TYPE Original Research
PUBLISHED 08 November 2022
DOI 10.3389/fmicb.2022.999183



OPEN ACCESS

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SPECIALTY SECTION

This article was submitted to Microbial Physiology and Metabolism, a section of the journal Frontiers in Microbiology

RECEIVED 20 July 2022 ACCEPTED 21 October 2022 PUBLISHED 08 November 2022

CITATION

Lovelace AH, Chen H-C, Lee S, Soufi Z, Bota P, Preston GM and Kvitko BH (2022) RpoS contributes in a host-dependent manner to *Salmonella* colonization of the leaf apoplast during plant disease. *Front. Microbiol.* 13:999183. doi: 10.3389/fmicb.2022.999183

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RpoS contributes in a host-dependent manner to Salmonella colonization of the leaf apoplast during plant disease

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Contaminated fresh produce has been routinely linked to outbreaks of Salmonellosis. Multiple studies have identified Salmonella enterica factors associated with successful colonization of diverse plant niches and tissues. It has also been well documented that *S. enterica* can benefit from the conditions generated during plant disease by host-compatible plant pathogens. In this study, we compared the capacity of two common S. enterica research strains, 14028s and LT2 (strain DM10000) to opportunistically colonize the leaf apoplast of two model plant hosts Arabidopsis thaliana and Nicotiana benthamiana during disease. While S. enterica 14028s benefited from co-colonization with plantpathogenic Pseudomonas syringae in both plant hosts, S. enterica LT2 was unable to benefit from Pto co-colonization in N. benthamiana. Counterintuitively, LT2 grew more rapidly in ex planta N. benthamiana apoplastic wash fluid with a distinctly pronounced biphasic growth curve in comparison with 14028s. Using allelic exchange, we demonstrated that both the N. benthamiana infectiondepedent colonization and apoplastic wash fluid growth phenotypes of LT2 were associated with mutations in the S. enterica rpoS stress-response sigma factor gene. Mutations of S. enterica rpoS have been previously shown to decrease tolerance to oxidative stress and alter metabolic regulation. We identified rpoSdependent alterations in the utilization of L-malic acid, an abundant carbon source in N. benthamiana apoplastic wash fluid. We also present data consistent with higher relative basal reactive oxygen species (ROS) in N. benthamiana leaves than in A. thaliana leaves. The differences in basal ROS may explain the host-dependent disease co-colonization defect of the rpoS-mutated LT2 strain. Our results indicate that the conducive environment generated by pathogen modulation of the apoplast niche can vary from hosts to host even with a common disease-compatible pathogen.

KEYWORDS

Salmonella enterica, produce-borne illness, Pseudomonas syringae, rpoS, apoplast, hydrogen peroxide, biomultiplier

Introduction

Fresh produce has become an increasing source for foodborne illnesses over the last several decades. In the United States, fresh produce associated foodborne outbreaks increased from 0.7% of reported outbreaks in the 1970s to 33% in 2011 (Bennett et al., 2014). From 2004 to 2015, fresh produce was the leading cause of food borne illness outbreaks (Fischer and Plunkett, 2015). Salmonella enterica is a common causal agent of foodborne illnesses and a significant number of cases of Salmonellosis have been linked to consumption of fresh or minimally processed produce (Hanning et al., 2009; Berger et al., 2010; Kozak et al., 2013; Bennett et al., 2015; Crowe et al., 2015; Fischer and Plunkett, 2015; Carstens et al., 2019). Plant-associated Salmonellosis outbreaks, while infrequent, are both economically disruptive and a recurrent public health concern (Bennett et al., 2015; Crowe et al., 2015). In response to the impacts of plantassociated Salmonellosis, multiple studies have sought to characterize the abiotic and biotic factors that contribute to S. enterica colonization of both edible and non-edible portions of crop plants as well as model plant hosts (Cooley et al., 2003; Barak et al., 2005, 2009, 2011; Noel et al., 2010; Meng et al., 2013; Garcia et al., 2014; Potnis et al., 2014, 2015; de Moraes et al., 2017, 2018; Jacob and Melotto, 2019; Montano et al., 2020). Mutational and comparative genetic studies have identified factors involved in S. enterica plant attachment, internalization, persistence, and translocation between inoculated to non-inoculated portions of plant hosts (recently reviewed by Zarkani and Schikora, 2021). Based on these studies, different plant hosts and plant host niches often pose distinct fitness requirements for S. enterica colonization. These studies have also shown that S. enterica colonization of plants is not limited to external plant surfaces. S. enterica can become internalized through roots and is able to actively bypass stomata to colonize the leaf apoplastic space (Gu et al., 2011; Karmakar et al., 2018; Rodrigues Oblessuc et al., 2019; Chahar et al., 2021).

The majority of genetic studies that have identified S. enterica plant colonization factors have focused on binary plant host-S. enterica interactions. However, interactions between S. enterica and plants are also modulated by plantassociated microbes (Poza-Carrion et al., 2013). It has been well documented that the activities of bacterial plant pathogens can allow S. enterica to more effectively colonize various plant niches (Wells and Butterfield, 1997; Meng et al., 2013; Pollard et al., 2014; Potnis et al., 2014, 2015; Deblais et al., 2021; Cowles et al., 2022). Plant pathogens can act as biomultipliers for S. enterica, creating favorable conditions that allow S enterica to reach higher loads in plant tissues than they can achieve independently (Barak and Schroeder, 2012). For instance, the plant soft rot necrotrophic pathogen Pectobacterium carotovorum secretes plant cell wall degrading enzymes that disrupt plant tissues and increase nutritional availability to S. enterica (Cox et al., 2013; George et al., 2016). George et al. (2018) used a TnSeq strategy to identify S. enterica 14028s genetic factors contributing to successful co-colonization of *P. carotovorum* tomato soft rot. However, for hemibiotrophic plant pathogens like *Pseudomonas syringae*, that translocate type III secretion systems effectors (T3Es) to subvert plant immunity and cause disease, *S. enterica* may also benefit from other forms of pathogen-associated niche modulation (Barak and Schroeder, 2012).

Plants have a distributed innate immune system structured around two interconnected tiers of immune receptors, Nucleotidebinding domain \underline{L} eucine-rich \underline{R} epeat receptors (NLRs) and Pattern Recognition Receptors (PRRs; Ngou et al., 2022). NLRs detect specific plant pathogen effector proteins delivered into the plant cytosol and activate plant immunity in response to plant pathogens. Conversely, Pattern Recognition Receptors (PRRs) are presented on the plant cell surface and are able to bind to cognate conserved "non-self" microbial molecular patterns common to both plant pathogens and non-plant pathogens. The archetypal PRR is Arabidopsis thaliana FLS2, which binds to broadly conserved epitopes of bacterial flagellin forming a kinase signaling complex to activate plant immune signaling responses (Zipfel et al., 2004; Chinchilla et al., 2006). It has been demonstrated that S. enterica activates PRR-mediated immune responses in diverse plants (Meng et al., 2013; Garcia et al., 2014; Jacob and Melotto, 2019). PRR-mediated immunity results in non-permissive conditions in the apoplast that are widely suppressive to bacterial proliferation (Velasquez et al., 2022). PRR-compromised A. thaliana displayed dysbiosis resulting in elevated populations of non-plant-pathogenic bacterial endophytes (Xin et al., 2016; Chen et al., 2020). For a given plant host, disease-compatible P. syringae translocate T3Es that collectively suppress PRR-mediated immune signaling, either evade or suppress signaling by host NLRs, and increase water and nutrient availability in the apoplast (Xin et al., 2018; Lovelace and Ma, 2022). S. enterica has been previously observed to benefit from apoplastic water soaking caused by hemibiotrophic Xanthomonas plant pathogens (Potnis et al., 2015). From a sociomicrobiology framework, the plant pathogen-modulated conducive apoplastic niche provides "common goods" allowing other bacteria co-occupying the apoplast to hitch a free ride and derive a reproductive benefit (Macho et al., 2007; Rufian et al., 2018; Chen et al., 2020). However, we still lack a detailed understanding of the metabolic and stressor landscape that distinguishes the conducive niche generated by plant pathogens from the suppressive non-permissive apoplastic niche, as well as to what degree plant pathogen-modulated apoplastic niches vary from host-to-host or pathogen-to-pathogen.

