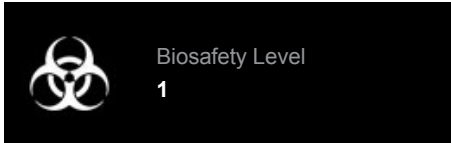




Product Sheet

pAD2 (ATCC® 87469™)

Please read this FIRST



Intended Use

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

Citation of Strain

If use of this culture results in a scientific publication, it should be cited in that manuscript in the following manner: pAD2 (ATCC® 87469™)

Shipping Information

Distributed: freeze-dried

American Type Culture Collection
PO Box 1549
Manassas, VA 20108 USA
www.atcc.org

800.638.6597 or 703.365.2700
Fax: 703.365.2750
Email: Tech@atcc.org

Or contact your local distributor

Description

Designation: pAD2

Distribution Host:

Distribution host: *Escherichia coli* unknown (ATCC 33694)

Propagation

Growth Conditions

Temperature: 37°C

Medium

ATCC® Medium 1227: LB Medium (ATCC medium 1065) with 50 mcg/ml ampicillin

Vector Information

Size (kb): 6.352

DESCRIPTION OF VECTOR:

Intact vector size: 6.352

Type of vector: phagemid

Cloning sites:

Polylinker sites:

Other unique sites: BamHI

Construction: pRS406

Host range: *Saccharomyces cerevisiae*; *Escherichia coli*

Features (with orientation and position when available):

marker(s): URA3, ->

replicon: f1, <-

other: LYS2 deleter cassette

replicon: pMB1

marker(s): ampR, <-

Vector: pAD2 (phagemid)

Construction: pRS406

Marker(s):URA3,ampR

Construct size (kb): 6.352

Features: marker(s): URA3

marker(s): ampR

other: LYS2 deleter cassette

replicon: f1

replicon: pMB1

References

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

Notes

Restriction digests of the clone give the following sizes (kb): KpnI--6.3;

KpnI/SstI--4.3, 2.0; ClaI--6.3.

- ATCC staff

Deleter vector for constructing designer yeast strains with a non-revertable deletion of a *lys2* auxotrophic marker.

- Yeast 14: 115-132, 1998

The two step selection process requires a *ura3* transformation host (this host can be created using pJL164 (ATCC 87471)). After transformation with the ClaI linearized vector, URA3 integrants are selected on *ura-* plates.

- Yeast 14: 115-132, 1998

The designer deletion strain is then recovered by selection on 5-FOA plates (loss of URA3 and LYS2 markers by a homologous recombination event).

- Yeast 14: 115-132, 1998



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Biosafety Level: 1

Appropriate safety procedures should always be used with this material. Laboratory safety is discussed in the current publication of the *Biosafety in Microbiological and Biomedical Laboratories* from the U.S. Department of Health and Human Services Centers for Disease Control and Prevention and National Institutes for Health.

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Additional information on this culture is available on the ATCC web site at www.atcc.org.

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