A pBBR1-based vector with IncP group plasmid compatibility for *Methylorubrum extorquens*

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Abstract
Plasmids are one of the most important genetic tools for basic research and biotechnology, as they enable rapid genetic manipulation. Here we present a novel pBBR1-based plasmid for *Methylorubrum extorquens*, a model methylotroph that is used for the development of C1-based microbial cell factories. To develop a vector with compatibility to the so far mainly used pCM plasmid system, we transferred the pBBR1-based plasmid pMiS1, which showed an extremely low transformation rate and caused a strong growth defect. Isolation of a suppressor mutant with improved growth led to the isolation of the variant pMis1_1B. Its higher transformation rate and less pronounced growth defect phenotype could be shown to be the result of a mutation in the promotor region of the rep gene. Moreover, cotransformation of pMis1_1B and pCM160 was possible, but the resulting transformants showed stronger growth defects in comparison with a single pMis1_1B transformant. Surprisingly, cotransformants carrying pCM160 and a pMis1_1B derivative containing a mCherry reporter construct showed higher fluorescence levels than strains containing only the pMis1_1B-based reporter plasmids or a corresponding pCM160 derivative. Relative plasmid copy number determination experiments confirmed our hypothesis of an increased copy number of pMis1_1B in the strain carrying both plasmids. Despite the slight metabolic burden caused by pMis1_1B, the plasmid strongly expands the genetic toolbox for *M. extorquens*.

Key words
cotransformation, expression system, *Methylorubrum extorquens* AM1, plasmid, plasmid copy number, Rep gene

1 | INTRODUCTION

*Methylorubrum extorquens* has great potential to become a universal production strain for the C1-based bioeconomy (Chen & Lan, 2020; Ochsner et al., 2014; Zhang et al., 2019). In the last decade, several *M. extorquens* strains with heterologously expressed metabolic pathways were described for the biotechnological production of various products including 1-butanol, 3-hydroxypropionic acid, mono- and dicarboxylic acids, mevalonate or α-humulene (Hu & Lidstrom, 2014; Liang et al., 2017; Lim et al., 2019; Schada von Borzyskowski et al., 2018; Sonntag et al., 2014, Sonntag, Kroner, et al., 2015; Sonntag, Müller, et al., 2015; Yang et al., 2017). The development of production strains, however, is limited by the organism's
restricted genetic accessibility. Although the bacterium has served as a model organism for methylotrophy for many decades and is therefore well described, only a few genetic tools have been developed. Plasmids are a key tool for the biotechnological applicability of a strain. The simultaneous use of two independent plasmids provides experimental flexibility and simplifies screenings and production strain developments. The compatibility of two or more plasmids in a single bacterial cell requires not only different selection markers but, more importantly, different origins of replication and partitioning systems (del Solar et al., 1998; Novick, 1987). As part of their characterization, plasmids are divided into Inc groups: plasmids from the same group are usually incompatible.

For M. extorquens AM1, besides a recently described set of mini chromosomes (Carrillo et al., 2019), derivatives of the pCM plasmid system (Marx & Lidstrom, 2001) were used almost exclusively as episomal vectors for heterologous gene expression. The expression vectors pCM80 (TcR) and pCM160 (kanR) are based on pDN19, a small IncP vector (Marx & Lidstrom, 2001). The authors of the respective study isolated the derivative pDN19X from M. extorquens AM1 transformed with pDN19. This derivative could be efficiently maintained and retransferred in M. extorquens AM1 and was used as a basis for the development of the pCM system (Marx & Lidstrom, 2001). In their work, the authors also showed that pBBR1MCS-2, a vector derived from Bordetella bronchiseptica pBBR1, could be transferred into M. extorquens with very low transformation efficiency and a significantly reduced transformant growth rate. Since this vector is known to be compatible with IncP group plasmids (Antoine & Locht, 1992; Kovach et al., 1994, 1995), we chose it as a starting point for the development of a plasmid with compatibility with the pCM vector system. The GC content of 64.6% seems to be applicable to M. extorquens AM1 (overall GC content of 68.5% [Vuilleumier et al., 2009]) and its diverse usability as a broad range vector can be also useful for certain applications. Here, we describe a pBBR1-derived plasmid with a mutation upstream of the rep gene that confers suitability for cotransformation with established pCM plasmids.

2 | MATERIALS AND METHODS

2.1 | Plasmids and bacterial strains

A list of all used plasmids and bacterial strains is given in Table 1. Reporter gene plasmids were constructed by subcloning synthesized DNA fragments (BioCat) into plasmid backbones. Used sequences and restriction enzymes are listed in Table 2. Analysis of sequences and creation of plasmid map was done with SnapGene (www.snapgene.com). Transformation of M. extorquens AM1 with plasmid DNA was performed as previously described (Toyama et al., 1998). If not stated differently, 100 ng of plasmid DNA was used for transformations.