Salmonella enterica subsp. enterica Typhimurium strains, 14028s and LT2 have been workhorse strains for the characterization of *S. enterica*, genetics, physiology, and host interactions and have been used routinely in studies of *S. enterica* plant colonization. In this study, we compared the competency of 14028s and LT2 (strain DM10000), to benefit from co-colonization of the leaf apoplastic space with the plant pathogen partner *P. syringae* pv. tomato DC3000 (*Pto*) in two

divergent model plant hosts, A. thaliana and Nicotiana benthamiana. Pto is a well characterized model plant pathogen that is natively disease compatible with A. thaliana Col-0 and is also disease compatible with the common laboratory accession of N. benthamiana when the effector gene hopQ is deactivated to prevent immune recognition by the Roq1 NLR receptor (Wei et al., 2007; Yao et al., 2013; Schultink et al., 2017). We determined that LT2 did not derive a benefit from co-colonization of the N. benthamiana leaf apoplast with Pto and linked that phenotype to a newly identified and presumed null allele of the rpoS sigma factor. We also determined that there are statistically elevated basal ROS levels in N. benthamiana leaves compared to Arabidopsis leaves. The differences in basal ROS may explain the host-dependent Pto infection-associated colonization phenotype of the rpoSmutated LT2 strain. In addition to identification of rpoS as a genetic factor contributing to *S. enterica* disease co-colonization in a host-dependent manner, our work is consistent with observations that the disease-associated conducive apoplastic niche can vary from host to host and pose distinct fitness requirements (Helmann et al., 2019).

Materials and methods

Plant growth and bacterial culturing conditions

Arabidopsis thaliana Col-0 seeds suspended in sterile 0.1% agarose were sown in SunGrow Professional potting mix and stratified in darkness for 1 day at 4°C before being grown in a growth chamber (Conviron A1000) with 14-h light (70 µmol) at 23°C. Plants were removed from the chamber at 4 weeks and kept at 12-h day and 12-h night conditions in the growth room prior to inoculation (4-5 weeks old) or apoplastic extractions (6–7 weeks). N. benthamiana were sown in the same potting mix amended with 1 g L⁻¹ Peter's 20-20-20 fertilizer and grown in a growth chamber with $12\,h\,day$ at 26°C (125 $\mu\text{mol})$ and 12 h night at 23°C. Two weeks after sowing, seedlings were transplanted into 6-inch pots and fertilized. Plants were removed from the chamber at 5 weeks and kept in the growth room prior to inoculations or apoplastic extractions (6-9 weeks old). Leaves from 4-week-old N. benthamiana plants were used for apoplastic fluid isolation. P. syringae pv. tomato strain DC3000 (Pto) strains, and S. enterica serovar Typhimurium strains used in this study are listed in Supplementary Table S1. All Pto strains were grown on King's B (KB) medium (per 11=20.0 g proteose peptone 3, 0.4 g MgSO₄·7H₂O, glycerol 10 ml, 2.0 g K₂HPO₄·3H₂O, with 18 g agar for solidified media) with $60\,\mu g\,mL^{-1}$ of rifampicin at 30°C. S. enterica strains were grown routinely in Lysogeny Broth (LB) medium (per 1 l = 10 g tryptone, 5 g yeast extract, 10 g NaCl, pH 7.5+ with 15 g agar for solidified media) at 37°C. To enumerate S. enterica populations from a mixed population, samples were grown at 42°C.

Plant inoculation and sampling procedures

To prepare S. enterica bacterial inocula, overnight cultures made from single colonies were incubated in LB at 37°C, pelleted using centrifugation, suspended in 0.25 mm MgCl₂, and were diluted to an optical density at $600\,\mathrm{nm}$ (OD₆₀₀) of 0.8 (approximately 5 × 108 CFU mL⁻¹), as determined using a Biospectrometer (Eppendorf, Hamburg, Germany). Pto inoculum was prepared as described in Lovelace et al. (2018) and diluted to OD_{600} = 0.8. Bacterial inocula for both individual and co-inoculations were further diluted in 0.25 mm MgCl₂ to the desired concentrations. S. enterica strains at $5 \times 10^5 \text{CFU} \, \text{mL}^{-1}$ (approximately LOG 3.5 CFU/cm²) were mixed with Pto or $Pto\Delta hrcC$ (a Type III Secretion System mutant) to a final Ptoconcentration of $5 \times 10^6 \, \text{CFU} \, \text{mL}^{-1}$ (approximately LOG 4.5 CFU/ cm²) and syringe-inoculated into four A. thaliana leaves per plant. S. enterica strains at 5×10^5 CFU mL⁻¹, were mixed with $Pto\Delta hopQ$ (disease compatible with *N. benthamiana*) or $Pto\Delta hrcC$ to a final concentration of 5×10^5 CFU mL⁻¹ and syringe inoculated into fully expanded N. benthamiana leaves. All inocula were serial diluted and plated on appropriate media with antibiotics to verify bacterial concentrations. All inoculated hosts were incubated for 3 days under high humidity (90-100% RH) at ambient room temperature. At 3 days post inoculation, four 4 mm diameter leaf discs (~0.5 cm² total) per treatment were collected with a 4 mm diameter biopsy punch. Leaf discs were macerated for 2 min in 0.1 ml of 0.25 mm MgCl2 using a SpeedMill Plus homogenizer (Analytik Jena, Jena, Germany) and the bacterial CFU/cm² leaf tissue was determined by serial dilution spot plating as described in Liu et al. (2015). Initial bacterial populations (CFU/cm² leaf tissue) from co-inoculations were determined from independent plant samples (Supplementary Figure S1). All plant inoculation assays were repeated three times with 3 plants per treatment. For co-inoculation assays, statistical differences between mean final bacterial populations were determined using a 2- tailed t-test for each strain (p < 0.05).

Extraction of apoplastic wash fluid and GC-MS analysis

Apoplastic wash fluid (AWF) was crude extracted using vacuum infiltration as described by O'Leary et al. (2014) with slight modifications. Whole A. thaliana plants or fully expanded N. benthamiana leaves were cut and placed into a 500 ml beaker with 300 ml of sterile distilled water. Repeated cycles of vacuum at 95 kPa for 2 min followed by slow release of pressure were applied until leaves were fully infiltrated. Excess water was blotted from plant tissue before leaves were rolled into 20 ml syringes which were placed into 50 ml conical tubes. Tubes were centrifuged at 1,000 rpm for 10 min at 4°C and the fractions were pooled and stored at $-80\,^{\circ}\text{C}$. AWF samples were filter sterilized using $0.2\,\mu\text{M}$ RapidFlow filters for subsequent experiments.

For metabolomic analyses, N. benthamiana apoplastic wash fluid (NbAWF) samples were collected as described by O'Leary et al. (2016). NbAWF samples were prepared for GC-MS analysis using a modified version of the method of Lisec et al. (2006). One hundred and fifty microliter samples of NbAWF were mixed with 700 µl of methanol, supplemented with 10 µg mL⁻¹ of ribitol, and then shaken at 70°C for 10 min, followed by centrifugation for 5 min at 11,000×g. Next, 700 μl of supernatant was removed and mixed sequentially by vortexing with 375 µl of cold chloroform and 500 μ l of cold ddH₂O. Samples were centrifuged at 2,200×g for 15 min, then 250 µl of supernatant was transferred to a fresh tube and dried in a vacuum concentrator without heat. The samples were derivatized in 29 µl of pyridine containing $20\,mg\,mL^{-1}$ methoxyamine and $50\,\mu l$ of N-Methyl-N-(trimethylsilyl) trifluoroaceamide (MSTFA) as described previously (Lisec et al., 2006). GC-MS analysis was performed as described previously (O'Leary et al., 2016). Area values of identified BAWF compounds from six biological replicates were averaged and normalized against a ribitol internal standard and standard deviation was calculated for each compound.

Bacterial growth assays

Bacterial inocula were generated as described above for all strains. Cell suspensions in 0.25 mm MgCl₂ were standardized to an OD_{600} of 0.8 (5 \times $10^{8}\,CFU\,mL^{-1}).$ Aliquots of 300 μl of bacterial inocula were diluted into 2.7 ml of NbAWF, AtAWF or M9 glucose medium (CSHL, 2010) to a final concentration of approximately $5 \times 10^7 \text{CFU} \, \text{mL}^{-1}$. To make macronutrient and micronutrient amended NbAWF, 1,000× concentrated macronutrients and micronutrients including sodium chloride, magnesium sulfate, ammonium sulfate, calcium chloride, glucose, potassium phosphate, potassium chloride, and sodium phosphate, were dissolved in distilled water and filter sterilized using 0.2 µM filters before being diluted to 1× in NbAWF. The same volume of water was used as a control. Aliquots of 400 µl were inoculated into 5 replicate wells of a Bioscreen honeycomb plate (Bioscreen Technologies, Bertinoro, Italy). The OD₆₀₀ was measured every hour for up to 36h in a Bioscreen C plate reader with low to medium shaking at 22°C (Bioscreen Technologies, Bertinoro, Italy). Raw absorbance readings were normalized by subtracting the initial absorbance readings from subsequent hourly readings.

