## Comparative analysis of pSMA198 found in *Streptococcus macedonicus* ACA-DC 198, the first streptococcal plasmid of the pCI305/pWV02 family of theta-replicating replicons

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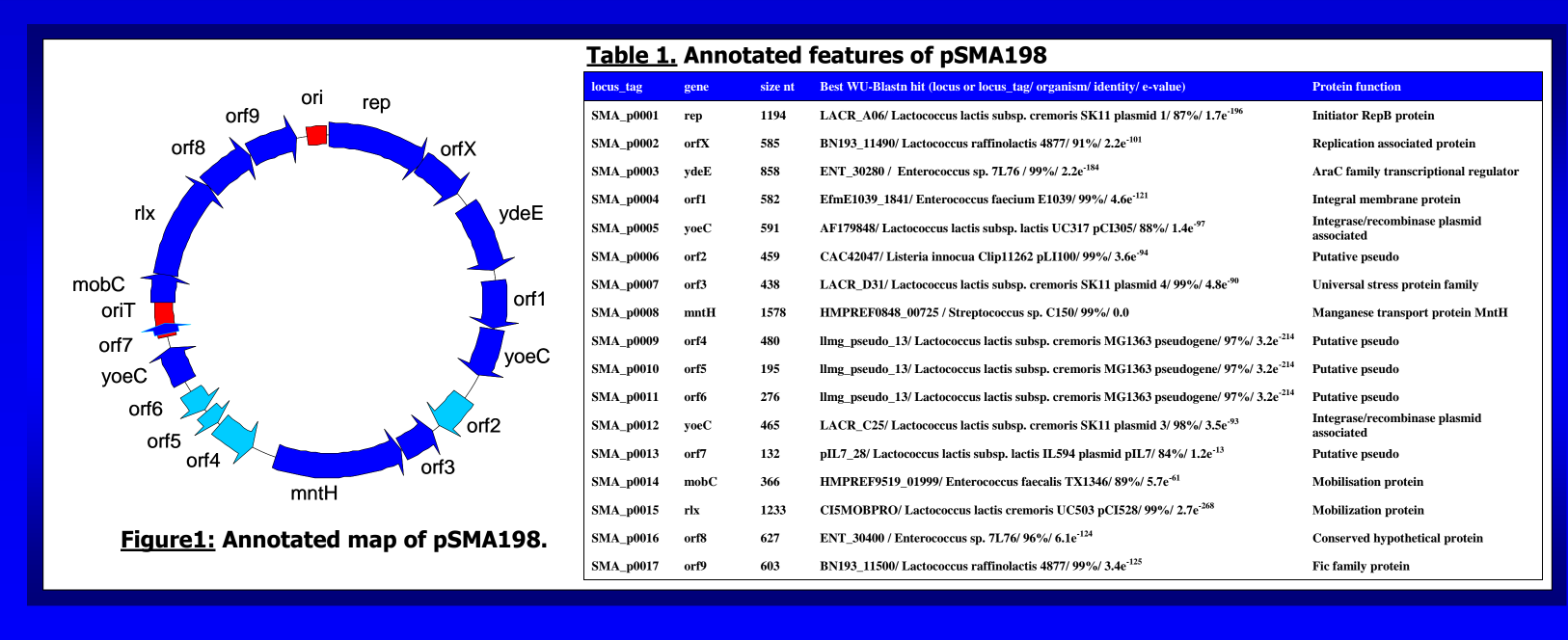
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## **Abstract**

