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Supplementary Data

Identification of dual receptor binding protein systems in lactococcal 936 group phages

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Supplementary Table S1. Measurements of morphological features of the examined phages.

	Head diameter	Tail length incl. bp	Tail width	(Total) bp length	(Total) bp width
p2	54.2 ± 1.9 (n=23)	160.6 ± 5.9 (n=23)	12.9 ± 0.7 (n=23)	12.7 ± 1.3 (n=23)	18.1 ± 1.0 (n=23)
4.2	55.4 ± 2.6 (n=19)	158.6 ± 4.0 (n=19)	12.9 ± 0.7 (n=19)	18.6 ± 1.6 (n=12)	17.0 ± 1.2 (n=12)
4R15L	55.4 ± 1.7 (n=17)	159.6 ± 5.0 (n=17)	12.8 ± 0.7 (n=17)	19.8 ± 1.4 (n=11)	17.0 ± 1.3 (n=11)
4R16L	54.0 ± 2.1 (n=20)	158.5 ± 5.6 (n=20)	12.3 ± 0.6 (n=20)	19.0 ± 2.3 (n=8)	17.2 ± 1.1 (n=8)

All measurements are presented in nm.

Supplementary Table S2. Oligonucleotides used in this study.

Oligonucleotide	Sequence	Target
RBP2F	agcagcccatggcacaccatcaccatcaccattcttctgtataaataaatactttttcagtc	Forward primer for cloning of <i>rbp2</i> in pNZ8048
RBP2R	agcagcaagcttttttaataaaagtagcttgcg	Reverse primer for cloning of <i>rbp2</i> in pNZ8048
RBP1F	agcagcccatggcgccaccatcaccatcaccattcttctgtatacaaaaatacgttttttagtcc	Forward primer for cloning of <i>rbp1</i> in pNZ8048
RBP1R	agcagctctagattacttgctagcagctcctccc	Reverse primer for cloning of <i>rbp1</i> in pNZ8048 and pTX8048
RBP1pTXF	agcagcggatccatgacgataacaaaatacgc	Forward primer for cloning of <i>rbp1</i> in pTX8048
RBP1pETMF	agcagcccatggcaataaataaatactttttcagtc	Forward primer for cloning of <i>rbp1</i> in pETM11
RBP1pETMR	agcagcggatccttacttgctagcagctcctccc	Reverse primer for cloning of <i>rbp1</i> in pETM11
BpF	agcagcccatggaaggaggcgtaatgcaccatcaccatcaccattcagtaagacagtataaaat	Forward primer for cloning of the baseplate region in pETM11
BpR	aggaggggatccttatttaataaagtagcttgc	Reverse primer for cloning of the baseplate region in pETM11
ΔRBP2R	aggaggggatccttacttgctagcagctcctccc	Reverse primer for cloning of the ΔRBP2 construct in pETM11
ΔRBP1R _i	gtatatttatttggcatttacatctctctttctac	Internal reverse primer used for the construction of the ΔRBP1 construct via SOEing PCR

Δ RBPF ₂	tagaaaaggaagatatgtaaatggcaataataataatatac	Internal forward primer used for the construction of the Δ RBPF1 construct via SOEing PCR
Δ HPR ₁	cgtatattttgtatcgtcattttatcctctattcccctcataaagg	Internal reverse primer used for the construction of the Δ HPR construct via SOEing PCR
Δ HPF ₂	cctttatggaggggaatagaggataaaatgacgataacaaaatatacg	Internal forward primer used for the construction of the Δ HPR construct via SOEing PCR
DitR	aggaggggatcctaaataaaatcaacttcttttg	Reverse primer for the cloning of the <i>dit</i> gene in pETM11
DitTalHPR	aggaggggatcctacatactctctttctacaatttgagc	Reverse primer for the cloning of the Dit and Tal complex in pETM11
TalF	agcagcccatggaaggaggcgtaatgcaccatcaccatcaccattggcagaatataattatag	Forward primer for cloning of the <i>tal</i> gene in pETM11
TalR	aggaggggatccttatcctctattcccctcata	Reverse primer for cloning of the <i>tal</i> gene in pETM11
pNZ8048F	caggagaaggacgatagca	Forward checking primer for pNZ8048 and pTX8048
pNZ8048R	tcttcttattctcgctttg	Reverse checking primer for pNZ8048 and pTX8048
pETM11F	gattacgacatcccactactg	Forward checking primer for pETM11 and pETM30
pETM11R	cgggctttgtagcagccggatc	Reverse checking primer for pETM11 and pETM30
pQE30F	cagggttattgtctcatgagcg	Forward checking primer for pQE30
pQE30R	cagctcaccgtctttcattgcc	Reverse checking primer for pQE30