For the competition assay, standardized inocula of each S. enterica strain and Pto were diluted in combination with either 0.25 mm $\,$ MgCl $_2$ or an inoculum partner for individual or co-inoculations, respectively, in M9 glucose medium or NbAWF to a final concentration of $5\times10^6\,{\rm CFU\,mL^{-1}}$ of each bacterium. Aliquots of 400 μl were inoculated into 5 replicate wells of a Bioscreen honeycomb plate and the growth (measured as ${\rm OD}_{600}$) was monitored until peak ${\rm OD}_{600}$ was achieved; 1 day for samples grown in NbAWF and 2 days for samples grown in minimal media. Serial dilutions were plated on appropriate plates with antibiotics to determine the initial and final population sizes of

both *S. enterica* strains and *Pto* measured as CFU mL⁻¹. Initial populations were measured from three aliquots of the original suspension and final populations were measured from individual wells in the Bioscreen honeycomb plate. Statistical differences between mean initial and final bacterial populations under single and co-inoculating conditions were determined using a 2-way ANOVA for each strain (p<0.05). At least two independent experiments were performed for all growth assays.

Gene fragment swap by allelic exchange

Salmonella enterica knock-out clones were generated in the LT2 strain background using the pR6KT2G suicide vector which allows for SacB-mediated sucrose counter-selection using methods defined by Stice et al. (2020) with modifications. The promoter and gene sequence of rpoS (STM14_3526) from 14028s was obtained from KEGG Gene using the organism code "seo." Flanks of 300 bp preceding and following the first 352 bp of *rpoS* were synthesized with attB1 and attB2 extensions for Gateway clonase recombination as double stranded DNA gblocks by Twist Bioscience (Supplementary Table S1). The synthesized gene fragment was cloned with BP clonase II into pR6KT2G according to the manufacturer's recommendations (Thermo Scientific, Waltham, MA, United States). The cleaned reaction mixture was electroporated into competent E. coli MaH1 pir+ cells and transformed cells were grown on LB amended with 10 µg mL⁻¹ gentamicin. Allelic replacement constructs were confirmed by BsrGI digest and sequencing before transformed into electrocompetent E. coli RHO5 pir+ cells.

The wild-type LT2 strain and RHO5 pR6KT2G::rpoS donor strain were mated on LB plates amended with 250 µg mL⁻¹ diaminopimelic Acid (DAP). Single crossover merodiploids were recovered from the mating mixture on LB plates amended with 10 μg mL⁻¹ gentamicin. Two merodiploid colonies were selected for counter selection in a liquid culture of 1 ml LB and 3 ml 1 M sucrose for 24 h at 37°C. Following counter selection, a portion of the diluted mixture was plated on LB plates amended with X-gluc (5-bromo-4-chloro-3-indolyl-beta-D-glucuronic acid, cyclohexylammonium salt). Candidate allelic replacement strains were not blue in color indicating eviction of the plasmid construct. Genomic DNA extractions were performed on candidate colonies using the Gentra Puregene kit according to the manufacturer's instructions (Qiagen, Hilden, Germany). Candidate colonies were screened using external primers (Supplementary Table S1) by PCR using Phusion HiFi polymerase according to the manufacturer's instructions (New England BioLabs, Ipswich, MA, United States). Candidates with the expected PCR fragment size were sequenced using external primers to confirm the gene deletion.

The resulting LT2 Δ rpoS¹⁻³⁵² strain was replaced with the rpoS 352 bp gene fragment from 14028s using the same homologous recombination procedure used to generate the mutants. Flanks of 300 bp preceding and following the 14028s rpoS gene fragment

were synthesized with *attB1* and *attB2* extensions for Gateway compatibility as double stranded DNA gblocks by Twist Bioscience (Supplementary Table S1). This gene fragment was cloned as described above and candidate knock-in strains were screened using external primers by PCR; the resulting gene fragment was digested using the AvaII restriction enzyme according to the manufacturer's instructions to distinguish 14028s and LT2 genotypes (New England BioLabs, Ipswich, MA, United States). Candidates with the expected PCR and digest fragment sizes were sequenced using the external primers to confirm the 14028s *rpoS* gene knock-in clones.

Hydrogen peroxide tolerance

LT2 and LT2 $rpoS_{14028s}$ strains were tested for viable bacterial population and growth in response to treatment with different concentrations of hydrogen peroxide. From overnight cultures prepared as described above, strains were pelleted by centrifugation and washed twice with fresh LB and adjusted to OD_{600} of 0.8 (approximately $5 \times 10^7 \, \mathrm{CFU} \, \mathrm{mL^{-1}}$). Aliquots of 200 μl bacterial inocula were treated with 0, 4, and 8 mm hydrogen peroxide in clear flat bottom plates on an orbital shaker at 1500 rpm at room temperature for 2 h. An equal volume of LB was used as media control. Serial dilutions were plated on LB plates to determine the population sizes of both S. enterica strains at the end point as CFU mL⁻¹ relative to the media control. Populations were measured from three aliquots of the original suspension and final populations were measured from individual wells. For measuring bacterial growth, plates were set up as described above and OD600 was measured using the SpectraMax iD3 reader (Molecular Devices, San Jose, CA, U.S.A) with the kinetic mode for 24h at 22°C. Two independent experiments were performed for CFU counts and growth curves.

Carbon source utilization

Sugars, sugar alcohols, and organic carboxylic acids identified in the NbAWF GC-MS profile were selected as carbon sources for testing of differential utilization between S. enterica LT2 and LT2 rpoS_{14028s} strains. Lactose was used as a negative control carbon source not utilized by S. enterica. Cultures of each S. enterica strain were generated from a single colony in LB at 37°C as described above. Cultures were pelleted by centrifugation and washed twice with fresh 1X M9 salt. The bacterial concentration was adjusted to $\mathrm{OD}_{600}\, of\, 0.8$ and was diluted 1 to 10 (approximately $5 \times 10^6 \, \text{CFU} \, \text{mL}^{-1}$). To prepare the 10 X concentration stocks, 4% of carbon sources including sucrose, D-glucose, galactose, D-fructose, L-malic acid, succinic acid, myo-inositol, and lactose were dissolved in distilled water and filter sterilized using 0.2 µM filters before dilution to 0.4% of each carbon source as well as a water only control. Aliquots of $200\,\mu l$ of inoculum with carbon source were inoculated into 4 replicates well at a 96-well flat bottom microplate with medium shaking at 22°C in Biotek Synergy II microplate reader (Winooski, VT). $\rm OD_{600}$ was measured every hour for up to 30 h and raw absorbance readings were normalized by subtracting the initial value of absorbance.

DAB and POX assays

Pto and derivatives were inoculated in A. thaliana and N. benthamiana to examine the in situ accumulation of hydrogen peroxide. Pto inocula were prepared as described above. For N. benthamiana, $1 \times 10^6 \text{CFU mL}^{-1}$ of $Pto\Delta hopQ$ (compatible with *N. benthamiana*) or $Pto\Delta hrcC$ were syringe-inoculated into fully expanded leaves. For A. thaliana, concentrations of 1 × $10^7 {
m CFU \, mL^{-1}}$ Pto or Pto $\Delta hrcC$ were used. 0.25 mm MgCl $_2$ was treated as buffer control in both host plants. All inoculated plants were incubated under high humidity (90-100% RH). 6 h and 24 h post inoculation, four 4mm leaf discs were collected for 3,3'-diaminobenzidine (DAB) staining. Leaf discs were collected in a 12-well microplate and submerged in DAB staining buffer as described in Daudi and O'Brien (2012). In short, leaf discs were gently vacuum infiltrated at 95 kPa for 2 min, then the plate was covered with aluminum foil and placed on a standard laboratory shaker for 4h at 80-100 rpm at room temperature. Following, the foil was removed, and DAB staining buffer was replaced with the bleaching buffer (ethanol: acetic acid: glycerol = 3:1:1). The plate was placed in a boiling water bath (~90-95°C) for 5-10 min until the chlorophyll had been removed. Leaf discs were taken out and excessive liquid was blotted off. Color intensity of brown precipitate was quantified using ImageJ (Schneider et al., 2012), 1/3 of the area from edges were omitted to exclude the woundingassociated signal from the cut edges of the discs. For each experiment, four plants were pooled as biological replicates and experiments were performed three times.

