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Interaction between *Salmonella* and Plants: Potential Hosts and Vectors for Human Infection

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Abstract

Fruits and vegetables are important for a healthy diet. However, when eaten raw and contaminated with human pathogens (HPs) they may cause a disease outbreak. Contamination with HPs can occur along the entire farm-to-fork production chain and Salmonella enterica is one of the most common foodborne pathogens. A range of biotic and abiotic environmental factors can influence the complex interactions between Salmonella and plants. Moreover, the outcome of experiments largely depends on the experimental design and parameters or methods employed, and on top, on the accompanying plant microbiome and the genetic equipment of the plant and the Salmonella strain. Particularly mobile genetic elements contribute to the diversification and adaptation of Salmonella to the plant environment. So far, little is known about the key processes and factors influencing the attachment and potential internalization of Salmonella in plants and the plant specific responses. It is therefore important to better understand the ecology of Salmonella in the soil and plant environment, in order to propose practicable recommendations for prevention of foodborne diseases. This also requires improved sensitivity and specificity of detection methods. In this chapter, we present the current knowledge, research needs, and methodology regarding the complex interactions between Salmonella and plants.

Keywords: Salmonella enterica, plant, biofilm, colonization mechanisms, interaction

1. Introduction

The natural microbiome of plants includes a wide diversity of microorganisms and is a key determinant of plant health and productivity, e.g., by supporting the uptake



© 2017 The Author(s). Licensee InTech. This chapter is distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/3.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited. [cc) BY of mineral nutrients in roots or suppressing pathogen growth and inducing the hostimmune system [1–3]. Due to its relevance, the plant microbiome (totality of microorganisms associated with the plant) is even called the second plant genome. Because of the tight interplay between plants and their epiphytic and endophytic microorganisms the terms holobiont and meta-organisms are used as well. The plant microbiome is important not only for plant growth and health, but is also positively influencing human health [4]. However, besides positive effects on human health plants can also be carriers of bacterial HPs.

Salmonella is one of the major causal agents of foodborne gastroenteritis and represents a major threat to public health. It is estimated that each year 93.8 million cases of salmonellosis occur globally (86% of which foodborne), with 155,000 deaths [5]. Consumption of raw plants is more and more recognized as a source for HPs and associated with disease outbreaks in several countries. The number of outbreaks linked to fresh produce, spices, and nuts surpassed those linked to foods of animal origin [6]. Sources of HPs in the production chain and factors contributing to the contamination of fruits and vegetables include for example the application of organic fertilizers such as animal manures, contaminated irrigation water, insect and animal vectors but also the use of contaminated seeds [7]. *Enterobacteriaceae* such as *Erwinia, Serratia*, and *Pantoea* belong to bacteria typically associated with the phyllosphere [8–10]. However, it is not completely understood how *Salmonella* persists in the plant environment and which environmental factors trigger its survival. In this chapter, we discuss factors influencing the survival of *Salmonella* in the agricultural environment as well as adaptations that allow successful colonization of plants, such as attachment, biofilm formation, and internalization.

2. Contamination of fresh produce

Besides contaminated animal products, *Salmonella* outbreaks are increasingly associated with fruits and vegetables. Already on the field, plants may be contaminated via soil or irrigation, especially if watered with surface water [11–14]. *Salmonella* has been shown to persist in various ecological niches in soil as well as in irrigation water and fertilizers [15–17]. In this context, the watering system and the agricultural practices seem to play a key role in the prevention of contaminated with human pathogens. For instance, lettuce plants were more likely to be contaminated with *Escherichia coli* when watered using overhead sprinklers when compared to subsurface drip or surface furrow irrigation [18]. Besides, even noncontaminated rain-sized water droplets could transfer HPs from contaminated soil or plants to other plants [19].

Organic fertilizers like manure, biogas plant digestates and sewage sludge offer an additional route for contamination of fresh produce. Similarly, animals like birds, game, mice, or insects can contribute to the contamination of fresh produce directly or indirectly via feces or irrigation water [7, 14, 20]. Often underestimated are soil particles, which can be carried by the wind over long distances and contribute to the transient of microbiome between plants [8]. Hence, wind-caused spread of HPs should also be considered. Contaminated plant residues might constitute additional risk if incorporated into soil before the planting of next crop.

The infection of plants is essentially dependent on the ability of HPs to survive and persist in the agricultural environment. *Salmonella*, for example, was shown to survive in soil for more than 200 days if the soil was fertilized [21, 22]. The survival of diverse bacteria newly introduced into soil has been subject of research for many years [23], and the mechanisms that govern this process, compared often to microbial invasion, were described in many studies (recently reviewed by [24]). In order to survive in the soil, HPs need to find an adequate ecological niche in which they can establish. Furthermore, their ability to do so and to survive for extended time increased when the indigenous microbial community was reduced as a result of, for example, sterilization [25]. In addition, the survival of microorganisms that successfully invaded the soil is highly dependent on the environmental heterogeneity [26–28].

Contamination of fresh produce with HPs like *Salmonella*, can occur before the harvest and also along the whole production chain [11, 14]. Since the epidemiological investigations start very often long time after the contamination or the harvest, it is very challenging to assess whether the contamination took place in the field or occurred "post-harvest" during the processing. Consequently in the majority of cases, the information available does not necessarily reveal the real causes of contamination [29].

3. Epidemiology of Salmonella in agricultural systems

Fresh produce contaminated with *Salmonella* can easily trigger a salmonellosis outbreak, and despite the difficulties with identification, in the past years fresh produce were repeatedly identified as the outbreak source. Among the outbreaks in the USA, *Salmonella* is the leading cause of the fresh produce-originated foodborne diseases [30]. The available data are depending on the procedures and records in particular countries. At least 12 large, fresh produce-related *Salmonella* outbreaks have been reported since 2010, an overview of international outbreaks with more than 100 associated cases is presented in **Table 1**.

Although fruits and vegetables were identified as source of human pathogens, it is not clear whether the plants were colonized in the field or during processing. *Salmonella* may live epiphytically or be internalized through wounds, the root system, stomata, or hydathodes (see below). Additionally, *Salmonella* can be entrapped in fruits or seeds after contamination of flowers [31, 32]. Moreover, large outbreaks can be destructive to consumer's confidence which results in economic losses [33, 34]. Therefore, the research on the ecology of HPs like *Salmonella* in relation to farming and harvesting practices is very important for human health and also for the economy.