Here we analyze pSMA198, the first plasmid isolated from Streptococcus macedonicus ACA-DC 198, and we attempt to clarify the route of its original acquisition. Based on the similarity profiles of the plasmid's replication initiation protein (Rep) and its origin of replication (ori), pSMA198 was found to be a novel member of the pCI305/pWV02 family of theta-replicating plasmids. The pCI305/pWV02 family consists of plasmids of narrow host range that are mainly found in lactococcal species. Comparative analysis of the pSMA198 revealed a high degree of similarity with plasmids pSK11b, pVF22 and pIL5 over its replication backbone, its mobilization backbone and most of its length, respectively. All these three plasmids have been isolated from Lactococcus lactis strains deriving from milk or its products supporting that S. macedonicus acquired pSMA198 from the latter species and that this acquisition took place in the dairy environment. Both pSMA198 and the chromosome of S. macedonicus exhibit a high degree of pseudogenes, indicating that they must have evolved under the same gene decay processes. Furthermore, we were able to determine chromosomal regions that may have originated from pSMA198, also supporting a long co-existence of the two replicons. In addition, pSMA198 is carried by S. macedonicus strains segregated in five different genotypes by pulsed-field gel electrophoresis (PFGE), showing that pSMA198's acquisition is not a recent event. We propose that our overall analysis of pSMA198 points towards the habituation of S. macedonicus ACA-DC 198 to the dairy environment.

## **Results and Discussion**

Streptococcus macedonicus ACA-DC 198 carries a novel plasmid of 12,728 bp assigned as pSMA198. The plasmid has a 35.0% G+C content, lower than that of the S. macedonicus chromosome (37.6%), indicating that it may have been acquired from another organism. Overall 17 CDSs were annotated on pSMA198 (Fig.1 and Table 1).



The first gene codes for a replication initiation protein (Rep). The rep gene showed 87% identity (e-value 1.7e-196) with the respective gene found on plasmid 1 of L. lactis subsp. cremoris SK11. Among the WU-Blast hits of Rep we identified the RepB proteins of the pCI305 and the pWV02 plasmids  $(78\% identity, e-value 7.4e^{-162} and 75\% identity, e-value 2.6e^{-152}, respectively) that are the prototypes of the pCI305/pWV02 family of the lactococcal$ theta-replicating plasmids. Multiple sequence alignment of the top Rep hits including the RepB proteins of pCI305 and pWV02 with ClustalW revealed a high degree of conservation over most of their length (Fig. 2).

