

The role of L-arabinose metabolism for *Escherichia coli* O157:H7 in edible plants

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Abstract

Arabinose is a major plant aldopentose in the form of arabinans complexed in cell wall polysaccharides or glycoproteins (AGP), but comparatively rare as a monosaccharide. L-arabinose is an important bacterial metabolite, accessed by pectolytic micro-organisms such as *Pectobacterium atrosepticum* via pectin and hemicellulose degrading enzymes. However, not all plant-associated microbes encode cell-wall-degrading enzymes, yet can metabolize L-arabinose, raising questions about their use of and access to the glycan in plants. Therefore, we examined L-arabinose metabolism in the food-borne pathogen *Escherichia coli* O157:H7 (isolate Sakai) during its colonization of plants. L-arabinose metabolism (*araBA*) and transport (*araF*) genes were activated at 18 °C *in vitro* by L-arabinose and expressed over prolonged periods *in planta*. Although deletion of *araBAD* did not impact the colonization ability of *E. coli* O157:H7 (Sakai) on spinach and lettuce plants (both associated with STEC outbreaks), *araA* was induced on exposure to spinach cell-wall polysaccharides. Furthermore, debranched and arabinan oligosaccharides induced *ara* metabolism gene expression *in vitro*, and stimulated modest proliferation, while immobilized pectin did not. Thus, *E. coli* O157:H7 (Sakai) can utilize pectin/AGP-derived L-arabinose as a metabolite. Furthermore, it differs fundamentally in *ara* gene organization, transport and regulation from the related pectinolytic species *P. atrosepticum*, reflective of distinct plant-associated lifestyles.

INTRODUCTION

Arabinose is an abundant aldopentose in plant material that is not found in animals. It is present almost entirely as polysaccharides (arabinans) and glycoproteins, as L-arabinofuranose (L-Araf) complexed in the RG1 and RG2 pectin components of plant cell walls [1], as side chains in hemicellulose [2] or in arabinogalactan-proteins, AGP [3]. Free monomeric L-arabinopyranose (L-Arap) is present in intracellular and apoplasmic compartments but is rare in comparison to fructose and sucrose [4]. L-arabinose metabolism is widespread in microbes, reflective of a beneficial function for accessing the carbohydrate directly from plants or as a dietary fibre in animal guts. Microbial metabolism of L-arabinose has been well characterized for fundamental understanding of

metabolic processes [5] and for biotechnological applications [6, 7].

L-arabinose metabolic systems comprise transport and metabolism genes and a master regulator. In *E. coli* and related bacteria, L-Arap is transported into the cell by an ABC transporter system where AraF is the periplasmic component that binds L-Arap with high affinity, AraH is the trans-membrane protein and AraG the ATP-binding component [8]. AraE is an H⁺ symporter with relatively low affinity (140–320 μm) for L-Arap [9]. Intracellular L-arabinose enters the pentose phosphate pathway in a three-step degradation pathway via AraA, an isomerase that converts it to L-ribulose; AraB, a ribulokinase that catalyses L-ribulose phosphorylation to L-ribulose-5-phosphate; and AraD, an epimerase that

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Abbreviations: AGP, arabinogalactan-proteins; ELISA, enzyme-linked immunosorbent assay; GFP, green fluorescent protein; L-Araf, L-arabinofuranose; L-Arap, L-arabinopyranose; Pba, *Pectobacterium atrosepticum*; PCWDE, plant cell-wall degrading enzymes; RG, rhamnogalacturonans; STEC, shigatoxigenic *Escherichia coli*; VTEC, verocytotoxigenic *Escherichia coli*.

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converts L-ribulose 5-phosphate to D-xylulose-5-phosphate [5]. Transport and metabolism genes are under the control of AraC, an activator that is triggered when complexed with L-Arap [10]. Expression of the *ara* genes is under catabolite control and cyclic AMP forms a dimeric complex with CRP to co-regulate with AraC in the absence of glucose [11]. *araF* in *E. coli* has two promoters, under the control of σ^{70} and σ^S , respectively [12, 13], and is repressed by a small RNA [14].

Shigatoxigenic/verocytotoxigenic *E. coli* (STEC/VTEC), predominately serotype O157:H7, is a food-borne pathogen that can be transmitted through the food-chain by edible plants and utilize plants as secondary hosts [15]. STEC can mobilize metabolic pathways that are specific for different plant tissues [16, 17], including L-arabinose metabolism, which was induced on exposure to crude extracts of spinach leaf lysates and spinach cell-wall polysaccharides [16]. Yet, *E. coli* do not encode cell-wall-degrading enzymes (PCWDE) in contrast to the related members within the *Enterobacteriaceae*. The phytopathogen *Pectobacterium atrosepticum* encodes pectinases that are considered virulence factors in plant disease [18]. This raises fundamental microbial ecology questions as to whether STEC are able to exploit an apparently inaccessible metabolite in plant-microbe interactions without the aid of cell-wall-degrading enzymes. Therefore, we tested the hypothesis that L-arabinose metabolism facilitates the ability of STEC to colonize plants, by comparing L-arabinose response of *E. coli* O157:H7 isolate Sakai *in vitro* and *in planta* to that of *P. atrosepticum*. Horticultural crop species (spinach and lettuce leaves, and broccoli microgreens) that are considered a risk of foodborne illness from STEC were used as the most relevant plant models. To understand the contribution of L-arabinose metabolism to colonization of plants by STEC (Sakai), we took a reductionist approach to quantify gene expression in response to plant extracts *in vitro*, which was validated by a qualitative approach of gene expression *in planta*.

METHODS

Bacteria

Bacterial strains were as follows: *E. coli* STEC isolate Sakai [19]; *E. coli* strain AAEC185A [20] used for cloning; *P. atrosepticum* isolate SCRI-1043 [21] and its derivative mutants *expI-* and *outD-* [22]. An *araBAD* knock-out mutant of STEC (Sakai) was constructed by allelic exchange, cloning the upstream and downstream flanking region with primers ECs0066No_for, ECs0066Ni_rev and ECs0066Ci_for, ECs0066Co_rev on *PstI* and *NotI* and *NotI* and *SalI* sites, respectively, into the exchange vector pTOF24 [23], with a tetracycline gene introduced into the *NotI* site for selection. The deletion mutation was confirmed by PCR and Sanger sequencing. An *araC* knock-out mutant of STEC (Sakai) was a generous gift for this study, made by lambda-Red recombination [24]. Bacteria were routinely grown with aeration in lysogeny broth (LB) or MOPS medium [25] at 37 or 18°C (STEC), or 27°C (Pba) supplemented with 0.2% glucose or glycerol where indicated, 10 μ m thiamine and MEM essential and non-essential amino

acids (Sigma M5550 and M7145) termed rich defined MOPS (RD-MOPS). Antibiotics were included to maintain transformed plasmids at 50 μ g ml⁻¹ kanamycin (Kan), 25 μ g ml⁻¹ chloramphenicol (Cam), 10 μ g ml⁻¹ tetracycline (Tet) or 50 μ g ml⁻¹ ampicillin (Amp).

