Inferring the Role of the Metabolism of Polyamines in the Phytopathogenic Bacteria *Pseudomonas Syringae*: A Meta-Analysis Approach

Leandro Solmi
Laboratorio de Estrés Biótico y Abiótico en Plantas-Instituto Tecnológico de Chascomús (INTECh), Consejo Nacional de Investigaciones Científicas y Técnicas-Universidad Nacional de General San Martín (CONICET-UNSAM), Buenos Aires (CP7130)

Heman Guillermo Rosli
Laboratorio de Interacciones Planta Patógeno-Instituto de Fisiología Vegetal (INFIVE)

Marina Alejandra Pombo
Laboratorio de Interacciones Planta Patógeno-Instituto de Fisiología Vegetal (INFIVE)

Santiago Stalder
Laboratorio de Estrés Biótico y Abiótico en Plantas-Instituto Tecnológico de Chascomús (INTECh), Consejo Nacional de Investigaciones Científicas y Técnicas-Universidad Nacional de General San Martín (CONICET-UNSAM), Buenos Aires (CP7130)

Franco Rubén Rossi
Laboratorio de Estrés Biótico y Abiótico en Plantas-Instituto Tecnológico de Chascomús (INTECh), Consejo Nacional de Investigaciones Científicas y Técnicas-Universidad Nacional de General San Martín (CONICET-UNSAM), Buenos Aires (CP7130)

Fernando Matías Romero
Laboratorio de Estrés Biótico y Abiótico en Plantas-Instituto Tecnológico de Chascomús (INTECh), Consejo Nacional de Investigaciones Científicas y Técnicas-Universidad Nacional de General San Martín (CONICET-UNSAM), Buenos Aires (CP7130)

Oscar Adolfo Ruiz
Laboratorio de Estrés Biótico y Abiótico en Plantas-Instituto Tecnológico de Chascomús (INTECh), Consejo Nacional de Investigaciones Científicas y Técnicas-Universidad Nacional de General San Martín (CONICET-UNSAM), Buenos Aires (CP7130)

Andrés Gárriz (✉ garriz@intech.gov.ar)
Laboratorio de Estrés Biótico y Abiótico en Plantas-Instituto Tecnológico de Chascomús (INTECh), Consejo Nacional de Investigaciones Científicas y Técnicas-Universidad Nacional de General San Martín (CONICET-UNSAM), Buenos Aires (CP7130)

Research Article
Abstract

*Pseudomonas syringae* is a phytopathogenic bacteria causing disease in a wide variety of economically important plant species. To succeed in plant invasion, these bacteria rely on virulence mechanisms that subvert plant immunity and create favorable conditions for growth. This process requires a precise regulation in the production of important proteins and metabolites. Among them, the family of compounds known as polyamines have attracted attention as they are involved in essential cellular processes. However, it is not known yet how phytopathogenic bacteria regulates polyamine homeostasis in the plant environment. In this work, we conducted a meta-analysis of publicly available transcriptomic data with the purpose to understand the regulation of the metabolism of polyamines and their links to pathogenicity in *P. syringae*. We demonstrated that polyamine synthesis is induced in the early stages of the disease, along with gene expression activation and energy production. On the contrary, the synthesis of these compounds is repressed whereas its transport is up-regulated at later stages, which correlates with the expression of virulence genes and the metabolism of nitrogen and carboxylic acids. We also showed that plant defense mechanisms partially hinder polyamine synthesis, which could reduce cell fitness in the plant environment. In addition, our analysis suggested that a proper bacterial response to oxidative conditions requires a reduction in polyamine production. The implications of these conclusions are discussed.

Introduction

Polyamines comprehend a family of biological compounds that are essential for all living organisms. They are positively charged at physiological pH and bind to polyanionic compounds such as RNA, DNA, proteins and cell membranes. This ability explains their important participation in RNA translation, DNA replication, membrane stability and cell stress tolerance. The most abundant polyamines in bacteria are the diamine putrescine (Put) and the triamine spermidine (Spd), even though polyamines with chemical similarities such as cadaverine and norspermidine might be also dominant in some species. In turn, the occurrence of the tetraamine spermine (abundant in mammals and plants) is rather rare.