For the peroxidase (POX) assay, 4 mm-size of leaf discs from A. thaliana and N. benthamiana were collected and washed in 1× Murashige-Skoog solution (Murashige and Skoog, 1962) on a rotary shaker for 1 h to remove the signal from damage. Pto or $Pto\Delta hrcC$ were used as inocula for A. thaliana, and $Pto\Delta hopQ$ and Pto\(Delta\)hrcC were used in N. benthamiana. Leaf discs were then individually placed in a 96-well clear flat bottom plate with 50 μl of 5 x 10⁷CFU mL⁻¹ bacterial suspension in 1× Murashige– Skoog solution and gently vacuum infiltrated for 2 min at 95 kPa. Leaf discs were submerged with bacterial inocula and incubated at 80-100 rpm shaking at room temperature for 20 h after the vacuum infiltration. After the incubation, leaf discs were removed from individual well and POX activity was measured as described in Mott et al., 2018 (Mott et al., 2018). Briefly, $50\,\mu l$ of 1 mg mL⁻¹ 5-aminosalicylic acid solution (pH 6.0) with 0.01% hydrogen peroxide was pipetted to each well and set for 2 min. 20 µl of 2 m NaOH solution was then added to each well to stop the reaction. Quantification of POX was measured at OD_{600} immediately.

Results

Salmonella enterica LT2 has a host-dependent defect in co-colonization of the Nicotiana benthamiana apoplast during Pseudomonas syringae disease

Salmonella enterica 14028s and LT2 strain DM10000 were assessed for their capacity to achieve increased loads in the leaf apoplast of A. thaliana and N. benthamiana when co-inoculated with a common host-compatible *P. syringae* (*Pto*) pathogen partner. Suspensions of S. enterica were standardized to $5 \times 10^5 \, \text{CFU} \, \text{mL}^{-1}$ and pressure inoculated directly into the leaf apoplast using a needless syringe. S. enterica strains were either inoculated singly or co-inoculated with host-compatible Pto or $Pto\Delta hopQ$ at inoculum concentrations used routinely in each plant host $(5 \times 10^6 \, \text{CFU} \, \text{mL}^{-1})$ for A. thaliana and $5 \times 10^5 \text{CFU}\,\text{mL}^{-1}$ for N. benthamiana respectively), or the same concentrations of the $Pto\Delta hrcC$ control strain, which lacks a functional Type III Secretion System (T3SS) and is incapable of translocating T3Es and causing disease. Three days post inoculation, the bacterial loads of S. enterica and Pto strains in leaves were determined by leaf tissue maceration and dilution plating under appropriate selective conditions.

Both S. enterica 14028s and LT2 benefitted from infection co-colonization in the A. thaliana apoplast based on their increased in load versus either independent S. enterica inoculation or co-inoculation with the disarmed, non-pathogenic, Pto \(\Delta hrc C \) T3SS- control strain (Figures 1A,B). Without the pathogenic Pto strain the S. enterica populations decreased slightly compared with the day 0 population (Supplementary Figure S1). This indicates that T3SS-dependent Pto infection resulted in conducive conditions in the A. thaliana apoplast that could be opportunistically exploited by both S enterica strains. S. enterica 14028s also benefited from $Pto\Delta hopQ$ infection co-colonization in N. benthamiana, as has been observed previously (Figure 1C; Meng et al., 2013). However, although S. enterica populations had increased by day 3 compared with the day 0 population in each treatment (Supplementary Figure S1), S. enterica LT2 achieved similar populations in N. benthamiana when co-inoculated with pathogenic $Pto\Delta hopQ$, non-pathogenic $Pto\Delta hrcC$ and when inoculated independently (Figure 1D). As only this combination of plant host and S. enterica strain resulted in Pto infection-independent colonization, this implicated both N. benthamiana plant host and LT2 strain factors in the phenotype.

The Salmonella enterica LT2 Pto infection-independent colonization is unlikely to be caused by nutritional competition with Pto

Based on the assumption that, as a specialist plant pathogen, *Pto* would be better equipped to utilize nutrients available in the

N. benthamiana apoplast than S. enterica, our initial hypothesis was that the LT2 Pto infection-independent colonization might result from Pto outcompeting S. enterica LT2 for a growthlimiting nutritional resource. To address this, we recovered N. benthamiana apoplastic wash fluid (NbAWF), using standard methods for apoplastic wash fluid extraction. We determined endpoint populations via dilution plating of the S. enterica strains and Pto both individually and while co-cultured in filter-sterilized ex planta NbAWF or M9 glucose medium as a control. Both S. enterica strains grew well in NbAWF fluid and M9 glucose medium both individually and when co-cultured with Pto (Figure 2). In fact, counter to our initial hypothesis, it was the Pto partner that reached lower endpoint population sizes in NbAWF when co-cultured with either S. enterica strain than when cultured individually (Figure 2B). This experiment did not support S. enterica-Pto nutritional competition as being a major mediator of LT2 Pto infection-independent colonization in N. benthamiana.

Salmonella enterica LT2, counter-intuitively, grows more rapidly in Nicotiana benthamiana AWF than Salmonella enterica 14028s and with a distinct biphasic growth curve

While culturing S. enterica in ex planta NbAWF we made the unexpected observation that LT2 grew more rapidly than 14028s and with a more pronounced biphasic growth curve (Figure 3A). Conversely, both S. enterica strains displayed largely similar growth behavior in filter-sterilized A. thaliana AWF and M9 glucose medium (Figures 3B,C). The biphasic growth of LT2 in NbAWF seemed consistent with diauxic growth associated with catabolite repression. Based on this hypothesis, we were curious whether biphasic growth in NbAWF could be nutritionally suppressed. We augmented NbAWF with components of M9 glucose medium in which both S. enterica strains grew similarly. The biphasic growth of LT2 in NbAWF was not suppressed by augmentation with sodium chloride, magnesium sulfate, ammonium sulfate, or calcium chloride (Supplementary Figure S2). However, augmentation with either glucose or potassium phosphate suppressed NbAWF biphasic growth with LT2 growth being more starkly suppressed than 14028s by these two compounds (Supplementary Figure S3). Compared to augmentation with glucose, augmentation with potassium phosphate had a temporary effect on biphasic growth suppression. To test whether the phosphate anion or potassium cation was responsible for transient suppression of biphasic growth we supplemented NbAWF with potassium chloride, potassium phosphate, or sodium phosphate. In both cases where a phosphate anion was used, S. enterica biphasic growth was transiently suppressed while augmentation with potassium chloride resulted in similar biphasic growth to that of the water control (Supplementary Figure S4). The observed growth suppression via