Gene expression analysis

Reporter plasmids were constructed in a pACYC-derived vector, carrying the *gfp+* gene, termed pKC026 [26], with promoter regions amplified from STEC (Sakai) or Pba (1043) genomic DNA with a proof-reading polymerase (Phusion, NEB) and inserted via the *XbaI* cloning site (Table S1, available in the online version of this article). A low copy vector *gfp+* reporter was generated, derived from pWSK29 27 by sub-cloning the *gfp+* gene plus the *ara* genes from previously generated plasmids into the *PstI* site (Table S1). Transformed STEC (Sakai) or Pba (1043) were grown at 18 or 27°C, respectively, in RD MOPS glycerol with aeration to OD₆₀₀ of 2.0 and sub-inoculated at 1:100 into 10 ml RD MOPS glycerol supplemented with 10 μ m to 10 mm L-arabinose. Cell density and GFP fluorescence were measured at 2 h intervals, from 150 μ l aliquoted into a black 96-well plate and measured in a GloMax Multi Detection System (Promega) machine (excitation 490 nm, emission 510–570 nm). Dose-response was measured at 0, 2, 4, 6, 8 and 24 h in RD MOPS glycerol and 0, 1, 2, 3, 4, 5 h in RD MOPS glucose. Time points for analysis were selected at maximum GFP emission, since the levels reached a maximum and then decreased presumably as arabinose was depleted and metabolism switched to glycerol. Measurements in plant extracts were taken at daily intervals over 5 days. Four sample reps were measured per test, and the experiment repeated three times. GFP is expressed as relative fluorescent units after subtraction from the vector-only control (pKC026 or pWSK29) data and normalized for cell density to OD₆₀₀ of 1.0. Arabinose concentrations were transformed (Log₁₀) for graph plotting and linear-regression analysis. Gene-expression analysis was measured directed by quantitative reverse transcriptase PCR (qRT-PCR) as previously [16] from STEC (Sakai) grown in minimal M9 medium (1 \times M9 salts, 1 mm MgSO₄, 0.3 mm CaCl₂) with 0.2% glycerol for 48 h at 18°C, and sub-inoculated 1:100 in minimal M9 supplemented with 0.2% L-arabinose or glucose, or with 0.2% oligo-/polysaccharides (1-5)- α -linked pectin backbone +/- α -L-arabinofuranosidase treatment; (1-5)- α -L-arabinobiose (Ara2); (1,5)- α -L-arabinoheptose (Ara7), (Megazyme, Bray, Ireland) at 18°C and sampled as indicated. Primers for STEC *araA* and *araD* (Table S1) were tested for efficiency and removal of DNA confirmed from direct PCR as described previously [16]. PCR products were amplified with iTaq Universal SYBR Green Supermix (Bio-Rad) measured on a StepOnePlus machine (Applied Biosystems), data normalised to *gyrB* (ECs4634) and expressed as 2^{- $\Delta\Delta$ CT} for fold-change compared to a control situation.

Plants and plant extracts

Spinach (*Spinacia oleracea*) cultivar Amazon (Sutton Seeds, UK) lettuce *cv.* Salinas (*Lactuca sativa*) (Tozer Seeds, UK),

Table 1. Expression of genes related to the processing, transport and utilization of arabinose

Gene name	Temp.	Spinach leaf lysates	Spinach root exudates	Spinach polysaccharides	Lettuce polysaccharides
<i>araA</i>	-3.21	NS	8.94	10.68	NS
<i>araB</i>	-4.94	NS	19.98	9.37	-2.18
<i>araC</i>	-5.76	NS	9.14	NS	-4.97
<i>araD</i>	NS	NS	3.58	NS	NS
<i>araE</i>	NS	NS	NS	NS	NS
<i>araF</i>	NS	NS	10.45	6.01	-2.70
<i>araG</i>	NS	NS	NS	2.16	NS
<i>araJ</i>	NS	2.22	-3.10	NS	NS

Fold-change gene-expression levels for *E. coli* O157:H7 (Sakai) exposed to temperature shift (37 to 18 °C), or plant extracts (all at 18 °C), data extracted from previous microarray measurements [16]. NS, not significantly different from the control.

tomato (*Solanum lycopersicum* cv. Moneymaker) (Thompson and Morgan, UK), broccoli (*Brassica oleracea* var. *italica*) (Unwins, UK), and *Nicotiana benthamiana* (Hutton stocks) were grown individually in 9 cm³ pots in compost for colonization assays in a glasshouse for 3 weeks in standard compost. Plants for root exudates or vermiculite for polysaccharide extracts were grown from surface sterilized seeds (2% calcium hypochlorite solution) in pots containing autoclaved autoclaved rockwool (Progrow), as described previously [16]. In brief, exudates were collected from 24 plants per batch by three aqueous extractions of the rockwool, and for the polysaccharide extracts, whole roots were excised from the plant and briefly washed in SDW to remove as much of the vermiculite as possible. A similar extract was made from vermiculite only as a negative control. Leaf and root extracts were frozen in liquid nitrogen and ground to a fine powder, for which 10 g was used for pectin (CDTA-treatment) and hemicellulose (NaOH treatment) fractions from an alcohol insoluble residue, which required a heating step (50 °C) to separate the fractions [28]. All lysates/polysaccharide extracts were stored at -20 °C.

ELISA was used to quantify arabinans, as described previously [29]. In short, 100 µl volumes of 1:10 plant polysaccharide extract (in 0.1 M NaHCO₃ buffer, pH 9.6) were incubated overnight at 4 °C to coat 96-well microtitre plates (NUNC, Maxisorb) blocked with 3% skimmed milk protein in TBS (1 h, RT) and washed three times with TBS (100 mM Tris, 150 mM NaCl, pH 7.5). Primary antibodies (PlantProbes) were added at 1:20 dilution in 3% skimmed milk protein in TBS (2 h, RT) and the plates washed three times in TBS. Secondary antibody, goat anti-rat horseradish peroxidase at 1:1000 was added (1 h, RT) and the plates washed three times in TBS, then developed with ABTS solution: 22 mg ABTS (Sigma Aldrich, St. Louis, USA) diluted in 100 ml of citrate buffer (50 mM sodium citrate, 0.05% H₂O₂, pH 4.0). Absorbance was measured at 405 nm on a microplate reader (Varioskan, ThermoFisher Scientific, USA).

Pba (1043) was grown in RD MOPS glycerol for 24 h at 27 °C, and sub-inoculated into fresh RD MOPS supplemented with

1% (w/v) spinach alcohol-insoluble polysaccharide extract plus 0.2% or 0.4% glycerol, over 24 h at 27 °C. STEC (Sakai) were grown in minimal M9 at 18 °C supplemented with oligosaccharides (Ara2, Ara7) exactly as per gene-expression analysis, and c.f.u. measurements taken at 0, 16, 32 and 48 h on selective MacConkey agar plates.

Plant colonisation assay

Three-week-old plants were used for leaf or root inoculations as per [4], inoculated with STEC (Sakai), diluted to OD₆₀₀ of 0.02 (equivalent to 10⁷ c.f.u. ml⁻¹) in SDW used to make bacteria suspensions. Leaves were dip-inoculated by submerging the foliar parts in 1 l bacterial suspension for 30 s, or roots were colonized by partially submerging plant pots in the suspension for 1 h. The pots were then transferred to the growth chamber until sampling. Plants were sampled at 0, 5, 10 and 14 days post-infection (days p.i.), aerial tissue removed aseptically from with a sterile scalpel, the compost removed by washing with SDW, and the roots transferred into 50 ml tubes, washed with PBS and the fresh weight determined. The tissue was macerated with a mortar and pestle, and tenfold dilutions plated on MacConkey+Kan agar for selective detection of STEC (Sakai) over the endemic microbiota. Data was analysed and plotted in Microsoft Excel, GraphPad Prism or RStudio, and significant differences calculated by one-way ANOVA for each tissue ($P > 0.05$).

