**Shigella sonnei** O-Antigen Inhibits Internalization, Vacuole Escape, and Inflammasome Activation

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**ABSTRACT** Two *Shigella* species, *Shigella flexneri* and *Shigella sonnei*, cause approximately 90% of bacterial dysentery worldwide. While *S. flexneri* is the dominant species in low-income countries, *S. sonnei* causes the majority of infections in middle- and high-income countries. *S. flexneri* is a prototypic cytosolic bacterium; once intracellular, it rapidly escapes the phagocytic vacuole and causes pyroptosis of macrophages, which is important for pathogenesis and bacterial spread. In contrast, little is known about the invasion, vacuole escape, and induction of pyroptosis during *S. sonnei* infection of macrophages. We demonstrate here that *S. sonnei* causes substantially less pyroptosis in human primary monocyte-derived macrophages and THP1 cells. This is due to reduced bacterial uptake and lower relative vacuole escape, which results in fewer cytosolic *S. sonnei* and hence reduced activation of caspase-1 inflammasomes. Mechanistically, the O-antigen (O-Ag), which in *S. sonnei* is contained in both the lipopolysaccharide and the capsule, was responsible for reduced uptake and the type 3 secretion system (T3SS) was required for vacuole escape. Our findings suggest that *S. sonnei* has adapted to an extracellular lifestyle by incorporating multiple layers of O-Ag onto its surface compared to other *Shigella* species.

**IMPORTANCE** Diarrheal disease remains the second leading cause of death in children under five. *Shigella* remains a significant cause of diarrheal disease with two species, *S. flexneri* and *S. sonnei*, causing the majority of infections. *S. flexneri* are well known to cause cell death in macrophages, which contributes to the inflammatory nature of *Shigella* diarrhea. Here, we demonstrate that *S. sonnei* causes less cell death than *S. flexneri* due to a reduced number of bacteria present in the cell cytosol. We identify the O-Ag polysaccharide which, uniquely among *Shigella* spp., is present in two forms on the bacterial cell surface as the bacterial factor responsible. Our data indicate that *S. sonnei* differs from *S. flexneri* in key aspects of infection and that more attention should be given to characterization of *S. sonnei* infection.

**KEYWORDS** O-Antigen, *Shigella*, host-pathogen interactions, inflammasomes, macrophages
and Asia, *S. flexneri* accounts for 66% of cases and *S. sonnei* 24% of cases (2). However, in areas with good socioeconomic conditions and a high gross domestic product per capita, such as North America and Europe, *S. sonnei* is responsible for up to 80% of infections (3). Transitional countries that have recently undergone socioeconomic improvements show a shift from *S. flexneri* to *S. sonnei* as the dominant species (4–6). As a number of large populous countries undergo this shift (e.g., Brazil, India, and China), *S. sonnei* is emerging as an important pathogen.

The pathogenesis of *S. sonnei* is poorly understood and generally assumed to be similar to *S. flexneri*. The growing importance of *S. sonnei* has led to a reevaluation of its pathogenesis and has revealed some important differences from *S. flexneri*. These include a novel adhesin (7, 8), an antibacterial type 6 secretion system (T6SS) (9), and a group 4 capsule (G4C), which protects it from serum-mediated killing (10). Both species have a homologous type 3 secretion system (T3SS) that promotes secretion of effectors into host cells.

Unlike other *Shigella* species which contain multiple serotypes, there is only one *S. sonnei* serotype. The genes encoding biosynthesis and export of the O-Ag are encoded on the pSS virulence plasmid and were horizontally acquired from *Plesiomonas shigelloides*. In all other *Shigella* spp., these genes are located on the chromosome (11). *S. sonnei* O-Ag is composed of two unusual sugars, 2-acetamido-2-deoxy-L-altruronic acid and 2-acetamido-2-deoxy-L-fucose, which are not present in the O-Ags of other *Shigella* spp. or indeed in many bacteria (12). Importantly, the G4C of *S. sonnei* is also composed of the O-Ag polysaccharide, linked to an unknown lipid anchor rather than the lipid A/core as in the lipopolysaccharide (LPS) (10). Therefore, the surface of *S. sonnei* is covered with two O-Ag layers.

Pyroptotic cell death is considered an important component of *S. flexneri* pathogenesis (13), allowing *S. flexneri* to escape macrophage-mediated killing, induce local inflammation, and invade epithelial cells from the basolateral side (14). In the canonical pathway for caspase-1 activation and pyroptosis, NOD and leucine-rich repeat containing proteins with CARD or PYD (NLRCs or NLRPs), AIM2-like receptors or Pyrin protein can respond to pathogen- and/or danger-associated molecular patterns. This leads to the assembly of the sensor, e.g., NLRC3 or NLRC4, and the adaptor protein, ASC, into a signaling platform, known as the inflammasome, which activates caspase-1 (15). In the noncanonical pathway, caspase-4 directly senses and is activated by cytosolic LPS (16). Active caspase-1 and active caspase-4 can cleave gasdermin-D (GSDMD) (17). Once cleaved, the N-terminal of GSDMD forms pores in the cell membrane to cause swelling and membrane rupture. The proinflammatory cytokines interleukin-1β (IL-1β) and IL-18 are also cleaved by active caspase-1 into their mature forms and released (18, 19).