4. Factors influencing the survival of Salmonella in soil

Successful establishment of human pathogenic bacteria in soil depends on a variety of biotic and abiotic factors (see **Figure 1** for an overview). Numerous studies, carried out under different conditions, showed that among them are weather or atmospheric conditions like tem-

Salmonella Serovar	Vector	Year	Country	Cases/serovar confirmed	Reference
S. Newport	Tomatoes	2015	USA	115/81	[112]
S. Poona	Cucumbers	2015-16	USA	907/907	[113]
Unknown	Onions, tomatoes	2015	USA	200/0	[114]
S. Enteriditis	Sprouts, beans	2014	USA	115/0	[113]
S. Newport	Cucumbers	2014	USA	275/0	[113]
S. Typhimurium	Cantaloupe	2012	USA	261/261	[115]
S. Braenderup	Mangoes	2012	USA, Canada	127/0	[113]
S. Newport	Mung beans	2011	Germany, The Netherlands	106/32	[116]
Unknown	Produce-based salads, broccoli salad	2011	Japan	1500/0	[117]
S. Agona	Fruit, papaya	2011	USA	106/0	[113]
S. I4,[5],12:i:-	Vegetables, sprouts alfalfa sprouts	5,2010	USA	140/0	[118]
S. Hvittingfoss	Vegetables, leafy greens, lettuce, fruit, tomatoes, olives	2010	USA	114/108	[119]

Table 1. International salmonellosis outbreaks associated with fresh produce.

perature, UV radiation, and moisture content of the soil [7, 35]. In general, temperature has an important effect on growth and decay rates of bacteria. Most studies examined the influence of temperature on survival of enteric bacteria under isothermal conditions, showing a generally reduced survival of Salmonella in soil with increasing temperature and, accordingly a better persistence in soil at lower temperatures [36, 37]. Semenov et al. [38] analyzed how temperature fluctuations affect Salmonella enterica serovar Typhimurium (S. Typhimurium) in cow manure and demonstrated increased decay rates with increasing amplitudes of daily oscillations. Besides temperature, water availability is a key factor for *Salmonella* to survive in the environment. Humidity in soil depends on rainfall and watering as well as on evaporation. Soil moisture also depends on soil properties like clay content or pore size. In general, it seems that survival of Salmonella in soil is promoted by high humidity while water shortage has a detrimental influence on persistence, probably due to drought stress [39-41]. The soil type and its physical and chemical characteristics have a strong influence on the fate of bacterial HPs. Those characteristics include texture and particle size distribution, which affect adsorption of *Salmonella* to soil particles. The soil type determines the extent of *Salmonella* leaching, if the bacteria are applied to the soil surface via contaminated slurry or manure as shown by Bech et al. [42]. In this study, percolation of S. Typhimurium was more pronounced in loamy

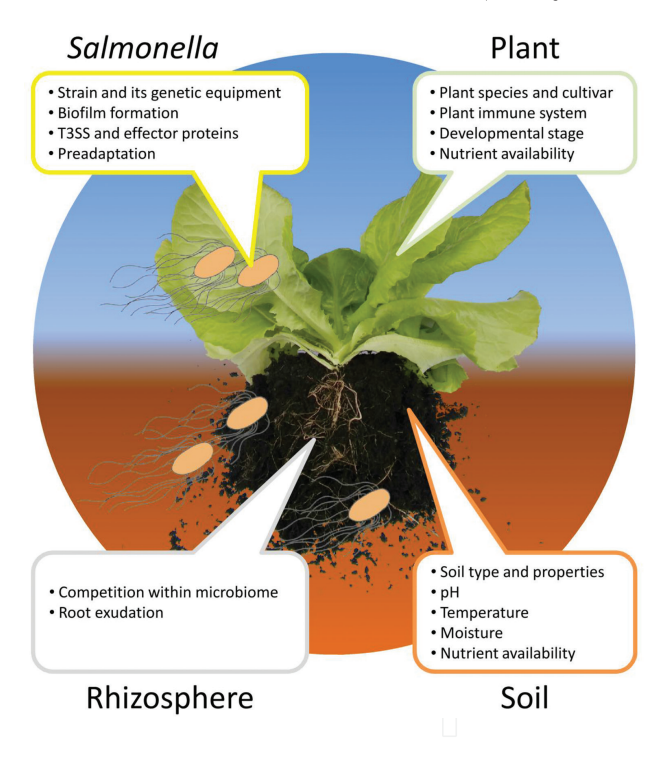


Figure 1. Factors influencing the survival of Salmonella in soil and its colonization of plants.

than in sandy soil with leaching bacteria reaching 1 m of depth at 10⁵ CFU/ml of leachate. Transport and survival of bacterial pathogens in soil is also influenced by amendment of fertilizers probably because of the presence of organic matter [43]. Leaching of *Salmonella* through soil was observed to reach greater depths after application of slurry than of manure [44]. In the same line, the application method of fertilizers can also have an effect on *Salmonella* survival in soil since an injection of manure or slurry or clumping of the applied fertilizer

aboveground protect bacteria in the soil from desiccation, UV, and high temperatures [43, 45, 46]. Agricultural practices like tillage that have an effect on the porosity of soil determine the extent of leaching [47] and the availability of oxygen. While a detrimental influence of aeration on survival of E. coli O157:H7 has been demonstrated, the oxygen availability does not influence the survival of Salmonella [48]. Soil pH is also an important factor for Salmonella survival and Salmonella can survive in the environment with neutral to acidic pH while alkaline pH has a detrimental effect on its persistence [49]. Another important determinant of Salmonella survival in soil is the availability of nutrients. In this environment, nutrients can only partly be used by bacterial HPs and are generally rather scarce. Salmonella is chemoheterotrophic and therefore depends on carbohydrates, lipids, and protein in its environment as sources for energy, nitrogen, and amino acids. Addition of organic fertilizers improves nutrient availability by addition of readily available carbon and nitrogen sources as well as other nutrients. But amendment of fertilizers also changes the microbiological properties of soil by introducing microorganisms to the soil microbial community. Moreover, the additional nutrients stimulate growth of copiotrophic soil bacteria which might compete for the nutrient resources [50]. So far, no clear correlations between the type of fertilizer and survival in soil have been identified [51]. But when survival in manure was compared to survival in manureamended soil, Salmonella usually survived better in soil [22]. This could be due to competition by the microbial flora of manure, which is more concentrated than in soil.

The soil microbial community and its composition have a great influence on the survival of *Salmonella* [52]. In the soil ecosystem, *Salmonella* has to compete with the indigenous microbial community for space and nutrients [24]. For example, it was shown that *Salmonella enterica* serovar Newport (*S.* Newport) survived about 10 weeks longer in sterilized soil compared to nonsterilized manure-amended soil [22]. Similarly, a better survival was found in γ -irradiated than in untreated soil [53]. These results indicate suppression by the native microbial community. Overall, results demonstrate the importance of the microbial community affecting the fate of *Salmonella* in soil. Plant pathogens, fungi, viruses, and animal pests present in the environment can degrade the plant material and increase the content of available nutrients or provide entry sites facilitating internalization into plants [7, 14, 54]. They may also serve as vectors [7, 55]. Effects of protists have been analyzed using protozoa showing that their presence can foster or reduce survival of different species. For example, *Salmonella enterica* serovar Thompson was accumulated in vesicles of *Tetrahymena* [56], while growth of protozoa can also decimate *S.* Typhimurium populations [37].

