

Licensing opportunity



Single plasmid systems for inducible dual protein expression and gene regulation in lactic acid bacterium *Lactococcus lactis* for microbial cell factories in industrial producing proteins

Field of use

Microbiology, molecular biology

Current state of technology

Stage of Development:
Available for demonstration

IPR status Know-how

Publication TBA

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Background

Tools for recombinant protein expression have been relatively well developed. *L. lactis* is therefore comparable to other well established bacterial expression systems, such as *Escherichia coli* and *Bacillus subtilis*. Advanced techniques for genetic engineering are required to develop *L. lactis* further as a microbial cell factory. Simultaneous expression of two or more proteins is beneficial for various applications, including the expression of multi-subunit proteins, the use of *L. lactis* as a mucosal delivery vehicle or as a multistep biocatalyst.

Description of the Invention

Here, plasmids for co-expression of two recombinant proteins in *L. lactis* have been developed and their effectiveness assessed by the expression of model proteins. Plasmids were further upgraded and a single plasmid CRISPR-Cas9 system has been developed. Duplication of the nisin promoter enabled the balanced, inducible expression of two model proteins in *L. lactis*, thus constituting a new tool for recombinant protein expression in this organism. A similar strategy resulted in a single plasmid CRISPR-Cas9 system that can be used, among other possible applications, for plasmid curing or CRISPRi-mediated gene regulation in *L. lactis*.

Plasmids will be applied in the future research in *L. lactis* for concomitant expression of therapeutic and reporter proteins, as well as for plasmid curing and gene silencing.

Main Advantages

- New tool for recombinant protein expression in *L. lactis* was developed.
- Duplication of the nisin promoter enabled the balanced, inducible expression of two model proteins in *L. lactis*.