*S. flexneri* can activate the NLRC4 and NLRP3 inflammasomes (20). The T3SS needle and rod proteins (MxiH and MxiI, respectively) are recognized by hNaip/mNaip1 and mNaip2 proteins, which interact with NLRC4 and promote caspase-1 activation (21, 22). NLRP3 senses decreased cytosolic potassium levels and activates caspase-1 (23). A T3SS effector, Ipah7.8, has been shown to be important for activation of both the NLRC4 and NLRP3 inflammasomes (20). In the case of *Shigella*, it is unclear whether pyroptosis benefits the host or the bacteria. *S. flexneri* is thought to use pyroptosis to escape the macrophage and infect epithelial cells. However, recent studies using *Salmonella* suggest that pyroptosis results in killing of bacteria by forming pore-induced intracellular traps (24) or GSDMD targeting of bacterial membranes (25). It is currently unknown whether *S. sonnei* activates the same inflammasomes as *S. flexneri* and whether this is beneficial for the host or bacteria.

In this study, we demonstrate for the first time that *S. sonnei* induces caspase-1-dependent pyroptosis of human macrophages. However, we observed that equivalent bacterial inocula induced much less cell death for *S. sonnei* than *S. flexneri*. We show this is due to the O-Ag of *S. sonnei*, which reduces internalization and vacuole escape, resulting in less cytosolic bacteria. Our studies reveal an important role for the *S. sonnei* O-Ag in regulating bacterial interactions with macrophages, with one consequence being a reduction in inflammatory cell death.
**RESULTS**

*S. sonnei* induces less macrophage cell death than *S. flexneri*. Previous research into the interactions of *Shigella* with macrophages has largely focused on *S. flexneri*, which robustly induces pyroptosis in macrophages (20). To investigate whether *S. sonnei* behaved in a similar manner, we infected primary human CD14⁺ monocyte-derived macrophages (hMDMs) and measured the uptake of propidium iodide (PI), as an indicator of membrane damage that precedes pyroptosis. Unexpectedly, *S. sonnei* induced 50% less PI uptake than *S. flexneri* (Fig. 1A).

Similar experiments in phorbol-12-myristate-13-acetate (PMA)-differentiated THP1 cells recapitulated the reduced PI uptake during *S. sonnei* infection compared to *S. flexneri* (Fig. 1B). In addition, a lactate dehydrogenase (LDH) release assay comparing lytic cell death showed *S. sonnei* induced less cell death than *S. flexneri* (Fig. 1C). To ensure the reduced cell death was not a unique feature of the widely used *S. sonnei* strain 53G, we included a recent clinical isolate, *S. sonnei* 381, alongside *S. sonnei* 53G and compared these to two different *S. flexneri* strains M90T (serotype 5a) and 2457T (serotype 2a). Notably, both *S. sonnei* strains induced lower PI uptake in macrophages (Fig. 1B).

There are fewer cytosolic *S. sonnei* than *S. flexneri*. Induction of macrophage cell death by *S. flexneri* requires the bacteria to be cytosolic, which entails two steps: internalization and vacuole escape. We hypothesized that differences in these processes between *S. flexneri* and *S. sonnei* might be responsible for the differences in cell death observed. To investigate why *S. sonnei* induced less cell death, we treated hMDMs or PMA-treated THP1 cells with 50 μM Z-VAD-fmk, a pan-caspase inhibitor, to inhibit cell death (Fig. S1A in the supplemental material) and performed a gentamicin protection assay to calculate the number of intracellular bacteria (Fig. 1D and E). *S. sonnei*-infected macrophages had reduced numbers of intracellular bacteria compared to *S. flexneri*.

As the earliest time point that can be measured in the gentamicin protection assay is 1 h 40 min postinfection, it is possible bacteria were already killed by this time point, which would misrepresent the relative efficiency of internalization (because internalized and killed bacteria would not be detected). To address this, we enumerated intracellular bacteria by differential staining at 40 min postinfection, which confirmed that fewer THP1 cells harbored intracellular bacteria when infected with *S. sonnei* than when infected with *S. flexneri* (Fig. 1F).

