The *Lactococcus lactis* KF147 nonribosomal peptide synthetase/polyketide synthase system confers resistance to oxidative stress during growth on plant leaf tissue lysate

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Abstract

Strains of *Lactococcus lactis* isolated from plant tissues possess adaptations that support their survival and growth in plant-associated microbial habitats. We previously demonstrated that genes coding for a hybrid nonribosomal peptide synthetase/polyketide synthase (NRPS/PKS) system involved in production of an uncharacterized secondary metabolite are specifically induced in *L. lactis* KF147 during growth on plant tissues. Notably, this NRPS/PKS has only been identified in plant-isolated strains of *L. lactis*. Here, we show that the *L. lactis* KF147 NRPS/PKS genes have homologs in certain *Streptococcus mutans* isolates and the genetic organization of the NRPS/PKS locus is conserved among *L. lactis* strains. Using an *L. lactis* KF147 mutant deficient in synthesis of NrpC, a 4′-phosphopantetheinyl transferase, we found that the NRPS/PKS system improves *L. lactis* during growth under oxidative conditions in *Arabidopsis thaliana* leaf lysate. The NRPS/PKS system also improves tolerance of *L. lactis* to reactive oxygen species and specifically H$_2$O$_2$ and superoxide radicals in culture medium. These findings indicate that this secondary metabolite provides a novel mechanism for reactive oxygen species detoxification not previously known for this species.

**KEYWORDS**

*Lactococcus lactis*, natural products, NRPS/PKS, Plant-associated bacteria, reactive oxygen species, secondary metabolites

1 | INTRODUCTION

Numerous bacterial species in multiple phyla possess the capacity to synthesize secondary metabolite peptides and carboxy acids by nonribosomal peptide synthetases (NRPS) and polyketide synthases (PKS) (Weissman, 2015). These secondary metabolites can include proteinogenic and nonproteinogenic amino acids and ketides assembled into diverse end-products with functions ranging from antibiotics to immunosuppressants (Siezen & Khayatt, 2008). NRPS/PKS systems have been described in detail for many industrially significant bacteria but there are few reports characterizing these systems and their resulting products by lactic acid bacteria (LAB) (Lin et al., 2015; Wu et al., 2010).

LAB are gram-positive, nonsporulating bacteria in the *Firmicutes* phylum and constitute a genetically diverse collection of species characterized by their fermentation of mono- and di-saccharides to lactic acid. Strains of *Lactococcus lactis* are among the most extensively characterized LAB and are used in a variety of applications including starter cultures in cheese production (Cavanagh, Fitzgerald, & McAuliffe, 2015), industrial product synthesis (Mierau et al., 2005), and for the delivery of therapeutics to the human gastrointestinal...
tract (Cano-Garrido, Seras-Franzoso, & Garcia-Fruitos, 2015; Wells & Mercenier, 2008). Although L. lactis has been most extensively investigated for its adaptation to dairy environments, it is generally regarded that plants are the ancestral habitat for this species (Cavanagh, Fitzgerald, & McAuliffe, 2015; Cavanagh, Casey et al., 2015). L. lactis KF147, a strain originally isolated from mung bean sprouts, is genetically distinct from dairy-associated isolates (Siezen et al., 2008). Its unique traits include the ability to metabolize a wide array of plant-derived carbohydrates and synthesize exopolysaccharides (Siezen et al., 2008).

The genome of L. lactis subspecies lactis KF147 contains a hybrid NRPS/PKS gene cluster (Siezen et al., 2008, 2010). These chromosomally-located genes have thus far only been identified in plant-derived lactococci (Siezen et al., 2011). However, the conditions in which those genes are expressed were not described. Because the production of secondary metabolites poses a significant energy burden on the cell, genes necessary for the production of those cell products are frequently cryptic in standard laboratory media (Rutledge & Challis, 2015). In fact, stringent transcriptional control and regulation of NRPS/PKS production is a frequent challenge that impedes the study of these systems and other secondary metabolites (Rutledge & Challis, 2015). Therefore, it was notable when we found that the L. lactis KF147 genes coding for the hybrid NRPS/PKS are induced during the growth of this strain in Arabidopsis thaliana leaf tissues (Golomb & Marco, 2015). The findings provided an opportunity to study the functional relevance of the NRPS/PKS system in L. lactis. Here, we examine the genetic relatedness of the NRPS/PKS to other lactococci and Streptococcus mutans strains and demonstrate the contribution of the secondary metabolite to L. lactis growth in oxidative conditions, most appreciably on plant tissues.