Intracellular and extracellular contents of polyamines are controlled by a tight regulation of their pathways of synthesis, catabolism, and transport (Figure 1). Depending on the initial substrate, two pathways are used for the formation of Put. One of them requires the decarboxylation of the amino acid arginine, whereas an alternative pathway decarboxylates ornithine instead. Decarboxylation of arginine is carried out by SpeA (arginine decarboxylase) and forms agmatine, this metabolite thus being later converted into Put by two consecutive enzymatic steps catalysed by agmatine deiminase (AguA) and N-carbamoylputrescine amidohydrolase (AguB). In turn, ornithine is decarboxylated by SpeC (ornithine decarboxylase) to obtain Put as the final product. Then, Put is transformed into Spd by SpeE (spermidine synthase) with the addition of amino propyl groups provided by decarboxylated S-adenosyl methionine, which in turn is the product of the enzymatic step catalysed by SpeD (S-adenosyl methionine decarboxylase). This anabolic pathway is quite conserved in bacteria and several studies demonstrated their significance for growth, stress tolerance and virulence. Despite the catabolic and transport
pathways have been less explored, there is also evidence suggesting that they contribute to polyamine homeostasis and cell fitness\textsuperscript{10–12}. Several routes have been described for the degradation of polyamines in bacteria. First, they can be converted into glutamyl-derivatives by the action of glutamyl-polyamine synthases and then oxidized to gamma-amino butyric acid (GABA) via glutamyl-polyamine oxidases. This involves the protein families PauA and PauB (as they are known in \textit{Pseudomonas aeruginosa}), respectively\textsuperscript{13}. Put might also be transaminated by SpuC using 2-oxo-glutaric acid or pyruvate as amino receptors to render gamma-amino butanal, which is then converted to GABA. In addition, the existence of a Spd dehydrogenase protein has been described in some species\textsuperscript{14,15}, but this enzyme is not widespread in bacteria. In turn, a combination of transport systems participates in the incorporation as well as the secretion of polyamines, and bacterial genomes generally show redundancy in the cognate genes\textsuperscript{16–20}. Among these transporters, PotABCD, PotFGHI, PotE, and SapBCDF are the most studied. PotABCD and PotFGHI are involved in Spd and Put incorporation, respectively, whereas PotE and SapBCDF seem to be involved mostly in polyamine secretion.

The roles played by these pathways in bacteria have been studied in greater extent in \textit{Escherichia coli} and \textit{P. aeruginosa}, but also in other human pathogens such as \textit{Streptococcus pneumoniae}, \textit{Vibrio cholerae}, and \textit{Salmonella enterica}. We currently count on a rather incomplete notion of the importance of polyamines in growth and virulence of phytopathogenic bacteria, even though some reports revealed they are essential in these organisms as well\textsuperscript{21}. In this trend, Spd is required for the synthesis of the toxin Phevamine A in \textit{Pseudomonas syringae}, which helps to suppress the oxidative burst induced in plant tissues upon recognition of the pathogen\textsuperscript{22}. In addition, the secretion of Put by phytopathogenic bacteria has recently attracted some attention. This process was demonstrated to occur in different bacterial species such as \textit{Ralstonia solanacearum} growing at the plant xylem, \textit{P. syringae} dwelling at the apoplastic compartment, and \textit{Dickeya zeae} growing under \textit{in vitro} conditions\textsuperscript{23–25}. On the basis of different reports demonstrating that polyamines have the ability to scavenge reactive oxygen species (ROS) and activate the antioxidant machinery\textsuperscript{26–29}, it was proposed that Put secretion by bacteria constitutes a mechanism to counteract the oxidative stress imposed by plants at the site of the infection, a function that remains to be corroborated. Alternatively, it was proposed that Put secretion in the \textit{D. zeae}/maize interaction is involved in cell-to-cell communication during the invasion of the host\textsuperscript{25}. More work is required to reach a better understanding of the roles that polyamines play in such microorganisms.

As the number of molecular studies examining phytopathogenic bacteria is constantly growing, meta-analysis of publicly available transcriptomic data represents a wonderful tool to comprehend the incidence of gene networks in pathogenesis\textsuperscript{30}. It is plausible that the comparison between these studies will lead to the recognition of common gene expression patterns and the identification of the metabolic pathways implicated in cell behavior. Under this premise, we speculated that this kind of approach could enable us to discern the signatures that characterize the regulation of the metabolism of polyamines in bacterial pathogens of plants and find the associations between the homeostasis of polyamines and bacterial virulence.
We focused this study on transcriptomic data obtained from the bacterial species *P. syringae*. This plant pathogen is worldwide distributed and more than 50 pathovars have been described, which altogether cause disease in the most important crops \(^{31,32}\). We also contemplated other plant and human pathogens to test whether the identified expression signatures could be extrapolated to non-related pathogenic systems. Our work demonstrates that polyamine synthesis is induced initially during plant colonization, a phenomenon that can be partially suppressed by plant defense responses. In addition, even though the expression of genes from the catabolism and transport of polyamines varied among studies, we were still able to conclude that polyamine incorporation is induced in later stages of infection, probably as an attempt to maintain the supply of these important metabolites when their synthesis is reduced. Finally, an interesting finding was that polyamine synthesis is repressed under oxidative stress, which disagrees with the accepted idea that polyamines are required for tolerance to these conditions. The implications of such findings are discussed.