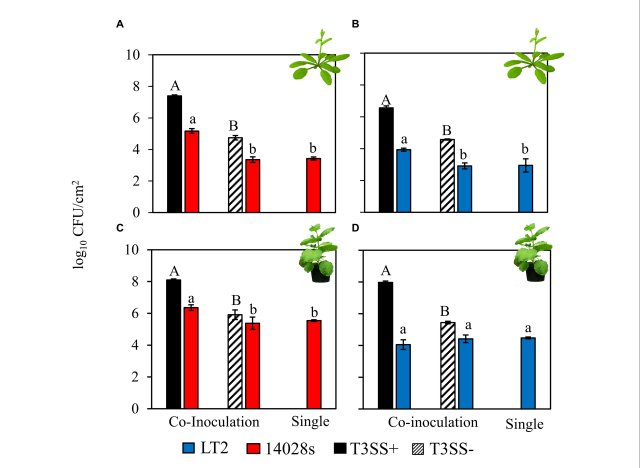


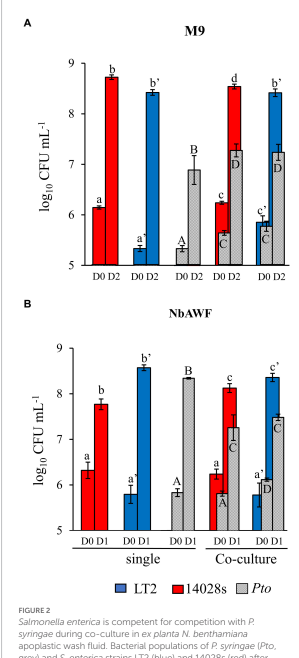
FIGURE 1
Salmonella enterica LT2 has a host-dependent defect in co-colonization of the Nicotiana benthamiana apoplast during P. syringae disease.
Bacterial populations of S. enterica strains (A,C) 14028s (red), and (B,D) LT2 (blue) with co-inoculation partner P. syringae (Pto) or $Pto\Delta hopQ$ with a functional Type III Secretion System (T3SS+, black) and $Pto\Delta hrCC$ without a functional T3SS (T3SS-, striped) Inocula were syringe infiltrated into model plant hosts, (A,B) A. thaliana CoI-0 at a concentration of 5×10^6 CFUmL⁻¹ for Pto strains and 5×10^6 CFUmL⁻¹ for S. enterica strains and (C,D)
N. benthamiana at a concentration of 5×10^6 CFUmL⁻¹ for all strains. Bacterial populations were measured as log colony forming units per cm² of leaf tissue (log₁₀ CFU/cm²) 3days post-inoculation. Data are means \pm SD (n=3 plants). Different letters indicate significant differences (2-tailed t-test for each strain at p<0.05). Three independent experiments were conducted with similar results.

nutrient augmentation is consistent with variation in the regulation of catabolite repression between the two strains.

Both the *Pto* infection-independent colonization and NbAWF biphasic growth phenotypes of LT2 are associated with mutations in the *rpoS* gene

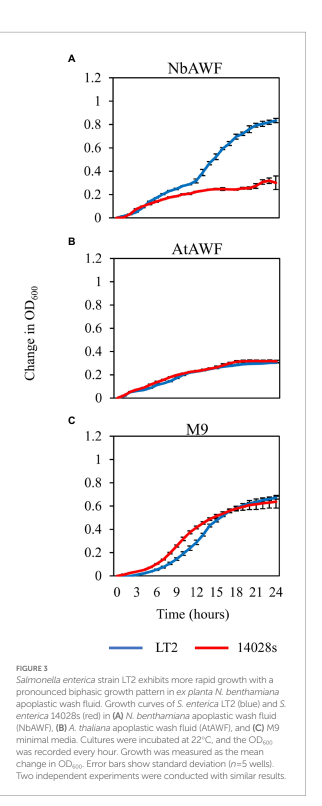
The infection-independent colonization phenotype of LT2 in *N. benthamiana* implicated both *S. enterica* strain and *N. benthamiana* host factors. A notable genetic difference between the LT2 and 14028s is the mutation of the LT2 *rpoS* sigma factor AUG start codon to the less-efficient UUG start codon (Swords et al., 1997; Wilmes-Riesenberg et al., 1997). This mutation results in lower RpoS protein levels and has been shown to be associated with reduced stress tolerance (Wilmes-Riesenberg et al., 1997; Jarvik et al., 2010). While confirming the *rpoS* sequences of 14028s

and LT2, we identified an additional 8-bp deletion mutation starting at nucleotide position 344 in the rpoS gene of LT2 DM10000. This 8-bp indel introduces a premature UGA stop codon at $rpoS_{LT2}$ nucleotide position 352 resulting in a predicted 117 aa truncated protein versus the 330 aa residue full-length *rpoS* ORF of 14028s and a likely null allele (Figure 4A). To our knowledge this 117 aa truncated allele has not been described previously and is not common among other LT2 isolates. We were curious whether these *rpoS* mutations contributed to our observed LT2 disease co-colonization defect and NbAWF growth phenotypes. We used two-step allelic exchange to delete the LT2 rpoS 1-352 region and subsequently replaced the deletion with the corresponding $rpoS_{14028s}$ coding sequence resulting in rpoS with an ATG start codon and the full-length open reading frame. Consistent with LT2 $rpoS_{14028s}$ having restored function compared to the native truncated rpoS_{LT2-117} allele, we observed that LT2 rpoS_{14028s} gained a dramatically improved tolerance to H₂O₂ (Figure 4B). When we tested the growth of the LT2 $rpoS_{14028s}$ strain



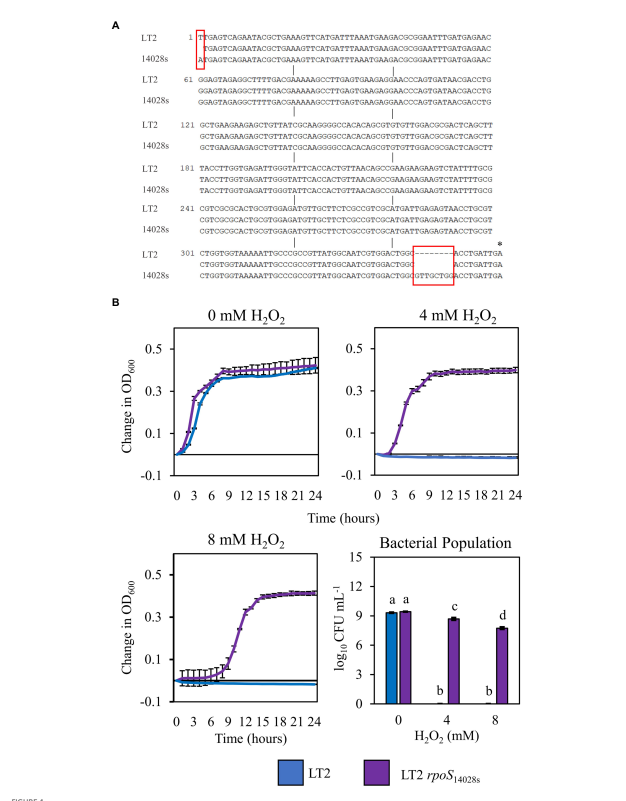
Salmonella enterica is competent for competition with P. syringae during co-culture in ex planta N. benthamiana apoplastic wash fluid. Bacterial populations of P. syringae (Pto, grey) and S. enterica strains LT2 (blue) and 14028s (red) after inoculation in (A) M9 glucose medium (M9) and (B) N. benthamiana apoplastic wash fluid (N4) Bacterial populations were measured as log colony forming units per mL of culture (\log_{10} CFU mL $^{-1}$) on day 0, 1, and 2. Data are means \pm SD (n=3-5). Different letters indicate significant differences (2-way ANOVA for each strain at p<0.05). Asterisk indicates significant difference between co-inoculation partners (2 tailed t-test at p<0.05). Two independent experiments were conducted with similar results.

in NbAWF, we observed that it had lost the pronounced biphasic growth phenotype and displayed growth behavior similar to 14028s (Figure 5A). In addition, we observed that LT2 $rpoS_{14028s}$ gained the capacity to benefit from Pto disease co-colonization in N. benthamiana (Figure 5B).



RpoS restoration alters utilization of L-malic acid, an abundant carbon source in *Nicotiana benthamiana* apoplastic wash fluid

Based on the previously observed capacity of glucose to suppress LT2 biphasic growth in NbAWF, we speculated that the



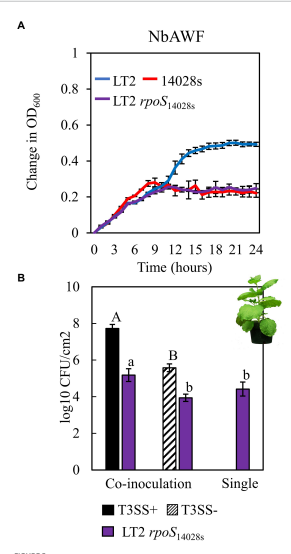


FIGURE 5 Restoration of the rpoS ORF in S. enterica LT2alters both ex planta and in planta phenotypes. (A) Growth curves of LT2 (blue), 14028s (red), and LT2 rpoS_{14028s} (purple) in N. benthamiana apoplastic wash fluid (NbAWF). Cultures were aliquoted into 5 replicate wells, incubated at 22°C, and the $\ensuremath{\mathsf{OD}}_{600}$ was recorded every hour. Growth was measured as the average change in measured OD_{600} with standard deviation error bars (n=5 wells). Two independent experiments were conducted with similar results. (B) Population of S. enterica LT2 rpoS_{14028s} in N. benthamiana, co-inoculated with P. syringae (Pto) with (T3SS+, black) and without (T3SS-, striped) a functional Type III Secretion System. $Pto\Delta hopQ$ and $Pto\Delta hrcC$ were used as T3SS+ and T3SS- respectively. Inocula were syringe-infiltrated at concentrations defined in Figure 1. Data are means \pm SD (n=3 plants). Bacterial populations were measured as log colony forming units per cm² of leaf tissue (log₁₀ CFU/cm²) 3days postinoculation. Data are means \pm SD (n=3 plants). Different letters indicate significant differences (2-tailed t-test for each strain at p<0.05). Three independent experiments were conducted with similar results.