In addition to the environment in which *Salmonella* is introduced, the bacterial characteristics are crucial for persistence. Firstly, the genetic disposition of the strain, for example, the presence of type III secretion system (T3SS), the ability to form biofilms, chemotaxis, or motility are important. Studies using strains with mutations influencing these characteristics usually resulted in reduced survival [57–59]. *Salmonella* can also produce an *O*-antigen capsule, which improves survival under desiccation stress [60]. Furthermore, the ability to form biofilms enhances environmental persistence of some *Salmonella* serovars [61]. Similarly, a biofilm-producing *Salmonella* strain survived chlorination significantly better than the biofilm-deficient mutant [62]. The conditions under which *Salmonella* are grown before their inoculation in the environment are also important since preadaptation influences the persistence [20]. Finally,

many studies on the survival of HPs in soil employed a relatively high inoculum, which under natural conditions occurs only locally, e.g., by fecal point contaminations. Inoculation resulted in a fast initial decline of inoculated bacterial populations with usually low numbers of *Salmonella* that survive for a long time after the inoculation.

In conclusion, studies analyzing the survival of *Salmonella* demonstrated complex interactions with the environment and a network of factors, which might play an important role in the persistence of *Salmonella*. Therefore, the very often contradictory results reflect the variability of strains, their survival strategies in a complex environment as well as differences in experimental setups used.

5. Attachment to plant surfaces and biofilm formation

Attachment and adhesion of Salmonella to plant surfaces are essential steps of plant colonization. Several bacterial elements such as fimbrial structures, nonfimbrial adhesins, flagella, cellulose, and lipopolysaccharides (LPS) are important bacterial factors for colonization [63, 64]. Although previous studies demonstrated that the attachment depends on plant and bacterial factors, no single factor was found to be essential, suggesting that bacteria use several parallel mechanisms to ensure attachment to different plants or to different plant cells under a wide variety of conditions [65]. Furthermore, the attachment of S. enterica to plant surfaces appears to be serovar-dependent [66]. For example, the strength of the attachment to basil, lettuce, or spinach leaves differed between S. enterica serovars. While S. Typhimurium, Salmonella enterica serovar Enteritidis, and Salmonella enterica serovar Senftenberg were efficient, other serovars including Salmonella enterica serovar Agona, Salmonella enterica serovar Heidelberg or Salmonella enterica serovar Arizonae showed less attachment [67]. Clear differences in attachment were also observed in leaves of different age, for example, S. Typhimurium showed a better attachment to older compared to younger lettuce leaves [68]. Additionally, S. enterica serovars were reported to actively move toward plant roots, attracted by root exudates [69]. There, they are able to efficiently attach and to form biofilms at natural openings or wounds [70, 71].

Several other studies provided evidence for biofilm formation by *Salmonella* on plant surfaces [72]. Within biofilms, bacteria are generally well-protected against environmental stresses, antibiotics, and disinfectants. The importance of biofilms for the attachment of *Salmonella* to plants and their role in the persistence in plants was recently described by Yaron and Romling [65]. Biofilm formation of *Salmonella* is influenced by environmental conditions and is reported to be maximal under reduced nutrient availability, aerobic conditions, low osmolarity, and mid temperatures [73], which are characteristic for the plant surface. In contrast, it was shown *in vitro* that *S*. Typhimurium cells grown at 37°C, the temperature in the animal host, do not produce cellulose and fimbriae [64]. Furthermore, the *red dry and rough (rdar)* and the *smooth and white (saw)* morphotypes, regulated by the *agfD* promoter and defined by a combination of traits such as the presence of thin aggregative fimbriae (tafi), cellulose, and *O*-antigen capsule, might affect the dispersal of *Salmonella* in an agricultural environment [74]. In contrast to the *saw* morphotype, the *rdar* morphotype, isolated from tomato, showed

better attachment to plant surfaces [74, 75]. Biofilm-producing *Salmonella* on parsley showed a higher resistance against disinfectants than the biofilm-deficient mutant. Furthermore, after a storage period of the plant, the cells that were able to produce the biofilm matrix were significantly more resistant to the disinfection treatment [62]. A screening of 6000 transposon mutants of *S*. Newport resulted in the identification of 20 mutants selected for reduced adherence to alfalfa sprouts [70]. Interestingly, these mutants contained insertions associated with genes, for example, for the surface-exposed aggregative fimbriae nucleator (*agfB*) and the general transcriptional regulator *rpoS*. The respective proteins have been reported to regulate the production of curly, cellulose, and other adhesins such as pili. Two other genes (STM0278 and STM0650) were identified as important factors for the colonization of alfalfa seedlings. Both play an important role in the formation of biofilms [76]. Furthermore, bacterial cellulose and curly were involved in the colonization of parsley with *S*. Typhimurium from irrigation water [77].

Although many factors influencing the colonization of plants were identified by *in vitro* experiments, a more detailed investigation of genes of *Salmonella* that are expressed during the colonization of plants is needed. New techniques for the isolation of mRNA from samples containing both plant and bacterial materials as well as for the quantitative PCR allow the analysis of the transcriptome and the identification of genes with related functions [78].

6. Internalization of Salmonella into plant tissues

An increasing number of salmonellosis outbreaks associated with plants shows that human pathogenic bacteria use plants as a niche for replication or as hosts and vectors for animal and human infection (Table 1). For a long time it was assumed that Salmonella rather survives on plant surfaces than colonizes the plant interior. This view has been challenged by recent reports. Today we know that Salmonella can actively enter and spread within the plant. Plants offer multiple entry possibilities for HPs; stomata, for example, were identified already a few years ago. Stomata are used for gas exchange between the surroundings and cells of the inner mesophyll layers, this is necessary for proper photosynthetic efficacy. They can close if a pathogen is recognized. Some pathogens, however, produce toxins (coronatine), which reopen stomata and therefore allow their use as gates for colonization of underlying tissues. Salmonella was shown to gather around the open stomata and enter the mesophyll tissue of lettuce leaves [58]. Similar to lettuce, a high incidence of internalization was observed in arugula leaves, while romaine and red-lettuce, as well as basil showed significantly lower internalization rates [79]. Interestingly, in this study parsley and tomato leaves showed only marginal internalization [79]. In addition to stomata, also hydratodes and trichomes allow an internalization of Salmonella into leaves [31, 80, 81]. Not only Salmonella or phytopathogenic bacteria use stomata as entry points, also other Enterobacteriaceae, for example E. coli, use similar strategies to access the plant's interior [82, 83]. Importantly to note is the fact that the preference to gather around open stomata manifests only in photosynthetically active leaves, and an artificial opening of the stomata at night has no effect on the bacterial behavior [58]. This observation is in line with the proposed hypothesis that those bacteria are in a direct competition for C- and N-sources with the native leaf microbiome [69, 71], and suggests a chemotaxis toward the newly synthesized products of the photosynthesis.