Internalized *S. flexneri* rapidly lyse the phagocytic/endosomal vacuole in order to access the cell cytosol and escape lysosomal degradation (26). To investigate how well *S. sonnei* escaped into the cytosol, we used chloroquine, an antibiotic that only accumulates in vacuoles at high enough concentrations to kill bacteria, allowing discrimination between cytosolic and vacuolar bacteria (27). *S. sonnei* showed a reduction in vacuole escape compared to *S. flexneri* (Fig. 1G). Taken together, these data indicated there are less cytosolic *S. sonnei* compared to *S. flexneri* at the same multiplicity of infection (MOI), which may result in the reduced macrophage cell death observed with *S. sonnei*. By increasing the *S. sonnei* MOI to obtain equivalent numbers of cytosolic bacteria to *S. flexneri* (Fig. S1B), *S. sonnei* and *S. flexneri* induced similar levels of cell death (Fig. 1H and J) and cell lysis (Fig. S1C). These findings confirm that cytosolic bacteria are required for induction of cell death in *S. sonnei* and *S. flexneri* and that *S. sonnei* does not access the cytosol as efficiently as *S. flexneri*.

The T3SS is required for vacuole escape but not internalization of *S. sonnei*. The T3SS of *S. flexneri* is required for bacteria to lyse the phagocytic vacuole and access the cytosol (28). Consistent with this, a *S. sonnei* T3SS mutant (∆mxiD) had an impaired ability to escape the vacuole (Fig. 2A) and reduced cell death measured by PI uptake (Fig. 2B) and LDH release (Fig. S1D). The *S. sonnei* T3SS was required to induce vacuole lysis and hence produce cytosolic bacteria.

It is unclear whether *Shigella* internalization into macrophages is predominantly T3SS-dependent invasion or phagocytic uptake. T3SS-mediated invasion of epithelial
cells by *S. flexneri* triggers extensive membrane recruitment to engulf the bacteria. To visualize *S. flexneri* and *S. sonnei* uptake, we performed scanning electron microscopy (SEM) on infected cells and were able to see membrane recruitment around attached *S. flexneri* but not *S. sonnei* (Fig. 2C and D). Since phagocytic uptake and T3SS-mediated...
invasion both involve membrane rearrangement, these would be difficult to distinguish visually. Instead, we performed gentamicin protection assays with wild-type (WT) and T3SS mutant strains to quantify the number of intracellular bacteria in macrophages. Our experiments showed that in both hMDMs and THP1 cells, internalization into macrophages was T3SS dependent for *S. flexneri* but not *S. sonnei* (Fig. 2E and F). This suggested that the majority of *S. flexneri* actively invaded macrophages, in contrast to *S. sonnei*, which were mainly internalized by phagocytic uptake.

*S. sonnei* and *S. flexneri* induce similar pyroptosis pathways in infected macrophages. Given that cytosolic bacteria induce cell death through inflammasome
activation, we characterized the inflammasome pathways activated by *S. sonnei*. Since the *S. flexneri* inflammasome activators MxiH, MxiI, and IpaH7.8 proteins are 100% identical between *S. sonnei* 53G and *S. flexneri* M90T, we hypothesized they would activate the NLRC4 inflammasome. At comparable levels of cytosolic bacteria, similar activation of caspase-1 and proteolytic cleavage of GSDMD and IL-18 were observed (Fig. 3A). The involvement of the inflammasome pathway was confirmed using ASC<sup>mRFP</sup> THP1 cells, which revealed that both bacteria induced comparable levels of cells with ASC-containing inflammasome foci during infection (Fig. 3B and C). Further, infected GSDMD-silenced THP1 cells (THP<sub>GSDMD-miR</sub>, validated in Fig. S2A and B) underwent reduced cell death, suggesting pyroptosis is the dominant type of cell death induced by *S. sonnei* and *S. flexneri* (Fig. 3D).

Cells deficient in caspase-4 showed reduced pyroptosis (Fig. 3E), however, loss of caspase-1 almost completely abolished pyroptosis (Fig. 3F; all knockout cells are validated in Fig. S2C to E), indicating the canonical pathway of pyroptosis predominates in *S. sonnei*- and *S. flexneri*-infected macrophages. Treatment with the NLRP3 inhibitor, MCC950 (29), did not markedly affect cell death (Fig. 3G and validated in Fig. S2F), suggesting that NLRP3 plays a minor role in pyroptosis. ASC-deficient THP1 cells showed a partial reduction in cell death levels compared to WT THP1 cells (Fig. 3H). Taken together, these results are consistent with NLRC4 activation contributing to pyroptosis during *S. sonnei* infection of human macrophages, which is similar to previous reports for *S. flexneri*.

The T6SS and LVP instability do not account for reduced cell death caused by *S. sonnei*. All *Shigella* spp. harbor a large virulence plasmid (LVP) that encodes the T3SS, its effectors and additional important virulence factors. The LVP of *S. sonnei* is less stable than *S. flexneri* due to the evolution of different toxin-antitoxin systems (30). We inserted an antibiotic resistance cassette onto the LVP to create a stabilized LVP and used this strain to test whether LVP loss affected the amount of cell death that was induced. The LVP stabilized *S. sonnei* induced similar cell lysis as WT *S. sonnei*, indicating that differences in plasmid retention was not responsible for the altered interaction with macrophages (Fig. 4A).