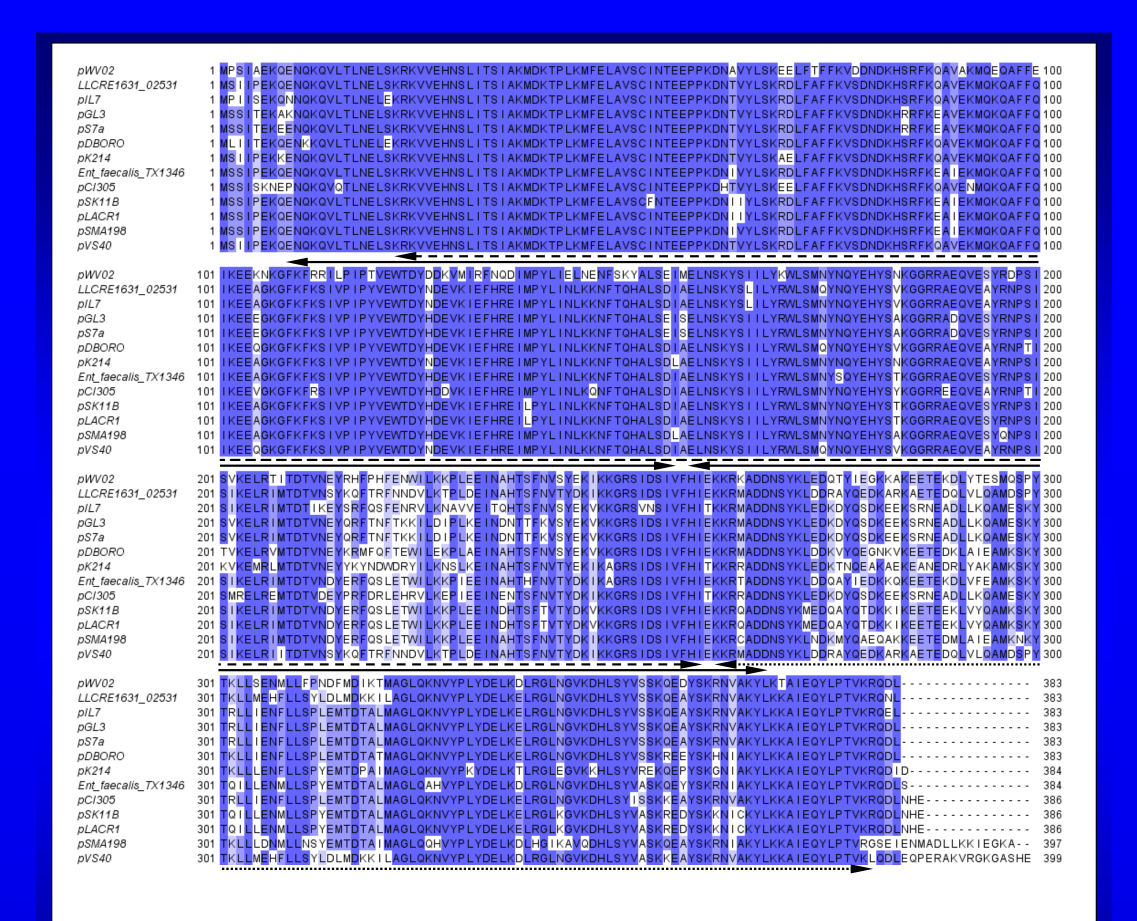


Figure 2: Multiple sequence alignment of RepB sequences relevant to the respective protein of pSMA198 performed with ClustalW. Double-headed arrows indicate the positions of the sequence signatures within the proteins as determined by InterProScan as follows: initiator Rep protein (dashed line), L. lactis RepB C-terminal (dotted line) and two consecutive winged helix-turn-helix transcription repressor DNA-binding (solid line).

Subsequently, we looked upstream of the rep gene in an attempt to identify the origin of replication (ori) of pSMA198. WU-Blastn similarity searches and multiple sequence alignment directed us towards a pCI305/pWV02 type of ori (Fig. 3). Indeed, we determined a segment spanning 242 nucleotides that contains an AT-rich region, three and a half direct repeats (DRs) of 22-bp iterons and two inverted repeats (IRs). This structure includes all the necessary elements for the binding of the required protein complexes, the melting of DNA, the initiation of the replication and the control of PCN. The pattern of the pSMA198 ori along with the similarity of its Rep with the lactococcal RepB shows that pSMA198 is certainly a member of the narrow host range pCI305/pWV02 family of replicons, which are normally found in *Lactococcus* species.