From the consumers' point of view, not only the internalization into leaves but also the translocation within the plant, e.g., toward fruits is important. In some crop plants, e.g., tomato, such translocation was detected [81]. The authors showed internalization into the tomato fruits when the entire plant was systemically colonized. Still, the colonization rates seemed rather low [81]. Nonetheless, in light of the persistent pathogenicity in animals after the passage through a plant host [84], the internalization mechanisms are of high interest. Some detailed mechanisms were already suggested. Erlacher and coworkers proposed one of those possible mechanisms: colonization of the niche below the cuticle layer of the epidermis [9]. Obviously such a behavior protects bacteria from the harsh conditions on the leaf surface (UV light, drought, and quick changes in temperature) but also from surface sterilization agents. Another strategy would be an intracellular lifestyle, which would resemble the strategy in the animal infection model. Until now, this possibility remains unverified, two reports postulated internalization into plant cells using Arabidopsis and tobacco systems [84, 85]. Yet, another helpful strategy is the efficient formation of biofilms, this strategy was discussed above and was reviewed by Yaron and Romling [65]. Only recently, it was discovered that particular Salmonella strains may avoid the recognition by the plant immune system [86], which would make them very well adapted colonizers (see below).

Many row eaten crop plants plants associated with salmonellosis outbreaks or food poisoning are usually grown in soil (lettuce, basil, parsley, etc.). In such cases the translocation from the potentially contaminated soil (through manure or irrigation water) via roots into the harvested and consumed plant parts is of enormous importance. Several reports assessed already this possibility and pointed at a very diverse picture with regard to pathogenic *E. coli* or *Salmonella*. Here the high heterogeneity with regard to colonization in the plant population is very remarkable [69, 87], usually about 20% of the plant population is colonized, however, this range may vary from 0 to 100% and strongly depends on plant species and bacterial strain [51, 69, 77, 88–90].

7. The function of T3SS and the role of plant immune system during the interactions between plant and *Salmonella*

Bacterial pathogens use T3SS and T4SS to inject so-called effector proteins directly into the cytoplasm of host cells. Those effectors are able to manipulate the host immune system and suppress the otherwise negative effects of defense responses. *Salmonella* uses two T3SS and more than 40 effectors in order to manipulate the immune system (perception mechanisms and signaling cascades) as well as the cytoskeleton of animal cells at different stages of the infection process [91]. Recent discoveries from others and our group imply that the

mechanisms used in animal and plant hosts may resemble each other [59, 85, 86, 92-95]. The inoculation with the wild-type Salmonella strains and mutants in one or both of the T3SSs showed that functional secretion systems are required for efficient plant colonization [59, 85, 93]. Two observations allow such a conclusion: (1) The mutants had lower proliferation rates when compared to the respective wild type, which suggests that a functional T3SS helps with the colonization of plants; and (2) T3SS mutants induced stronger immune response of the host plant. Similar to animals or humans, plants respond to colonization of pathogenic bacteria inducing numerous immune responses, among others are oxidative burst and enhanced expression of Pathogenesis Related (PR) genes. Both were observed after inoculation with Salmonella and both were stronger if the inoculation was performed using mutants in T3SS [59, 85, 86]. Those results suggest that the wild-type strain is able to suppress the immune response. It is very plausible to think that this suppression is due to functional T3SS-dependent effector proteins. We know only little about their function in plant cells, since only two effectors (SseF and SpvC) were evaluated in this respect. SseF together with SseG are translocated into animal cells and are responsible for the establishment of the reproduction niche [96]. In plants, SseF induces the hypersensitive response (HR) [94]. Important is the fact that silencing of the suppressor of SGT1 eliminates the response to SseF, suggesting that this effector is recognized in R protein-dependent manner, which is the usual recognition method of pathogen effectors during the effector-triggered immunity (ETI). SpvC is a phosphothreonin lyase which dephosphorylates activated MAP kinases. Those kinases build a core compound in the signaling cascade leading from the perception of the pathogen on the cell surface to the transcriptional response at the chromatin level. Especially the trio MPK3, MPK4, and MPK6 plays an important role in plants [97], and is activated (phosphorylated) during the response to Salmonella [84]. SpvC interacts actively with the MPK6 and dephosphorylates this kinase, consequently abolishing the signal transduction [95]. A comprehensive overview of the reports regarding the plant immune responses to HPs was published only recently and is an excellent compendium of the current knowledge [98].

8. Salmonella changes its physiology in contact with plant host

During the interaction between *Salmonella* and crop plants, not only the plant reacts to the presence of the bacteria, also *Salmonella* adapts to the conditions represented by a plant organism. Recent results show that bacteria modify their physiology and motility in order to adjust to the physiological conditions occurring in plants. Several authors evaluated the transcriptional changes of bacteria when in contact with plants or plant-originated products [99, 100]. Interestingly, the analysis of the transcriptome, revealed a partial overlap between bacteria from macerated lettuce or cilantro leaves and bacteria from intestine, suggesting that those bacteria might be better adapted to the exploitation of plant material than estimated [100]. Similar results were observed for the pathogenic *E. coli* O157:H7, which seem to change its enzymatic and metabolomic equipment in order to utilize plant compounds [101, 102]. In addition, the bacteria upregulate a plethora of genes related to attachment, antimicrobial

resistance and response to oxidative stress [101]. Very striking was the fact that although plant filtrates or root exudates contain numerous amino acids, which are available to the bacteria as C and N sources, *E. coli* induced many amino acid synthesis pathways probably to supplement the missing compounds [102].

9. Detection, characterization and quantification of *Salmonella* in environmental samples

Salmonella is rarely detected in crop plants. For example, in a previous study *Salmonella* could not be detected in more than 170 plants but in their environment [103]. This suggests that environmental factors are affecting the prevalence of *Salmonella* in the field, or that the sensitivity of the currently used detection system is not sufficient.

Traditional methods for the detection and identification of HPs often rely on cultivationdependent techniques followed by biochemical and serological identification, which is typically time-consuming and laborious [104]. Furthermore, in response to environmental stresses Salmonella can enter a physiological state where the cells remain viable, but are no longer culturable on typically used growth media. Salmonella in this VBNC state are often only detectable by methods depending on nucleic acids. This highlights the importance of the complementary use of cultivation-dependent and -independent detection methods for the diagnosis and prevention of food contamination and foodborne diseases. In the recent decades, there have been increasing efforts to develop and improve molecular methods for the rapid detection and characterization of pathogens in animals and animal products [105-108]. These methods, which include immunological as well as biosensorand nucleic acid-based assays (e.g., ELISA, PCR, microarrays, next generation sequencing) have an improved sensitivity and specificity but are also time-, cost-, and labor-demanding. Typically, to further increase the sensitivity of these methods nonselective or selective enrichment steps are employed. One of the most challenging problems is the sample preparation, which is strongly depending on the sample matrix, associated inhibitory compounds, and the bacterial load.