**Results**

**Bacteria repress polyamine metabolism under apoplastic-like conditions.** As a first approach to evaluate the relationship between the homeostasis of polyamines and bacterial virulence, we looked for works comparing the transcriptome of cells growing in *hrp*-inducing medium (HIM) and rich media. HIM composition resembles nutritional conditions of plant apoplast such as low pH and reduced nitrogen/carbon ratio and consequently induces the expression of the *hrp/hrc* genes. The *hrp/hrc* cluster encodes type 3 secretion system (T3SS) components translocating effector proteins into plant cells and are essential for pathogenesis \(^{33,34}\). Two works following these criteria were found. Vandelle *et al* \(^{35}\) used a multistrain-whole genome platform to evaluate early responses in three biovars of *P. syringae pv. actinidiae* (Psa1, 2 and 3) as well as in *P. syringae pv. tomato* DC3000 (Pst), and Nobori *et al* \(^{36}\) examined the transcriptional profile of *Pst* by RNAseq. Figure 2 shows the expression levels of genes involved in synthesis, catabolism, and transport of polyamines when cells grow in HIM compared to rich media. This analysis evidenced the down-regulation of the anabolic pathway, even though the extension of the repression as well as the genes being repressed varied among species and biovars. It is worthy to note that either the expression of *speE* or *speD* was diminished, suggesting that the synthesis of Spd (and seemingly not of Put) is blocked in response to apoplast-mimicking conditions. In turn, relatively minor changes in the expression of genes involved in the degradation and transport of polyamines were evidenced. In this trend, down-regulation of polyamine catabolism and transport was more evident in *Psa3(2)* as well as in *Pst*, which agrees with the similarities between these strains reported by Vandelle *et al* \(^{35}\) in general gene expression. Interestingly, *Psa3* biovars strongly up-regulate the T3SS genes compared to other biovars and are responsible for severe outbreaks of bacterial canker in kiwi fruits in important production areas \(^{35}\). Whether the repression of the polyamine metabolism is associated to the formation of T3SS and aggressiveness of plant pathogens should be studied in further research.

**Polyamine metabolism is induced in bacteria during plant invasion.** We then expanded our investigation by analysing transcriptomic profiles in *Pseudomonas* species during their growth in plant tissues. For this
purpose, five studies using *P. syringae* pathovars were selected. Lovelace *et al* \(^{37}\) and Nobori *et al* \(^{36,38}\) evaluated gene expression in *Pst* invading Arabidopsis, whereas Yu *et al* \(^{39}\) and McAtte *et al* \(^{40}\) focused their research in *P. syringae* pv. *syringae* B728a (*Pss*) and *Psa* colonizing *Phaseolus vulgaris* and Kiwifruit, respectively. It is important to note that the basal conditions to which all samples were compared in these works were different. Thus, whereas Lovelace *et al* \(^{37}\) and Nobori *et al* \(^{36,38}\) used cells grown in the rich medium King’s B and McAtte *et al* \(^{40}\) used the rich medium LB, Yu *et al* \(^{39}\) on the contrary considered bacteria cultivated in minimal nutrient conditions. Therefore, in the light of the observations described in the previous section, the changes in gene expression caused by minimal media should be considered to interpret the results correctly.

In contrast to the expression profiles observed during growth under *in vitro* conditions, this analysis demonstrates an early induction of polyamine biosynthesis in *Pst* (during the first 6 h after the inoculation, 6 hai) (Figure 3A). Thus, genes in charge of the synthesis of Put (*SpeA, SpeC, AguA and AguB*) and Spd (*SpeD, SpeE*) resulted up-regulated. Interestingly, genes related to Put synthesis were not differentially expressed later at 48 hai and the synthesis of Spd was suppressed in both *Pst* and *Pss*. Therefore, we speculate that a rapid induction of polyamine synthesis is important at the start of plant invasion in these strains. Besides, either the repression of this pathway is required later during the development of disease or rather it is provoked by the activation of the plant defense system. In turn, the analysis of expression of genes from the catabolism and transport of polyamines under *in planta* conditions is rather inconclusive, as many of them showed contrasting expression patterns among the dataset included. The only consistent observation was an early repression of *spuH* and induction of *potD1*. It could be plausible that these pathways are more sensitive to different experimental setups and/or that their regulation strongly depends on the bacterial genotype. In fact, we observed that expression profiles shown by *Psa* differs considerable from those observed in other strains, as polyamine biosynthesis was not regulated in the early period of plant invasion, and most of the genes involved in polyamine catabolism and transport were induced throughout this process.

In the interest of confirming if polyamine synthesis is important during early plant invasion stages, we searched for phenotypes in mutants of the cognate genes at the Fitness Browser database described at Helmann *et al* \(^{41,42}\) (https://fit.genomics.lbl.gov/cgi-bin/myFrontPage.cgi). In agreement with this, we found that the transposon-mediated disruption of the *speE* gene in *Pss* leads to a mild negative fitness score when growing in the apoplast of green beans (data not shown), suggesting that at least at the start of pathogenesis the anabolism of Spd results important.