Plant colonization for microscopy

N. benthamiana leaves ~6 cm length were infiltrated with STEC (Sakai) at 10⁷ c.f.u. ml⁻¹ or Pba (1043) 10⁶ c.f.u. ml⁻¹ in 0.5 × Murashige and Skoog (MS) medium (Sigma Aldrich) and maintained in an environmental cabinet with 16 h daylength and a continuous temperature of 21 °C. Samples were imaged 4–11 days post-infiltration for STEC (Sakai) and 1–4 days post-infiltration for Pba (1043). Broccoli seed was surface sterilized and germinated on purple capillary matting (Grofelt, UK) watered with bacteria at 10³ c.f.u. ml⁻¹ in 0.5 × MS as described previously [30] and imaged 5–11 days

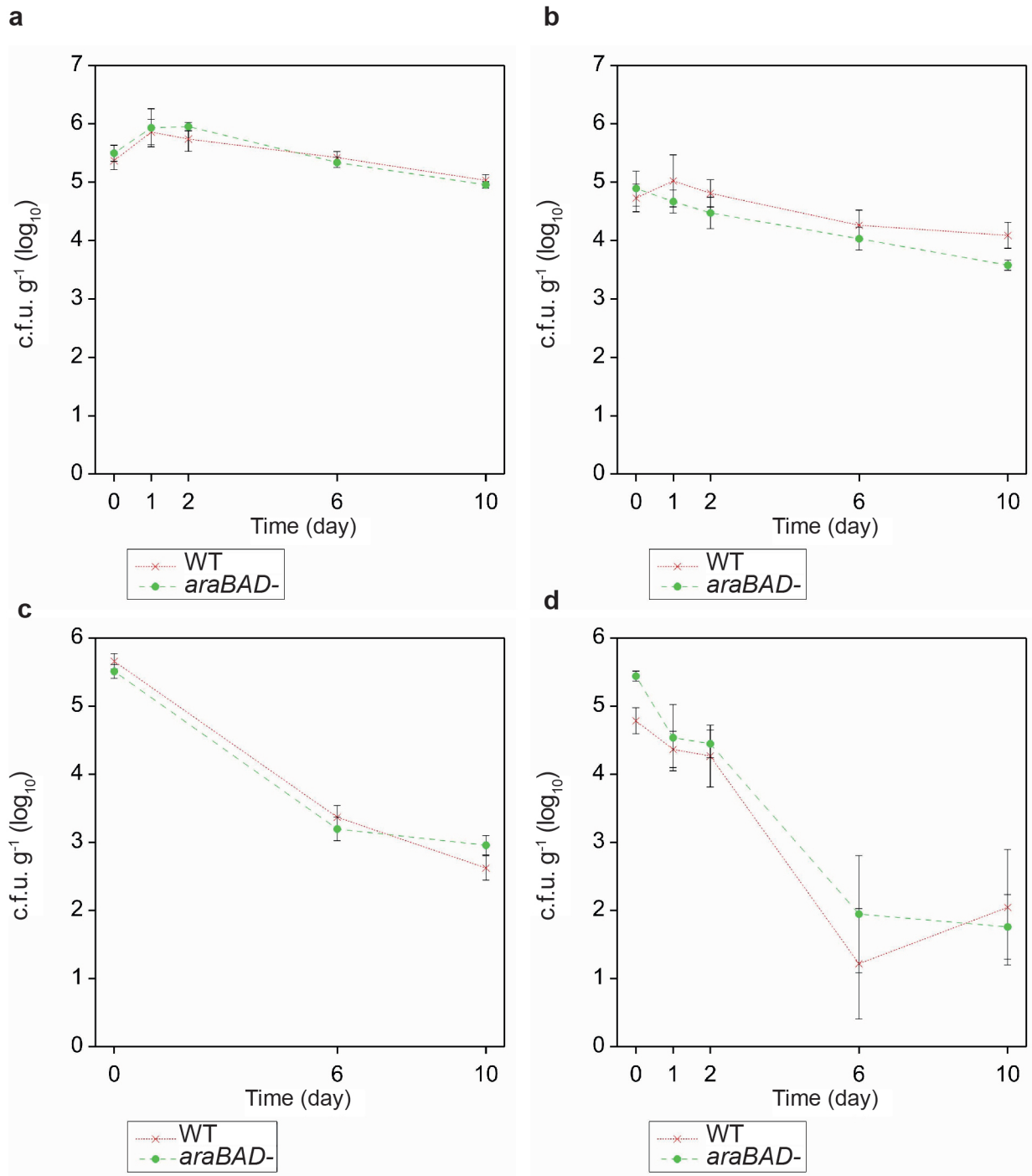


Fig. 1. Colonization of spinach and lettuce plants by Sakai WT and Sakai *araBAD*⁻. Number of bacteria recovered from lettuce or spinach leaves or roots. Boxplots show the median and interquartile range, overlaid with individual data points. STEC (Sakai) WT (cyan) or *araBAD* mutant (orange) were enumerated from four (lettuce) or five (spinach) plant samples and the experiment was repeated three times for spinach leaves. Samples were taken on day 0, 1, 2, 6 and 10 or for spinach leaves on day 0, 6, 10.

post-sowing. Inoculated cotyledons were harvested for antibody labelling 6 days post-sowing, with methods adapted from [31] using microtubule-stabilizing buffer (MTSB; 50 mM PIPES, 5 mM EGTA, 5 mM MgSO₄, 4.5 mM KOH adjusted to pH 7.0 with 10M KOH) and blocking buffer (2%

Albumin fraction V BSA in MTSB). Individual cotyledons were cut and placed in SDW, before incubation in blocking buffer for 2 h at 37°C, followed by overnight incubation at 4°C with α-O157 (*E. coli* antisera O157 monovalent, rabbit-derived, MAST ASSURE, M12030) diluted 1:500 in blocking

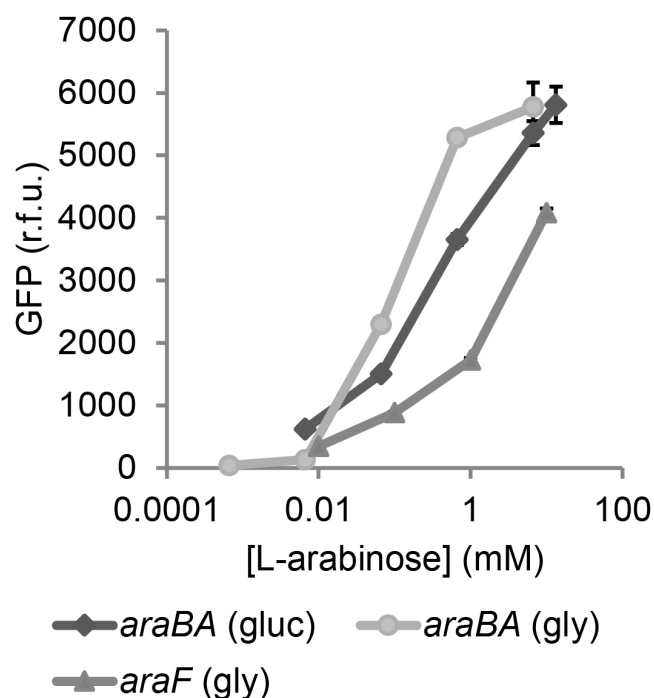


Fig. 2. Dose-response expression of STEC (Sakai) *araBA::gfp+* GFP measurements from STEC (Sakai) transformed with pACYC*araBA::gfp+* / pJM058 (*araBA*_{STEC}) or pACYC*araF*_{STEC}::*gfp+* / pJM066 (*araF*_{Pba}) grown at 18 °C in RD MOPS glucose (gluc) or RD MOPS glycerol (gly) supplemented with different concentrations of L-arabinose of 0, 0.01, 0.1, 1 or 10 mM. Maximal GFP expression levels are presented from the mean of triplicate samples at 5 h for pJM058 (*araBA*_{STEC}) in glucose; 8 h pJM058 (*araBA*_{STEC}) in glycerol; or 6 h pJM066 (*araF*_{Pba}) glycerol.