So far, knowledge is scarce regarding the specific and reliable detection of *Salmonella* in complex and often heterogenous plant- and environmental-matrices (e.g., vegetables, spices, soil samples, manure, biogas digestates) as well as the appropriate extraction and purification techniques. For iceberg lettuce, carrot- and cucumber-peelings, qPCR detection limits of 10³ bacterial cells per gram were reported [109]. Since the infectious dose of *Salmonella* was reported to be less than 10³ cells [110], small numbers have to be detected reliably. Besides direct extraction of total DNA from the sample material, a preceding enrichment step in the respective culture media can be performed. This enrichment has the advantage to increase the sensitivity of detection and additionally to reactivate cells in the dormant VBNC state. After extraction of DNA from the respective samples, *Salmonella* can be detected by qPCR or PCR-Southern blot hybridization, e.g., by detection of the *invA* gene [104, 111]. Alternatively or additionally to DNA-based methods, RNA-based methods

can be used. Apart from a more laborious sample preparation, RNA-based methods have the advantage that in contrast to DNA-based methods only living and active *Salmonella* are detected, i.e., *Salmonella* relevant for a potential infection of humans. Especially the detection of mRNA of pathogenicity determinants could be appropriate to prove the viability and potential virulence of HPs.

Microarrays and next-generation sequencing technologies offer intriguing possibilities regarding the rapid and accurate detection as well as genetic characterization of *Salmonella* in environmental matrices. However, the costs and technical requirements for the analysis of large data sets still limit their practicability in the day-to-day qualitative and quantitative detection. The further development of rapid, reliable, and cost-effective high-throughput detection methods will very likely contribute to the understanding of the ecology of *Salmonella* in the plant environment and consequently help to reduce or prevent infections mediated by plant-associated HPs.

10. Conclusions

Today the notion that human pathogenic bacteria such as Salmonella might persist on or within plants in low numbers is widely accepted. The research on the interactions between crop or model plants and *Salmonella* is obviously driven by its medical aspects and the need for better prevention methods. We already know various features of the interactions but many are still not fully understood. New techniques that use high-throughput analyses and unbiased approaches are useful. Numerous national survey agencies started to use next-generation sequencing for the epidemiological analysis of salmonellosis outbreaks and have therefore direct access to the genome sequences of particular serovars. They are also able to monitor the genomic changes, for example, reception of new plasmids or pathogenicity islands, which are important prerequisites in virulence of the bacterial strain. Similarly, the full range of "omic" approaches is being used in model systems providing very detailed data on both partners in the Salmonella-plant interaction at biochemical, physiological, and transcriptional levels. The study of those interactions harbors even more potential, it permits the characterization of the different infection mechanisms and the different strategies available for *Salmonella* in contact with diverse hosts. New and more efficient prevention strategies greatly depend on our understanding of these mechanisms. Therefore, the new findings might significantly improve our possibilities to diminish the number of future outbreaks.

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References

- [1] Berendsen RL, Pieterse CM & Bakker PA. The rhizosphere microbiome and plant health. *Trends Plant Sci* 2012;**17**:478–486.
- [2] Buee M, De Boer W, Martin F et al. The rhizosphere zoo: An overview of plant-associated communities of microorganisms, including phages, bacteria, archaea, and fungi, and of some of their structuring factors. *Plant Soil* 2009;**321**:189–212.
- [3] Turner TR, James EK & Poole PS. The plant microbiome. *Genome Biol* 2013;14:10.
- [4] Berg G, Grube M, Schloter M et al. The plant microbiome and its importance for plant and human health. *Front Microbiol* 2014;**5:491**.
- [5] Majowicz SE, Musto J, Scallan E et al. The global burden of nontyphoidal *Salmonella* gastroenteritis. *Clin Infect Dis* 2010;**50**:882–889.
- [6] Brandl MT, Cox CE & Teplitski M. Salmonella interactions with plants and their associated microbiota. *Phytopathology* 2013;**103**:316–325.
- [7] Brandl MT. Fitness of human enteric pathogens on plants and implications for food safety. *Annu Rev Phytopathol* 2006;44:367–392.
- [8] Rastogi G, Sbodio A, Tech JJ et al. Leaf microbiota in an agroecosystem: spatiotemporal variation in bacterial community composition on field-grown lettuce. *Isme Journal* 2012;6:1812–1822.
- [9] Erlacher A, Cardinale M, Grube M et al. Biotic stress shifted structure and abundance of *Enterobacteriaceae* in the lettuce microbiome. *PLoS ONE* 2015;**10**:e0118068.
- [10] van Overbeek LS, van Doorn J, Wichers JH et al. The arable ecosystem as battleground for emergence of new human pathogens. *Front Microbiol* 2014;**5**:17.
- [11] Olaimat AN & Holley RA. Factors influencing the microbial safety of fresh produce: A review. *Food Microbiol* 2012;**32**:1–19.

- [12] Allende A & Monaghan J. Irrigation water quality for leafy crops: A perspective of risks and potential solutions. *Int J Environ Res Public Health* 2015;**12**:7457–7477.
- [13] Li B, Jackson SA, Gangiredla J et al. Genomic evidence reveals numerous Salmonella enterica serovar Newport reintroduction events in suwannee watershed irrigation ponds. *Appl Environ Microbiol* 2015;81:8243–8253.
- [14] Jacobsen CS & Bech TB. Soil survival of *Salmonella* and transfer to freshwater and fresh produce. *Food Res Int* 2012;**45**:557–566.
- [15] Barak JD & Liang AS. Role of soil, crop debris, and a plant pathogen in Salmonella enterica contamination of tomato plants. *PLoS One* 2008;3:e1657.
- [16] Duffy EA, Lucia LM, Kells JM et al. Concentrations of *Escherichia coli* and genetic diversity and antibiotic resistance profiling of salmonella isolated from irrigation water, packing shed equipment, and fresh produce in texas. *J Food Protect* 2005;1:70–79.
- [17] Miles JM, Sumner SS, Boyer RR et al. Internalization of *Salmonella enterica* serovar montevideo into greenhouse tomato plants through contaminated irrigation water or seed stock. *J Food Protect* 2009;4:696–914.
- [18] Fonseca JM, Fallon SD, Sanchez CA et al. *Escherichia coli* survival in lettuce fields following its introduction through different irrigation systems. *J Appl Microbiol* 2011;**110**:893–902.
- [19] Monaghan JM & Hutchison ML. Distribution and decline of human pathogenic bacteria in soil after application in irrigation water and the potential for soil-splash-mediated dispersal onto fresh produce. J Appl Microbiol 2012;112:1007–1019.
- [20] Semenov AM, Kuprianov AA & van Bruggen AH. Transfer of enteric pathogens to successive habitats as part of microbial cycles. *Microb Ecol* 2010;60:239–249.
- [21] Islam M, Morgan J, Doyle MP et al. Persistence of salmonella enterica serovar Typhimurium on lettuce and parsley and in soils on which they were grown in fields treated with contaminated manure composts or irrigation water. *Foodborne Pathog Dis* 2004;**1**:27–35.
- [22] You Y, Rankin SC, Aceto HW et al. Survival of Salmonella enterica serovar Newport in manure and manure-amended soils. Appl Environ Microbiol 2006;72:5777–5783.
- [23] Acea MJ, Moore CR & Alexander M. Survival and growth of bacteria introduced into soil. Soil Biol Biochem 1988;20:509–515.
- [24] Mallon CA, Elsas JD & Salles JF. Microbial invasions: the process, patterns, and mechanisms. *Trends Microbiol* 2015;23:719–729.
- [25] Mallon CA, Poly F, Le Roux X et al. Resource pulses can alleviate the biodiversity–invasion relationship in soil microbial communities. *Ecol Soc Am* 2015;96:915–926.
- [26] Fierer N & Jackson RB. The diversity and biogeography of soil bacterial communities. Proc Natl Acad Sci USA 2006;103:626–631.

