The influence of plant species, tissue type and temperature on the capacity of Shigatoxigenic *Escherichia coli* to colonise, grow and internalise into plants.

**Running title**: STEC growth characteristics in plants

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Abstract (247 / 250 words)

Contamination of fresh produce with pathogenic *Escherichia coli*, including Shigatoxigenic *E. coli* (STEC), represents a serious risk to human health. Colonisation is governed by multiple bacterial and plant factors that can impact the probability and suitability of bacterial growth. Thus, we aimed to determine whether the growth potential of STEC for plants associated with foodborne outbreaks (two leafy vegetables and two sprouted seed species), is predictive for colonisation of living plants as assessed from growth kinetics and biofilm formation in plant extracts. Fitness of STEC was compared to environmental *E. coli*, at temperatures relevant to plant growth. Growth kinetics in plant extracts varied in a plant-dependent and isolate-dependent manner for all isolates, with spinach leaf lysates supporting the fastest rates of growth. Spinach extracts also supported the highest levels of biofilm formation. Saccharides were identified as the major driver of bacterial growth, although no single metabolite could be correlated with growth kinetics. The highest level of *in planta* colonisation occurred on alfalfa sprouts, though internalisation was 10-times more prevalent in the leafy vegetables than in sprouted seeds. Marked differences in *in planta* growth meant that growth potential could only be inferred for STEC for sprouted seeds. In contrast, biofilm formation in extracts related to spinach colonisation. Overall, the capacity of *E. coli* to colonise, grow and internalise within plants or plant-derived matrices were influenced by the isolate type, plant species, plant tissue type and temperature, complicating any straight-forward relationship between *in vitro* and *in planta* behaviours.
Importance (149 / 150 word)

Fresh produce is an important vehicle for STEC transmission and experimental evidence shows that STEC can colonise plants as secondary hosts, but differences in the capacity to colonise occur between different plant species and tissues. Therefore, an understanding of the impact of these plant factors have on the ability of STEC to grow and establish is required for food safety considerations and risk assessment. Here, we determined whether growth and the ability of STEC to form biofilms in plants extracts could be related to specific plant metabolites or could predict the ability of the bacteria to colonise living plants. Growth rates for sprouted seeds (alfalfa and fenugreek) exhibited a positive relationship between plant extracts and living plants, but not for leafy vegetables (lettuce and spinach). Therefore, the detailed variations at the level of the bacterial isolate, plant species and tissue type all need to be considered in risk assessment.
Introduction

Contamination of fresh produce from Shigatoxigenic *Escherichia coli* (STEC) presents a serious hazard as a cause of food-borne illnesses, diarrhoea and enterohemorrhagic disease. Fresh produce is a major vehicle of transmission of STEC, with foods of plant origin accounting for the majority of *E. coli* and *Shigella* outbreaks in the USA (47). Fresh produce is often eaten raw or minimally processed and contamination of the produce can occur at any point along the food chain from farm to fork, with major outbreaks e.g. from spinach (27) and sprouted seeds (5). STEC has been shown to interact with plants and can use them as secondary hosts (15, 22), which has implications for pre-harvest contamination, as well as persisting on post-harvest produce (26, 28, 30).

Colonisation of host plants by *E. coli* is governed by a range of environmental, bacterial and plant factors. Initial contact and attachment of bacteria on plant tissue is defined by motility, adherence factors and plant cell wall components (58, 59), while establishment is influenced by a range of plant biotic (40, 61) and abiotic factors (14, 56). The ability of bacteria to grow in the presence of plant material is a key factor in assessing risk, and although proliferation is well known to be influenced by physio-chemico factors (4, 53), risk assessments for STEC on fresh produce tend to consider plants as a homogenous whole (11, 19, 48).

STEC preferentially colonise the roots and rhizosphere of fresh produce plants over leafy tissue and have been shown to internalise into plant tissue, where they can persist in the apoplastic space as endophytes (12, 66). The apoplast contains metabolites, such as solutes, sugars, proteins and cell wall components (50) and as such provides a rich environment for many bacterial species, both commensal bacteria and human pathogens (18, 25). The rate of
STEC internalisation is dependent on multiple factors including the plant species and tissue (67) and how plants are propagated (16, 17). Specificity in the response of STEC to different plant species and tissue types has been demonstrated at the transcriptional level (8, 34). Therefore, there is a need to take into account specificity of the STEC-plant interactions that could impact risk.

Determination of the growth potential of a bacterial population takes into account the probability of growth together with the suitability of the growing population for a particular environment (20). It is used as a measure in risk assessment, e.g. for growth of STEC in water (64). In plant hosts, bacterial growth potential is governed by several factors, including bacterial growth rates, initial adherence and colony establishment, which is often in biofilms, as well as plant-dependent factors including metabolite availability and plant defence responses (23). Therefore, the aim here was to determine if in vitro growth kinetics and biofilm formation of STEC in plant extracts, together with plant metabolite analysis, could be related to colonisation of plants that are associated with food-borne outbreaks, and hence inform on growth potential of STEC in planta. Use of genetically distinct E. coli isolates (two STEC, two environmental and one laboratory isolate) enabled assessment of bacterial phenotypic variation within plants or plant-derived matrices to be compared. Growth kinetics and biofilm formation were quantified in different tissue extracts of two leafy vegetables, lettuce and spinach, and two sprouted seeds, fenugreek and alfalfa sprouts. Growth kinetics was related to metabolomics of the extracts. Quantification of in planta colonisation and internalisation allowed a correlation analysis for the two STEC isolates.
Results