buffer. The samples were rinsed twice in MTSB and incubated at 37 °C with Alexa Fluor 568 nm goat anti-rabbit (Invitrogen A11011) diluted 1:1000 in blocking buffer followed by three 5 min washes and mounting in MTSB for imaging. For LM6 labelling, cotyledons were fixed in 100% methanol for 20 min at 37 °C followed by the addition of fresh methanol at 60 °C for 3 min, then gradual addition of water until the methanol was diluted to 20%. The cotyledons were then transferred to fresh SDW prior to antibody labelling as described above with LM6 (PlantProbes 1,5-arabitan, 1:50) and α -O157 (1:500) followed by secondary antibodies Alexa Fluor 488 nm goat anti-rat (Invitrogen A11006, 1:100) for 1 h and Alexa Fluor 568 nm goat anti-rabbit (Invitrogen A11011, 1:100) for a further 2 h.

Confocal microscopy

N. benthamiana leaves were infiltrated with sterile distilled water (SDW) and secured to microscope slides with double-sided adhesive tape prior to imaging of the abaxial surface as described previously [32]. Broccoli cotyledons were mounted on slides under cover-slips, in SDW for STEC (Sakai) or dry for Pba colonized leaves (since addition of water affects distribution of Pba rendering images meaningless), and imaged via either surface. Images were collected at Nyquist resolution

using a Nikon A1R confocal laser scanning microscope fitted with either an NIR Apo 40×0.8W water dipping lens or a CFI Plan Fluor 10×0.3 lens, and GaAsP detectors. Images represent false-coloured maximum intensity projections, unless otherwise stated, produced, linearly adjusted and re-sized using the splines option, where necessary for publication, using NIS-elements AR software. GFP, Alexa Fluor 488 nm (green) and chlorophyll (blue) were excited at 488 nm with emissions collected at 500–530 nm and 663–737 nm, respectively, and Alexa Fluor 568 nm (magenta) excited sequentially at 561 nm with emission at 570–620 nm.

RESULTS

Arabinose metabolism genes are induced on exposure to spinach extracts

Whole transcriptomic analysis of *E. coli* O157:H7 (Sakai) from a previous study had shown induction of arabinose-associated genes on exposure to spinach extracts (Table 1, extracted from [16]). The differential expression profiles were confirmed using a different approach (qPCR) and showed that the isomerase, *araA* was induced on exposure to spinach cell-wall polysaccharides but not lettuce (Fig. S1a). In contrast, *araA* expression was reduced after 1 h following infiltration into either spinach leaves (16.82-fold+/-4.68) or lettuce leaves (14.08-fold+/-1.89) relative to the no-plant control condition (Fig. S1b). Thus, STEC (Sakai) response to L-arabinose appears dynamic and is impacted by tissue type, raising the hypothesis that L-arabinose metabolism impacts STEC (Sakai) colonization of plants.

Arabinose metabolism is not essential for colonization of plants

To determine any essentiality of arabinose metabolism *in planta*, a knock-out mutant of the *araBAD* operon was made in STEC (Sakai) and tested in plant colonization ability primarily on spinach leaves and confirmed on the original plant models, spinach roots, lettuce roots and leaves. Both the WT and the *araBAD* mutant strains showed essentially the same patterns of colonization, on both species and in both tissue types, with a decrease from the initial starting inoculum over time, which was greatest on the foliar tissue, as expected from a high inoculation dose. There was no significant difference between the WT and *araBAD* mutant (>95%) and any reductions in colonization by the *araBAD* mutant strain were marginal, e.g. on lettuce roots at day 10 (Fig. 1). Thus, the arabinose metabolism genes are not essential for STEC (Sakai) colonization of spinach or lettuce, leaves or roots.

STEC (Sakai) *ara* genes are transcriptionally responsive *in vitro*

Arabinose metabolism was not essential for STEC colonization of plant tissue, yet it represents a major potential nutrient source in plant hosts. Therefore, expression of STEC (Sakai) *ara* metabolism and transport genes was examined in detail under *in vitro* conditions. Transcriptional GFP reporter plasmids were generated for metabolism (*araBA*)

Table 2. Expression of STEC and pba *ara* genes in native and non-native backgrounds show differences in regulatory control

Plasmid/isolate	STEC (Sakai)	Pba (1043)	STEC Δ <i>araC</i> (Sakai)
pACYC <i>araBAD</i> _{STEC} :: <i>gfp</i> + (pJM058)	11151 (\pm 493)	15566 (\pm 93)	15809 (\pm 224)
pACYC <i>araF</i> _{STEC} :: <i>gfp</i> + (pJM066)	4086 (\pm 64)	35215 (\pm 560)	275 (\pm 3)
pWSK <i>araBA</i> _{Pba} :: <i>gfp</i> + (pJM067)	1882 (\pm 138)	10213 (\pm 182)	NT
pWSK <i>araF</i> _{Pba} :: <i>gfp</i> + (pJM068)	3278 (\pm 260)	25075 (\pm 1804)	NT
pACYC <i>araBA</i> _{Pba} :: <i>gfp</i> + (pJM064)	5050 (\pm 124)	*	186 (\pm 3)
pACYC <i>araF</i> _{Pba} :: <i>gfp</i> + (pJM065)	4868 (\pm 81)†	†	1747 (\pm 20)†

GFP was measured in response to 10 mM L-arabinose at maximal expression times (4 or 6 h), for cultures grown in RD MOPS glycerol at 27 °C. Blue background indicates native expression and red indicates non-native expression.

(representative experimental replicate, total four reps).

*>600000 r.f.u.

†High background: 1100–1300 r.f.u. in the absence of arabinose.

NT, not tested.

and high-affinity transport (*araFGH*) genes of STEC (Sakai). Responsiveness to L-Arap was measured at 18 °C, a temperature relevant to colonization of plants, and to show utility of the reporters as biosensors. Growth of STEC (Sakai) transformed with any of the reporter plasmids (Table S1) in defined medium with/out L-Arap was not impaired compared to the vector only control (pKC026) (Fig. S2). GFP levels from *araBA*_{STEC} (pACYC*araBA*_{STEC}::*gfp*+, pJM058) increased in a dose-responsive manner to the concentration of L-Arap when cultures were grown in either glycerol ($R^2=0.933$, $t=7.481$, $P=0.0017$) or glucose ($R^2=0.973$, $t=12.012$, $P=0.0003$) (Fig. 2). Expression of *araF*_{STEC} (pACYC*araF*_{STEC}::*gfp*+, pJM066) increased in a similar manner ($R^2=0.891$, $t=4.674$, $P=0.0185$), although expression was ~1.5-fold lower than for *araBA*_{STEC} (Fig. 2). The specificity of the reporters was confirmed by growing STEC (Sakai) transformed with *araBA*_{STEC} (pJM058) or *araF*_{STEC} (pJM066) reporters in medium supplemented with xylose, which resulted in no expression of *araBA*_{STEC} (pACYC*araBAD*_{STEC}::*gfp*+, pJM058). As expected *araF*_{STEC} (pACYC*araF*_{STEC}::*gfp*+, pJM066) was induced in xylose as it is a low-affinity xylose transporter, but at a reduced level compared to arabinose (Table S2). Utility of *araBA*_{STEC} (pACYC*araBAD*_{STEC}::*gfp*+, pJM058) was also demonstrated during biofilm formation of non-pathogenic *E. coli* [33]. Therefore, the promoter constructs could effectively be used as specific biosensors for the presence of L-arabinose.