- [27] Martiny JB, Bohannan BJM, Brown JH et al. Microbial biogeography: Putting microorganisms on the map. *Nat Rev Microbiol* 2006;**4**:102–112.
- [28] Horner-Devine MC, Lage M, Hughes JB et al. A taxa–area relationship for bacteria. *Nature* 2004;**432**:750–753.
- [29] Suslow TV, Oria MP, Beuchat LR et al. Production practices as risk factors in microbial food safety of fresh and fresh-cut produce. *Compr Rev Food Sci Food Saf* 2003;**2**:38–77.
- [30] Hanning IB, Nutt JD & Ricke SC. Salmonellosis outbreaks in the united states due to fresh produce: Sources and potential intervention measures. *Foodborne Pathog Dis* 2009;6:635–648.
- [31] Gu G, Cevallos-Cevallos JM & van Bruggen AH. Ingress of *Salmonella enterica* Typhimurium into tomato leaves through hydathodes. *PLoS One* 2013;8:e53470.
- [32] Guo X, Chen J, Brackett RE et al. Survival of salmonellae on and in tomato plants from the time of inoculation at flowering and early stages of fruit development through fruit ripening. *Appl Environ Microbiol* 2001;67:4760–4764.
- [33] WBG. The economic impact of ebola on sub-Saharan Africa: Updated estimates for 2015. 2015;1–17.
- [34] Butler D. The next time the world is ill-prepared for the next epidemic or pandemic. But the horror of the ebola outbreak in West Africa may drive change. *Nature* 2015;**524**:22–25.
- [35] Santamaría J & Toranzos GA. Enteric pathogens and soil: a short review. *Int Microbiol* 2003;6:5–9.
- [36] Arrus KM, Holley RA, Ominski KH et al. Influence of temperature on *Salmonella* survival in hog manure slurry and seasonal temperature profiles in farm manure storage reservoirs. *Livest Sci* 2006;102:226–236.
- [37] García R, Baelum J, Fredslund L et al. Influence of temperature and predation on survival of *Salmonella enterica* serovar Typhimurium and expression of *invA* in soil and manure-amended soil. *Appl Environ Microbiol* 2010;**76**:5025–5031.
- [38] Semenov AV, van Bruggen AH, van Overbeek L et al. Influence of temperature fluctuations on *Escherichia coli* O157:H7 and *Salmonella enterica* serovar Typhimurium in cow manure. *FEMS Microbiol Ecol* 2007;**60**:419–428.
- [39] Bernstein N, Sela S & Neder-Lavon S. Effect of irrigation regimes on persistence of *Salmonella enterica* serovar Newport in small experimental pots designed for plant cultivation. *Irrigation Sci* 2007;26:1–8.
- [40] Or D, Smets BF, Wraith JM et al. Physical constraints affecting bacterial habitats and activity in unsaturated porous media—a review. *Adv Water Resour* 2007;**30**:1505–1527.
- [41] Holley RA, Arrus KM, Ominski KH et al. *Salmonella* survival in manure-treated soils during simulated seasonal temperature exposure. *J Environ Qual* 2006;**35**:1170–1180.

- [42] Bech TB, Johnsen K, Dalsgaard A et al. Transport and distribution of *Salmonella enterica* serovar Typhimurium in loamy and sandy soil monoliths with applied liquid manure. *Appl Environ Microbiol* 2010;**76**:710–714.
- [43] Horswell J, Hewitt J, Prosser J et al. Mobility and survival of *Salmonella Typhimurium* and human adenovirus from spiked sewage sludge applied to soil columns. *J Appl Microbiol* 2010;**108**:104–114.
- [44] Semenov AV, van Overbeek L & van Bruggen AH. Percolation and survival of *Escherichia coli* O157:H7 and *Salmonella enterica* serovar Typhimurium in soil amended with contaminated dairy manure or slurry. *Appl Environ Microbiol* 2009;75:3206–3215.
- [45] Hutchison ML, Walters LD, Moore T et al. Fate of pathogens present in livestock wastes spread onto fescue plots. *Appl Environ Microbiol* 2005;**71**:691–696.
- [46] Nicholson FA, Groves SJ & Chambers BJ. Pathogen survival during livestock manure storage and following land application. *Bioresour Technol* 2005;96:135–143.
- [47] Hruby CE, Soupir ML, Moorman TB et al. Effects of tillage and poultry manure application rates on *Salmonella* and fecal indicator bacteria concentrations in tiles draining Des Moines Lobe soils. *J Environ Manage* 2016;**171**:60–69.
- [48] Semenov AV, van Overbeek L, Termorshuizen AJ et al. Influence of aerobic and anaerobic conditions on survival of *Escherichia coli* O157:H7 and *Salmonella enterica* serovar Typhimurium in Luria-Bertani broth, farm-yard manure and slurry. *J Environ Manage* 2011;92:780–787.
- [49] Bennett DD, Higgins SE, Moore RW et al. Effects of lime on *Salmonella enteritidis* survival in vitro. *J Appl Poult Res* 2003;**12**:65–68.
- [50] Ding GC, Radl V, Schloter-Hai B et al. Dynamics of soil bacterial communities in response to repeated application of manure containing sulfadiazine. *Plos One* 2014;9(3):e92958.
- [51] Franz E, van Diepeningen AD, de Vos OJ et al. Effects of cattle feeding regimen and soil management type on the fate of *Escherichia coli* O157:H7 and *Salmonella enterica* serovar Typhimurium in manure, manure-amended soil, and lettuce. *Appl Environ Microbiol* 2005;71:6165–6174.
- [52] Moynihan EL, Richards KG, Brennan FP et al. Enteropathogen survival in soil from different land-uses is predominantly regulated by microbial community composition. *Appl Soil Ecol* 2015;89:76–84.
- [53] Goberna M, Podmirseg SM, Waldhuber S et al. Pathogenic bacteria and mineral N in soils following the land spreading of biogas digestates and fresh manure. *Appl Soil Ecol* 2011;49:18–25.
- [54] Ge C, Lee C, Nangle E et al. Impact o f phytopathogen infection and extreme weather stress on internalization of *Salmonella* Typhimurium in lettuce. *Int J Food Microbiol* 2014;**168–169**:24–31.