Two main defense responses are triggered in plants against the attack of potential pathogens \(^{43}\). The first layer of defense is activated upon the recognition of pathogen associated molecular patterns (PAMPs, such as the flagellin peptide flg22 and chitin) and is known as PTI (PAMP-triggered immunity). Pathogens introduce effectors inside plant cells to counteract PTI and promote virulence \(^{44}\). Resistant plants can detect directly or indirectly some of these effectors initiating a second defense response, which has been coined as Effector-triggered immunity (ETI) or nucleotide-binding Leu-rich repeat (NLR)-
triggered immunity (NTI)\textsuperscript{45}. Our analysis showed that the pre-induction of PTI (by treating plants with flg22 before inoculation) hinders the induction of polyamine biosynthetic genes and leads to the upregulation of genes from the transport systems SapBCDF and PotABCD (Figure 3B, flg22 vs. mock). In agreement with this observation, the effectorless polymutant \textit{PstD36E} (unable to supress plant immune system) demonstrated a relatively minor induction of polyamine biosynthesis compared to WT \textit{Pst}, whereas components of the \textit{sapBCDF/potABCD} operons were also induced. These results suggest that plant defense mechanisms affect polyamine synthesis in bacteria, which conduces to the activation of polyamine incorporation to overcome their limitation. In addition to that, our analysis also indicates that the translocation of bacterial effectors helps to counteract the negative effects of plant defense responses on polyamine homeostasis. In turn, the activation of ETI (evaluated by plant inoculation with strains expressing either the avirulence genes \textit{avrRPT2} or \textit{avrRps4}) originate slight effects on bacterial polyamine metabolism, as just a higher induction of genes coding for polyamine transporters in the \textit{Pst-AvrRpt2} strain were verified. It should be considered though that ETI causes a rapid induction of plant defense mechanisms that strongly reduces pathogen’s growth. Thus, it is possible that the effects on bacterial polyamine metabolism could be masked on such drastic environmental conditions.

\textbf{Different networks are coordinated with polyamine homeostasis during plant invasion.} The broad diversity of conditions used in the datasets included in this study enabled us to perform a co-expression analysis with the purpose of identifying gene modules being co-ordinated with the homeostasis of polyamines. We first analysed the data collected by Nobori \textit{et al}\textsuperscript{36,38} using \textit{Pst}/Arabidopsis as a model system. In this case, four genes coding for polyamine biosynthetic enzymes (\textit{speA}, \textit{speC}, \textit{speE} and \textit{speD}), one for a transporter (\textit{potF}) and four involved in polyamine catabolism (\textit{pauA1}, \textit{pauA2}, \textit{pauA4} and \textit{spuC}) were found to establish significant co-expression relationships with other gene nodes. An evaluation of these groups of genes as well as GO term enrichment analysis identified over-representation of gene categories intimately related to nitrogen compound biosynthesis, carboxylic acid metabolism, energy generation, primary metabolic processes, transmembrane transport, gene expression and macromolecular biosynthetic processes (Figure 4A and 4C). Interestingly, genes from different polyamine pathways established contrasting relationships with these nodes. Thus, whereas polyamine anabolic genes showed positive co-expression scores with membrane transport, amino acid synthesis and regulation of gene transcription/translation, those in charge of polyamine catabolism showed negative scores, suggesting the existence of regulatory mechanisms that avoid simultaneous activation of both pathways. It is interesting to note also that polyamine biosynthesis shows positive correlation with genes involved in energy generation (ATP synthase subunits and electron transport proteins) but negative correlation with genes involved in flagellum formation (Figure 4A). This indicates that energy generation is strongly associated with polyamine synthesis, and that polyamines are not required (or they might be even counter-productive) for the activation of bacterial movement (Figure 4A). We then conducted a dendrogram clustering analysis to better understand the links in the regulation of these gene groups. Thus, the identified biosynthetic genes are grouped together in cluster III (Figure S1A), which includes genes induced at the beginning of plant invasion (6 hai) and repressed at later stages (48 hai). Interestingly, these genes are also repressed in samples taken from PTI-induced plants or in the
hypovirulent strain *Pst*-D36E, as well as in *in vitro* experiments using both rich or minimal media. In turn, the polyamine catabolic genes were ordered in cluster II. Even though this group is also mildly induced in the early periods of plant invasion and repressed *in vitro*, the outstanding feature of these genes is a strong induction at later stages of the infection. Altogether, these results agree with our previous assumption that polyamine biosynthesis is required to ensure bacterial fitness at the start of plant invasion, which would be related to the effects of polyamines on gene expression regulation and energy production. Polyamine accumulation, on the contrary, would not be required at subsequent stages, where the catabolic pathway is activated to reduce their contents.