2 | EXPERIMENTAL PROCEDURES

2.1 | Bacterial strains and culture conditions

Lactococcus lactis subsp. lactis KF147 (Kelly, Davey, & Ward, 1998) was provided by NIZO food research (Ede, The Netherlands) and was maintained as a frozen glycerol stock at −80°C. L. lactis was routinely grown at 30°C without agitation on M17 broth (BD, Franklin Lakes, NJ) supplemented with 0.5% d-glucose (GM17). Arabidopsis thaliana Col-1 leaf lysate (ATL medium) was prepared as previously described (Golomb & Marco, 2015). Escherichia coli DH5α was grown at 37°C on LB (Fisher Scientific, Waltham, MA). When necessary, the culture medium was supplemented with ampicillin at 100 μg/ml for E. coli and erythromycin at 5 μg/ml for L. lactis. For reactive oxygen species tolerance experiments, optical density (OD) at 600 nm was used to examine L. lactis growth in GM17 in a Synergy 2 microplate reader (BioTek, Winooski, VT) in 200 μl volumes. L. lactis numbers were also determined by plating serial dilutions of the cultures onto GM17 agar and incubation at 24 hr prior to viable cell enumerations. L. lactis used for growth experiments was prepared from stationary phase cells incubated in GM17 for approximately 18 hr. Cells were collected by centrifugation (10,000g, 3 min) and washed twice in phosphate-buffered saline [PBS, 137 mmol/L NaCl, 2.7 mmol/L KCl, 4.3 mmol/L Na2HPO4, 1.4 mmol/L KH2PO4 (pH 7)] prior to use.

2.2 | Sequence analysis of the L. lactis KF147 NRPS/PKS genes

Annotated NRPS/PKS genes were retrieved from the complete genome sequence of KF147 (Siezen et al., 2010). Homology searches were performed, using BLASTp against the NCBI nr database (http://blast.ncbi.nlm.nih.gov/Blast.cgi).

2.3 | Construction of the L. lactis nrpC deletion mutant BAL1

Standard molecular biology techniques were performed as previously described (Sambrook, Fritsch, & Maniatis, 1989). Double-crossover homologous recombination was used to generate a markerless nrpC deletion mutant of KF147, using the suicide vector pRV300 (Leloup, Ehrlich, Zagorec, & Morel-Deville, 1997). All primers used for mutant construction are shown in Table S1. The upstream flanking region of nrpC was PCR amplified, using primers A and B and the downstream flanking region was PCR amplified, using primers C and D. The resulting amplicons were combined by splicing-by-overlap extension (SOEing) PCR (Horton, Cai, Ho, & Pease, 1990). The resulting PCR product was digested with SalI and SacII, cloned into pRV300, and transformed into E. coli DH5α to yield pNRPC-KO. pNRPC-KO was next introduced into KF147 by electroporation as previously described (Golomb & Marco, 2015). Single-crossover mutants were selected on GM17 agar plates supplemented with erythromycin. A single-crossover mutant grown on GM17 broth in the absence of erythromycin for approximately eight passages was sufficient to facilitate excision of the plasmid. Double-crossover mutants were identified by replica plating onto GM17 agar supplemented with erythromycin. Erythromycin-sensitive colonies were screened by PCR for a nrpC deletion. PCR products with primers A and D that resulted in a product size of 1125 bp indicated the absence of nrpC and pRV300 from the chromosome (Table S1). The PCR products were also sequenced (UC Davis DNA Sequencing Facility (http://dnaseq.ucdavis.edu) for confirmation. A single nrpC deletion mutant, strain BAL1, was used in subsequent experiments.

2.4 | ROS tolerance

The capacity of L. lactis to tolerate Reactive oxygen species (ROS) was examined in multiple ways including (1) growth in H2O2 (2) paraquat (methyl viologen dichloride hydrate), (3) xanthine with xanthine oxidase (XO), and (4) under respiratory conditions in the presence of hemin. For these studies, GM17 was supplemented to a final concentration with either 0.5 mmol/L H2O2 (Macron Fine Chemicals, Center Valley, PA), 40 mmol/L paraquat (Sigma-Aldrich, St. Louis, MO), or 5 mmol/L xanthine (Sigma-Aldrich) in the presence of 0.3 U/ml XO (EMD Millipore, Billerica, MA). For the growth in the presence of xanthine, controls were supplemented with xanthine but not XO. In ATL, 10 mmol/L paraquat was used. For growth under
respiratory conditions, *L. lactis* was incubated with continuous aeration in GM17 supplemented with porcine hemin at a final concentration of 10 μg/ml (Chem-Impex International, Wood Dale, IL) and where indicated, 20 mM/L paraquat.