E. coli growth rates in plant extracts

To relate growth potential to colonisation of STEC in fresh produce plants, in vitro growth rates were first measured in plant extracts. Primary modelling of in vitro growth data in plant extracts successfully fitted 86.7 % (117 of 135) growth curves with a non-linear Baranyi model (SM1). Mis-fits were improved by manually truncating the growth curves to before the observed decrease in cell density that occurred in stationary phase, resulting in $R^2_{\text{adj}} = 0.996$ (Fig. S1, Table S1a). Comparison of the maximum growth rates ($\mu$) showed highest growth rates in spinach extracts, with fastest growth in leaf lysates at 18 °C or apoplast at 25 °C (Fig. 1A), while in lettuce the fastest growth occurred in apoplastic extract at all temperatures tested (Fig. 1B). All isolates grew consistently faster in fenugreek sprout extracts than in alfalfa, and either sprout extract supported faster growth than defined medium (RDMG) (Fig. 1C). The E. coli O157:H7 isolates showed differential responses in the different extracts and their growth rates were as fast or faster than the environmental isolates in almost all extracts. The lowest growth rates occurred for the laboratory-adapted isolate MG1655. The plant extract tissue-type as well as the bacterial isolate significantly impacted $\mu$, from a two-way ANOVA at 18 °C ($F(4, 7363) = 76.3; p < 0.0001$ and $F(8, 7363) = 436.4; p < 0.0001$, for bacterial isolate and extract type, respectively) and at 20 °C ($F(4, 8387) = 160.3; p < 0.0001$ and $F(8, 8387) = 416.1; p < 0.0001$, for bacterial isolate and extract type, respectively).

Growth was almost always highest at 25 °C, although with exceptions, e.g. for E. coli O157:H7 isolate ZAP1589 in lettuce extracts. Growth characteristics were similar at both 18 and 20 °C, but $\mu$ were in general lower at 20 °C than at 18 °C. This counterintuitive result was
reproducible and occurred in all growth experiments. It meant that secondary modelling for
temperature was not possible. It was possible, however, for temperature-effects of growth in
the defined medium without plant extracts, which produced a linear distribution for temperature
for all five *E. coli* isolates ($R^2 = 0.996$ to 1) (SM2), indicating the effect was due to the plant
extracts and not a systemic error.

Metabolite analysis of fresh produce plant extracts

To establish the impacts of different plant components on the growth of the *E. coli* isolates,
metabolite analysis was determined for the extracts. Detection of absolute levels of mono- and
disaccharides (sucrose, fructose, glucose, arabinose) showed the highest abundance in
fenugreek sprout extracts, followed by lettuce apoplast and lettuce leaf lysates (Table 2).
Sucrose was the most abundant sugar in all species and cultivars, except for alfalfa, which had
high levels of fructose and glucose. Arabinose was only detected in the apolastic fluid of
spinach and lettuce, accounting for 0.36 % and 0.23 % of all sugars, respectively. A two-way
ANOVA showed significant differences for tissue types ($F (7, 60) = 16.5; p < 0.0001$).
The levels of amino acids and other metabolites were determined from identification of 116
polar metabolites, of which 60 were assigned and mapped onto a simplified polar metabolite
pathway for plants to visualise metabolite availability for the bacteria (Fig. S2). The abundance
ratio of each compound against the internal standard ribitol, generated a response ratio (RR)
to allow semi-quantitative comparison (Table S2). Differences occurred between species and
tissue types in a similar pattern to the mono- and disaccharides (Table 2), and for 12
metabolites including fructose, glucose and sucrose, there were significantly different RR (two-
way ANOVA and Tukey multiple comparison, $F (7, 854) = 37.2, p < 0.0001$). Small amounts of
arabinose could be found in all tissues with no significant differences between host species or
tissue types. Grouping metabolites by structure (Fig. 2A) for monosaccharides,
polysaccharides, amino acids, organic acids and other metabolites, showed that the highest
total saccharides were present in fenugreek sprouts, while alfalfa was higher in
monosaccharides and amino acids. The organic acids in spinach apoplast consisted mainly of
oxalic acid, which was almost double the amount in spinach leaf lysates. The percentage
composition showed that the majority of metabolites in all lettuce extracts are polysaccharides,
compared to mainly of organic acids in all spinach extracts.

Significant variation of the metabolite content occurred between plant tissues, as well as for
and individual metabolites (two-way ANOVA assuming a parametric distribution, F (420, 854) =
43.15; p < 0.001). A principal components analysis (PCA) showed that the first five
components accounted for ~ 85 % of variance, and 50 % of the variance for all detectable
polar metabolites (n=116) was attributed to PC1 and 2 (Fig. 2B). This was supported by
significant positive correlation for leaf lysates and apoplast extracts of lettuce and spinach (R²
> 0.97), a weak correlation for the root lysates based on species (R² 0.542 – 0.757), with no
significant correlation between any species for the tissues.

The influence of plant extract metabolites on E. coli growth

To relate any specific plant metabolites to bacterial growth, a correlation analysis was carried
out between the plant extracts growth rates for two E. coli O157:H7 isolates (Sakai and
ZAP1589) and the assigned metabolites. Several organic acids positively associated with
maximal growth rates (µ), although there was a temperature-dependent effect. Metabolites
associate with growth at 18 °C for isolate Sakai were galactosyl glycerol, threonic acid, and
oxoproline (p ~ 0.04); at 20 °C, malic acid, fumaric acid and quinic acid (p = 0.014 – 0.048); and at 25 °C oxalic acid (p = 0.009), aspartic acid (p = 0.038), glutamic acid (p = 0.046), coumaric acid (p = 0.011) and uridine (p = 0.011). Chlorogenic acid (trans-5-O-caffeoyl-D-
quininate) was consistently associated with growth for all temperatures (p 0.04 at 18 °C, p 0.004 at 20 °C, and p 0.04 at 25 °C). E. coli isolate ZAP1589 gave similar results, although there was also a bacterial isolate effect as there were no significant associations at 20 °C. Therefore, no single metabolite was identified as the major factor influencing E. coli growth rate, with a significant impact from growth temperature.