enhanced biphasic growth of LT2 compared with 14028s and LT2 $rpoS_{14028s}$ in ex~planta NbAWF could be caused by rpoS-dependent differential utilization patterns for a carbon source found in NbAWF. Analysis by GC–MS of NbAWF detected 69 compounds

including sugars and sugar derivatives, amino acids and amino acid derivatives, and organic acids (Supplementary Table S2). We tested LT2 and LT2 rpoS_{14028s} for differential utilization of seven carbon sources identified in NbAWF (fructose, glucose, sucrose, galactose, myo-inositol, malic acid, and succinic acid) as the sole carbon sources in M9 medium. We also tested lactose as a negative control which was not detected in NbAWF and is not utilized by S. enterica. Of these compounds, there was a notable difference in the growth behavior of the two strains in M9 L-malic acid medium with LT2 showing a decreased lag phase compared with LT2 rpoS_{14028s} (Figure 6). Of note, malic acid was identified as the apoplastic metabolite with the second highest relative concentration in NbAWF after sucrose (Supplementary Table S2). Thus, the *rpoS*-associated variations in biphasic growth in NbAWF may be mediated by variation in the utilization malic acid present in NbAWF.

Evidence for higher basal ROS in Nicotiana benthamiana compared to Arabidopsis thaliana leaves

Reactive oxygen species play important roles in plant immune signaling (Shapiguzov et al., 2012; Qi et al., 2017). Based on the rpoS-association for infection co-colonization in N. benthamiana but not A. thaliana, and the stark rpoS-dependent H₂O₂ tolerance phenotype of LT2, we were curious whether there might be differences in the ROS concentrations of *N. benthamiana* leaves compared with A. thaliana leaves. We used two distinct assays to quantify ROS in leaf discs of both hosts. 1) Diaminobenzidine (DAB) staining in which H₂O₂-mediated oxidation of DAB results in the production of a brown precipitate in leaf tissue that can be quantified by image analysis and 2) the peroxidase (POX) assay in which the H₂O₂-mediated oxidation of 5-aminosalyclic acid by apoplastic plant peroxidases results in a color change that is measured spectrophotometrically. We used these techniques to monitor ROS levels after treatment with disease-compatible *Pto* pathogen (T3SS+) or non-pathogenic *PtoΔhrcC* (T3SS-) compared with a MgCl2 buffer control. Although, the 24h treatment with the *N. benthamiana* disease-compatible $Pto\Delta hopQ$ resulted in less ROS signal than treatment with the $Pto\Delta hrcC$ strain based on the DAB assay, ROS signal was significantly higher in N benthamiana than A. thaliana in every treatment and time point tested (Figure 7). This is consistent with N. benthamiana maintaining higher relative basal ROS than A. thaliana. It is possible that the elevated basal ROS of N. benthamiana combined with the reduced ROS tolerance of LT2 might directly contribute to the LT2 Pto infection-independent colonization phenotype.

Discussion

In this study, we observed that *S. enterica* subsp. *enterica* Typhimurium LT2 strain DM10000 is unable to benefit from co-colonization of the *N. benthamiana* apoplast with a

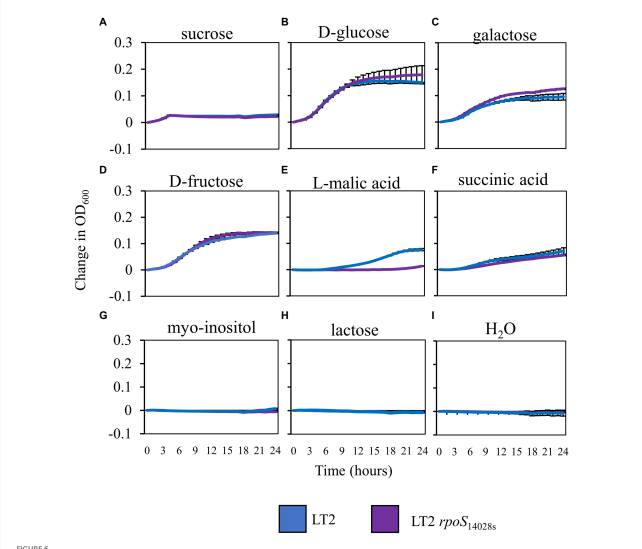
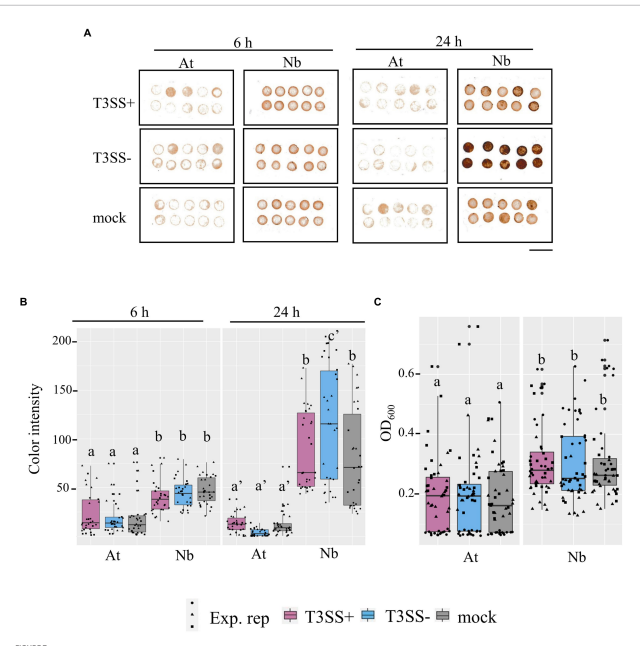


FIGURE 6
Salmonella enterica growth in M9 with NbAWF carbon sources. Growth curves of S. enterica LT2 (blue) and LT2 $rpoS_{14028s}$ (purple) in M9 medium supplemented with different carbon sources LT2 (blue) and LT2 $rpoS_{14028s}$ (purple) in M9 medium (A) sucrose, (B) D-glucose, (C) galactose, (D) D-fructose, (E) L-malic acid, (F) succinic acid, (G) myo-inositol, (H) lactose, and mock control (I) H_2O . Cultures were aliquoted into 4 replicate wells, incubated at $22^{\circ}C$, and the OD_{600} was recorded hourly. Growth was measured as the average change in measured OD_{600} with standard deviation error bars (n=4 wells). Two independent experiments were conducted with similar results.

disease-compatible *P. syringae* plant pathogen partner. We demonstrated that the *N. benthamiana* LT2 *Pto* infection-independent colonization phenotype was linked to a presumed null allele of the *rpoS* sigma factor. The *rpoS*_{LT2-117} mutation was also associated with increased sensitivity to hydrogen peroxide, pronounced biphasic growth *in ex planta* NbAWF, and altered utilization of L-malic acid. In addition, we present evidence for higher relative levels of basal ROS in *N. benthamiana* than in *A. thaliana* which, when considered in the context of the low hydrogen peroxide tolerance of LT2 strain DM10000, could potentially mediate the LT2 *Pto* infection-independent colonization phenotype.