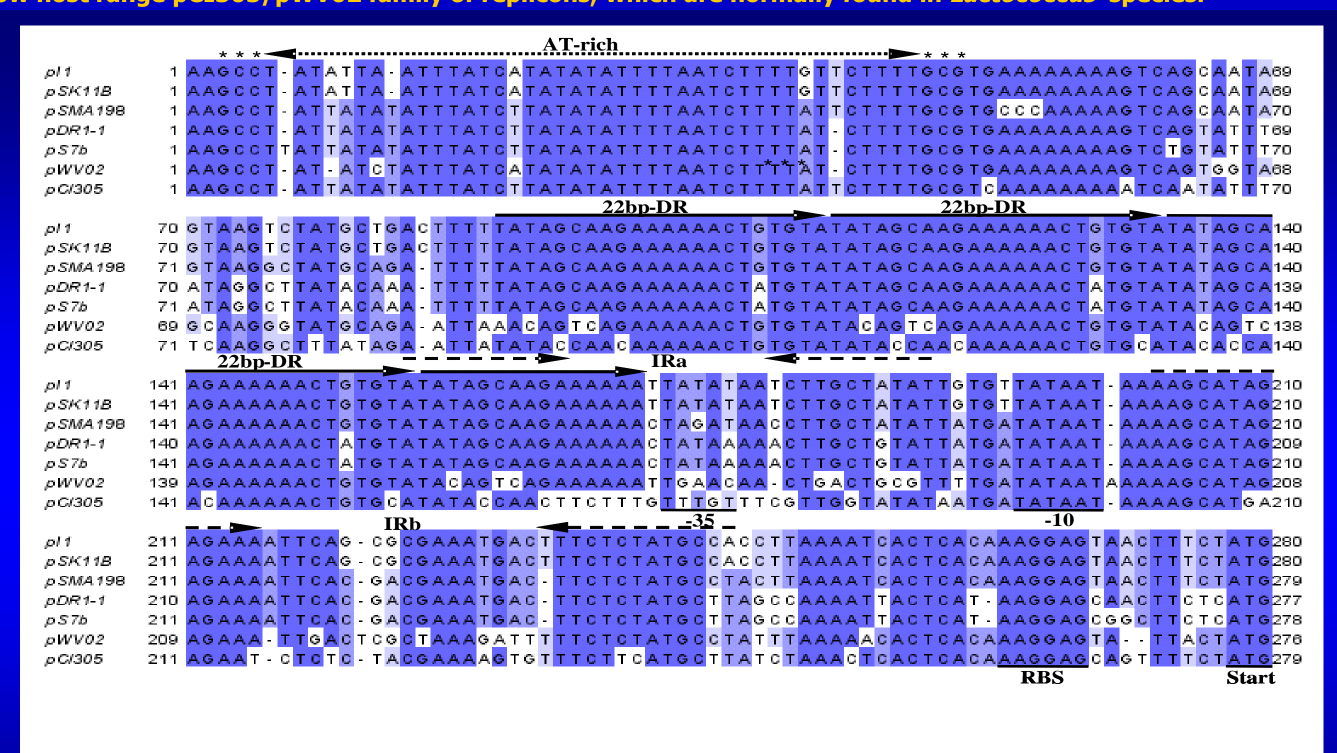
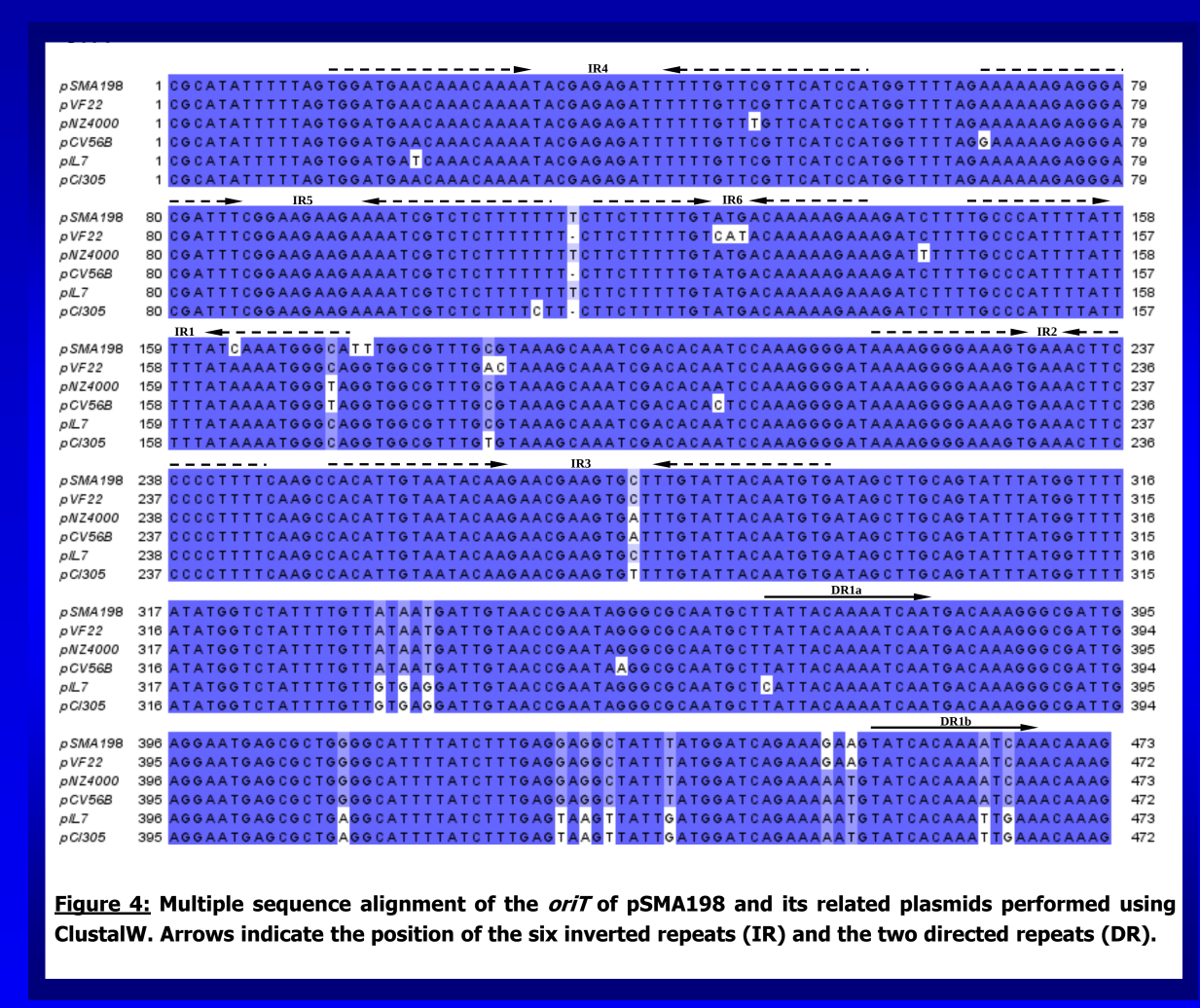