The main metabolite groups were then investigated as groups that could influence bacterial growth, by generating defined ‘artificial’ growth media comprising the main plant extract metabolites. The six most abundant metabolites were selected from lettuce apoplast or sprout extracts to represent contrasting metabolite profiles (Table 3). Each of the major groups of saccharides (SA), organic acids (OA) or amino acids (AA) were assessed independently by dilution, to restrict their effect, and at temperatures relevant to lettuce (18 °C) and sprouts (25 °C). Maximal growth rates were similar in the sprout and lettuce extract artificial medium (Fig. 3), although reduced compared to the ‘complete’, natural extracts (Fig. 1). Growth rates were significantly reduced when the concentration of the saccharide group (SA) was reduced for both artificial media (all p < 0.0049), while restriction of the amino acids (AA) or organic acids (OA) had no impact (Fig. 3). The SA-dependent effect occurred for all E. coli isolates, although there were also significant isolate dependencies (two-way ANOVA, F (16, 28637) = 39.5; p < 0.0001 at 25 °C; two-way ANOVA, F (4, 9544) = 401.3; p < 0.0001 at 18 °C).
The influence of plant extracts on *E. coli* biofilm formation

On host tissue *in planta*, bacterial colonies are more likely to be present in biofilms rather than as single cells. Therefore, the influence of the plant extracts of the leafy vegetables was tested for *E. coli* biofilm ability in isolation, i.e. on polystyrene surfaces. Spinach leaf lysates and root lysates were the only extracts that induced biofilm for all isolates, albeit minimal for isolate MG1655 (*p* < 0.0011, compared to isolate MG1655) (Table 4). The remaining extracts were not as conducive for biofilm formation, with the exception of one of the environmental isolates (JHI5025). This was not explained by different growth rates since this isolate did not exhibit the fastest growth rates in the extracts compared to the others (Fig. 1) and presumably therefore reflect increased adherence in the presence of the plant extracts. A qualitative risk ranking was determined for implementation of biofilm formation as a risk factor for the *E. coli* O157:H7 isolates (Sakai and ZAP1589) that identified spinach roots as the highest risk (from highest to lowest): spinach roots > spinach leaves > lettuce roots > lettuce leaves > spinach apoplast > lettuce apoplast.

*E. coli* O157:H7 colonisation and internalisation *in planta*

*E. coli* O157:H7 colonisation of leafy vegetables and sprouts was quantified to determine whether growth kinetics and biofilm formation in the extracts were predictive of *in planta* colonisation. Colonisation of the *E. coli* O157:H7 isolate (ZAP1589) was quantified on spinach and lettuce, and for both isolates (ZAP1589 and Sakai) on sprouted seeds. Our previous *in planta* data for lettuce and spinach plants showed that the highest levels of *E. coli* isolate Sakai occurred on spinach roots (67). Inoculation of spinach and lettuce with the high dose (*10^7* cfu ml^-1^) of *E. coli* isolate ZAP1589 also resulted in higher levels of bacteria on the roots compared
to leaves, with similar levels on spinach and lettuce roots, e.g. 2.53 ± 0.97 and 2.69 ± 0.88 log (cfu g⁻¹) at day 14, respectively (Fig. 4A, B). *In planta* colonisation of sprouted seeds by the two *E. coli* O157:H7 reference isolates was quantified for plants grown under conditions that mimic industry settings (hydroponics at 25 °C, three days) (Fig. 4C-F). A low inoculation dose of 10³ cfu ml⁻¹ was used and total viable counts on day 0 were estimated by MPN since they fell below the direct plating detection threshold. Total counts of isolate Sakai increased by 4.5 log (cfu g⁻¹) on alfalfa sprouts and 3 log (cfu g⁻¹) on fenugreek sprouts, between 0 and 2 dpi. Viable counts for isolate ZAP1589 were generally lower on both sprouted seeds compared to isolate Sakai, but still reached 6.00 ± 0.253 log (cfu g⁻¹) on alfalfa 2 dpi.

Internalisation was also assessed since endophytic behaviour is a feature of *E. coli* O157:H7 colonisation of fresh produce plants and growth potential could be reflected by growth in the apoplast washings. Internalisation of isolate ZAP1589 occurred to higher levels in spinach roots compared to lettuce roots (Fig. 4A, B), although the prevalence was similar in both plant species (60 % and 58.3 % of plants contained endophytic bacteria). In contrast, internalisation in sprouts only occurred on three occasions in all the experiments: isolate Sakai in alfalfa (1.07 log (cfu g⁻¹)) and fenugreek (1.53 log (cfu g⁻¹)) on day 1, and isolate ZAP1589 in alfalfa (1.87 log (cfu g⁻¹)) on day 2. The prevalence was 7.1 % (1/14 samples positive), although the viable counts were close to the limit of detection by direct plating. Therefore, internalisation of *E. coli* O157:H7 isolates Sakai and ZAP1589 appeared to be a rare event on sprouted seeds, although they colonised the external sprout tissue to higher levels than on lettuce or spinach.
Correlating in planta colonisation with plant extract growth rate kinetics

To relate growth kinetics in extracts with in planta growth, growth rates were estimated for in planta growth. This was possible for sprouted seeds since colonisation levels increased over time (Fig. 4). Alfalfa plants supported significantly faster growth rates for both E. coli O157:H7 isolates compared to fenugreek, at 2.23 ± 0.213 log cfu g⁻¹ per day (R² = 0.720) and 1.50 ± 0.0913 log cfu g⁻¹ (R² = 0.863) for Sakai on alfalfa and fenugreek sprouts, respectively, and for isolate ZAP1589, rates of 2.24 ± 0.159 log cfu g⁻¹ (R² = 0.822) and 0.710 ± 0.116 log cfu g⁻¹ (R² = 0.464) per day on alfalfa and fenugreek sprouts, respectively. The difference in growth rate between the isolates on fenugreek sprouts was significant (p < 0.0001). Although in planta growth rates for E. coli isolates Sakai were estimated on spinach tissues (leaves, roots or internalised in leaf apoplast) or lettuce (leaves, roots) from low inoculation dose (10³ cfu ml⁻¹), (67) these were non-significant since growth over the 10 day period was minimal or completely constrained, with a high degree of plant-to-plant variation. Growth rate estimates were not made when a high starting inoculum was used since the colonisation levels decreased over time (Fig. 4).