Even though the T6SS of *S. sonnei* has only been described to have antibacterial activity (9), T6SSs from other bacteria (e.g., *Francisella tularensis* [31, 32]) have activity within macrophages. We therefore created a *S. sonnei* T6SS mutant (ΔtssB) to determine whether there was any contribution by the T6SS to cell death but found no difference in LDH release (Fig. 4A), indicating that the T6SS was not responsible for the altered interaction with macrophages. Altogether, these results ruled out the loss of LVP or a contribution by the T6SS in the reduced cell death observed for *S. sonnei*.

*S. sonnei* O-Ag prevents internalization into macrophages. *S. sonnei* O-Ag is incorporated into the G4C as well as being attached to the lipid A/core of LPS (Fig. 4B). The incorporation of the O-Ag into LPS and G4C is genetically separable, which we exploited to investigate their respective roles in the interaction with macrophages. The G4C of *S. sonnei* reduces bacterial invasion of epithelial cells by impairing T3SS activity (10) and could therefore play a similar role in macrophage internalization. We confirmed that *S. sonnei* ΔG4C invaded HeLa cells more efficiently (see Fig. S3B). Uptake and pyroptosis induced by *S. sonnei* ΔG4C was statistically similar to WT bacteria, although we did observe slightly greater cell death with the ΔG4C mutant (Fig. 4C and D). This was consistent with predominantly phagocytic uptake of *S. sonnei* by macrophages.

We then deleted the O-Ag synthesis operon (genes *wbg*<sub>T</sub> to *wbg*<sub>2</sub>) (33) to create a strain devoid of all O-Ag (both LPS and G4C linked) (Fig. 4B). This strain (ΔO-Ag) demonstrated increased internalization and cell death compared to *S. sonnei* Δg4c (Fig. 4C and D) and WT *S. sonnei*. In contrast, an LPS O-Ag-deficient strain (ΔwaaL), which retains the G4C, showed equivalent internalization as WT *S. sonnei* (see Fig. S3A). Therefore, the presence of the *S. sonnei* O-Ag per se, rather than specifically the O-Ag
in the capsule or attached to the lipid A/core, impedes macrophage internalization, and its complete removal enhances bacterial internalization.

We have shown that S. sonnei cell death is T3SS dependent due to the requirement for cytosolic bacteria. The T3SS tip accessibility has previously been shown to be

**FIG 3**  S. sonnei and S. flexneri induce similar pyroptosis pathways when normalized for numbers of cytosolic bacteria. (A) Immunoblots were performed on Shigella-infected THP1 cells to visualize cleavage of caspase-1, GSDMD, and IL-18 at 3 h postinfection. (B and C) ASC-IRFP THP1 cells expressing nontargeting miRNA or GSDMD-targeting miRNA were infected with S. sonnei (MOI of 20) or S. flexneri (MOI of 5) for 3 h. Cell death was measured by PI uptake over a 3-h time course and is plotted as the AUC. ***, P < 0.001** (by one-way ANOVA, n = 3). (D) THP1 cells left untreated or treated with 5 μM MCC950 were infected with S. sonnei (MOI of 20) or S. flexneri (MOI of 5) for 3 h. Cell death was measured by PI uptake over a 3-h time course and is plotted as the AUC. ****, **P < 0.0001** (by one-way ANOVA with Tukey’s multiple-comparison test, n = 4). (E) Control THP1 cells (Casp1 KO) were infected with S. sonnei (MOI of 20) or S. flexneri (MOI of 5). Cell death was measured by PI uptake over a 3-h time course and is plotted as the AUC. ****, **P < 0.001**; ***, P < 0.0001** (by one-way ANOVA with Tukey’s multiple-comparison test, n = 3). (F) Control THP1 cells (Casp1 KO) were infected with S. sonnei (MOI of 20) or S. flexneri (MOI of 5). Cell death was measured by PI uptake over a 3-h time course and is plotted as the AUC. ****, **P < 0.001**; ***, P < 0.0001** (by one-way ANOVA with Tukey’s multiple-comparison test, n = 3). (G) Control THP1 cells (Casp1 KO) were infected with S. sonnei (MOI of 20) or S. flexneri (MOI of 5). Cell death was measured by PI uptake over a 3-h time course and is plotted as the AUC. ****, **P < 0.001**; ***, P < 0.0001** (by one-way ANOVA with Tukey’s multiple-comparison test, n = 3). (H) Control THP1 cells (Casp1 KO) were infected with S. sonnei (MOI of 20) or S. flexneri (MOI of 5). Cell death was measured by PI uptake over a 3-h time course and is plotted as the AUC. ****, **P < 0.001**; ***, P < 0.0001** (by one-way ANOVA with Tukey’s multiple-comparison test, n = 3).
enhanced upon removal of the G4C and further exposed by removal of the O-Ag (10).