The host-microbe and microbe-microbe interactions that influence the outcomes of three-way human enteric pathogen, plant host, plant pathogen interactions are complex. Multiple studies have observed that *S. enterica* plant colonization is impacted by plant genotype and that different *S. enterica* factors influence fitness and colonization of different plant hosts (Barak et al., 2011; de Moraes et al., 2017; Jacob and Melotto, 2019; Montano et al., 2020). Highlighting that not all plant pathogen-modulated niches are equally exploitable by *S. enterica*, only a subset of *Xanthomonas* tomato pathogens conferred a co-colonization fitness advantage to *S. enterica* serovars Enteritidis and Baildon (Potnis et al., 2015). Recent work by Cowles et al. (2022) determined that the *Xanthomonas* T3E AvrHah1, associated with leaf water soaking, is both necessary and sufficient for increased persistence of 14028s during *Xanthomonas* disease co-colonization of tomato leaves. Deblais et al. (2021), demonstrated that *S. enterica* LT2 (strain JSG626) persistence on tomato leaves was enhanced by the



Evidence for higher basal ROS in N. benthamiana levels compared to A. thaliana leaves. (A,B) in situ H₂O₂ in N. benthamiana and A. thaliana inoculated with P. syringae (Pto) with (T3SS+) and without (T3SS-) a functional Type III Secretion System. The detected H_2O_2 level within inoculated tissues of N. benthamiana (Nb) and A. thaliana (At) were measured with DAB staining at both an earlier time point (6h) and later time point (24h). T3SS+ strains, Pto and $Pto\Delta hopQ$, were used in A. thaliana and N. benthamiana, respectively. $Pto\Delta hrcC$ was used as T3SS- treatment for both plants. Whole leaves were syringe-infiltrated with bacterial suspension ($1 \times 10^7 \text{CFUmL}^{-1}$ for *A. thaliana* and $1 \times 10^6 \text{CFUmL}^{-1}$ for *N. benthamiana*) or 0.25mm MgCl₂ treatment as mock treatment. 4mm diameter of leaf discs were collected at each time point post inoculation. Scale bar, 10mm. (C) Peroxidase (POX) assay shows peroxidase levels in N. benthamiana than in A. thaliana. Leaf disks from Col-0 (At) and N. benthamiana (Nb) were vacuum infiltrated with bacterial suspension $(5\times10^7\text{CFUmL}^{-1})$ and incubated for 20h. T3SS+ and T3SS- strains used for both plants were described as above. Absorbance of OD_{600} was measured as POX activity. Graphs show data from three experimental replicates (Exp. rep) with means \pm SE, n=16 discs from 4 plants. Different letters indicate statistically significant differences as analyzed by 1-way ANOVA (Tukey test, p<0.05).

bacterial tomato pathogen Clavibacter michiganensis subsp. michiganensis but not by Xanthomonas gardneri. Deblais et al. (2021) also noted that LT2 strain JSG626 displayed better persistence on tomato than six other S. enterica strains in preliminary testing. It would be interesting to characterize

whether these strains also vary in their capacity to benefit from disease co-colonization.

Amongst the set of identified genetic polymorphisms between 14028s and LT2, the LT2 rpoS UUG start codon mutation is both well known and well characterized due to its role in reduced stress

tolerance and LT2 virulence attenuation (Fang et al., 1992; Swords et al., 1997; Wilmes-Riesenberg et al., 1997). We determined that the replacement of LT2 DM10000 rpoS with the 14082s rpoS full length coding sequence greatly enhanced tolerance to hydrogen peroxide and allowed LT2 DM10000 to benefit from disease co-colonization in the N. benthamiana apoplast. Previous studies have identified both RpoS and oxidative stress tolerance as contributing to S. enterica colonization of legume sprouts (Barak et al., 2005; Wahlig et al., 2019). Barak et al. (2005), reported that transposon-inactivation of rpoS resulted in decreased attachment by S. enterica Newport to alfalfa sprouts based on decreased RpoSmediated production of curli and cellulose. However, our use of direct leaf apoplastic infiltration likely abrogates attachment requirements as a component of S. enterica plant colonization. Wahlig et al. (2019) inferred that an observed decrease in LT2 H₂O₂ tolerance also mediated reduced colonization of Medicago trunculata sprouts by LT2 but not 14028s in a manner dependent on the functional MtFLS2 flagellin immune receptor. Conversely Zarkani et al. (2019) reported no major difference in the colonization and persistence of 14028s and LT2 on tomato leaves using either direct infiltration or dip inoculation. In addition, in two independent studies by de Moraes et al. (2017) and Montano et al. (2020), transposon insertions in rpoS were not identified among S. enterica factors contributing to 14028s colonization of either tomato fruit pericarps or lettuce leaves, respectively.

While comparing our observations to previously published work, there are two issues worth noting. First, the majority of comparative and mutational genetic studies that have identified *S. enterica* plant colonization factors have focused on independent *S. enterica* colonization, while co-colonization of the plant pathogen-modulated apoplastic niche very likely poses distinct fitness requirements. Second, in our work we identified a novel and presumed null allele of *rpoS* in LT2 strain DM10000 which may confound direct comparison with published results using *S. enterica* LT2. Considering that *rpoS* mutations can be selected unintentionally during laboratory culture (Notley-McRobb et al., 2002; Ferenci, 2003), it may be beneficial to characterize the *rpoS* integrity of other *S. enterica* strains used in plant-colonization experiments.

To our knowledge, the rpoS $_{LT2-117}$ allele has not been previously reported and is not common among LT2-derived strains. While the LT2 rpoS $_{UUG}$ allele is associated with levels of H_2O_2 tolerance comparable to that of a WT rpoS allele, null mutations in rpoS are associated with a dramatic loss in H_2O_2 tolerance and catalase activity (Wilmes-Riesenberg et al., 1997). The rpoS $_{LT2-117}$ allele is presumably also a null mutation based on its H_2O_2 sensitivity phenotype and degree of ORF truncation (Figure 4). In addition to regulation of catalase expression and H_2O_2 tolerance, the RpoS sigma factor regulates tolerance to acid stress, high osmolarity, high temperature, and promotes survival during extended starvation (Wilmes-Riesenberg et al., 1997). RpoS also plays a major role in the regulation of *S. enterica* metabolic transition into stationary phase (Levi-Meyrueis et al., 2014). Paradoxically, although rpoS is upregulated during stationary phase and plays a

role the regulation of starvation survival genes, rpoS inactivating mutations can also provide a selective advantage under nutrient limitation conditions (Notley-McRobb et al., 2002; Ferenci, 2003). The antagonistic pleiotropy of rpoS mutants providing both a selective advantage under nutrient limitation and a selective disadvantage in response to physiological stressors has been studied extensively in E. coli but is also supported in S. enterica (Notley-McRobb et al., 2002; Ferenci, 2003; Robbe-Saule et al., 2007; Levi-Meyrueis et al., 2014). In E. coli, mutations in rpoS can be readily recovered by selecting strains for improved growth on a non-preferred carbon source such as succinate in the absence of physiological stressors (Notley-McRobb et al., 2002; Dong et al., 2009). Under these selective regimes rpoS alleles are often recovered with reduced activity rather than full null alleles presumably because these alleles provide the strains with limited stress tolerance (Ferenci, 2003). The relevancy of rpoS antagonistic pleiotropy for S. enterica plant colonization either with or without plant pathogen partners in a natural setting is unclear. Presumably, physiological stressors associated with colonization of plant tissues would provide purifying selection against rpoS-inactivating mutations in S. enterica, but improved capacity for growth on non-preferred carbon sources common in plant-associated niches might also provide rpoS mutants with a selective advantage under certain conditions. Although the LT2 DM10000 rpoS_{LT2-117} allele was presumably unintentionally selected during laboratory culture, S. enterica Enteritidis and Typhi clinical isolates with rpoS-deactivating mutations and catalase defects have been previously identified indicating that these mutations are not solely an artifact of the laboratory (Robbe-Saule et al., 2003; Betancor et al., 2009).

Our work also indicates that phenotypes based solely on behavior in ex planta apoplastic wash fluid should be interpreted cautiously. The pronounced biphasic growth of LT2 DM10000 in ex planta NbAWF is dissimilar to the behavior of the strain in planta during N. benthamiana disease co-colonization. Ideally, NbAWF represents a partial metabolic snapshot of the water extractable fraction of the apoplastic compartment present at the time of sampling. However volatile and/or reactive compounds may not survive processing, filter sterilization, and storage. Apoplastic wash fluid cannot fully capture the dynamic complexity of active microbe-plant biotic interactions. We determined that NbAWF contains a wide diversity of potential carbon and nitrogen sources that S. enterica could utilize and serves as an effective growth medium for both 14028s and LT2. The co-occurrence of the rpoS-associated LT2 DM10000 N. benthamiana infectionindependent colonization phenotype and the pronounced ex planta NbAWF biphasic growth phenotype seems to be an extraordinary, although artifactual coincidence. Although RpoS in S. enterica has been more extensively studied in regards to its role in stress tolerance, over 45% of the RpoS regulon is associated with both the positive and negative regulation of metabolic genes (Levi-Meyrueis et al., 2014). The suppressive effect of glucose or phosphate augmentation on the NbAWF biphasic growth phenotype is consistent with rpoS-dependent alteration to diauxic

growth utilization patterns of a non-preferred carbon source in NbAWF. Exogenous phosphate has previously been shown to suppress catabolite control of the expression of csr genes which regulates motility, carbon storage, and virulence in S. enterica, which could explain why we observe suppression of biphasic growth with exogenous phosphate in NbAWF (Teplitski et al., 2006). Although our tests for rpoS-dependent altered utilization of NbAWF identified carbon sources was far from exhaustive, we did identify rpoS-dependent alteration in the utilization of L-malic acid in M9 medium with decreased growth lag observed in the LT2 strain with the *rpoS*_{LT2-117} allele (Figure 6). Dong et al. (2009) identified that E. coli selected for spontaneous rpoSdeactivating mutations also had shortened generation times when grown on malate as a sole carbon source. Although the connection between altered utilization of L-malic acid and NbAWF biphasic growth is solely correlative, it is notable that we identified malic acid as the second highest relative concentration metabolite in NbAWF, while Gao et al. (2020) did not identify malic acid among A. thaliana apoplastic wash fluid metabolites in which the two strains grew similarly.