2.2 | Media and culture conditions

Escherichia coli DH5α (NEB) was used for plasmid amplification and cloning. E. coli cultures were grown in LB (lysogeny broth) medium (Bertani, 1951). M. extorquens AM1 (Peel & Quayle, 1961)

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**TABLE 1** Plasmids and bacterial strains used in this study

| Name                        | Relevant features                                                                 | Reference                      |
|-----------------------------|-----------------------------------------------------------------------------------|--------------------------------|
| **Bacterial strains**       |                                                                                   |                                |
| Escherichia coli DH5α       | F- Δφ80lacZΔM15, Δ[locZYA-argF]U169, recA1, endA1, hsdR17 (rK-, mK+), phoA,       | ATCC                           |
|                             | supE44, λ-, thi-1 gyrA96 relA1                                                     |                                |
| Methylorubrum extorquens AM1| CmR, gram-negative, facultatively methylotrophic, obligate aerobic α-proteobacterium | Peel and Quayle (1961)         |
| **Plasmids**                |                                                                                   |                                |
| pCM160                      | Constitutive expression vector for M. extorquens; KanR, pmxaF (IncP)               | Marx and Lidstrom (2001)       |
| pCM80                       | Constitutive expression vector for M. extorquens; TcR, pmxaF (IncP)                | Marx and Lidstrom (2001)       |
| pMis1                       | *Pseudomonas putida* expression vector, KanR                                       | Mi et al. (2014)               |
| pMis4                       | *P. putida* expression vector; pMis1 derivative with mutated rep gene, KanR        | Mi et al. (2016)               |
| pBBR1MCS-2                  | Broad-host-range expression vector, KanR                                            | Kovach et al. (1995)           |
| pMis1_1B                    | pMis1 derivative, KanR                                                             | This study                     |
|                            | pCM160-based mCherry reporter plasmid                                              | This study                     |
| pMis1_1B_PmxaF_mCherry      | pMis1_1B based mCherry reporter plasmid, promoter rhoPBad was exchanged with PmxaF | This study                     |
| Plasmids/Sequences | Restriction enzymes used | Backbone |
|--------------------|--------------------------|----------|
| pMis1_1B_PmaxA_mCherry | EcoRI + XbaI | pMis1_1B |
| pCM160_mCherry | SphI | pCM160 |

**TABLE 2** Synthetic sequences used for cloning of reporter plasmids. DNA was synthesized by BioCat (Heidelberg, Germany). Underlined nucleotides mark restriction sites for cloning.
cultivations were performed in a liquid minimal medium (Peyraud et al., 2009) with 123 mM methanol and a final concentration of 12.6 µM CoCl₂ (Kiefer et al., 2009; Sonntag et al., 2014). Solid medium contained 1.5% [w/v] agar-agar. Antibiotics were used in concentrations of 50 µg/ml for kanamycin or 10 µg/ml for tetracycline hydrochloride. All media components were purchased from Carl Roth or Merck.

*M. extorquens* AM1 main cultures were inoculated to an OD₆₀₀ value of 0.1 with precultures grown for 48 h at 30°C and 180 rpm on a rotary shaker.

### 2.3 Growth monitoring and fluorescence assays

For the high-resolution monitoring of growth and mCherry fluorescence signals, *M. extorquens* AM1 cultures were grown in a BioLector® microreactor system (m2p-labs GmbH). Main cultures were grown in Flowerplate® wells at 30°C, 1000 rpm, and 85% humidity. The final cultivation volume was 1 mL. The level of mCherry reporter protein was measured via its fluorescence signal at 580/610 nm [ex/em]. Growth was monitored via scattered light signals.

### 2.4 Determination of relative plasmid copy numbers

The relative plasmid copy number (PCN) of pMis1_1B_mCherry was determined by comparing plasmid-specific real-time PCR fluorescence signals in strains containing two plasmids (=sample) to strains containing one plasmid (=control). The main cultures of respective strains were inoculated and cultivated as described above. Cells were harvested in the mid to late exponential growth phase after 45 h of cultivation. DNA was isolated with a QIAamp® DNA Mini kit (Qiagen) following the manufacturer’s instructions for bacterial cells. Real-time PCR primers pairs (Table 3) were designed for amplification of a specific region of the resistance markers on the plasmids (=target) as well as for amplification of a genomic reference sequence (=Ref), all yielding amplicons of 140–143 nucleotides in length. Real-time PCR experiments were performed using QuantiTect® SYBR® Green PCR Kit in a PikoReal™ System (Thermo Scientific) according to the manufacturer’s instructions. The following PCR protocol was used: 15 min at 95°C followed by 45 cycles of 94°C/15 s, 50°C/30 s, and 72°C/30 s. The specificity of primers was confirmed by melting curves and gel electrophoresis. The dynamic range of reaction was validated by testing a tenfold serial dilution of DNA template ranging from 50 ng to 5 pg. The efficiency (E) was subsequently calculated with Equation (1) (Higuchi et al., 1993; Rasmussen, 2001).