Comparison of the in planta and extract growth rate estimates were made for both E. coli O157:H7 isolates on sprouted seeds (at 25 °C) or in spinach and lettuce (at 18 °C) (Fig. 5). A positive correlation occurred for growth rate estimates in the sprouted seeds (R² = 0.516), although this was not significant. Since in planta growth in spinach or lettuce tissues was minimal, there was no correlation with growth rates in corresponding extracts. Therefore, the restrictions in bacterial growth that occurred with living plants meant that growth rates in...
extracts could not be extrapolated to *in planta* growth potential for leafy vegetables, but did bear a positive relationship for sprouted seeds.
The potential for food-borne bacteria to grow in fresh produce food commodities is a key consideration in quantitative risk assessment. Factors that influence bacterial growth are the plant species and tissue, the bacterial species or isolate, and the surrounding environment. The growth potential of a bacterial population consists of proportion of the growing sub-population and the suitability of the environment for growth, and it provides a quantitative description of probability of growth (20). Therefore, the factors that influence growth potential of STEC in edible plants include plant-dependent and physio-chemico factors, as well as bacterial isolate-specific responses. Metabolically active components of plants can be extrapolated from plant extracts for bacterial growth dynamic measurements coupled with metabolite analysis. They also represent a bacterial growth substrate in their own right that could arise during the post-harvest production process e.g. from cut surfaces. A number of studies show growth of food-borne bacteria on plant extracts during the production process (31, 51, 52) and growth potential for E. coli O157:H7 has been evaluated in water (64). Here, maximum growth rates in plant extracts were strongly influenced by the plant tissue type and species, as well as the E. coli isolate tested and overlaid by temperature-dependent effects. In planta growth rates, however, was markedly different between the sprouted seeds and leafy vegetables, with a growth restriction evident in the leafy vegetables. The plant-dependent factors that could account for this difference include plant age, defence response, growth conditions and associated microbiomes. As such growth rates in the extracts could not be used to infer in planta growth potential for spinach or lettuce. In contrast, proliferation on sprouted seeds did bear a positive relationship to growth rates in extracts, although it was also dependent on the plant species and on bacterial isolate tested.
Saccharides were shown to be the major driving force for *E. coli* growth, which is unsurprising given their role in central metabolism (37). Although the levels of the most abundant sugars, glucose, fructose and sucrose (the disaccharide of glucose and fructose) could explain the high growth rates in sprout extracts, similarly rapid growth did not occur in lettuce leaf lysate extract, despite an abundance of sugars, indicating that plant species-specific inhibitory compounds exist. This is supported by the occurrence of more rapid growth rates in spinach leaf extracts compared to lettuce. Plant-dependent factors that could influence bacterial growth potential include the innate defence response (29) and antimicrobial activity of plant secondary metabolites (65). Plant development stage is an important factor since sprouted seeds, which were abundant in glucose and fructose, are at a distinct developmental stage to mature plants, and young plants of a variety of species can serve as preferential secondary plant hosts for STEC (68).

Bacterial growth rates were not significantly impacted by manipulation of the major amino or organic acids from the extracts, although the phenolic acid, chlorogenate (*trans*-5-O-caffeoyl-D-quinate) was positively associated with growth. This contrasts to reports of its ability to inhibit fatty acid synthesis in *E. coli* isolate MG1655 (33) and prevent *E. coli* growth (69), but may be explained by differences in concentration between the extracts and exogenous application. Oxalate levels were relatively high in spinach, in keeping with previous reports that show an average as high as ~1000 mg / 100 g fresh weight (44) and correlated with growth for isolate Sakai at 25 °C. Amino acids levels were substantially higher in sprouted seed extracts compared to the leafy vegetables, which is likely a reflection of different developmental stages of the plants (3). It was notable that the artificial media did not support equivalent growth rates to the ‘complete’, natural extract media, indicating that other, minor nutrients in the extracts...
were utilised for maximal bacterial growth and also need to be accounted for in growth dynamics.

Bacteria including STEC, tend to form biofilms in association with plant tissue (10, 67, 68). Here, a risk ranking could be inferred from biofilm formation in the extracts, with spinach roots ranked highest. Curli is an important biofilm component for STEC associated with plants (6), but other biofilm components are likely to be responsible for the biofilm formation in extracts, since isolate Sakai did not form biofilms in spinach apoplast extract in vitro although does produce curli during endophytic colonisation and biofilm formation in leaves (67). This indicates that specific in planta cues induce different biofilm components. Alternative biofilm components that may be involved include Type 1 fimbriae, which was shown to be expressed by the environmental isolates JHI5025 and JIH5039 at 20 °C and promoted binding to spinach roots (36).

Internalisation of STEC into apoplastic spaces in plants presents a hazard as pathogens cannot be removed by conventional sanitation methods. However, growth potential for internalised E. coli O157:H7 could not be inferred from growth in apoplast extracts since endophytic proliferation was prevented or reduced in the apoplast (67). As the apoplast is a habitat for plant-associated endophytes (60) and phytopathogens (57), it appears that for E. coli additional factors such as the plant defence response need to be considered. The increased likelihood of internalisation into tissues of leafy vegetables compared to sprouted seeds for the E. coli O157:H7 isolates could be due to multiple factors including plant age, the competing microbiota and access to nutrients. Plant dependent factors have also been shown to impact colonisation of lettuce cultivars by STEC (54).
In planta colonisation of *E. coli* O157:H7 isolate Sakai was significantly higher than isolate ZAP1589, in both leafy tissue types and on both sprouted seed species (67). In contrast, growth rates in the plant extracts and in artificial media overlapped, albeit with specific extract-specific differences. Since isolate ZAP1589 was found to be flagellate but non-motile, this may reflect a role for flagella in plant colonisation (59). ZAP1589 growth rates on sprouted seeds were similar to the rates reported for other *E. coli* O157:H7 isolates on 2-day old alfalfa sprouts (7). Growth rates of both *E. coli* O157:H7 isolates in the extracts was, in general, as high as the environmental isolates, indicating similarities in fitness levels for STEC and environmental *E. coli* in the plant environment. As anticipated, almost all growth rates were lowest for the laboratory adapted K-12 isolate, and biofilm formation was essentially absent.