Pba (SCRI1043) *ara* genes are transcriptionally responsive *in vitro*

P. atrosepticum (Pba) and STEC O157:H7 isolates share a high proportion of their genomes [34] and since *ara* genes are widespread in bacteria, we hypothesized that L-arabinose metabolism and transport would be conserved between both species. The transcriptional activity of *araBA* and *araF* from Pba (strain 1043) were similarly quantified using the low copy number GFP reporter system (Supplementary Material), in response to L-Arap. The dose response of *araBA*_{Pba} (pJM067) and *araF*_{Pba} (pJM068) was similar to that of STEC (Sakai), and

both Pba (1043) constructs produced maximal levels of GFP after 4 h in RD MOPS glycerol supplemented with 10 mM L-Arap. Expression from the *araF*_{Pba} reporter was ~2.5-fold higher than from the *araBA*_{Pba} reporter (Table 2). The response to xylose was similar to the STEC (Sakai) constructs, although the *araF*_{Pba} transporter reporter expression level in xylose was 40% of the level in L-arabinose, showing some lack of specificity.

Regulatory control of arabinose metabolism and transport differs between Pba and STEC

Genetic organization of the *ara* loci are different between the species. In STEC arabinose metabolism and regulation are linked and are *in cis*, while transport is *in trans* and dependent on AraC (and CRP) binding. In Pba (1043) metabolism, transport and regulation are linked *in cis*, but the epimerase is located *in trans*, and *araC* is embedded within the transport polycistronic operon at the 3'-end (Fig. S3a). The STEC *araBA* genes are co-transcribed and the epimerase, *araD* is adjacent but assumed to be transcribed separately via a canonical ribosome binding site, while *araC* is divergently transcribed from *araBA* [35]. Six REP sequences are present 5' to the *araD*_{STEC} CDS. The AraC regulatory protein sequences share 57% overall amino acid identity, with conservation in all L-Arap binding residues, 8/10 beta barrels, one of the dimerization alpha helices and the DNA-binding / H-T-H domain (Fig. S3b)

To compare the STEC and Pba L-arabinose regulatory networks, exchange experiments were performed, where the L-arabinose biosensor plasmids were transformed into the non-native host. This artificial situation was used for a direct comparison between the species. The pACYC-based Pba reporters transformed into STEC (Sakai) allowed direct comparison as growth not inhibited in the presence of L-Arap. For each pair of strain backgrounds, expression was highest in Pba (1043), at levels up to 8.6-fold for the *araF*_{Pba} reporter (Table 2). The least amount of difference was for *araBAD*_{STEC} where the levels of expression were relatively similar, but still

Table 3. GFP measurements from *araBAD_{STEC}* in STEC (Sakai) inoculated in different media types

Medium	Day 0	Day 1	Day 2	Day 5
arabinose 6.661 mm	14.293	20234.293	58610.325	64072.291
arabinose 0.666 mm	10.193	10329.918	24307.419	28889.353
arabinose 66.609 μm	19.933	5752.784	1342.746	1733.018
arabinose 6.661 μm	29.340	576.825	292.451	415.993
arabinose 0.666 μm	17.607	< 0	45.755	330.483
RD MOPS only	30.287	42.179	< 0	< 0
0.2% glucose	38.473	23.344	< 0	99.814
Spinach leaf lysate	< 0	< 0	< 0	ND
Tomato leaf lysate	243.600	< 0	1494.266	< 0
Lettuce leaf lysate	< 0	< 0	< 0	< 0

Expression from STEC (Sakai) transformed with pJM058 (pACYC*araBAD_{STEC}::gfp+*).

*Values that were less than 0 after normalization are indicated by '< 0'; ND – no data.

higher in the Pba (1043) background. There was a difference between each pair of *ara* operons, such that the *araF_{Pba}* promoter was stronger than *araBA_{Pba}*, whereas the opposite was true for the STEC (Sakai). Furthermore, *araF_{STEC}* was over-induced in the Pba (1043) background, reversing the trend seen in the native STEC background. Hence, in Pba (1043), arabinose transport via the high affinity AraF transporter is particularly highly induced, even when the *araF* promoter is non-native, which is the opposite of the situation observed in STEC.

Regulatory control via AraC was examined with an *araC* deletion mutant for STEC (Sakai). Under *in vitro* conditions, the *araBAD_{STEC}* reporter construct was still induced in this background, to levels marginally higher than the WT (Table 2), (as expected due to negative feedback regulation of *araBA*) whereas *araF_{STEC}* was completely repressed in the absence of *araC*. In contrast, the Pba reporter *araBA_{Pba}* was repressed in the absence of *araC* in the STEC (Sakai) background, whereas *araF_{Pba}* was still expressed to ~50% of that in the WT STEC (Sakai) background (Table 2). Taken together, the data shows differences in regulatory control and potentially specificity in AraC-dependent regulation, with some level of constitutive expression of *araF_{Pba}*.

STEC and Pba *ara* genes are expressed in *in planta* and in some plant extracts

Since L-arabinose is normally complexed in plant tissue as L-Araf, *in vitro* plant extracts were used for quantification of gene expression in the absence of other factors. Expression from STEC (Sakai) transformed with the *araBA_{STEC}* reporter was assessed *in vitro* in media supplemented with a range of different plant extracts, to determine whether L-arabinose dependent-activity from the promoters could be quantified. However, detection of *araBA_{STEC}* was absent or minimal over 5 days of exposure in the plant extracts compared to the medium with L-Arap added (Table 3), which indicated

that the levels of L-arabinose were either below 0.6 μm or components in the plant extracts masked GFP detection by the plate reader. Therefore, a qualitative approach using microscopy was taken to determine if *ara* gene expression could be detected *in planta* over longer time periods.

Expression was assessed qualitatively from bacteria inoculated into the internal tissue of leaves and from leaf surfaces. Two plant models were chosen that are known to support high levels of colonization and in an attempt to override differences in the plant-microbe interactions, i.e. potential PCWDE production by Pba. *N. benthamiana* was used for infiltrated bacteria, since unlike spinach or lettuce, it allows unrestricted growth of internalized *E. coli* O157:H7 (Sakai) and Pba (1043) [36]; surface-associated bacteria were assessed from inoculation of broccoli microgreens, as a representative edible crop species that supports high growth of both isolates [30]. Colonies formed in both species after 11 days in the *N. benthamiana* apoplast, and they both expressed metabolism, *araBA* and transport, *araF* genes in native backgrounds (Fig. 3a–d). Some variation in expression was evident at the single cell level in all four constructs. Both STEC and Pba formed extensive colonies on the plant cell-wall margins of broccoli cotyledons after 6 days, and both expressed metabolism and transport reporter genes (Fig. 3e–h). Variation at the single cell level was evident (quantification on plant leaves was not possible). No GFP was detected from the empty vector (pKC026) from either STEC or Pba in broccoli or *N. benthamiana* (Fig. S4a, b). Thus, both species expressed L-arabinose metabolism and transport genes *in planta*, under different model systems.