We therefore hypothesized that the O-Ag was impeding T3SS-mediated invasion. To test this, we created a T3SS mutant in the O-Ag-deficient strain (ΔO-AgΔmxID). In keeping with our hypothesis, this strain had wild-type levels of internalization (Fig. 4E).
but impaired activation of cell death because it is unable to escape the vacuole. To further investigate the role of the O-Ag in shielding the T3SS, we complemented the O-Ag mutant (ΔO-Ag) with either the *S. sonnei* O-Ag synthesis operon (pSS) or the *S. flexneri* 5a O-Ag synthesis and modification operons (pSf5a) (34–36) (Fig. 4E). Both complemented strains impeded internalization of *S. sonnei* (Fig. 4F) and, as a consequence, reduced the level of cell death similar to those observed with WT *S. sonnei* (Fig. 4F). Interestingly, complementation with pSf5a produced a *S. flexneri*-like O-Ag ladder that migrated differently on SDS-PAGE than when expressed in *S. flexneri*. To determine whether this was due to different modal length of O-Ag controlled by WzzB, we introduced the *wzzB*Sf onto the pSf5a complementation plasmid (pSf5a/wzzB). In this strain the modal length of the O-Ag resembled that of the WT *S. flexneri*; however, a low level of expression was observed. The levels of internalization and cell death were not reduced to the levels of the WT *S. sonnei* and instead resembled the levels of the O-Ag mutant.

**DISCUSSION**

*S. flexneri* is known to induce pyroptosis in macrophages. This is considered a key step in the pathogenesis of *Shigella* since it allows bacteria to infect epithelial cells from the preferred basolateral side and leads to bacterial dissemination. In addition, pyroptosis creates an inflammatory response causing the recruitment of neutrophils, which disrupts the epithelial cell barrier and allows more *Shigella* to traverse the epithelial layer (37).

Here, we present evidence that *S. sonnei* does not use the same mechanisms during infection as *S. flexneri* (summarized in Fig. 5). In line with previous reports, we found that *S. flexneri* induces rapid pyroptosis upon internalization of infected macrophages (20, 22). However, *S. sonnei* induced markedly less macrophage cell death, which was
the result of a decreased number of cytosolic bacteria through a combination of fewer internalized S. sonnei and impaired vacuole escape. The requirement for cytosolic bacteria in the induction of inflammasomes was consistent for both S. sonnei and S. flexneri. Additional host responses are also likely to be affected by the reduced number of cytosolic bacteria for S. sonnei compared to S. flexneri.

Once S. sonnei and S. flexneri cytosolic numbers were normalized, pyroptosis proceeded via similar pathways and to similar levels. For both species, cell death was predominantly dependent on GSDMD and caspase-1, indicating the canonical inflammasome pathway is induced by Shigella. There may be a minor contribution to cell death for the noncanonical pathway, since immunoblots indicated that caspase-4 was activated by infection by both S. sonnei and S. flexneri and that caspase-4 deficiency or NLRP3 inhibition led to less pyroptosis over time than control cells. However, this difference was minor compared to that observed for ASC or caspase-1-deficient cells. NLRC4 has a caspase-recruitment and activation domain (CARD), which can enable its interaction with and activation of caspase-1 directly, bypassing the need for ASC (38–40). This suggests that the NLRC4 inflammasome has a prominent role in the cell death of S. sonnei-infected THP1 cells. These results are in line with those shown previously for S. flexneri, which suggest both the NLRP3 and NLRC4 inflammasomes are involved in S. flexneri-mediated macrophage death (20).

Interestingly, S. sonnei was able to reduce internalization into macrophages in an O-Ag-dependent manner. The O-Ag contributes to host immune evasion, and its role in evasion of complement mediated killing is well characterized (41). There are also examples of the O-Ag affecting cellular interactions, including impeding recognition and internalization by epithelial cells (Salmonella Typhimurium [42]) and macrophages (Burkholderia cenocepacia [43]). The modal length of the O-Ag from Salmonella Typhimurium or S. flexneri serotype 2a is important for T3SS-mediated invasion into macrophages and epithelial cells, respectively (44). Similarly, glucosylation of the S. flexneri serotype 5a O-Ag, which reduces the O-Ag length by half, enhances its invasiveness (45).