The ability of *E. coli* isolates to metabolise different carbon sources varies and could contribute to the isolate-dependent variations in growth rates. Although less than 50 % of *E. coli* isolates can metabolise sucrose (37), *E. coli* O157:H7 isolate Sakai encodes the sucrose transport genes (1) and sucrose degradation genes were expressed by this isolate on exposure to spinach extracts (8). The sucrose translocator from *S. enterica* serovar Typhimurium was expressed by a related epiphyte *in planta* (43). In contrast, fructose and glucose are sufficient sole carbon source-metabolites for *E. coli* and their role in bacterial metabolism is well characterised (37). An *E. coli* fructose metabolism gene has also been expressed in a related epiphyte *in planta* (32).

Growth rates normally positively correlate with temperature (55), as was observed for growth rates in the defined medium without plant extracts, which exhibited a linear distribution from 18°C to 25°C. However, maximal growth rates in the extracts were influenced in a non-linear manner by temperature. Similarly, a non-linear effect was reported in a meta-study on growth...
of STEC on lettuce (38). Since *E. coli* Sakai exhibits distinct metabolic responses to different plant tissues (8), it is possible that a temperature-dependent effect on metabolite content similarly impacted bacterial metabolism and resultant growth. This may explain the different organic acid-growth correlations that occurred at 20 °C 'vs' 18 °C. The implications are that a linear approximation, e.g. such as a Ratkowsky model, is not sufficient to describe *E. coli* growth in plant extracts, although it has been used to model growth on plants (39, 55).

In conclusion, growth potential *in planta* was described in part, by growth rates in plant extracts, but only for sprouted seeds. On the other hand, biofilm formation in plant extracts showed some relation to *in planta* colonisation in leafy vegetables. Plant species- and tissue-type dependent differences in metabolites meant that no single metabolite could be correlated with growth, and the only positive association was with the combined group of saccharides. The marked differences in *in planta* colonisation between the sprouted seeds and leafy vegetables reinforces the higher risk associated with very young plants, grown under conditions conducive for bacterial growth (68). Therefore, although this data can inform hazard identification and risk analyses, it is evident that important specificities within each plant-microbe system need to be considered, and it is not possible to take a generalised view of STEC-plant colonisation.
Materials and Methods

Bacteria and media

The bacterial isolates panel comprised five isolates: two *E. coli* O157:H7 isolates, two environmental *E. coli* isolates and an *E. coli* K-12 isolate (Table 1). *E. coli* ZAP1589 is a Stx negative derivative, generated from isolate H110320350. Regions flanking *stx* genes were amplified using specific primers: No-*stx1* (5’-ttgctggtctcgtacccggg), Ni-*stx1* (5’-cgctcttgccgccggtggaacgg), Co-*stx1* (5’-tcccttgcaccacctgcgtacg), Ci-*stx1* (5’-ccgttccagccgccgcaagacg), No-*stx2* (5’-ttgctggtctcgtacccggg), Ni-*stx2* (5’-cgctcttgccgccggtggaacgg), Co-*stx2* (5’-tcccttgcaccacctgcgtacg), Ci-*stx2* (5’-ccgttccagccgccgcaagacg) for *stx1* and *stx2*, respectively. The PCR fragments were cloned into pTOF25 and verified by sequencing. The kanamycin resistant gene from pTOF2 (41) was cloned into the *stx1*-deletion construct and tetracycline resistance gene from pTOF1-TcR (63) was cloned into the *stx2*-deletion construct. The plasmids were transformed into isolate H110320350 for allelic exchange to delete *stx1* and *stx2* sequentially, these were confirmed absent by PCR using primers:

- *stx1* (5’-ATAATGCATCGCTTGACTAC and 5’-AGAAGACCACAGTGAGTCATC) and *stx2* (5’-GGCAGTCTGAAACTGCCTCC and 5’-TCGCCAGTTATCTGACATTCT). Motility of isolate ZAP1589 and isolate H110320350 was tested on motility agar (0.7 %), and presence of the H7 flagella was confirmed by agglutination with the monoclonal H7 antibody.
Bacteria were cultured overnight in Lysogeny-broth medium (LB) at 37 °C (2), with shaking at 200 rpm. Prior to experimentation an aliquot of the overnight culture was inoculated 1:100 in rich defined 3-(N-morpholino)propanesulfonic acid (MOPS) medium (45) with 0.2 % glycerol and essential and non-essential amino acids, termed ‘rich defined MOPS glycerol’ (RDMG), for 24 h at 18 °C and 200 rpm. Bacteria were collected by centrifugation, washed in phosphate buffered saline (PBS) and adjusted to the required starting optical density (OD) 600 nm. Media was supplemented with 30 μg ml⁻¹ kanamycin, if required. Defined artificial ‘lettuce apoplast’ or ‘sprout extract’ media was generated by adding each group of constituents (Table 3) to a base minimal MOPS medium (MMM) lacking a carbon source and amino acids. Each component group was added at the defined concentration to represent the concentrations and composition present in lettuce apoplast or sprout extracts and by dilution of one major group at a time at: 1:50 saccharides (SA), 1:10 amino acids (AA) or 1:20 organic acids (OA), while the other groups were at 1:1. The pH of the sprout defined medium was 7.2 and lettuce apoplast defined medium 7.05. Viable counts were determined from 10-fold dilutions plated on MacConkey (MAC) agar, incubated overnight at 37 °C and counted manually the next day. All experiments were conducted in triplicate. Viable counts and OD₆₀₀ nm were plotted in Excel 2010.