To see whether *ara* gene regulatory control observed *in vitro* held *in planta*, expression of *araBA_{STEC}* (pJM058) transformed into Pba (1043) and *araBA_{Pba}* (pJM064) transformed into STEC was examined from surface-associated bacteria on broccoli leaves. Both constructs were expressed *in planta* in

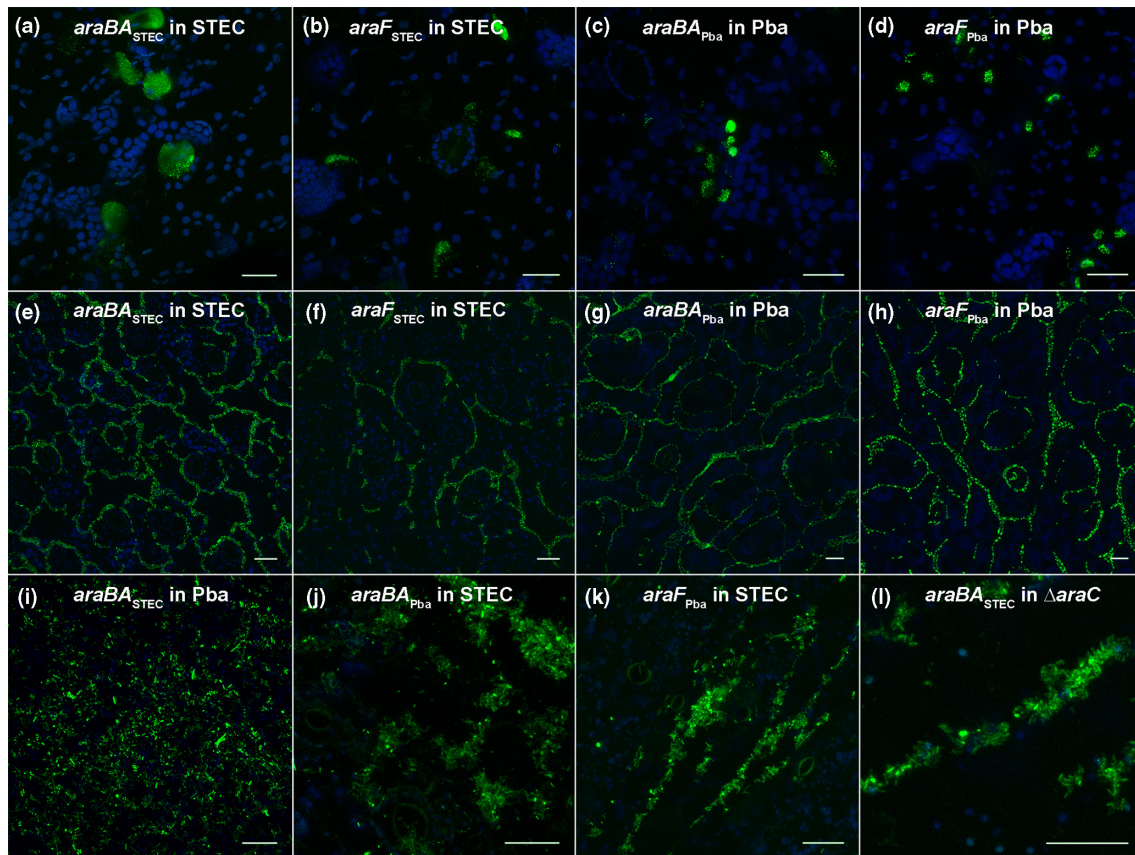


Fig. 3. *In planta* expression of arabinose metabolism and transport genes GFP reporter expression of *araBA* and *araF* plasmid constructs in STEC (Sakai), Pba (1043) or STEC (Sakai $\Delta araC$) backgrounds. Reporter constructs transformed in native backgrounds were examined from bacteria infiltrated into *N. benthamiana* leaves and colonies imaged at day 11 (a), day 4 (b, c) or day 1 (d) after inoculation. Reporter constructs transformed in native or non-native backgrounds were examined on broccoli cotyledons germinated from inoculated (imbibed) seeds. Bacterial colonies of STEC (e, f, j and k), Pba (g–i) or $\Delta araC$ (l) were present on the surfaces of cotyledons at day 5 (l), day 6 (e, j, k), day 7 (i), day 10 (g, h) and day 11 (f). Chloroplasts in the epidermal and mesophyll cells are shown (blue). All scale bars represent 25 μm .

the non-native backgrounds, although morphological differences occurred in Pba cells presumably due to GFP toxicity (Fig. 3i–k). Variation in expression at the single cell level was evident and in STEC: GFP levels from most cells was low with only a small number expressing a high level. A similar pattern occurred for STEC hosting *araF*_{Pba}. GFP was only detected from the *araBA*_{STEC} reporter in the STEC $\Delta araC$ mutant background (Fig. 3l), in keeping with quantitative data (Table 2). There was no detectable GFP from *araF*_{STEC} or either of the Pba constructs *araBA*_{Pba} or *araF*_{Pba} in this background (Fig. S4d–f), confirming their responsiveness to L-arabinose.

Since Pba (1043) encodes PCWDE, the responsiveness of the *araBA*_{Pba} reporter was quantified *in vitro* from incubation with a crude preparation of leaf cell-wall polysaccharides (spinach). Glycerol was added as a carbon source to facilitate bacterial growth and increased glycerol concentration used to promote induction of any PCWDE [37], although their expression was not measured per se. Growth of Pba (1043) reached comparable densities in all substrates and was not

affected by variable glycerol concentration. GFP levels for *araBA*_{Pba} reached ~ 450 r.f.u. in the presence of cell-wall polysaccharide extract with 0.4% glycerol, compared to ~ 150 r.f.u. in their absence, with no difference between 6- and 24 h incubation (Fig. 4a). This level of expression was \sim tenfold lower compared to *in vitro* response to L-Arap (Table 2), indicative of the difference in response to free L-Arap and complexed L-Araf. In contrast, no expression was detected from the STEC (Sakai) *araBA*_{STEC} reporter in cell-wall polysaccharides. However, incubation with purified L-Araf induced expression of STEC (Sakai) metabolism genes *araA* and *araD* in the presence the debranched form of pectin or as oligomers as arabinobiose and arabinoseptose, while the fully branched form of pectin did not (Fig. 4b). Furthermore, STEC (Sakai) showed marginal growth in minimal media supplemented with just arabinobiose or arabinoseptose, reaching 6.716 (± 0.240) and 7.024 (± 0.724) Log_{10} c.f.u. ml^{-1} (respectively) after 48 h at 18 °C from an inoculum of 6 Log_{10} c.f.u. ml^{-1} , compared with 8.643 (± 0.058) or 8.134 (± 0.365) Log_{10} c.f.u. ml^{-1} in medium supplemented with L-Arap or glucose