The accumulation of apoplastic ROS and H₂O₂ is a well established component of the plant immune response (Qi et al., 2017). Consistent with this, we observed statistically elevated DAB staining in N. benthamiana at 24h after treatment with non-pathogenic Pto \(\Delta hrcC. \) However, notably we measured statistically higher ROS signal in N. benthamiana than in A. thaliana with all treatments and timepoints (Figure 7). This could be indicative either of higher ROS production or of lower antioxidant activity in N. benthamiana relative to A. thaliana. Significant differences in defense responses between cultivars of lettuce, such as reactive oxygen species burst and callose deposition, were found to have a positive correlation with apoplastic persistence of 14028s as well as E. coli O157:H7 (Jacob and Melotto, 2019). While disease compatible pathogens can suppress aspects of immune signaling, this suppression is quantitative. For instance, Pto collectively requires catalases to detoxify external H₂O₂ to cause disease which would presumably be unnecessary if apoplastic H₂O₂ were completely suppressed during a compatible disease interaction (Guo et al., 2012). Likewise, the rpos_{LT2-117} -associated hydrogen peroxide sensitivity and catalase deficiency of LT2 DM10000 may be adequate to explain the *N. benthamiana* disease co-colonization defect.

The variation in *Pto* infection-dependent colonization competence of *S. enterica* LT2 in *A. thaliana* or *N. benthamiana* indicates a dissimilarity of the pathogen-modulated apoplastic environments between these two hosts. Apoplastic plant defenses are most typically studied in the context of specialist plant pathogens. However, these specialists, in addition to their active mechanisms to suppress aspects of the plant immune response, are likely well adapted to evade or tolerate intrinsic or immune-associated stressors of the apoplast. Conversely, human enteric pathogens like *S. enterica* may provide a clearer window into the selective pressures of the naïve non-permissive and the pathogen-modulated conducive apoplastic niches. The physiological and

transcriptional responses of Salmonella spp. and their close relatives to various stressors have been extensively studied and may therefore be more readily interpretable in the context of apoplastic plant defenses (Rychlik and Barrow, 2005; Shen and Fang, 2012). In our work, we identified a plant host-dependent contribution for the global regulatory sigma factor RpoS to infection co-colonization. Characterizing the contributions of S. entericaRpoS-regulated genes such as catalases and other oxidative stress tolerance genes would help to clarify the potential role of basal ROS as the mechanistic mediator of the N. benthamiana Pto infectionindependent colonization phenotype. In addition, previous studies have demonstrated the utility of conducting in planta transcriptomics of S. enterica (Jacob et al., 2021) and the use of TnSeq to identify genetic factors contributing to S. enterica disease co-colonization of tomato during soft rot disease (George et al., 2018). Future work based on S. enterica transcriptomics or TnSeq during disease co-colonization with T3SS-dependent plant pathogens could provide powerful new insights into the landscape of the pathogen-modulated conducive apoplastic niches in diverse plant hosts and how plant pathogens may impact S. enterica colonization of diverse plants.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary material, further inquiries can be directed to the corresponding author.

Author contributions

AL: Conceptualization, methodology, validation, formal analysis, investigation, writing, review and editing, visualization, and funding. H-CC: Conceptualization, methodology, validation, formal analysis, investigation, writing, review and editing, and visualization. SL: Formal analysis and investigation. ZS: Formal analysis, investigation, data curation, and writing. PB: Methodology, resources, and supervision. GP: Conceptualization, review and editing, supervision, project administration, and funding. BK: Conceptualization, resources, writing review and editing, supervision, project administration, and funding. All authors contributed to the article and approved the submitted version.

Funding

This work was supported by grants from the United States Department of Agriculture: USDA-NIFA 2018-07750 awarded to AL, National Science Foundation IOS 1844861 to BK and University of Georgia College of Agriculture and Environmental Science Interdisciplinary Seed Grant Program: Ensuring Safe Food and Water to BK. ZS received support through a CARA Fellowship.

Acknowledgments

We would like to thank Anna Karls for providing 14028s, Diana Downs for providing LT2 DM10000 as well as experimental guidance for working with *S. enterica*. We would also like to thank Wayne Parrot and Ron Walcott for use of equipment, Jovana Mijatović for assistance with data analysis and visualization, and Govindaraj Dev Kumar for critical review and helpful comments on the preparation of this manuscript.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.999183/full#supplementary-material

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SUPPLEMENTARY FIGURE ST

Starting population of Salmonella enterica LT2, 14028s and the coinoculated Pseudomonas syringae partner strains in Arabidopsis thaliana and Nicotiana benthamiana. Bacterial populations of S. enterica strains (A,C) 14028s (red), and (B,D) LT2 (blue) with co-inoculation partner P. syringae (Pto) or $Pto\Delta hopQ$ with a functional Type III Secretion System (T3SS+, black) and $Pto\Delta hrcC$ without a functional T3SS (T3SS-, striped). Inocula were syringe infiltrated into model plant hosts, (A,B) A. thaliana Col-0 at a concentration of $5\times10^{\circ}$ CFU mL⁻¹ for S. enterica strains and (C,D) N. benthamiana at a concentration of $5\times10^{\circ}$ CFU mL⁻¹¹ for all strains. Bacterial populations were measured as log colony forming units per cm² of leaf tissue (log₁₀ CFU/cm²) immediately after the inoculation. Data are means±SD (n=3 plants). Different letters indicate significant differences (2-tailed t-test for each strain at p<0.05). Three independent experiments were conducted with similar results.

SUPPLEMENTARY FIGURE S2

Salmonella enterica biphasic growth in N. benthamiana apoplastic wash fluid is unaltered after augmentation with specific macro and micronutrients. Growth curves of S. enterica LT2 (blue) and S. enterica 14028S (red) in N. benthamiana apoplastic wash fluid augmented with (A) 10mM sodium chloride, (B) 5mM magnesium sulfate, (C) 10mM ammonium sulfate, (D) 1mM calcium chloride, and (E) water. Cultures were aliquoted into 5 replicate wells, incubated at 22°C, and the OD600 was recorded every 2h. Growth was measured as the average change in measured OD600 with standard deviation error bars (n=5 wells). Two independent experiments were conducted with similar results.

SUPPLEMENTARY FIGURE S3

Salmonella enterica strain biphasic growth in N. benthamiana apoplastic wash fluid is suppressed by glucose and phosphate augmentation. Growth curves of S. enterica LT2 (blue) and S. enterica 14028s (red) in N. benthamiana apoplastic wash fluid augmented with (A) 25mM glucose, (B) 10mM potassium phosphate, and (C) water. Cultures were aliquoted into 5 replicate wells, incubated at 22°C, and the OD_{600} was recorded every 2h. Growth was measured as the average change in OD_{600} . Error bars show standard deviation (n=5 wells). Two independent experiments were conducted with similar results.

SUPPLEMENTARY FIGURE \$4

Salmonella enterica biphasic growth in N. benthamiana apoplastic wash fluid is suppressed by the phosphate anion. Growth curves of S. enterica LT2 (blue) and S. enterica 14028s (red) in N. benthamiana apoplastic wash fluid supplemented with (A) 10mM potassium chloride, (B) water, (C) 10mM potassium phosphate, and (D) 10mM sodium phosphate. Cultures were aliquoted into 5 replicate wells, incubated at 22°C, and the OD_{600} was recorded every hour. Growth was measured as the average change in measured OD_{600} with standard deviation error bars (n=5 wells). Two independent experiments were conducted with similar results.

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