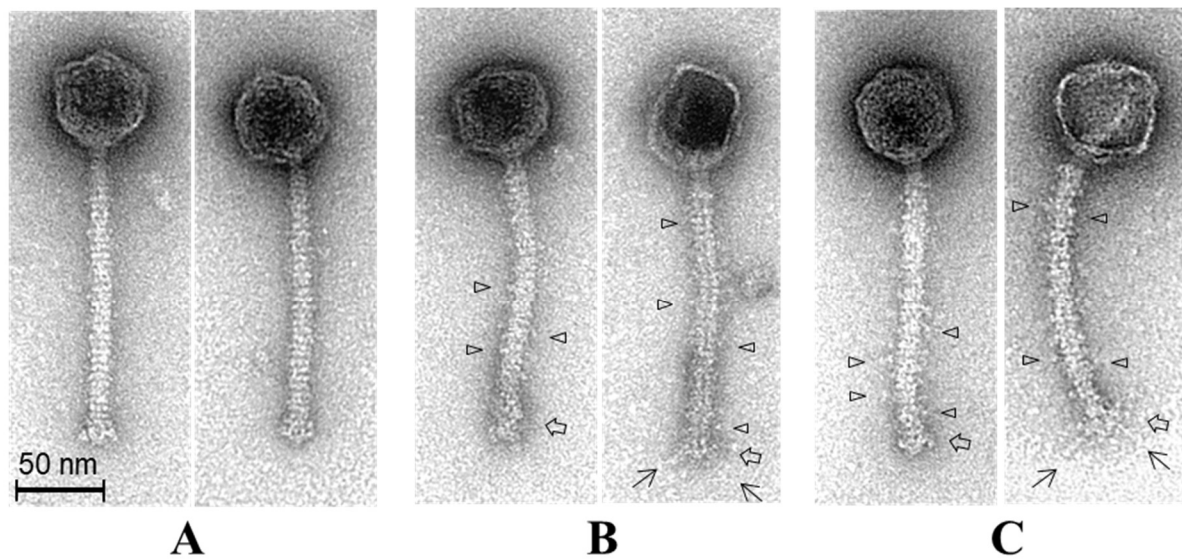


Figure S1. Representative micrographs of phages p2 (A), PhiR15L (B) and PhiR16L (C) stained with 2% uranyl acetate. ⇨ highlights the enlarged baseplate of phages PhiR15L (B) and PhiR16L (C). ▷ highlights some of the globular appendages which appear to coat the tail of the phages PhiR15L (B) and PhiR16L (C). PhiR15L and PhiR16L phage particles with empty heads (particles on right side in B & C) also show elongated appendages protruding from the baseplate (indicated with →).