### 2.5 Statistical analysis

Significant differences in *L. lactis* growth rates were determined using an unpaired, two-tailed Student t-test. Differences were considered significant if *p* < .05.

## 3 RESULTS AND DISCUSSION

### 3.1 The NRPS/PKS locus contains distinct functional features and is conserved among plant-associated *L. lactis* strains

The *L. lactis* KF147 genes coding for the NRPS/PKS system (llkf_1211 to llkf_1222) span over 40 kb (ca. 1.5% of the chromosome) and consist of a two-component response regulator (npkKR), biosynthetic enzymes (nrpA, pk1ABC, npkS, and npkB), a 4′-phosphopantetheinyl transferase (PPTase, npC), and a putative ATP-binding cassette (ABC) transporter (llkf_1217 and llkf_1218) (Figure 1). The complete locus spans from llkf_1209 to llkf_1235 and includes the two accessory genes llkf_1224 annotated as an AvrD macrolide biosynthetic protein and llkf_1125 annotated as a phytoene dehydrogenase (Figure 1). Finally, there are multiple hypothetical proteins (Figure 1).

Flanking one end of the NRPS/PKS locus in *L. lactis* KF147 is a DNA integration/recombination/inversion protein (llkf_1230) (Figure 1). This protein is similar to lactococcal and streptococcal phage integrases (30%-93% amino acid similarity) and shares a C-terminal catalytic domain with the *Bacillus subtilis* ICEBs1 conjugal transposon integrase. The integrase is required in *B. subtilis* for ICEBs1 excision and integration (Lee, Auchtung, Monson, & Grossman, 2007). Another feature of the ICEBs1 transposon is the formation of repeated nucleotide sequences (Wozniak & Waldor, 2010). This is similar to the action of *L. lactis* Tn6098 (Machielisen, Siezen, van Hijum, & Vlieg, 2011). Using Tandem Repeat Finder (Benson, 1999), we identified a 21 bp repeat (ACTATAATAAAAGATGGAGTG) between llkf_1225 and llkf_1226 in KF147 (and NCDO 2118), which might have been the insertion site for the NRPS/PKS (Figure 1). Notably, the surrounding hypothetical proteins do not share homology with other ICEBs1 genes and the G+C contents of the genes in the locus (35.3%) do not differ from the chromosome (34.9%). Therefore, these genes may have been acquired by horizontal gene transfer from a related bacterial species and constitute a genomic island. Alternatively, the NRPS/PKS system and flanking genes might have been present in an ancestral lactococcal genome and subsequently lost during adaptation to new habitats such as milk.

The same NRPS/PKS system is also present in the published genomes of *L. lactis* strains NCDO 2118 (completed, isolated from frozen peas) (Oliveira et al., 2014) and YF11 (draft, isolated from fermented corn) (Cavanagh, Fitzgerald, & McAuliffe, 2015; Du et al., 2013). Genes coding for the NRPS/PKS system were also detected in *L. lactis* Li-1, KF134, KF146, and KF196 according to comparative genome hybridization (Siezen et al., 2011) and confirmed by BLASTp analysis (data not shown). Each of these strains was isolated from (fermented) plant foods and are classified to the *L. lactis* subspecies lactis. Due to the high sequence conservation (>99%) amongst the NRPS/PKS systems in these strains, we expect them to synthesize the same secondary metabolite.

Inspection of the genes flanking the NRPS/PKS locus in *L. lactis* strains KF147 and NCDO2118 showed that they are inserted relative to other *L. lactis* strains between genes encoding acetolactate synthase (als, upstream) and two hypothetical LpxTG membrane proteins (lpxAB) followed by the kdpDEABC potassium transport system (downstream) (Figure 1 and Table 1). *L. lactis* strains that lack the NRPS/PKS system but share the same flanking gene arrangement as KF147 were isolated from other sources including strain IO-1 (a kitchen sink isolate) (Kato et al., 2012), CV56 (a vaginal isolate) (Gao et al., 2011), and KLDS 4.0325 (a dairy isolate) (Yang, Wang, & Huo, 2013) (Figure 1). Importantly, even within the *lactis* subspecies, there is considerable variation in the NRPS/PKS flanking regions because not all strains display this genetic composition. In this regard, the dairy-associated
### TABLE 1  BLASTp results of *L. lactis* KF147 NRPS/PKS genomic island