Plant extracts and metabolite analysis

Lettuce (Lactuca sativa) var. All Year Round and spinach (Spinacia oleracea) var. Amazon were grown individually in 9 cm³ pots in compost for microbiological assays, or in vermiculite for metabolite analysis, in a glasshouse for three weeks. Fenugreek (Trigonella foenum-graecum) and alfalfa (Medicago sativa) seeds were soaked in sterile distilled water (SDW) for 3 h at room temperature (RT), surface sterilized with 3 % calcium hypochlorite (20,000 ppm ml⁻¹ active chlorite) for 15 min, washed five times with SDW and soaked for 2 h in SDW at RT.
Sprouts were transferred aseptically on distilled water agar (DWA) (0.5 % agar) and sprouted for two (alfalfa) or five (fenugreek) days at 25 °C in darkness. Leaf apoplastic washings were collected as described previously (Methods SM3), optimised for spinach and lettuce to minimize cytoplasmic contamination (35). All tissue extracts were made as described previously (8). In brief, vermiculite was gently washed off the roots with tap water and rinsed with SDW. Leaves and roots were separated with a sterile scalpel, macerated in liquid nitrogen with a pestle in a mortar and stored at -20 °C until use and pre-processed for sample clarification by mixing 1 g with 20 ml SDW, soaked on a shaker for 4 h, centrifuged at 5000 rcf for 15 min, and the supernatant heated to 50 °C for 30 min. The extract was centrifuged at 5000 rcf for 15 min and filter sterilised through a 0.45 μm filter for root tissue or 0.1 μm filter for leaf tissue. Sprouts were macerated in liquid nitrogen, processed as described above without a washing step to remove vermiculite, and filter sterilised through a 0.22 μm filter. Apoplast extracts were filtered sterilised through a 0.1 μm filter (Durapore, Merck, Germany). Extracts were made from ~ 5 plants per sample for leaves and roots and up to 24 plants for apolastic washings or for sprouts. 10 ml plant extract samples were used for GC-MS analysis as described in Methods SM4. Lysates were prepared for HPLC described previously by (62).

Growth rate parameterisation

Representative edible species associated with food-borne outbreaks were used: two leafy greens (lettuce, spinach) and two sprouted seeds (fenugreek, alfalfa). Plant tissues used represented edible, non-edible and internalised tissues of the leafy greens from total lysates of leaves or roots, and apoplastic washing recovered from leaves, respectively, while total sprout lysates were used to represent edible sprouts. A panel of five *E. coli* was assessed (Table 1) to
compare relative fitness of two STEC O157:H7 Stx- isolates to two environmental isolates from plant roots and soil. A K-12 faecal-derived and laboratory-adapted isolate was included for reference. Growth was assessed at three temperatures (18, 20 and 25 °C) to represent relevant growth temperatures of field-grown leafy greens in northern temperate zones and sprouted seeds grown under controlled conditions. Growth kinetics were measured from optical densities derived from a plate reader (as described by others (20)).

**Bacterial growth rates**

Bacterial growth rates were determined using a pre-warmed plate reader Bioscreen C plate reader (Oy Growth Curves Ab Ltd, Finland), set to different temperatures. The *E. coli* isolates were grown as described above, adjusted to an OD$_{600}$ of 0.05 in PBS (~ 2.1 x 10$^7$ cfu ml$^{-1}$) and inoculated at a 1:10 dilution in plant extracts (at 1:20 w/v in dH$_2$O) or defined media (Table 3), in 200 μl total volume, in multi-well plates. Growth for the *E. coli* isolates was measured at 18, 20 and 25 °C in 100-microwell plates (Honeycomb, Thermo Fisher, USA). Wells were randomised in duplicate on the plate with negatives included. All growth curves in extracts were repeated three times with four replicates on plates. Measurements were recorded every 15 min for 48 hours and multi-well plates were shaken for 60 seconds pre- and post-measurement. Results were exported from plate reader proprietary software as tab-delimited files. For model fitting, 12 replicates of each isolate and medium type were averaged and converted to viable counts log (cfu h$^{-1}$) (Methods SM5). A conversion factor of 4.2 x 10$^8$ cfu ml$^{-1}$ was applied so that all growth curves could be modelled using DM-Fit (Methods SM1). Secondary modelling was applied for different temperature as described (Methods SM2). A 2-
way ANOVA was carried out for multiple comparisons (isolate / extract type) in Prism v6 (GraphPad Software Inc., USA).

Biofilms

Bacterial biofilms were measured as described previously (42). Bacteria were grown aerobically in LB at 37 °C for 12 h, sub-cultured (1:1000 v/v) in RDMG for 18 h at 18 °C, diluted in PBS to OD$_{600}$ of 0.05 and inoculated into plant extracts as per the growth rates determination in a 96 well polystyrene plate and incubated statically for 48 h at 18 °C. The washed wells were stained with 0.1 % crystal violet solution and solubilised with 95 % ethanol. The solution was transferred into a fresh plate and absorbance measured at 590 nm with a plate reader (Multiskan Go, Thermo Scientific, USA). Results were exported with the software SkanIt™ (Thermo Scientific, USA) to Microsoft Excel 2010 for analysis. A 2-way ANOVA was carried out for multiple comparisons (isolate / extract type) in Prism v6 (GraphPad Software Inc., USA).

Plant colonisation assay

Lettuce and spinach plants (~ 3 weeks old) were transferred to a growth chamber (Snijders) at 21 °C; 75 % humidity and 16 h light – 8 h dark cycle (400 μE/m2.s (30.000 lux)) three days prior to inoculation and were not watered for ~ 18 h prior to inoculation. Roots were inoculated by placing pots in a plastic box containing a 1 litre suspension of E. coli Sakai or ZAP1589, diluted to OD$_{600}$ of 0.02 (equivalent to $10^7$ cfu ml$^{-1}$) in SDW, which partially submerged pots. After 1 h inoculation, the pots were transferred to the growth chamber until sampling. Sprouts were inoculated with $10^3$ cfu ml$^{-1}$ bacteria in 0.5 l SDW for 1 h, rinsed with 0.5 x Murashige and Skoog (MS) basal medium (no sucrose), and transferred to petri dishes containing distilled
water agar (DWA) (0.8 % agar) and incubated for up to three days at 25 °C. Negative controls were incubated with SDW without bacteria.