- [55] Caldwell KN, Anderson GL, Williams PL et al. Attraction of a free-living nematode, *Caenorhabditis elegans*, to foodborne pathogenic bacteria and its potential as a vector of *Salmonella* Poona for preharvest contamination of cantaloupe. *J Food Protect* 2003;66:1964–1971.
- [56] Brandl MT, Rosenthal BM, Haxo AF et al. Enhanced survival of *Salmonella enterica* in vesicles released by a soilborne *Tetrahymena* species. *Appl Environ Microbiol* 2005;**71**:1562–1569.
- [57] Cooley MB, Miller WG & Mandrell RE. Colonization of *Arabidopsis thaliana* with *Salmonella enterica* and enterohemorrhagic *Escherichia coli* O157:H7 and competition by *Enterobacter asburiae*. *Appl Environ Microbiol* 2003;**69**:4915–4926.
- [58] Kroupitski Y, Golberg D, Belausov E et al. Internalization of *Salmonella enterica* in leaves is induced by light and involves chemotaxis and penetration through open stomata. *Appl Environ Microbiol* 2009;**75**:6076–6086.
- [59] Schikora M, Neupane B, Madhogaria S et al. An image classification approach to analyze the suppression of plant immunity by the human pathogen *Salmonella Typhimurium*. *BMC Bioinformatics* 2012;**13**.
- [60] Gibson DL, White AP, Snyder SD et al. *Salmonella* produces an O-antigen capsule regulated by AgfD and important for environmental persistence. *J Bacteriol* 2006;**188**:7722–7730.
- [61] Vestby LK, Moretro T, Langsrud S et al. Biofilm forming abilities of *Salmonella* are correlated with persistence in fish meal and feed factories. *BMC Vet Res* 2009;5.
- [62] Lapidot A, Romling U & Yaron S. Biofilm formation and the survival of *Salmonella* Typhimurium on parsley. *Int J Food Microbiol* 2006;**109**:229–233.
- [63] Wiedemann A, Virlogeux-Payant I, Chausse AM et al. Interactions of *Salmonella* with animals and plants. *Front Microbiol* 2014;5:791.
- [64] Tan MSF, White AP, Rahman S et al. Role of fimbriae, flagella and cellulose on the attachment of *Salmonella Typhimurium* ATCC 14028 to plant cell wall models. *PLos One* 2016;**11**:13.
- [65] Yaron S & Romling U. Biofilm formation by enteric pathogens and its role in plant colonization and persistence. *Microb Biotechnol* 2014;7:496–516.
- [66] Ongeng D, Geeraerd AH, Springael D et al. Fate of *Escherichia coli* O157:H7 and *Salmonella enterica* in the manure-amended soil-plant ecosystem of fresh vegetable crops: a review. *Crit Rev Microbiol* 2015;41:273–294.
- [67] Berger CN, Shaw RK, Brown DJ et al. Interaction of *Salmonella enterica* with basil and other salad leaves. *ISME J* 2009;**3**:261–265.
- [68] Kroupitski Y, Pinto R, Belausov E et al. Distribution of *Salmonella* Typhimurium in romaine lettuce leaves. *Food Microbiol* 2011;**28**:990–997.
- [69] Klerks MM, Franz E, van Gent-Pelzer M et al. Differential interaction of *Salmonella enterica* serovars with lettuce cultivars and plant-microbe factors influencing the colonization efficiency. *ISME J* 2007;1:620–631.

- [70] Barak JD, Gorski L, Naraghi-Arani P et al. *Salmonella enterica* virulence genes are required for bacterial attachment to plant tissue. *Appl Environ Microbiol* 2005;**71**:5685–5691.
- [71] Klerks MM, van Gent-Pelzer M, Franz E et al. Physiological and molecular responses of Lactuca sativa to colonization by Salmonella enterica serovar Dublin. Appl Environ Microbiol 2007;73:4905–4914.
- [72] Steenackers H, Hermans K, Vanderleyden J et al. *Salmonella* biofilms: An overview on occurrence, structure, regulation and eradication. *Food Res Int* 2012;**45**:502–531.
- [73] Gerstel U & Romling U. The *csgD* promoter, a control unit for biofilm formation in *Salmonella* Typhimurium. *Res Microbiol* 2003;**154**:659–667.
- [74] Cevallos-Cevallos JM, Gu G, Danyluk MD et al. Adhesion and splash dispersal of *Salmonella enterica* Typhimurium on tomato leaflets: effects of rdar morphotype and trichome density. *Int J Food Microbiol* 2012;160:58–64.
- [75] Cevallos-Cevallos JM, Gu G, Danyluk MD et al. *Salmonella* can reach tomato fruits on plants exposed to aerosols formed by rain. *Int J Food Microbiol* 2012;**158**:140–146.
- [76] Barak JD, Gorski L, Liang AS et al. Previously uncharacterized Salmonella enterica genes required for swarming play a role in seedling colonization. *Microbiology* 2009;155:3701–3709.
- [77] Lapidot A & Yaron S. Transfer of *Salmonella enterica* serovar Typhimurium from contaminated irrigation water to parsley is dependent on curli and cellulose, the biofilm matrix components. *J Food Prot* 2009;72:618–623.
- [78] Holmes A, Birse L, Jackson RW et al. An optimized method for the extraction of bacterial mRNA from plant roots infected with *Escherichia coli* O157:H7. *Front Microbiol* 2014;5:286.
- [79] Golberg D, Kroupitski Y, Belausov E et al. Salmonella Typhimurium internalization is variable in leafy vegetables and fresh herbs. Int J Food Microbiol 2011;145:250–257.
- [80] Barak JD, Kramer LC & Hao LY. Colonization of tomato plants by Salmonella enterica is cultivar dependent, and type 1 trichomes are preferred colonization sites. Appl Environ Microbiol 2011;77:498–504.
- [81] Gu G, Hu J, Cevallos-Cevallos JM et al. Internal colonization of *Salmonella enterica* serovar Typhimurium in tomato plants. *PLoS One* 2011;6:e27340.
- [82] Berg G, Erlacher A, Smalla K et al. Vegetable microbiomes: Is there a connection among opportunistic infections, human health and our 'gut feeling'? *Microb Biotechnol* 2014;7:487–495.
- [83] Erlacher A, Cardinale M, Grosch R et al. The impact of the pathogen *Rhizoctonia solani* and its beneficial counterpart *Bacillus amyloliquefaciens* on the indigenous lettuce microbiome. *Front Microbiol* 2014;5:175.
- [84] Schikora A, Carreri A, Charpentier E et al. The dark side of the salad: *Salmonella Typhimurium* overcomes the innate immune response of Arabidopsis thaliana and shows an endopathogenic lifestyle. *PLoS One* 2008;**3**:e2279.