Figure 3: Multiple sequence alignment of the *ori* of pSMA198 and its related plasmids performed using ClustalW. Arrows indicate the position of the AT-rich region, the 22-bp direct repeat (DR) iterons and the two inverted repeats (IR). The promoter (-35, -10 and the RBS) and the start codon of the rep gene are underlined.

Even though pSMA198 is not a self-transmissible plasmid, a cis-acting origin for transfer (oriT) that would allow its mobilization in the presence of a true conjugative plasmid was also predicted upstream of mobC (Fig. 4). The oriT exhibited a region of six consecutive IRs and two DRs. In addition we determined an identical nick site to those proposed previously for plasmids pS7a and pS7b, eight bases after the end of IR3. The nick site is where the mobilization nucleases cleave duplex DNA during transfer. Once more, these structures were highly conserved among several lactococcal plasmids including pCI305.



The relation of pSMA198 to other plasmids was further investigated (Fig. 5). pSMA198 showed highest identity over the entire length of its replication (ori-rep-orfX) and mobilization (oriT-mobC-rlx-orf8-orf9) backbones to plasmids pSK11b (78% identity, e-value 8.4e-253) and pVF22 (96% identity, e-value 0.0), respectively. Plasmid pSK11b has been isolated from L. lactis subsp. cremoris SK11, which is a widely used industrial starter in cheese making, and plasmid pVF22 has been isolated from the raw milk cheese strain L. lactis subsp. lactis biovar. diacetylactis DPC3901. Interestingly, the similarity between pSMA198 and each of the two plasmids mentioned above was basically restricted to the loci under investigation (i.e. the replication or the mobilization backbones) (Fig. 5). This led us to look for the plasmid that would have the highest identity with the complete sequence of pSMA198. The plasmid identified was pIL5 that has been isolated from L. lactis subsp. lactis IL594, which is also a cheese starter. pIL5 exhibited 97% identity (e-value 0.0) over approximately the three quarters of the pSMA198 sequence (Fig. 5). It should be emphasized that apart from the closest similarity hits mentioned above, the overriding majority of top hits in all similarity searches for the different features annotated on pSMA198 at the protein or nucleotide level originated from L. lactis dairy strains. For example, nine out of the ten top hits for the replication backbone derived from strains isolated from milk or its products.

