

pMR101 87115[™]

Description

Clone type: Vector

Host: Escherichia coli JM101 (ATCC 33876)

Storage Conditions

Product format: Freeze-dried

Intended Use

This product is intended for laboratory research use only. It is not intended for any animal or human therapeutic use, any human or animal consumption, or any diagnostic use.

BSL₁

ATCC determines the biosafety level of a material based on our risk assessment as guided by the current edition of *Biosafety in Microbiological and Biomedical Laboratories* (*BMBL*), U.S. Department of Health and Human Services. It is your responsibility to understand the hazards associated with the material per your organization's policies and procedures as well as any other applicable regulations as enforced by your local or national agencies.

Certificate of Analysis





For batch-specific test results, refer to the applicable certificate of analysis that can be found at www.atcc.org.

Vector Information

Construct size (kb): 4.099999904632568

Intact vector size: 4.100

Vector name: pMR101 (phagemid)

Type of vector: phagemid

Construction: pTM201/NS3-3, pET8c, pACYC177, pET11d

Host range: Escherichia coli

Vector information: Other unique sites: EcoNI, Sall, BstEII, PstI, BglII, Xbal

Cloning sites: Ncol; BamHI

Markers: kanR

Operator: lac, <-, 3788-3804

Promoters: T7 (phi10), <-, 3808-3827

Replicon: M13, →, 11-467; p15A, →, 1585-1587

Growth Conditions

Medium:

ATCC Medium 1236: LB Medium (ATCC medium 1065) with 25 mcg/ml kanamycin

Temperature: 37°C

Notes

Restriction digests of the clone give the following sizes (kb): BamHI--4.1; BstEII/PstI--3.0, 1.08; SalI--4.1.

- ATCC staff

There is an error in the reference in figures 1 and 2. The restriction sites,

Xbal and Bglll, are drawn in reverse order on the maps for pMR101.

- personal communication



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Expression vector (T7-based) with a kanR marker and a P15A replicon compatible with ColE1-derived plasmids. Particularly useful for co-transformation with ColE1-based ampR T7 expression vectors and the production of two proteins in the same cell.

- Gene 144: 59-62, 1994

Use of 5'Ncol and 3'BamHI cloning sites is similar to that of other expression systems, which facilitiates transfer of genes into these pMR vectors.

- Gene 144: 59-62, 1994

If used in an Escherichia coli strain that expresses T7 polymerase under the control of the lacUV5 promotor (such as BL21(DE3)), addition of IPTG can result in high levels of recombinant protein production.

- Gene 144: 59-62, 1994

Material Citation

If use of this material results in a scientific publication, please cite the material in the following manner: pMR101 (ATCC 87115)

References

References and other information relating to this material are available at www.atcc.org.

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