We then performed the same analysis using the dataset published by Yu *et al* \(^3^9\). This work compared gene expression in the strain *Pss* B728a during the infection of *P. vulgaris* 48 hai (apoplastic and epiphytic growth) and growth under different stressful conditions *in vitro* (Figure 4B). A much broader list of genes from the different pathways of the metabolism of polyamines was identified showing significant co-expression scores. GO term enrichment analysis indicated that categories belonging to degradative pathways and membrane transport were over-represented (Figure 4D). Interestingly, even though most of these genes were linked to others through positive correlations, they were clustered in different groups (Figure S1B) and many of these clusters included genes from different polyamine metabolism pathways. Given that only samples taken 48 hai were considered in this dataset, it can be hypothesised that at this stage of infection precise regulatory mechanisms modulates specific genes from the different pathways of the polyamine metabolism rather than regulating the entire route. Importantly, these regulatory mechanisms also operate in response to stressful conditions. The biosynthetic gene *speA* is mostly linked by negative edges to genes from secondary metabolism and regulators of gene transcription and translation (Figure 4B). This observation suggests that, as concluded above, polyamine synthesis may not be relevant for gene expression as it is at the start of the process. *speA* was grouped in cluster I, genes that are mildly repressed in response to low nitrogen availability but remained unaltered under other growing conditions (Figure S1B). Thus, it is conceivable that bacteria face a reduction in nitrogen through the development of the disease and respond by repressing the expression of *speA* to limit the use of nitrogen in favour of the synthesis of other essential compounds. Cluster II contains the operon formed by the Put/Spd transport system PotFGHI, the Put exporter PotE(1), the catabolic oxidase PauB3 and the Spd synthase SpeE. These genes established positive correlations with gene expression regulators and transmembrane proteins, which shows a mild induction when cells are grown under osmotic conditions but not *in planta*, suggesting that they play a role in this type of stress by result dispensable in the later stages of plant colonization. In turn, cluster III includes the largest amount of polyamine metabolism genes. All these genes belong to the catabolic and transport pathways and establish positive correlation with a large amount of genes up-regulated under low N conditions. It agrees with the idea that the induction of polyamine catabolism and transport helps to optimize the use of nitrogen under such stressful conditions. Interestingly, these genes also showed positive edges with many genes that conform the *hrp/hrc* operons, indicating that their induction in response to low N contents accompanies the activation of bacterial virulence. At last, cluster IV grouped genes up-regulated *in planta*, but not under other stressful conditions. In this cluster, we identified genes belonging to the
transporter systems PotABCD and PotFGHI, confirming the idea that transport of polyamines might play an important role during later periods of the infection.

**Polyamine metabolism is shut-down during the activation of the anti-oxidative response.** Polyamines have been described as essential metabolites to cope with oxidative stress in bacteria \(^{46-50}\). However, our co-expression analysis shown in the previous section did not show a significant correlation with genes involved in redox homeostasis. The only exception is a positive correlation between the gene coding for the catalase KatG and the glutamyl-Put synthase PauA2 in Figure 4A. Interestingly, it was demonstrated that KatG (along with KatB) is required for full virulence in \(Pst\) \(^{51}\) and the induction of this gene in response to \(H_2O_2\) in \(E.\ coli\) is dependent on polyamines \(^{50}\). Therefore, we wondered if the activation of the detoxifying machinery in bacteria growing under oxidative stress is accompanied by modifications in the metabolism of polyamines. With this in view, we looked first for transcriptomic data studying bacterial gene expression in response to \(H_2O_2\), and selected works that stressed cells for a short period to evaluate the immediate response. In this case, we also included species from other genera to test the existence of widespread regulatory mechanisms. As shown in Figure 5, our analysis that the metabolism of polyamines remains almost unaltered in response to stress, except for the repression of Put or Spd synthesis shortly after the addition of \(H_2O_2\) to the culture media. A stronger repression of polyamine synthesis and transport occurs in \(E.\ coli\) and \(S.\ enterica\) (Figure S2), suggesting that in these cases an earlier regulation of the intracellular concentration of these metabolites might be crucial for the antioxidant response. In fact, oxidative stress hallmark genes such as \(oxyR\), \(soxS\) and \(soxR\) resulted more highly activated than in \(Pseudomonads\) at the time points analysed. Even so, we conceive that despite suitable levels of polyamines might be required to activate essential component of the antioxidant machinery as reported previously, a rapid repression of polyamine metabolism occurs as part of the stress response. It would be interesting to explore, then, how phytopathogenic bacteria adapt the metabolism of polyamines to cope with the oxidative stress imposed by plants without affecting cell fitness and pathogenesis.

**Discussion**

The growing number of publicly available transcriptomic datasets represents a powerful source for the identification of the main features involved in particular cellular processes, such as growth, development, and stress resistance \(^{30}\). In the case of pathogenic microbes, these studies may provide a good deal of information concerning common mechanisms participating in the activation of virulence, and at the same time, they offer the possibility to evaluate the relative importance of specific pathways. In this work, we explored the importance of the metabolic routes maintaining the homeostasis of polyamines for pathogenesis and oxidative stress response in bacterial species from the genera \(Pseudomonas\), with an emphasis in \(P.\ syringae\).

We first compared gene expression levels in cells growing in minimal media in relation to rich media. Minimal media are thought to mimic apoplastic conditions, mainly low N/C ratio and acidic pH, which
lead to the induction of virulence-associated hrp/hrc operons coding for the T3SS. The studies selected for this work offered a time-lapse analysis spanning 8 h of culture and indicate the existence of a regulatory network suppressing polyamine homeostasis under minimal nutrient environments. These results curtail the importance of polyamines to support growth in minimal media (so that the synthesis of other metabolites is preferred) and besides, suggest that the mechanisms governing polyamine homeostasis differ from those inducing bacterial virulence. Even though this observation contradicts our previous gene expression analysis in Pst using qRT-PCR (where the polyamine biosynthetic gene speC and the catabolic enzymes pauA3 and pauB2 resulted mildly up-regulated in minimal medium), it should be considered that only samples taken 6 hai were analysed in that work and that variations in gene expression before or after that time could have been missed.