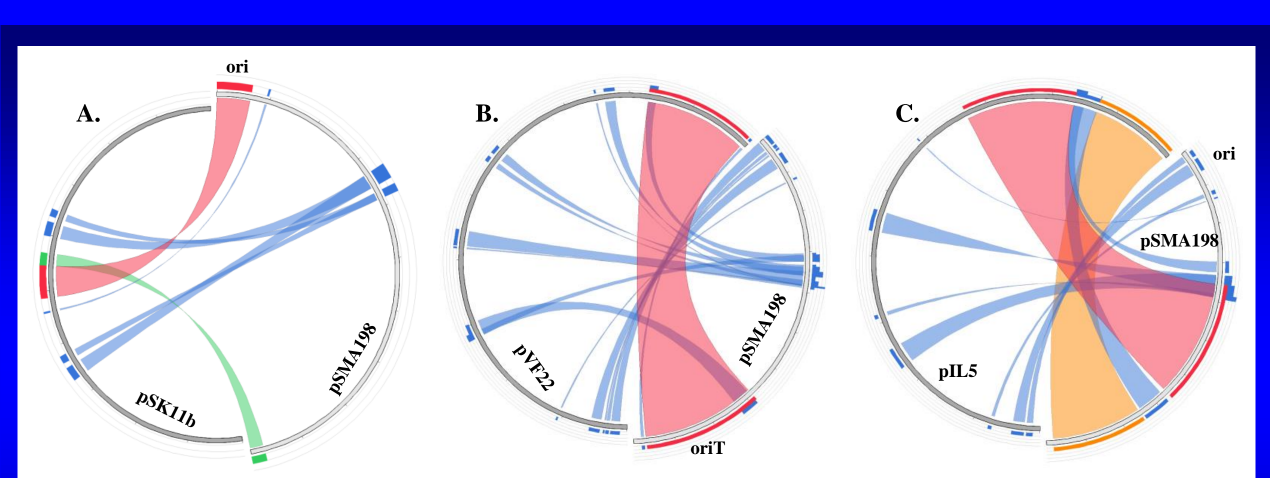
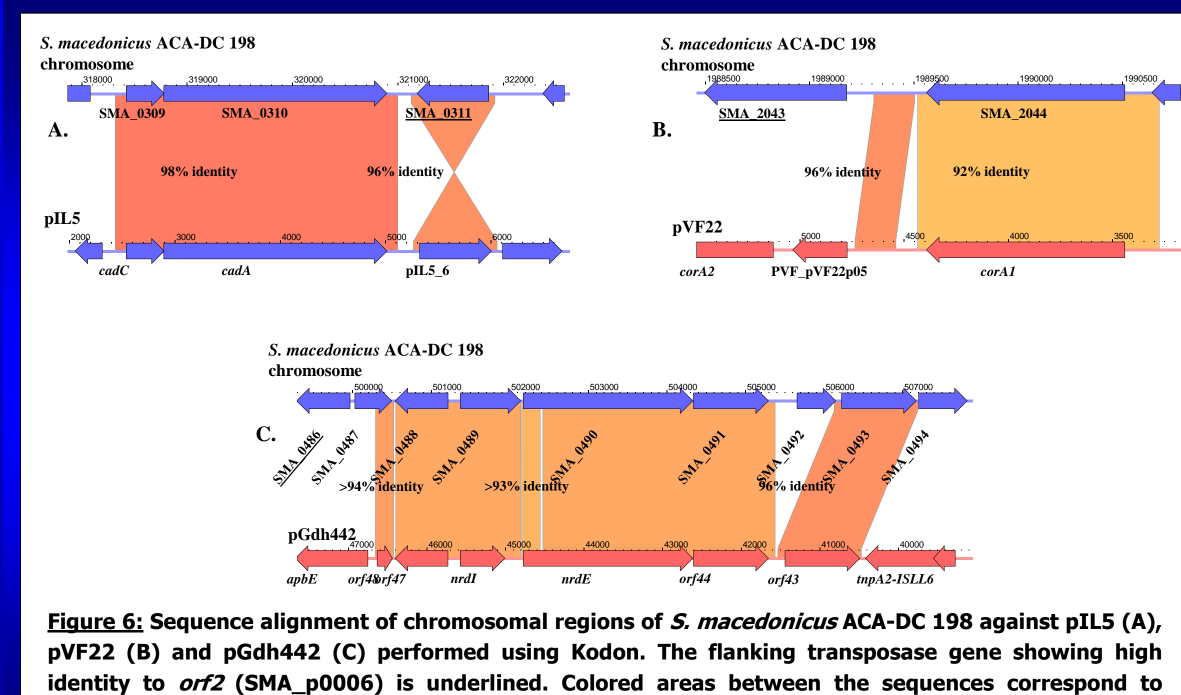