Unexpectedly, in our study the internalization of S. sonnei into macrophages was independent of its T3SS. This is in contrast to S. flexneri, which exhibits significant T3SS-mediated invasion into macrophages. This suggests that macrophage internalization is a combination of bacterium-driven invasion and phagocytic uptake for S. flexneri, whereas S. sonnei internalization is almost exclusively due to phagocytic uptake. S. sonnei O-Ag is incorporated into both the G4C and the LPS of S. sonnei. Only when all of the O-Ag layers of S. sonnei are removed can S. sonnei efficiently invade macrophages. The accessibility of Ipab was previously shown to increase upon removal of the G4C, and a further increase was observed for an O-Ag-deficient strain, indeed suggesting that the lipid A/core-linked O-Ag also contributes to shielding of the T3SS (10). The ability of the serotype 5a O-Ag synthesis and modification operon from S. flexneri to prevent the internalization of O-Ag-deficient S. sonnei indicates that the composition of the saccharides is not important for this phenotype. Furthermore, the inability of S. flexneri O-Ag when regulated by wzzB to complement for internalization of cell death suggests the modal length of the O-Ag is important. However, this strain also produced a small amount of O-Ag, and we cannot discount this as the reason for the failure to complement. Our data, and previously published data regarding the accessibility of the T3SS, support the conclusion that the O-Ag acts as a physical barrier to T3SS-mediated invasion rather than being antiphagocytic.

The results presented here, combined with previous investigations, indicate that S. sonnei and S. flexneri use different infection mechanisms. These mechanisms are also different from related Gram-negative enteric pathogens such as Salmonella spp. or enteropathogenic Escherichia coli, which also activate distinct inflammasome pathways in human macrophages (46–49). Increasing evidence points to S. sonnei being more adapted to an extracellular lifestyle since, compared to S. flexneri, it invades epithelial cells and macrophages poorly. This may partly explain the dominance of S. sonnei in developed countries, where improved living conditions, including reduced overcrowd-
ing and hence the person-to-person spread of pathogens, fails to lower \textit{S. sonnei} infection rates. These studies highlight that further investigation into \textit{S. sonnei} is required in order to implement appropriate measures to reduce infection rates.

**MATERIALS AND METHODS**

**Bacterial strains and growth.** Unless otherwise stated, all \textit{Shigella} strains (see Table S1 in the supplemental material) were routinely grown in tryptone soy broth (TSB) at 37°C with shaking at 200 rpm. Antibiotic selection was used when necessary as follows: 100 \( \mu \)g/ml ampicillin, 50 \( \mu \)g/ml kanamycin (Kn), 12.5 \( \mu \)g/ml chloramphenicol (Cm), 100 \( \mu \)g/ml erythromycin, 50 \( \mu \)g/ml streptomycin (Sm), and 10 \( \mu \)g/ml gentamicin (Gm).

**Cloning and mutagenesis.** \textit{S. sonnei} LVP\textsuperscript{stable}, \textit{ΔwaaL}, \textit{ΔtssB}, and \textit{ΔO-Ag} strains were constructed as follows (primer sequences are in Table S2 in the supplemental material). For \textit{S. sonnei} LVP\textsuperscript{stable} nucleotides (nt) 82936 to 83715 and nt 83716 to 84215 were amplified using primers 1 and 2 and primers 3 and 4. The chloramphenicol cassette was amplified from pKD3 using primers 21 and 22. Overlapping PCR was used to construct the mutagenesis fragment 82936–83715-Cm-83716–84215; note that the P1–P2 fragment was inserted in the opposite orientation). This fragment was further amplified by PCR with primers 1 and 4. Then, 2 \( \mu \)g of PCR product was electroporated into \textit{S. sonnei} using primers 6 and 22. Overlapping PCR was used to construct the mutagenesis fragment consisting of 5’ tssB-Kn-3’ tssB. This fragment was further amplified by PCR with the primers 6 and 9. Then, 2 \( \mu \)g of PCR product was electroporated into \textit{S. sonnei} 53G/pKD46 induced with 1 mM L-arabinose for 45 min to express Lambda Red recombinase genes. The electroporation was plated on TSB supplemented with Cm. Genomic insertion of \textit{cat} was verified by PCR using primers 5 and 22.

For the \textit{S. sonnei} \textit{ΔtssB} mutant, 500-bp fragments flanking tssB were amplified using primers 6 and 7 and primers 8 and 9. The kanamycin cassette was amplified from pKD4 using primers 21 and 22. Overlapping PCR was used to construct the mutagenesis fragment consisting of 5’ tssB-Kn-3’ tssB. This fragment was further amplified by PCR with the primers 6 and 9. Then, 2 \( \mu \)g of PCR product was electroporated into \textit{S. sonnei} 53G/pKD46 induced with 1 mM L-arabinose for 45 min to express Lambda Red recombinase genes. The electroporation was plated on TSB supplemented with Kn. Genomic insertion of \textit{kan} was verified by PCR using the primers 10 and 22.