A point to be considered is that the plant apoplast is richer in nutrients and that it is sufficient to support the growth of pathogens, at least throughout the early stages of infection. This could explain why in planta growth generates much broader modifications of bacterial transcriptomes than in vitro conditions, which not always mirrors to those observed in in vitro experiments. Thus, expression data from in planta studies should reflect the modulation of the polyamine metabolism and its link with pathogenesis in a more accurate manner. Different studies evaluating the transcriptional profiles of Pseudomonas species in planta were selected to examine the regulation of polyamine pathways during plant colonization. In this trend, our analysis found substantial changes with respect to the metabolic behaviour showed by cells in vitro, as the synthesis of polyamines was induced during the first stages of infection (up to 6 hai) and repressed later with the advance of pathogenesis (48 hai). At the earliest periods considered in this work, it is conceived that plants have already recognized the nature of the pathogen and triggered the first line of defense, whereas the microbe in turn is deploying virulence strategies to resist them. Thus, we concluded that an initial rise in the contents of polyamines via de novo synthesis is required to support cell fitness at this stage. If this is true, mutant strains perturbed in polyamine synthesis should be affected in their virulence. Even though this kind of mutants have not been deeply tested in Pseudomonads yet, a genome-wide fitness profiling made on Pss showed that deletion of ΔspeE conduces to mild negative fitness scores when growing in green pepper. The fact that no other mutant strains affected in genes contributing to polyamine biosynthesis or transport show negative scores could be explained on the bases of the existence of redundancy in these pathways. Thus, this would assure the supply of these metabolites during first steps of plant invasion even if these pathways are partially affected by the plant defense systems. In this trend, works in Ralstonia solanacearum (which exclusively depends on the SpeC pathway for Put synthesis) demonstrated that a ΔspeC mutant is hypovirulent in tomato.

The importance of maintaining polyamine contents early in the interaction can be explained through the participation of these compounds in essential molecular mechanisms. For instance, it has been shown that polyamines are required to assure correct ribosome ensemble and general transcription efficiency. In relation to this, our co-expression analysis based on the work made by Nobori et al., where most of the samples correspond to cells growing for 6 h both in vitro and in planta, indicated that genes involved
in polyamine biosynthesis were coexpressed with ribosome components as well as regulators of transcription/translation. On the contrary, polyamine catabolic genes showed a negative coexpression relationship with the same group of genes. We also distinguished the same pattern of coexpression with genes involved in ATP formation, carboxylic acid and nitrogen metabolism. Thus, polyamine accumulation would be associated to the synthesis of important compounds and a higher rate of energy generation and gene expression. An interesting observation derived from this analysis is the fact that polyamine biosynthetic genes were negatively correlated with genes participating in the formation of the flagella, which suggests that bacterial movement could be affected by polyamines in *Pseudomonads*. In agreement with this, it was recently demonstrated that Put accumulation by disrupting Spd formation or the Put aminotransferase pathway promotes the transition towards the formation of sessile biofilms in *P. aeruginosa*\(^{54}\). However, the link between polyamines and cell movement seems to depend on the bacterial species and the polyamine involved. In this trend, even though deletion of *speA* and *speC* in *Yersinia pestis* also perturbs biofilm formation\(^ {55}\), the lack of *speA* in *Dictyella zeae* and *Proteus mirabilis* decreases swimming and swarming mobilities\(^ {7,25}\). It would be interesting to corroborate the significance of polyamine contents for cell movement in *P. syringae*.