Figure 5: Sequence alignment in a circular fashion of pSMA198 against the lactococcal pSK11b (A), pVF22 (B) and pIL5 (C) of dairy origin. Local alignments produced by BLAST are presented using ribbons whose color corresponds to four quartiles of the alignment's bitscore (red: top 25%, orange: second 25%, green: third 25% and blue: worst 25%). In order to aid orientation, the position of the *ori* or *oriT* of pSMA198 has been added in the figures.

The possibility of genetic exchange between pSMA198 and the chromosome of S. macedonicus ACA-DC 198 was also examined. We were led to this hypothesis due to the existence of genes or pseudogenes of elements that would facilitate such an exchange in both the plasmid and the chromosome. In addition, the reduced size of pSMA198 as compared to some of its related plasmids, e.g. pVF22 and pIL5 (12 kb vs. 22 and 23 kb, respectively) indicates loss of genetic material, some of which could have moved to the chromosome. In the attempt to identify such regions, we employed two different strategies. We investigated for the presence of chromosomal genes showing high identity to the genes found on pSMA198 or its related plasmids (Fig. 6).



different levels of identity that is depicted within the areas

Our findings demonstrate that pSMA198 is a novel member of the pCI305/pWV02 family of theta-replicating plasmids. The pCI305/pWV02 replicon has been shown to be of narrow host range, mainly replicating in Lactococcus spp. pSMA198 is the first streptococcal plasmid to be described within this family. Based on the levels of identity of the pSMA198 replication backbone, which may reflect its evolutionary history, to the respective backbones of other plasmids, our data supports that S. macedonicus acquired pSMA198 from L. lactis. This exchange took place most probably in the milk environment as the overriding majority of closest related plasmids to pSMA198 (i.e. pSK11b, pVF22 and pIL5) are of dairy origin. This is in agreement with the isolation source of S. macedonicus ACA-DC 198 and the fact that S. macedonicus strains have dairy products as their primary ecological niche. The acquisition of pSMA198 by S. macedonicus ACA-DC 198 seems not to be a recent event. Prominently, the fact that the chromosome of S. macedonicus ACA-DC 198 and pSMA198 exhibit a high percentage of pseudogenes, indicates that they may have both evolved under the same gene decay processes. In addition, the potential exchange of genetic material between the chromosome and the plasmid designates a long co-existence of the two replicons. We propose that our overall analysis of pSMA198 points towards the habituation of S. macedonicus ACA-DC 198 to the dairy environment.

## **Bibliography**

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• Papadimitriou K., S. Ferreira, N. C. Papandreou, E. Mavrogonatou, P. Supply, B. Pot, and E. Tsakalidou (2012) Complete genome sequence of the dairy isolate Streptococcus macedonicus ACA-DC 198. J Bacteriol 194:1838-9. • Papadimitriou K., T. Plakas, R. Anastasiou, S. Ferreira, P. Supply, P. Renault, N. C. Papandreou, B. Pot, and E.Tsakalidou (In preparation) Analysis of the lactococcal plasmid pSMA198 found in Streptococcus macedonicus ACA-DC 198 points towards the habituation of the strain to the dairy environment.