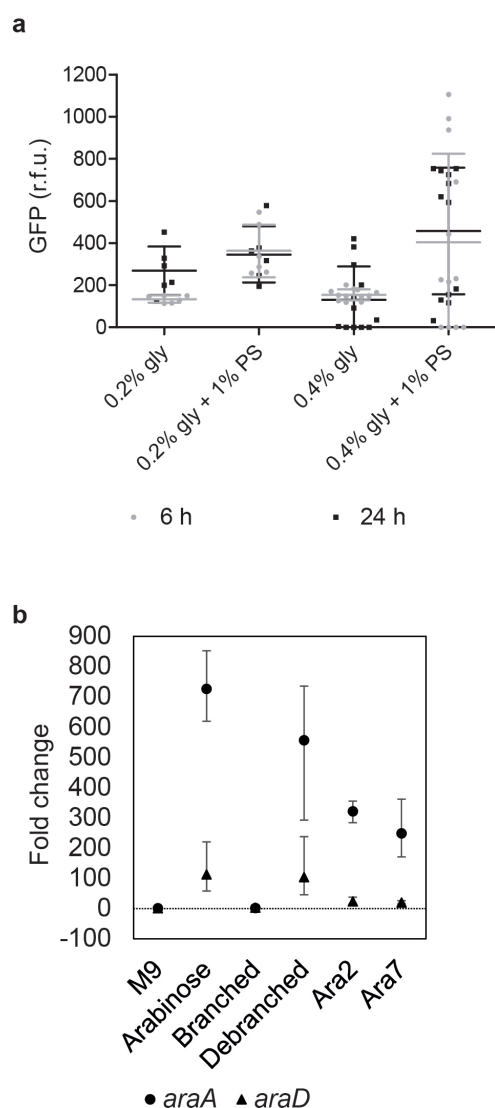


Fig. 4. Expression of *araBA_{Pba}* in the presence of plant cell-wall polysaccharides. Reporter expression for *araBA_{Pba}* (pJM067, pWSKaraBA_{Pba}::gfp+) in Pba (1043) following incubation in 1% (w/v) spinach alcohol-insoluble polysaccharide (1% PS) extract at two glycerol concentrations (0.2, 0.4%), for 6 or 24 h at 27°C (a). Fluorescence measurements, normalized for cell density, are expressed as average r.f.u. (+/-SD) relative to the control plasmid, from triplicate samples \times two or four independent experiments (for 0.2% or 0.4% glycerol, respectively). Gene expression measured directly from STEC (Sakai) *araA* and *araD* (b) after one h in M9 minimal medium only (M9); or supplemented with 0.2% L-arabinose (arabinose); (1-5)- α -linked pectin backbone which contains (1-3)- α -linked and possibly (1,2)- α -linked L-arabinofuranosyl residues (branched); (1-5)- α -linked backbone treated with α -L-arabinofuranosidase (debranched); (1-5)- α -L-arabinobiose (Ara2); and (1,5)- α -L-arabinoheptose (Ara7). Gene expression is expressed as averaged fold-change (+/-SD) relative to the reference gene (*gyrB*), from three biological \times three technical replicates.

(respectively). No growth occurred in a BSA-supplemented control medium. Therefore, STEC (Sakai) was responsive to L-Araf in oligosaccharides *in vitro* and able to utilize them for growth.

Arabinan complement of horticultural species

L-arabinose is abundant present in plant cell-wall pectin and glycoproteins as L-Araf, while L-Arap is much less so and a minor glycan in plants in comparison to glucose, sucrose and fructose [4]. In crude leaf cell-wall polysaccharide extracts of spinach and lettuce, spinach contained fivefold more L-Arap compared to lettuce leaf extracts [16]. A screen of pectin-enriched fractions (CDTA treatment) from 15 plant horticultural species and cultivars, with different tissues (leaves, root or sprouted seeds) with antibody probes confirmed that arabinans as L-Araf were more abundant of spinach than in lettuce across a range of cultivars [28]. It also showed that broccoli microgreens were comparable to sprouted seeds for linear pentasaccharides of L-Araf(1-5)- α -L-arabinan, LM6 (Table S3). Closer examination of broccoli microgreen-leaf showed (1-5)- α -L-arabinan (LM6) was dominant in the pectin-enriched fraction together with homogalacturonan (JIM7), and AGP (LM2) as the dominant glycoprotein (Fig. 5a). Processed arabinans in RG-I were detected in broccoli as a minor component (LM16). NaOH-derived enrichments detected xyloglucans as a principle component of hemicellulose (LM15, LM25). The distribution of (1-5)- α -L-arabinan (LM6) was visualized in broccoli cotyledons by antibody staining (Fig. 5b), with or without inoculated STEC (Sakai). (1-5)- α -L-arabinan was distributed along cell margins (LM6, green) and coincided with STEC (Sakai) localization (α -O157 antibody, magenta).

DISCUSSION

In STEC (Sakai), L-arabinose metabolism is active in plants but non-essential for its colonization: *ara* genes were responsive *in vitro* to leaf extracts and *in planta* during colonization for a range of plant models, but the gene cluster knock-out was not affected in spinach or lettuce colonization over a 10 day period. Therefore, it appears that STEC (Sakai) is able to take advantage of L-arabinose as an abundant plant-derived metabolite in an opportunistic manner. Use of different plant models to demonstrate either surface or internal plant colonization showed that STEC (Sakai) responded to L-arabinose once it establishes *in planta* colonies. Repression of metabolic genes on initial introduction into plants suggests that a period of adaption occurs before the L-arabinose system is induced *in planta*. The lack of dependency on L-arabinose metabolism by STEC (Sakai) in plant colonization is indicative of metabolic redundancy, in line with the non-essentiality of the *ara* gene cluster for growth in rich media [38], and in keeping with *E. coli* as a heterotroph [39].

For microbes that cross biological kingdoms, metabolic diversity is an essential asset. The *Enterobacteriaceae* tend to have mosaic genomes contributing to generalism, also described for plant or animal-associated pseudomonads [40]. Intestinal microbiota within animal hosts have access to and can metabolize ingested plant material, e.g. of *E. coli* KduI facilitates conversion of pectin-derived D-galacturonate in gnotobiotic mice [41]. In nature, fermentation of hexoses occurs through the Entner Duodoroff (ED) pathway, which we found was induced in STEC (Sakai) on *in vitro* exposure to plant extracts [16]. Discovery of non-phosphorylative pathway for pentoses, including L-arabinose,