We also demonstrated that pre-induction of plant defense responses leads to a minor induction of polyamine biosynthesis and to the up-regulation of polyamine transport genes. This also occurs when the effectorless D36E mutant strain is analysed, emphasising the importance of the T3SS function in avoiding the effects of plant defense mechanisms on pathogen´s metabolism. We speculate that a rise in the expression of polyamine transporters under these conditions is a consequence of lower expression of the anabolic genes. The mechanisms underlying the reduction in the synthesis of polyamines could be either 1) a bacterial response to the deleterious environment imposed by the plant defense system, or 2) that plant defense responses directly target the synthesis of these compounds to affect cell fitness. In relation to this, it is known that one of the main defense mechanisms deployed by plants against endophytic bacteria is the accumulation of reactive oxygen species at the apoplast\(^ {39,52}\), provoking oxidative stress conditions to bacteria dwelling at this niche. With this in mind, we explored transcription datasets obtained from different bacterial species exposed to oxidative stress and found that, even though at different extents, polyamine biosynthetic genes were down-regulated in bacteria. Hence, the reduction in polyamine synthesis might be associated with the induction of the antioxidative machinery that helps to cope with plant defense responses. This seems to contradict the accepted paradigm that polyamines help to cope with this stress\(^ {11,48,56,57}\). In this trend, secretion of polyamines by *R. solanacearum* at the tomato xylem was proposed by Lowe-Power *et al*\(^ {23}\) as a possible mechanism to subdue oxidative stress. These authors demonstrated that Put is accumulated at this compartment presumably as a product of the activation of bacterial metabolism, and that a Δ*speC* mutant resulted completely avirulent. In relation to this, we described in a previous work that *Pst* can also secrete large amounts of Put when growing in apoplastic washing fluids, although we could not find a link between this phenomenon and bacterial virulence. Our next experiments will try to use *Pst* mutants unable to synthesize polyamines to further test their roles in stress tolerance and its link with pathogenesis.
As mentioned above, our study showed that bacterial polyamine biosynthesis is repressed 48 hai, which is also associated to the induction of polyamine transporters and catabolic genes. A similar scenario is observed in later stages of pathogenic interactions established by other bacterial species. For instance, *Xanthomonas axonopodis pv. glycines* reduces the expression of *speD* but up-regulates other polyamine biosynthetic and transport genes after 72 hai in soybean \(58\), whereas polyamine synthesis was repressed in *Xanthomonas oryzae pv. oryzae* after 5 days while polyamine transport was activated \(59\) (data not shown). Importantly, hierarchical clustering of co-expressed gene analysis demonstrated that transcription profiles from samples taken 48 hai are grouped with those obtained from cells growing *in vitro* under low nitrogen conditions or in minimal medium, as well as in experiments using primed plants (pre-infiltrated with flg22) or the D36E strain. Based in this, it is reasonable to conclude that bacteria are facing more restrictive conditions at this stage. Besides, the induction of some of the polyamine metabolic genes at 48 hai is correlated with the induction of the T3SS. This observation supports the idea that the modulation of specific genes from the metabolism of polyamines, but not all, would be coordinated with the induction of virulence mechanisms. More work is needed to understand the regulatory mechanisms acting regulating genes from the polyamine metabolism throughout the infection and their impact on pathogenesis.

Altogether, these findings contribute to the hypothesis that polyamine synthesis in phytopathogenic *Pseudomonas* species is important during the early periods of plant invasion, which is accompanied by the activation of nitrogen and carboxylic acid metabolism, general gene expression and energy generation. Afterwards, this pathway is repressed in later stages with the consequent induction of the transport systems (Figure 6), coordinated with the expression of virulence genes and degradative enzymes. This metabolic shift enables the survival of cells in the plant environment. Whether this behaviour could be extrapolated to other bacterial species should be evaluated with the generation of specific mutant strains perturbed in the different branches of the polyamine metabolism.

**Materials And Methods**

**Identification of orthologs of the polyamine metabolism genes in bacterial genomes.** To identify groups of orthologs in the different bacterial species considered in this work, we followed two approaches. Thus, we explored the Ortholog Group Member database at Pseudomonas Genome DB (https://pseudomonas.com), or alternatively, applied the reciprocal best hit method using the NCBI’s blastp software \(60\). In those cases where this pipeline failed in finding potential orthologs in a defined database, we still included in our analysis the best hit if the query coverage was higher than 80% and E-value was below \(1 \times 10^{-10}\). A list of the genes considered for each strain is shown in Table S1.

**Selection of transcriptomic datasets from the public domain.** Microarray and RNAseq data were downloaded from the NCBI Gene Expression Omnibus (GEO). GEO repository was queried using keywords such as “Pseudomonas”, “Phytopathogenic bacteria”, “pathogenic bacteria”, “oxidative stress”, and “plant infection”. In total, nine analyses were selected. Seven of these works were performed in three different pathovars of *P. syringae* and other *Pseudomonads* such as *P. putida* and *P. aeruginosa*, whereas two
considered the human pathogens *E. coli* and *S. enterica* (see Table 1 for a brief description). We explored data obtained from *P. syringae* growing under *in vitro* and *in planta* conditions to unravel the links between polyamine metabolism and pathogenesis, whereas those works evaluating bacterial responses to H$_2$O$_2$ were used to corroborate the potential role of these metabolites in the stress response. In this last case, we excluded studies analyzing gene expression beyond 1 h of treatment, as we were interested in the early response. In addition, works that utilize concentrations of H$_2$O$_2$ higher than 15 mM were also ruled-out as these concentrations are significantly higher than the physiological amounts faced by bacteria in plant tissues. Because of the experimental heterogeneity associated to these datasets and the complexity underlying their normalization, we did not re-analyze them using a unified pipeline, but rather attempted to identify transcriptional patterns in a broader sense by comparing the expression levels reported by the authors in each case. Thus, gene expression levels were obtained from the DEGs (Differentially Expressed Genes) listed in each study, and genes were included in our analysis if the fold change (log$_2$FC) resulted >|0.5| and showed p-adjusted values <0.05.
Table 1
Description of the datasets used for the meta-analysis in this work. Bacterial species/strains, Plant host and Growth conditions are listed