| Gene          | Length (bp) | Accession          | Annotation                          | Organism               | Annotation                          | Similarity (%) | Coverage (%) |
|---------------|-------------|--------------------|-------------------------------------|------------------------|-------------------------------------|----------------|--------------|
| *als*         | 1665        | WP_012897772.1     | Acetolactate synthase               | *L. lactis*            | Acetolactate synthase               | 99             | 100          |
| Ilkf_1209     | 198         | WP_012897773.1     | Hypothetical protein                | *Virgibacillus pantothenticus* | Hypothetical protein               | 53             | 92           |
| Ilkf_1210     | 243         | WP_012897774.1     | Hypothetical protein                | No hits                |                                     |                |              |
| npkK          | 1776        | WP_012897775.1     | Histidine kinase                    | *S. mutans*             | Histidine kinase                    | 65             | 99           |
| npkR          | 603         | WP_012897776.1     | DNA-binding response regulator      | *S. mutans*             | Two-component system response regulator | 78             | 100          |
| npkA          | 17136       | WP_012897777.1     | Nonribosomal peptide synthetase     | *S. mutans*             | Nonribosomal peptide synthetase     | 69             | 99           |
| pksA          | 1266        | WP_012897778.1     | PKS acyl-CoA transferase            | *S. mutans*             | PKS acyl-CoA transferase            | 74             | 100          |
| pksB          | 4533        | WP_012897779.1     | PKS biosynthesis protein            | *S. mutans*             | PKS biosynthesis protein            | 68             | 99           |
| pksC          | 4716        | WP_012897780.1     | PKS biosynthesis protein            | *S. mutans*             | PKS biosynthesis protein            | 73             | 100          |
| Ilkf_1217     | 861         | WP_012897781.1     | ABC transporter, ATP-binding protein | *S. mutans*             | ABC transporter ATP-binding protein | 84             | 99           |
| Ilkf_1218     | 738         | WP_012897782.1     | MFS transporter permease            | *S. mutans*             | ABC transporter permease            | 78             | 100          |
| npkS          | 6441        | WP_012897784.1     | Polyketide synthase                 | *S. mutans*             | Polyketide synthase                 | 67             | 99           |
| npkB          | 696         | WP_012897785.1     | Thiosterase                         | *S. mutans*             | Thiosterase                         | 72             | 100          |
| npkC          | 669         | WP_012897786.1     | 4′-phosphopantetheinyl transferase  | *S. mutans*             | 4′-phosphopantetheinyl transferase  | 70             | 100          |
| Ilkf_1223     | 192         | WP_012897787.1     | Hypothetical protein                | No hits                 |                                     |                |              |
| Ilkf_1224     | 945         | WP_012897788.1     | Macrolide biosynthetic protein, AvrD family | *S. mutans*             | Macrolide biosynthetic protein, AvrD family | 61             | 96           |
| Ilkf_1225     | 1506        | WP_012897789.1     | Phytoene dehydrogenase              | *Bacillus thuringiensis* | Hypothetical protein               | 47             | 99           |
| Ilkf_1226     | 651         | WP_012897790.1     | Hypothetical protein                | *Enterococcus sp.*     | ATP-binding protein                 | 38             | 95           |
| Ilkf_RS06220  | 372         | WP_017864463.1     | Hypothetical protein                | *Burkholderia pseudomallei* | Hypothetical protein               | 33             | 36           |
| Ilkf_1230     | 1350        | WP_012897791.1     | DNA integration/ recombination/inversion protein | *Enterococcus durans* | Integrase                           | 33             | 91           |
| Ilkf_1231     | 1818        | WP_017864465.1     | Hypothetical protein                | *Enterococcus faecalis* | Hypothetical protein               | 34             | 97           |
| LLK_RS06235   | 1677        | WP_03603395.1      | Hypothetical protein                | *Paenibacillus polymyxa* | Hypothetical protein               | 31             | 98           |
| Ilkf_RS06240  | 1851        | WP_03603397.1      | Hypothetical protein                | *Desulfobulbus japonicus* | Hypothetical protein               | 32             | 96           |
| Ilkf_1234     | 135         | WP_017864467.1     | Hypothetical protein                | *L. lactis*             | Transposase                         | 85             | 70           | (Continues)
L. lactis strain IL1403 does not share the same gene arrangement. In IL1403, als is located on a distal part of the chromosome and this strain lacks kdpDEABC.