- [85] Shirron N & Yaron S. Active suppression of early immune response in tobacco by the human pathogen *Salmonella Typhimurium*. *PLoS One* 2011;6:e18855.
- [86] Garcia AV, Charrier A, Schikora A et al. *Salmonella enterica* flagellin is recognized via FLS2 and activates PAMP-triggered immunity in *Arabidopsis thaliana*. *Molecular Plant* 2014;7:657–674.
- [87] Gorbatsevich E, Sela Saldinger S, Pinto R et al. Root internalization, transport and inplanta survival of *Salmonella enterica* serovar Newport in sweet basil. *Environ Microbiol Rep* 2013;5:151–159.
- [88] Erickson MC, Webb CC, Diaz-Perez JC et al. Infrequent internalization of *Escherichia coli* O157:H7 into field-grown leafy greens. J Food Prot 2010;73:500–506.
- [89] Bernstein N, Sela S & Neder-Lavon S. Assessment of contamination potential of lettuce by *Salmonella enterica* serovar Newport added to the plant growing medium. *J Food Prot* 2007;**70**:1717–1722.
- [90] Franz E, Visser AA, Van Diepeningen AD et al. Quantification of contamination of lettuce by GFP-expressing *Escherichia coli* O157:H7 and *Salmonella enterica* serovar Typhimurium. *Food Microbiol* 2007;24:106–112.
- [91] Niemann GS, Brown RN, Gustin JK et al. Discovery of novel secreted virulence factors from *Salmonella enterica* serovar Typhimurium by proteomic analysis of culture supernatants. *Infect Immun* 2011;**79**:33–43.
- [92] Hernandez-Reyes C & Schikora A. *Salmonella*, a cross-kingdom pathogen infecting humans and plants. *FEMS Microbiol Lett* 2013; 343(1):1–7.
- [93] Schikora A, Virlogeux-Payant I, Bueso E et al. Conservation of *Salmonella* infection mechanisms in plants and animals. *PLoS One* 2011;6:e24112.
- [94] Ustun S, Muller P, Palmisano R et al. SseF, a type III effector protein from the mammalian pathogen *Salmonella enterica*, requires resistance-gene-mediated signalling to activate cell death in the model plant *Nicotiana benthamiana*. *New Phytol* 2012;**194**:1046–1060.
- [95] Neumann C, Fraiture M, Hernandez-Reyes C et al. The *Salmonella* effector protein SpvC, a phosphothreonine lyase is functional in plant cells. *Front Microbiol* 2014;**5**:548.
- [96] Deiwick J, Salcedo SP, Boucrot E et al. The translocated Salmonella effector proteins SseF and SseG interact and are required to establish an intracellular replication niche. *Infect Immun* 2006;74:6965–6972.
- [97] Pitzschke A, Schikora A & Hirt H. MAPK cascade signalling networks in plant defence. *Curr Opin Plant Biol* 2009;**12**:421–426.
- [98] Melotto M, Panchal S & Roy D. Plant innate immunity against human bacterial pathogens. *Front Microbiol* 2014;5:411.
- [99] Deng X, Li Z & Zhang W. Transcriptome sequencing of *Salmonella enterica* serovar Enteritidis under desiccation and starvation stress in peanut oil. *Food Microbiol* 2012;**30**:311–315.

- [100] Goudeau DM, Parker CT, Zhou Y et al. The *Salmonella* transcriptome in lettuce and cilantro soft rot reveals a niche overlap with the animal host intestine. *Appl Environ Microbiol* 2013;79:250–262.
- [101] Kyle JL, Parker CT, Goudeau D et al. Transcriptome analysis of *Escherichia coli* O157:H7 exposed to lysates of lettuce leaves. *Appl Environ Microbiol* 2010;**76**:1375–1387.
- [102] Crozier L, Hedley PE, Morris J et al. Whole-transcriptome analysis of verocytotoxigenic Escherichia coli O157:H7 (Sakai) suggests plant-species-specific metabolic responses on exposure to spinach and lettuce extracts. Front Microbiol 2016;7:1088.
- [103] Micallef SA, Rosenberg Goldstein RE, George A et al. Occurrence and antibiotic resistance of multiple Salmonella serotypes recovered from water, sediment and soil on mid-Atlantic tomato farms. *Environ Res* 2012;114:31–39.
- [104] Hein I, Flekna G, Krassnig M et al. Real-time PCR for the detection of *Salmonella* spp. in food: An alternative approach to a conventional PCR system suggested by the FOOD-PCR project. *J Microbiol Methods* 2006;66:538–547.
- [105] Park SH, Aydin M, Khatiwara A et al. Current and emerging technologies for rapid detection and characterization of *Salmonella* in poultry and poultry products. *Food Microbiol* 2014;38:250–262.
- [106] Mandal PK, Biswas AK, Choi K et al. Methods for rapid detection of foodborne pathogens: an overview. *Am J Food Technol* 2011;6:87–102.
- [107] Law JWF, Ab Mutalib NS, Chan KG et al. Rapid methods for the detection of foodborne bacterial pathogens: principles, applications, advantages and limitations. *Front Microbiol* 2015;5:19.
- [108] Zhao X, Lin CW, Wang J et al. Advances in rapid detection methods for foodborne pathogens. J Microbiol Biotechnol 2014;24:297–312.
- [109] Vojkovska H, Kubikova I & Kralik P. Evaluation of DNA extraction methods for PCR-based detection of *Listeria monocytogenes* from vegetables. *Lett Appl Microbiol* 2015;60:265–272.
- [110] Blaser MJ & Newman LS. A review of human salmonellosis: I. Infective dose. *Rev Infect Dis* 1982;4:1096–1106.
- [111] Nastasi A, Mammina C & Mioni R. Detection of *Salmonella* spp. in food by a rapid PCRhybridization procedure. *Microbiologica* 1999;22:195–202.
- [112] MDH. Update: Tomatoes identified as source of Salmonella outbreak in restaurant chain. 2015.
- [113] CDC. *Salmonella* outbreaks. Available at http://www.cdc.gov/Salmonella/outbreaks. html. Accessed July 2016:177–188.
- [114] CDHD. Health department investigating recent salmonella cases. 2015.

- [115] ISDH. Chamberlain farm produce, inc. 2012.
- [116] RKI. Salmonella Newport-Ausbruch in Deutschland und den Niederlanden, 2011. *Epidemiologisches Bulletin* 2012;**20**:177–188.
- [117] TJP. Available at http://www.japantimes.co.jp/news/2011/02/24/news/salads-caused-hokkaido-food-scare. Accessed July 2016. 2011.
- [118] FDH. Fei number 1000515256. 2011.
- [119] IDPH. Summary of S. ser. Hvittingfoss outbreak April-June 2010. 2010.





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