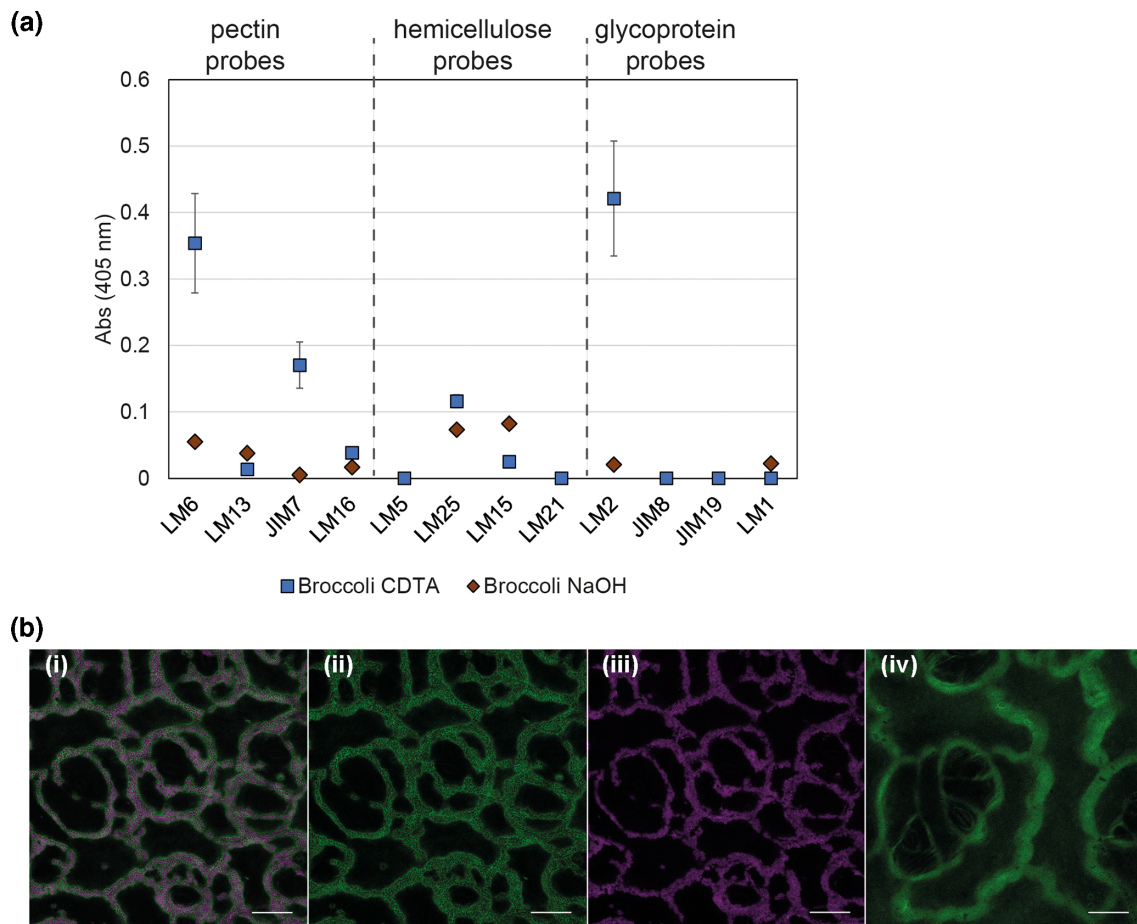


Fig. 5. Glycan complement of broccoli in plant cell-wall polysaccharide extracts. Alcohol insoluble residues for broccoli microgreen enriched for pectin by CDTA extraction (blue) or hemicellulose by NaOH extraction (brown) (a). ELISA quantification for indicated primary antibodies probes with secondary antibody conjugated with HRP, is shown as absorbance at 405 nm, averaged (+/-SD) from four extractions \times two technical repeats. One-way ANOVA showed significant differences between antibodies for both extractions (pectin $P=1.11 \times 10^{-4}$; hemicellulose $P=0.043$). Detection of (1-5)- α -L-arabinans in broccoli cotyledon tissue (b). Broccoli cotyledons, 7 days after germination in the presence of *E. coli* STEC Sakai (WT), treated with LM6 primary antibody, labelled with Alexa Fluor 488 nm (green) and α -O157 primary antibody, labelled with Alexa Fluor 568 nm (magenta) secondary antibody showing single-layer confocal images of inoculated (i, ii, iii) or non-inoculated tissue (iv). All scale bars represent 25 μ m.

may also facilitate colonization of plant niches as it has an equivalent role to the ED pathway [42]. Furthermore, promiscuity in pentose transporters broadens metabolite options when glycans become scarce [43], e.g. on leaf surfaces.

Examination of a range of plant species showed that arabinans (as *L-Araf*) were differently distributed in the pectin and AGP components between plant species and tissues. *L-Arap* is present in leaf cell-wall polysaccharides at relatively low levels compared to other glycans but also with still plant-species and tissue differences [4, 16]. STEC (Sakai) is responsive to *L-Araf* in debranched pectin and as arabinan oligomers as well as *L-Arap*, but not in branched pectin. We previously showed that STEC (Sakai) ECP targeted (1-5)- α -L-arabinans present in RG1 of pectin for adherence to plant cells [28], which may account in part at least, for observed differences in colonization. This raises a potential link between fimbrial-dependent attachment and metabolism to facilitate plant colonization, as has been seen in

uropathogenic *E. coli* with type 1 fimbriae [39]. Since *E. coli* do not encode pectinases and cannot access *L-Araf* directly, there are two potential sources for STEC *in planta*: from turn-over of cell-wall components, e.g. during plant growth and development from activity of plant-derived α -L-arabinofuranosidase glycoside hydrolases [3], and/or from pectinase activities of the endemic microbiome [44]. All plant inoculation experiments for STEC (Sakai) included endemic microbiomes in our work incurring microbial ecological interactions, whether competitive, neutral or mutualistic. As such, for STEC to have sufficient ecological fitness to invade established microbiomes, capacity for diverse metabolite utilization is critical.

P. atrosepticum is a member of the *Enterobacteriaceae* that differs from *E. coli* in that it encodes PCWDE, making them a useful comparison pair for metabolite use. *Pba* *L*-arabinose metabolism appears to be linked to PCWDE expression through the response regulator ExpM (RssB in *E. coli*) [45] since *araA*, *C* and

H were significantly induced > twofold after 12 h inoculation on potato tubers in a *Pba expM* mutant [22]. The quorum-sensing dependent *RsmA-rsmB* negative regulatory system (*CsrA-csrB* in *E. coli*) links metabolism to production and secretion of PCWDE through RpoS as a transcriptional activator of the *RsmA* [22]. *ExpM/RssB* negatively regulates RpoS that when phosphorylated, facilitates degradation of the sigma factor by ClpXP protease activity, and phosphorylation of *ExpM/RssB* is controlled (in part) by the sensor kinase *ArcB*, which is itself inactivated during carbon starvation [46]. Hence *ExpM* activity alleviates *RsmA*-dependent repression of the PCWDE. Therefore, the elevated RpoS levels that would occur in the *expM* mutant or in the presence of inactive, unphosphorylated *ExpM* induce expression of alternative metabolic pathways, including L-arabinose. The *ExpM*-dependent effect appeared specific for arabinose since the xylose genes were not similarly induced in *Pba*, and the *ara* genes were not impacted by other key virulence regulator mutants [22, 47].

Despite the taxonomic similarity of STEC (Sakai) and *Pba* (1043), differences exist in the genetic organization, expression profiles and regulation of L-arabinose metabolism and transport. L-arabinose transport appears to be more sensitive in *Pba* compared to STEC, since *in vitro* expression of *araF* was ~threefold higher than the metabolic genes *araBA*, whereas the converse occurred in STEC. There was also evidence for constitutive, or non-AraC dependent expression of *Pba* L-arabinose transport (via *AraF*), whereas STEC *araF* induction was completely dependent on AraC. Although the AraC amino acid sequences for L-arabinose binding and DNA-binding domains were conserved in both species, there is sufficient species-dependent specificity such that negative feedback of *araBA* by AraC was lost when the *Pba* genes were expressed in the STEC *araC* mutant, and *araF*_{STEC} was over-induced in the *Pba* background. Although it was not possible to make direct comparisons between *Pba* and STEC in *in planta*, expression of the biosensor reporters echoed the *in vitro* expression patterns for both species, thus provided a faithful representation.

STEC and *Pba* have different primary reservoirs (ruminants and potatoes, respectively), and as such, STEC can be considered an invader of plant niches [48]. Within the same environment, e.g. plants niches, different traits will contribute to their relative fitnesses. Although both species possess metabolic flexibility, nuanced regulatory differences and the link to PCWDE expression account for their different strategies for L-arabinose metabolism. In STEC, the system is primed toward opportunistic access, weighted to catabolic activation, while in *Pba*, the system is primed towards direct access through high-affinity transport. This has enabled STEC to exploit such an abundant plant metabolite during colonisation of plants, but not in a dependent manner.

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Conflicts of interest

The authors declare that there are no conflicts of interest.

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