For the \textit{S. sonnei} \textit{ΔwaaL} mutant, 500-bp fragments flanking waaL were amplified using primers 11 and 12 and primers 13 and 14. The kanamycin cassette was amplified from pKD4 using primers 21 and 22. Overlapping PCR was used to construct the mutagenesis fragment consisting of 5’ waaL-Kn-3’ waaL. This construct and pSEVA612S were digested with BamH and EcoRI, ligated, and transformed into \textit{E. coli} CC118. The resulting plasmid, pSEVA\textit{ΔwaaL-Kn}, was conjugated into \textit{S. sonnei} 53G. Briefly, 20 \( \mu \)l of helper \textit{E. coli} 1047 pRK2013 was incubated for 2 h at 37°C with 20 \( \mu \)l of the donor strain \textit{E. coli} CC118-lopir pSEVA\textit{ΔwaaL} on Luria-Bertani (LB) agar. Then, 40 \( \mu \)l of the receiver strain (\textit{S. sonnei} 53G/pACBSR) was added, and the plate was incubated for 4 h at 37°C. Conjugants were selected on TSB agar supplemented with Cm and Sm. Individual colonies were grown in TSB supplemented with Sm and 0.4% (wt/vol) L-arabinose (Sigma) for 8 h to induce expression of the I-SceI endonuclease from pACBSR and then plated on Kn plates. Genomic deletion of \textit{waaL} was verified by PCR using primers 15 and 22. The strains were passaged several times in liquid TSB to remove pACBSR, and bacteria sensitive to Sm were selected.

\textit{S. sonnei ΔO-Ag} mutant was constructed by amplifying 500-bp fragments upstream of \textit{wbgT} and downstream of \textit{wbgZ} using primers 16 and 17 and primers 18 and 19. The kanamycin cassette was amplified from pKD4 using primers 21 and 22. Overlapping PCR was used to construct the mutagenesis fragment consisting of 5’ \textit{wbgT}-kan-3’ \textit{wbgZ}. This construct and pSEVA612S were digested with BamH and EcoRI, ligated, and transformed into \textit{E. coli} CC118-lopir. The resulting plasmid, pSEVA\textit{ΔO-Ag-Kn}, was conjugated into \textit{S. sonnei} 53G, as described above. Genomic deletion of \textit{ΔO-Ag} was verified by PCR using primers 20 and 22.

Complementation vectors were constructed using standard molecular biology techniques. The 53G O-Ag operon was amplified with primers 23 and 24. The PCR product and pSEVA471 were digested with BamH and ligated to create pO-Ag\textsuperscript{53G}. The M90T \textit{gtt} operon was amplified with primers 25 and 26. The PCR product and pSEVA471 were digested with KpnI and BamH and ligated to create pSEVA471-gtt. The M90T O-Ag operon was amplified with primers 25 and 26. The PCR product and pSEVA471 were digested with KpnI and BamH and ligated to create pSEVA471-gtt. The resulting plasmid, pSEVA\textit{ΔO-Ag-Kn}, was conjugated into \textit{S. sonnei} 53G, as described above. Genomic deletion of \textit{ΔO-Ag} was verified by PCR using primers 20 and 22.

**Cell culture and infection.** THP-1 cells were maintained in Roswell Park Memorial Institute (RPMI) medium supplemented with 10% heat-inactivated fetal bovine serum (FBS), 5 mM HEPES, 5 mM sodium pyruvate, 100 \( \mu \)g/ml penicillin, and 100 \( \mu \)g/ml streptomycin. Cells were seeded at 7.5 × 10^5 cells/ml 72 h prior to infection in complete RPMI plus 100 ng/ml PMA. At 24 h prior to infection, the medium was replaced with phenol-red free, PMA-free complete RPMI medium. HeLa cells were maintained in Dulbecco modified Eagle medium (1,000 mg/liter glucose) supplemented with 10% FBS. Cells were seeded at 1 × 10^5 cells/ml 24 h prior to infection. All cell lines were incubated at 37°C and 5% CO\textsubscript{2}. Cells were infected with the indicated MOI and centrifuged for 10 min at 600 × g to synchronize infection. At 30 min postcentrifugation, Gm (150 \( \mu \)g/ml) was added directly to wells for the remainder of the experiment. Where indicated, inhibitors—Z-VAD-fmk (50 \( \mu \)M; R&D Systems) or MCC950 (5 \( \mu \)M; Tocris Bioscience)—were added to cells 1 h prior to infection. To induce NL3P3-driven caspase-1 activation, the cells were primed with ultrapure O111:B4 LPS (250 ng/ml; Invivogen) for 3 h and then treated with nigericin (20 \( \mu \)M; Sigma) for 45 min. To induce caspase-4 activation, unprimed cells were transfected with LPS (5 \( \mu \)g/ml)/H9262 using Lipofectamine 2000 (1% [wt/vol]; Invitrogen).

Infected HeLa cells were washed and fixed in 2.5% glutaraldehyde for analysis by SEM at an accelerating voltage of 25 kV using a JEOL JSM-5300 scanning electron microscope (JEOL, Herts, UK).
Generation of cell lines. The THP1 GSDMD<sup>miR</sup> cell line was described previously (50); THP1 Casp1 knockout (KO), THP1 Casp4-KO, and THP1 ASC KO cells were all kindly provided by Veit Hornung (51).