Lettuce and spinach roots were sampled at 0, 5, 10 and 14 days post infection (dpi), aseptically removed from aerial tissue with a sterile scalpel, the compost removed by washing with SDW, and the roots were transferred into 50 ml tubes, washed with PBS and the fresh weight determined. Sprouts were sampled at 0, 1, 2 dpi, where half were used to enumerate the total viable counts of *E. coli* and stored in PBS until further use (~ 30 min), and surface-associated bacteria were removed from the other half of the samples by surface sterilization with 200 ppm Ca(ClO)₂ for lettuce/spinach roots or 20,000 ppm Ca(ClO)₂ for sprouts, for 15 min. Surface decontamination of sprout tissue required at least 15,000 ppm of Ca(ClO)₂ to eradicate external *E. coli*, but endophytes appeared to be protected from the active chlorite since endemic internalised bacteria occurred on recovery media after surface decontamination with 20,000 ppm Ca(ClO)₂. The root/sprouts were washed five times with PBS to ensure removal of all loosely adherent bacterial cells and residual chlorine. Surface sterilisation was validated as described (67). Any samples containing surface-associated bacterial colonies were removed from subsequent analysis. Roots/sprouts were macerated using mortar and pestle in 2 ml PBS and ~ 50 mg sterile sand. The supernatant was diluted once for spinach and lettuce (1:1), three times for fenugreek (1:3) or four times for alfalfa (1:4) with PBS and 100 μl plated on MAC plates using a spiral plater (WASP, Don Whitley Scientific, UK) and incubated for 24 h at 37 °C. Plates were counted using a counting grid (WASP, Don Whitley Scientific, UK), multiplied by the dilution factor and converted to cfu ml⁻¹. The experiment was repeated three times with five replicate samples per time point, and sprout samples comprised multiple (> 15) sprouts. The limit of detection from direct plating was 20 cfu ml⁻¹, below which
values were manually levelled to <1 log (cfu ml⁻¹) for lettuce and spinach root data. Since the level of inoculation of sprouts for day 0 was below the detection limit, the numbers were semi-quantified by most probable number (MPN) method for 3 tube assay as described by Oblinger and Koburger (46). Samples were diluted 6-fold in buffered peptone water (BPW) and incubated overnight at 37°C, and positive samples confirmed by plating triplicate 100 μl samples on MAC agar and incubating overnight at 37°C.
Acknowledgments

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Conflict of interest disclosure

The authors declare no conflicts of interest.
References


Tables and Figures

Tables

Table 1  Bacterial isolates used in this study

<table>
<thead>
<tr>
<th>Isolate Name</th>
<th>Serotype</th>
<th>ST</th>
<th>Stx</th>
<th>Source</th>
<th>Reference</th>
<th>Genome$</th>
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<tr>
<td>MG1655</td>
<td>OR:H48</td>
<td>98</td>
<td>n/a</td>
<td>faecal/lab</td>
<td>(21)</td>
<td>NC_000913.1</td>
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<td>JHI5025</td>
<td>nd</td>
<td>2055</td>
<td>n/a</td>
<td>soil</td>
<td>(24)</td>
<td>ERS1939526</td>
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<td>JHI5039</td>
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<td>2303</td>
<td>n/a</td>
<td>root</td>
<td>(24)</td>
<td>ERS1939531</td>
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<td>Sakai</td>
<td>O157:H7</td>
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<td>negative</td>
<td>sprout / clinical</td>
<td>(9)</td>
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<td>ZAP1589</td>
<td>O157:H7</td>
<td>11</td>
<td>negative</td>
<td>leek / clinical</td>
<td>(49)*</td>
<td>PRJNA248042</td>
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</table>

ST = sequence type, Stx = Shiga toxin presence, nd = not determined, n/a = not applicable.
Isolate Sakai used here is the stx-inactivated derivative (9). * Isolate ZAP1589, derived from H110320350 (Perry et al., 2013) has both stx-encoding regions removed, and is H7 positive but non-motile. $ GenBank, ENA or BioProject accession numbers are provided for whole genomes.

Table 2  Quantification of saccharides from plant extracts

Concentrations of mono- and disaccharides determined by HPLC (μg mg⁻¹). ND – not detected.
Table 3  Composition of defined artificial media supplements

Concentration (μg ml⁻¹) as determined by HPLC and GC-MS for the major six components in sprout extracts (alfalfa and fenugreek combined), lettuce apoplast, used to generate defined 'artificial' media.

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>Sprouts</th>
<th>Lettuce apoplast</th>
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<tbody>
<tr>
<td>Saccharides (SA)</td>
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<td>Sucrose</td>
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<table>
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<th>glucose</th>
<th>fructose</th>
<th>sucrose</th>
<th>arabinose</th>
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<tr>
<td>fenugreek</td>
<td>24.5 ± 3.1</td>
<td>24.9 ± 3.7</td>
<td>75.6 ± 6.3</td>
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<td>alfalfa</td>
<td>35.4 ± 0.8</td>
<td>35.8 ± 18.6</td>
<td>3.5 ± 0.3</td>
<td>ND</td>
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<tr>
<td>lettuce apoplast</td>
<td>19.4 ± 1.8</td>
<td>23.4 ± 2.8</td>
<td>53.4 ± 20.7</td>
<td>0.226 ± 0.001</td>
</tr>
<tr>
<td>lettuce leaf lysates</td>
<td>10.7 ± 0.3</td>
<td>14.6 ± 0.4</td>
<td>50.1 ± 3.1</td>
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<tr>
<td>lettuce root lysates</td>
<td>9.9 ± 0.1</td>
<td>20.0 ± 0.9</td>
<td>22.5 ± 0.4</td>
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<tr>
<td>spinach apoplast</td>
<td>11.8 ± 2.0</td>
<td>8.0 ± 1.7</td>
<td>38.3 ± 7.0</td>
<td>0.211 ± 0.049</td>
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<td>spinach leaf</td>
<td>21.9 ± 2.9</td>
<td>6.1 ± 0.8</td>
<td>32.8 ± 2.6</td>
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<td>spinach root</td>
<td>17.4 ± 1.2</td>
<td>9.00 ± 0.9</td>
<td>29.4 ± 1.5</td>
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### Table 4

Biofilm formation for reference *E. coli* isolates in plant tissue extracts. Biofilms were formed on polystyrene multiwall plates following incubation in spinach (Sp.) and lettuce (Lt.) extracts (apoplast; leaf; root) and rich defined MOPS medium with glycerol (RDMG) at 18°C, for 48 hrs in static conditions. The average (+ variance) density of crystal violet at OD₅₉₀ nm is presented. P value summaries are provided per isolate for each extract type vs' RDMG (ns p > 0.05; * p ≤ 0.05; ** p ≤ 0.01; *** p ≤ 0.001; **** p ≤ 0.0001).