\[
E = 10^{\frac{1}{slopes}}
\]

The amount of starting material for the final experiments was 500 pg of template DNA. All reactions were performed in biological triplicates. The relative ratio (R) was calculated with the Pfaffl efficiency-corrected model with averaged controls (Equation 2) (Pfaffl, 2004).

\[
R = \frac{(E_{target})^{ΔC_P}_{target(MEAN control-sample)}}{(E_{Ref})^{ΔC_P}_{Ref(MEAN control-sample)}}
\]
RESULTS AND DISCUSSION

3.1 Increased transformation efficiency of pMis variant pMis1_1B

We tested different variants of pBBR1 (Antoine & Locht, 1992) for transformability in M. extorquens AM1, as pBBR1MCS-2 was transferred into M. extorquens AM1 already, albeit with very low efficiency (Marx & Lidstrom, 2001). Plasmids pMis1 and pMis4 are derivatives of pBBR1MCS-2 carrying a rhamnose-inducible promoter (Mi et al., 2016). In pMis4, a mutation in the rep gene leads to a G159S modification in the Rep protein. This mutation prevented the plasmid burden caused by pMis1 in Pseudomonas putida (Mi et al., 2016). The named plasmids and a pCM160 control were transferred into M. extorquens AM1, and an aliquot of 100 µl of the transformation mix was spread on selective agar plates. Although transformation of M. extorquens AM1 with pCM160 yielded almost a lawn of colonies, other plasmids yielded a low number of transformants after 192 h of incubation: On the pBBR1MCS-2 transformation plate, only 2 very small colonies were visible, indicating a severe growth defect of the respective transformants. Transformation with pMis1 yielded in a sole, big colony, and transformation with pMis4 in a small number of big colonies. The respective plasmids of these colonies were isolated and 30 fmol of pDNA was retransformed into M. extorquens AM1 along with pCM160 as a control. Only the plasmid isolated from the pMis1 transformant showed a clear increase in transformation efficiency compared to pMis1 and was named pMis1_1B (Figure 1a). Colonies of pMis1_1B transformants were clearly visible after 96 h of incubation, with inconsistent colony size, whereas colonies of pMis1 transformants were not visible even after 240 h of cultivation.

Complete sequencing of pMis1_1B revealed an insertion of one cytosine residue 22 bp upstream of the rep gene (Figure 1b). This insertion upstream of the rep region has likely altered the plasmid copy number (PCN). For pBBR1-based plasmids, the Rep protein is known to play a crucial role in PCN control by binding to the origin of replication (ori) (Antoine & Locht, 1992; del Solar et al., 1998). Consequently, modifications in the rep gene region have been shown to influence the PCN (Mi et al., 2016; Tao et al., 2005; Wadood et al., 1997). Interestingly, the currently used pCM system is based on a similar experiment with the plasmid pDN19 (Marx & Lidstrom, 2001). pDN19 was transferred into M. extorquens AM1 with low efficiency, but a mutated derivative (pDN19X) could be isolated from a transformant. pDN19X was efficiently maintained and showed enhanced transformation efficiency. The described single-point mutation in traJ led to changes in transformability even though all
functions were provided by helper plasmids. The authors, therefore, speculated that transformability was not enhanced by changes in TraJ but by a change in PCN. This confirms that the PCN is an extremely important factor for *M. extorquens* plasmids.