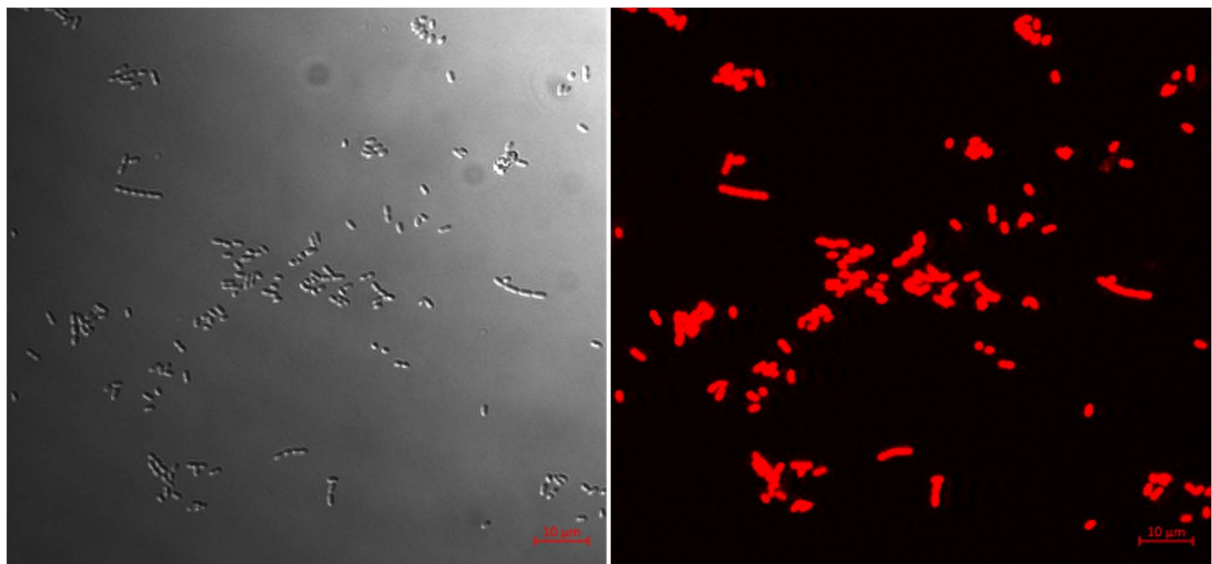


Figure S2. Fluorescent binding assays using mCherry tagged RBP2_{Phi4.2}. Protein was added at a concentration of 50 μg/ml. Scale bars correspond to 10 μm. Cells were visualized using differential interference contrast (DIC) microscopy (panel on the left), and fluorescent confocal microscopy (panel on the right) at the mCherry excitement wavelength of 514 nm.

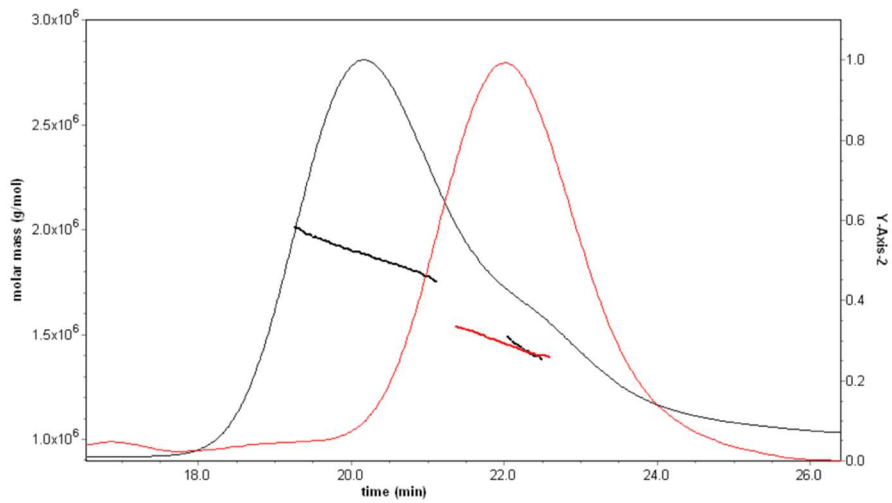


Figure S3. SEC/MALS/RI analysis of the full baseplate (black curve) and the Δ RBP1 (red curve) complexes. The molar mass (left axis), and the UV280nm absorbance (right axis) are plotted as a function of the column elution time. The column used was a 24-ml Superose 6 HR10/30 column (GE Healthcare, Cork, Ireland).