<table>
<thead>
<tr>
<th>Treatment / Isolate</th>
<th>Sakai</th>
<th>ZAP1589</th>
<th>JHI5025</th>
<th>JHI5039</th>
<th>MG1655</th>
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<tr>
<td>Fructose</td>
<td>1443.4</td>
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<td>Glucose</td>
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<td>Asparagine</td>
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<td>Malic acid</td>
<td>n/a</td>
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<td>2,3-dihydroxy-propanoic acid</td>
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<tr>
<td>Isolate</td>
<td>Sp. apoplast</td>
<td>Sp. leaf</td>
<td>Sp. root</td>
<td>Lt. apoplast</td>
<td>Lt. leaf</td>
</tr>
<tr>
<td>--------------------</td>
<td>--------------</td>
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<tr>
<td></td>
<td>0.002 ± 0.001 (ns)</td>
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<td>0.013 ± 0.000 (ns)</td>
<td>0.001 ± 0.002 (ns)</td>
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<td></td>
<td>0.071 ± 0.000 (***)</td>
<td>0.128 ± 0.001 (****)</td>
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**Figure Legends**

**Figure 1**  Maximum growth rates (μ) of reference *E. coli* isolates in plant extracts.

Maximum growth rates (μ) were calculated using the Baranyi model for the reference *E. coli* isolates in spinach (A) or lettuce (B) apoplast (circles), leaf lysates (triangles) and root lysates (diamonds) extracts, or in alfalfa (circles) or fenugreek (triangles) sprouts lysate extracts (C) with RDMG (diamonds) as no-plant extract control, at 18, 20 or 25 °C. Each point is the average rate (n = 12), with standard errors indicated by bars. P value summaries from multiple comparison analysis by isolate ‘vs’ MG1655 or by extract type ‘vs’ RDMG are provided in Table S1b and Table S1c, respectively.

**Figure 2**  Plant extract metabolomics and grouping

The 60 assigned metabolites from all species and tissues are separated into amino acids, organic acids, mono- and polysaccharides and others (A) by their mean total response ratio (with SD indicated by bars). (B) Score plot of principal component 1 (31 % variance) and component 2 (19 %) for all 116 polar metabolites, for alfalfa (ALF) in red, fenugreek (FEN) in blue, spinach (SAP, SLL, SRL) green and lettuce (LAP, LLL, LRL) black.

**Figure 3**  Maximum growth rates (μ) in artificial media mimicking plant extracts.

Maximum growths rates (μ) calculated using the Baranyi model for the *E. coli* isolates at 18 °C and 25 °C in media mimicking (A) lettuce apoplast or (B) sprout lysates (a mixture of alfalfa and fenugreek sprout metabolites) with specified dilutions. The base minimal MOPS medium (MMM) was supplemented with saccharides (SA), organic acids (OA) or amino acids (AA) at the dilution specified. Each point is the average rate with standard errors indicated by bars.
Figure 4  Total and internalised counts for *E. coli* O157:H7 *in planta*.

The number of *E. coli* isolate ZAP1589 recovered from inoculation (10^7 cfu ml⁻¹) of (A) spinach (var. Amazon) or (B) lettuce (var. All Year Round) roots at 0, 5, 10 and 14 dpi.. The number of *E. coli* isolate ZAP1589 recovered from alfalfa (C) or fenugreek (D), and *E. coli* isolate Sakai recovered from alfalfa (E) or fenugreek sprouts (F), from inoculation at 10^3 cfu ml⁻¹, sampled at 0, 1 and 2 dpi.. Averages (lines) and individual samples counts are shown for the total (black) or internalised population (red) (n = 15: ~ 1.5 g per sample for sprouts, individual plants for spinach & lettuce). Sprout d0 data was assessed by MPN (level of detection = 0), otherwise minimum counts were manually levelled to the direct plating detection limit of 10 cfu g⁻¹ on d1.

Figure 5  Comparison of *in planta* and extract growth rates for *E. coli* isolates Sakai and ZAP1589

Growth rates for *in planta* estimates were plotted against estimates for plant extract extracts, on a Log_{10} cfu day⁻¹ basis for *E. coli* isolates Sakai and ZAP1589, normalised per g fresh weight for plant tissues or per ml for plant extracts. Estimates for sprouted seeds (alfalfa – Alf; fenugreek – Fen) were obtained for growth at 25 °C, and at 18 °C for spinach (Sp.) or lettuce (Lt.) tissues (apoplast – A; leaves – L; roots – R).

Supplemental Figure 1  Manual correction of growth rate misfits in DMFIT.

Example of a correction with *E. coli* isolate JHI5039 grown in lettuce leaf lysate, 18 °C. A) DMFIT could not fit a non-linear curve on data (n = 193) with a decrease in the stationary phase (R^2_{adj} = 0.001). B) Data was cut off manually (n = 49) to achieve better fits (R^2_{adj} = 0.996). A complete list of fits including data points are in Supplemental Table 3.
Supplemental Figure 2  Simplified polar metabolic pathways in plants

Interaction between major polar pathways (colour coded) in green leafy plants. Metabolism of carbohydrates degradation (green) is linked to amino acid degradation (dark blue and purple), which feed into the TCA cycle (red). The arrows pointing outside are entries into the non-polar fatty acid pathway. The glutamate group (orange) leads into the urea cycle. The light blue cycle described the acyl chain synthesis. Modified from the metabolomic pathway in Solanum, based on Dobson, et al. (13).