### 3.2 | Compatibility of pMis1_1B with the established pCM plasmids

For testing the compatibility of pMis1_1B with the pCM system, *M. extorquens* AM1 was transformed with pMis1_1B and pCM80 simultaneously. The cotransformation did lower the transformation efficiency by 2-fold compared to transformation of pMis1_1B alone (Figure 2a). Furthermore, colonies of cotransformants were only clearly visible after 8 days of growth, indicating an increased metabolic burden caused by the presence of both plasmids. Detailed growth monitoring in liquid growth medium confirmed these results: While *M. extorquens* AM1 harboring pMis1_1B already showed a reduced growth rate compared to the pCM160-containing strain or the plasmid-free strain, cotransformation with both plasmids strongly reduced the growth rate (Figure 2b). Plasmid-induced reduction of growth rates under methylotrophic growth conditions due to metabolite limitations have already been described for *M. extorquens* AM1 (Kiefer et al., 2009). We, therefore, tested increased amounts of several media components but were unable to identify any medium-related limitation (data not shown). Only a reduction in the kanamycin concentration resulted in an increased growth rate, but most probably reduced plasmid maintenance stability or copy number, as revealed by fluorescence quantification experiments using an mCherry reporter derivative of pMis1_1B (Figure A1). Mutations in the rep-gene region were already successfully used to increase the copy number of pBBR-based broad host range plasmids and the growth rate of corresponding transformants (Mi et al., 2016; Tao et al., 2005). A more directed screening of rep gene expression level variants might identify a more suitable expression level associated with less plasmid burden.

### 3.3 | Characterization of pMis1_1B as pCM coexpression vector

Despite the lower growth rates of the cotransformants, pMis1_1B is nevertheless a promising candidate for simultaneous gene expression from two plasmids in *M. extorquens* AM1. To investigate the plasmid stability and general expression levels from pMis1_1B, we used pMis1_1B_PmxaF_mCherry containing the reporter gene under the control of the strong native constitutive promotor PmxaF. Expression of mCherry did not change the growth behavior of the respective strains.
We investigated the mCherry expression levels from respective pCM and pMis1_1B constructs in strains with one or two plasmids, respectively. The previously described growth defect of the cotransformants was also visible in this experiment (Figure 3a). However, there was a clear synergistic effect in terms of expression: The mCherry signal originating from pMis1_1B_P_mxaF_mCherry was substantially increased if pCM80 was additionally present. The maximal mCherry fluorescence signal of the cotransformants even exceeded the values from pCM160_mCherry single transformants. One explanation for this finding could be an increased PCN. We therefore, determined the relative PCN of pMis1_1B_mCherry for strains containing two plasmids and strains containing only a single plasmid by real-time PCR. Since the efficiency of the used primer pairs varied from 1.73 to 1.96, the E-corrected Pfaffl-method was used (Pfaffl, 2004). Although the values determined for each of the three replicate strains containing two plasmids showed a strong variation, a clear increase in PCN was detectable when both plasmids are present in M. extorquens AM1 (Figure 3b). Thus, the presence of pCM80 led to a strong increase in the pMis1_1B copy number.

4 | CONCLUSIONS

The pBBR1 derivative pMis1_1B, which we characterized in this study represents a novel plasmid for M. extorquens, which is compatible with the widely used pCM system. Even though the growth rate of M. extorquens AM1 was affected when both plasmids were present in the cells, pMis1_1B could be a powerful tool for certain applications. For example, it provides enormous facilitation for the combinatorial testing of multiple enzymes during the development of synthetic pathways. Moreover, a new plasmid system that is not as stably maintained as pCM could be useful for applications where an expression plasmid is only needed transiently as for the expression of Cre-recombinase in a Cre/IoxP recombination system or for CRISPR gene editing. With pMis1_1B, a new useful tool has been added to the expanding genetic toolbox for M. extorquens.

AUTHOR CONTRIBUTIONS

Laura Pöschel: Conceptualization (equal); investigation (equal); methodology (equal); visualization (lead); writing—original draft (lead); writing—review and editing (equal); Elisabeth Gehr: Conceptualization (equal); investigation (equal); methodology (equal); writing—original draft (supporting); Markus Buchhaupt: Conceptualization (equal); methodology (equal); funding acquisition (lead); project administration (lead); writing—original draft (supporting); writing—review and editing (equal).

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CONFLICT OF INTEREST

None declared.

DATA AVAILABILITY STATEMENT

The plasmid sequence of pMis1_1B is available in the NCBI GenBank under accession number OP441404: https://www.ncbi.nlm.nih.gov/nuccore/OP441404. The plasmid itself has been deposited to Addgene. All other data generated or analyzed during this study are included in this published article.

ETHICS STATEMENT

None required.

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APPENDIX

**Figure A1** Effect of variation of kanamycin concentration on the growth of *Methylorubrum extorquens* AM1 pCM80/pMis1_1B_p_mxaF_mCherry cotransformants. The maximal mCherry fluorescence signal is a measure of pMis1_1B_p_mxaF_mCherry abundance, while the maximal slope of the scattered light signal is an indicator of growth rate. Three independent biological replicates were measured; standard deviations are indicated by error bars.

**Figure A2** Comparison of scattered light measurements of *Methylorubrum extorquens* AM1 cotransformants containing pCM80 combined with either pMis1_1B or pMis1_1B_p_mxaF_mCherry. Measurements of biological triplicates are shown.