3.2 | The L. lactis NRPS/PKS is homologous to S. mutans TnSmu2

Protein alignments of the L. lactis KF147 NRPS/PKS system revealed a high degree of amino acid similarity (65%–84%) and coverage (99%–100%) to an NRPS/PKS system in multiple strains of Streptococcus mutans on the TnSmu2 genomic island (Table 1) (Wu et al., 2010). The L. lactis KF147 NRPS/PKS system is most similar to S. mutans LJ23, a strain for which the complete genome sequence is available (Aikawa et al., 2012). None of the genes downstream of llkf_1224 had homologs in S. mutans, with the exception of the gene encoding the putative macrolide biosynthetic protein AvrD (llkf_1224). This gene is in the same location relative to the NRPS/PKS system in S. mutans LJ23 and shares 61% amino acid identity with 96% coverage (Table 1).

Although the function of the NRPS/PKS in S. mutans LJ23 has not been directly investigated, the same system in S. mutans UA140 was recently characterized (Wu et al., 2010). This NRPS/PKS product was the first to be characterized for a LAB species and shown to confer pigmentation and improve oxygen and hydrogen peroxide stress tolerance (Wu et al., 2010). The S. mutans UA140 NRPS/PKS locus is identical to that of S. mutans LJ23 (personal communication, Fengxia Qi) and therefore is also conserved with L. lactis KF147 (Figure 1 and Table 1).

3.3 | ATL and paraquat in combination delay growth of the NRPS/PKS deletion mutant

We previously found significant increases in L. lactis KF147 NRPS/PKS gene transcript levels during growth in Arabidopsis leaf tissue lysate (ATL) compared to in GM17 (Golomb & Marco, 2015). Therefore, we investigated the importance of the NRPS/PKS system in conferring optimal growth rates to L. lactis in ATL by constructing strain BAL1, a markerless, nrpC deletion mutant of L. lactis KF147. The nrpC gene was targeted for deletion because it encodes the PPTase responsible for transferring the 4′-phosphopantetheine moiety to the carrier protein domain of nonribosomal peptide synthetases and polyketide synthases. Removal of the PPTase prevents substrate attachment and synthesis of the NRPS/PKS product (Beld, Sonnenschein, Vickery, Noel, & Burkart, 2014). Static incubation in ATL revealed that the NRPS/PKS system did not provide a specific advantage to L. lactis during exponential phase growth in this medium, as both KF147 and BAL1 exhibited the same growth rates and no differences in final cell yields were observed (data not shown).

Because the NRPS/PKS system from L. lactis KF147 is homologous to a system in S. mutans strains UA140 and LJ23, we reasoned that the requirement for the NRPS/PKS product could depend on the extent to which reactive oxygen species are released by the plant tissues. To study the possible function of the NRPS/PKS system in tolerance...
of oxidative stress on plants, we supplemented ATL with 10 mmol/L paraquat, a redox cycling compound that generates superoxide radicals in the presence of reducing agents (Lock & Wilks, 2010). Under those conditions, both KF147 and the ΔnrpC mutant BAL1 exhibited no growth for 24 hr (Figure 2). By 72 hr, wild-type KF147 had grown to high cell densities, whereas BAL1 viable cell numbers declined by 10-fold (Figure 2). These results indicate that paraquat undergoes redox cycling by reducing agents in ATL and generates superoxide radicals against which the NRPS/PKS product provides protection.

3.4 | *L. lactis* KF147 NRPS/PKS provides limited advantages during growth in standard laboratory culture medium

No differences in growth rate or final cell density were observed between *L. lactis* KF147 and BAL1 during static incubation in GM17 at 30°C, the standard laboratory culture medium and growth conditions for this species (data not shown). Moreover, unlike *S. mutans* UA140, there was no visible change in cellular pigmentation between KF147 and the ΔnrpC mutant.