<table>
<thead>
<tr>
<th>Authors</th>
<th>Study accession</th>
<th>Bacterial strain</th>
<th>Plant host</th>
<th>Growth conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nobori et al'36,38</td>
<td>GSE103442, GSE138901</td>
<td><em>P. syringae pv. tomato</em> DC3000</td>
<td><em>Arabidopsis thaliana</em> Col-0</td>
<td><em>In vitro</em> (minimal and rich media) and <em>In planta</em></td>
</tr>
<tr>
<td>Yu et al'39</td>
<td>GSE42544</td>
<td><em>P. syringae pv. syringae</em> B728a</td>
<td><em>Phaseolus vulgaris</em> cultivar Bush Blue Lake 274</td>
<td><em>In vitro</em> (minimal and rich media, stressful conditions) and <em>In planta</em></td>
</tr>
<tr>
<td>McAtte et al'40</td>
<td>PRJNA472664</td>
<td><em>P. syringae pv. actinidiae</em> ICMP 18884</td>
<td><em>Actinidia chinensis</em> Planch. var. chinensis “Hort16A”</td>
<td><em>In planta</em></td>
</tr>
<tr>
<td>Vandelle et al'35</td>
<td>GSE164472</td>
<td><em>P. syringae pv. actinidiae</em> biovars 1, 2 and 3</td>
<td>None</td>
<td><em>In vitro</em> (minimal and rich media)</td>
</tr>
<tr>
<td>Bojanovic et al'64</td>
<td>GSE85475</td>
<td><em>P. putida</em> KT2440</td>
<td>None</td>
<td><em>In vitro</em> (H₂O₂ amended media)</td>
</tr>
<tr>
<td>Jozefczuk et al'65</td>
<td>GSE20305</td>
<td><em>E. coli</em> MG1655</td>
<td>None</td>
<td><em>In vitro</em> (H₂O₂ amended media)</td>
</tr>
<tr>
<td>Chang et al'66</td>
<td>GSE3090</td>
<td><em>P. aeruginosa</em> PA01</td>
<td>None</td>
<td><em>In vitro</em> (H₂O₂ amended media)</td>
</tr>
<tr>
<td>Liu et al'67</td>
<td>GSE 155479</td>
<td><em>S. enterica</em> subs. enterica serovar Enteritidis ATCC 13076</td>
<td>None</td>
<td><em>In vitro</em> (H₂O₂ amended media)</td>
</tr>
</tbody>
</table>

Co-expression analysis.

Datasets uploaded by Nobori et al'36,38 and Yu et al'39 were used to calculate Pair-wise gene expression Pearson correlation across all the samples to generate a similarity matrix, which then were the inputs for the construction of co-expression networks (using R/WGCNA version 1.34) as described by Sharma et al'61. Hierarchical cluster trees were created by setting gene co-expression module seizure of 30, deepSplit at level 1 and tree mergeCutHeight at 0.20. Figures were created using Cytoscape'62 and enriched GO terms were identified using BiNGO plugin'63.

Declarations
Acknowledgments

L. Solmi, H. G. Rosli, M. A. Pombo, F. R. Rossi, F. M. Romero, O. A. Ruiz and A. Gárriz, are members of the Research Staff of the Consejo Nacional de Investigaciones Científicas y Técnicas (CONICET, Argentina). This work was partly funded by the project “Universidades Agregando Valor” #8550 from the Secretaría de Políticas Universitarias, Argentina. The authors would like to thank Josefina Gárriz for his diligent proofreading of this manuscript.

Author Contributions

L.S and A.G. designed the study; S.S., L.S. and A.G. prepared figures; A.G. wrote the main text. All authors analyzed and reviewed the manuscript.

Competing interests

The author declares no competing interests.

References


### Figures

**Figure 1**

Schematic representation of polyamine metabolism pathways. Biosynthetic enzymes are represented in white rectangles, while black and grey rectangles depict the enzymes involved in polyamine catabolism and transport, respectively. Adapted from Schneider and Wendisch and created with BioRender.com
Figure 2

Heatmap representing polyamine metabolism gene expression levels in *Pseudomonas syringae* during growth in apoplastic-like conditions compared to rich media (|log$_2$FC|>0.5, p adj<0.05).
Figure 3

Polyamine metabolism gene expression in *P. syringae* during growth in plant tissues. Heatmap representing polyamine metabolism gene expression levels in *P. syringae* during **A)** plant invasion in comparison to basal conditions and **B)** in Flg22-treated plants compared to mock, as well as non-virulent strains compared to WT Pst (|log₂ FC|>0.5, p-adj<0.05)
Figure 4

mRNA co-expression networks in *P. syringae* using the datasets from Nobori *et al* (2018, 2020) (A) and Yu *et al* (2013) (B). Red and green edges denote negative and positive correlations, respectively, for highly correlated genes ($R^2 > 0.75$ for Nobori *et al*, 2018, 2020; and $R^2 > 0.99$ for Yu *et al*, 2013)). Functions are color-coded. Pearson’s correlation coefficients were used. Bar graphs in C) and D) depict GO-enrichment analysis of the genes included in A and B, respectively.
Figure 5

Heatmap showing polyamine metabolism and oxidative stress gene expression in *Pseudomonas* species during exposure to H$_2$O$_2$ in comparison to basal conditions (|log$_2$FC|>0.5, p adj<0.05).
Figure 6

Legend not included with this version

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- FigureS1.pptx
- FigureS2.jpg
- TableS1.docx