Isolation of primary hMDMs. Leukocytes cones were obtained from the NHS blood and transfusion service (from anonymous healthy donations), as previously described (46). Blood from each donor was diluted 1:4 with phosphate-buffered saline (PBS), transferred into a LeucoSep tube (Greiner Bio-One), and centrifuged at 1,000 g × g for 20 min at room temperature (slow acceleration and deceleration to prevent disturbance of the layers) to obtain the buffy coat containing white blood cells. This was separated and washed three times with RPMI. Cell were washed with MACS buffer (50 mg/ml bovine serum albumin [BSA] and 2 mM EDTA in PBS). CD14<sup>+</sup> cells were isolated by MACS using biotinylated anti-CD14<sup>+</sup> antibody and anti-biotin microbeads according to the manufacturer’s protocol (Miltenyi Biotec). Monocytes were cultured in complete RPMI plus 20 ng/ml recombinant human macrophage colony-stimulating factor (M-CSF) for 7 days to promote differentiation into hMDMs. The medium was replaced with complete RPMI lacking antibiotics and M-CSF 24 h prior to infection.

Internalization and vacuole escape assays. To prevent cell death, cells were treated with Z-VAD-fmk (50 μM) 1 h prior to infection. The cells were infected with bacteria, as described above. For internalization assays, the cells were washed with serum-free RPMI and lysed with TritonX-100 (0.5%) at 1.5 h postinfection. For vacuole escape assays, the cells were treated 30 min postinfection either with 200 μg/ml chloroquine and 150 μg/ml gentamicin or with 150 μg/ml gentamicin alone for 1 h and then lysed with Triton X-100 (0.5%). Serial dilutions were prepared, plated on LB agar, and incubated overnight at 37°C.

PI uptake assays. Cells and bacterial strains were prepared as described above. Prior to infection, the cells were supplemented with 5 μg/ml PI (Invitrogen). For time course assays, fluorescence was measured at 630 nm every 10 min with a POLARStar 623 Omega plate reader (BMG Labtech) (50). Uninfected controls treated with Triton X-100 (0.05%) were used to calculate the percent uptake.

LDH assays. Infections were performed as described above. At 3 h postinfection, supernatants were harvested. An LDH assay was performed according to the kit instructions (CytoTox 96 nonradioactive cytotoxicity assay; Promega). The absorbance was measured at 490 nm, and values are expressed as percentages of the 100% lysis control. All values are normalized to the uninfected control.

Immunoblot. Infections were performed as described previously, except that prior to infection the cells were washed with PBS and infections were done in Opti-MEM plus 5 mM sodium pyruvate. Supernatants were precipitated in acetone (1:4 [vol/vol]) overnight at −20°C, the acetone was aspirated, and the samples were left to air dry. Cells were lysed in radioimmunoprecipitation assay buffer (120 mM Tris [pH 8.0], 300 mM NaCl, 2% NP-40, 1% sodium deoxycholate, 2 mM EDTA) supplemented with complete protease inhibitor and 1 mM phenylmethylsulfonyl fluoride. Laemmli buffer and 5% 2-mercaptoethanol were added to the lysates. Precipitated supernatants were resuspended in respective cell lysates to create pooled samples. Mouse anti-hcaspase-1 (AdipoGen), mouse anti-caspase-4 (Santa Cruz Biotechnology), goat anti-hIκBα (R&D Systems), and rabbit anti-hIL18 (MBL International) were used at 1:1,000 dilutions, and mouse anti-hGSDMD (Santa Cruz Biotechnology) was used at a 1:500 dilution.

Immunofluorescence microscopy. Cells were seeded and infected as described previously. To calculate the percentage of THP1 cells infected, in/out staining was performed as follows. At 40 min after the addition of bacteria (T = 0), the cells were washed three times with cold PBS. Rabbit anti-sonnei (1:100; phase 1 and 2 sera; Fisher Scientific) or rabbit anti-flexneri (1:500; serotype 5a sera; PHE) diluted in 2% BSA–PBS was added to the cells. The cells were then incubated with antibodies on ice for 30 min. Next, the cells were washed with cold PBS and incubated on ice with donkey anti-rabbit-Alexa 594. S. flexneri serotype 5a antibody (Abcam), followed by anti-rabbit horseradish peroxidase, and developed with chemiluminescence, or fixed and silver stained as previously described (52).

Statistical analysis. The number of independent repeats performed for each experiment was determined (indicated by "n" in the figure legends). One-way analysis of variance (ANOVA) or a Student t test was performed to compare means, as implemented in GraphPad Prism 8. Errors bars represent the standard errors of the means throughout.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/mBio.02654-19.

FIG S1, TIF file, 1.5 MB.

FIG S2, TIF file, 1.5 MB.

FIG S3, TIF file, 1.5 MB.
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