To provide a more oxidative environment, KF147 and the BAL1 mutant were grown in the presence of paraquat. Wild-type KF147 and the ΔnrpC mutant BAL1 were both unaffected by the presence of paraquat in GM17 (data not shown). In conditions conducive for *L. lactis* respiratory metabolism (i.e. hemin supplementation and aeration), sufficient reducing agents (e.g. NADH dehydrogenases NoxA and NoxB) might be present and could induce paraquat redox cycling and subsequent superoxide formation. Under respiratory conditions without paraquat, no change in growth was observed between KF147 and BAL1 (Figure 3a) whereas in the presence of paraquat, both strains exhibited a biphasic growth pattern (Figure 3b). During the first 2.5 hr, both strains exhibited significantly decreased growth rates (μ = 0.744 ± 0.024 hr⁻¹ and μ = 0.671 ± 0.016 hr⁻¹ for KF147 and BAL1, respectively) (Figure 3b) compared to exposure to respiratory conditions in the absence of this compound (p < .0005 for both) (Figure 3a). The growth impairment was greater for BAL1 than for wild-type KF147 in the presence of paraquat (p < .0005). After 2.5 hr, the growth rates of both strains slowed considerably (μ = 0.225 ± 0.013 hr⁻¹ and μ = 0.144 ± 0.008 hr⁻¹ for KF147 and BAL1, respectively) and the deletion mutant was again more impaired (p < .0005) (Figure 3b). Similarly, exposure of *L. lactis* KF147 and BAL1 to the superoxide-generating reagents xanthine oxidase (XO) and xanthine under static conditions in GM17 also resulted in a subtle but significant decrease in growth rate for BAL1 compared to KF147 (Table 2).

The addition of H₂O₂ to GM17 also resulted in a significantly decreased growth rate for BAL1 as compared to KF147 (Table 2). Both *L. lactis* and *S. mutans* lack catalase (Price, Zeyniyev, Kuipers, & Kok, 2011), and therefore the NRPS/PKS in these strains appears to also provide an alternate mechanism for protection against H₂O₂. Overall, although the growth impairment of BAL1 was significant under oxidative conditions in GM17, the effect was not as great as in ATL. This difference might have been due to the relatively reduced expression of the NRPS/PKS gene cluster in GM17 and/or to a lack secondary metabolite precursors in the laboratory culture medium.

![Figure 2](image1.png)  
**FIGURE 2** Growth of *L. lactis* KF147 and BAL1 in ATL with paraquat. *L. lactis* KF147 and BAL1 were inoculated into 10 ml ATL containing 10 mmol/L paraquat and incubated at 30°C without aeration. The average ± SD of three replicate cultures is shown.

![Figure 3](image2.png)  
**FIGURE 3** Growth of *L. lactis* KF147 and BAL1 in GM17 under respiratory conditions with and without paraquat. *L. lactis* KF147 and BAL1 were inoculated into GM17 under respiratory conditions (continuous aeration and 10 μg/ml hemin) (a) or respiratory conditions and 20 mmol/L paraquat (b), Bacteria were incubated at 30°C in microtiter plates and OD was measured at 600 nm. The average ± SD of at least eight replicate cultures is shown.
TABLE 2 Growth rates (hr⁻¹) of L. lactis KF147 and BAL1 exposed to ROS

| ROS     | KF147       | BAL1       | p     |
|---------|-------------|------------|-------|
| Xanthine & XO | 0.581 ± 0.036 | 0.535 ± 0.009 | .022  |
| H₂O₂   | 0.465 ± 0.017 | 0.386 ± 0.026 | <.0005|

4 | CONCLUSIONS

Although numerous LAB species are annotated to contain NRPS/PKS systems, few have been biochemically characterized and these are limited to Lactobacillus reuteri TMW1.656 (Lin et al., 2015) and S. mutans UA159 (Joyner et al., 2010), and UA140 (Wu et al., 2010). Remarkably, the L. lactis KB147 NRPS/PKS system is similar in both gene structure and function to the NRPS/PKS in S. mutans UA140 and LJ23 and not to orthologous systems in other lactic acid bacteria. Both of the latter species are members of the Streptococcaceae, however, they inhabit distinct ecological niches and it is unclear as to in which species the NRPS/PKS system originated (or perhaps as a third as of yet unknown species). In addition to the NRPS/PKS system, the genetic locus also contains a gene annotated as avrD (likf_1224). It is notable that this protein is also produced by certain plant pathogens and is recognized by plant host resistance (R) proteins to result in plant production of an oxidative burst and generation of superoxide radicals (Lamb & Dixon, 1997; Spoel & Dong, 2012). This finding is consistent with our results showing that the production of the NRPS/PKS appears to be dispensable for L. lactis during growth in standard laboratory media, but relevant to the plant environment, possibly either to counter responses to pathogenic bacteria (oxidative burst) or to survive generally, aerobic conditions on leaf tissues. In conclusion, tolerance to ROS compounds due to expression of an NRPS/PKS product represents a novel strategy for L. lactis ROS detoxification, possibly during growth on living plants, and with potential application to support the viability of this organism in culture production and preservation.

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CONFLICT OF INTEREST

None declared.

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Additional Supporting Information may be found online in the supporting information tab for this article.