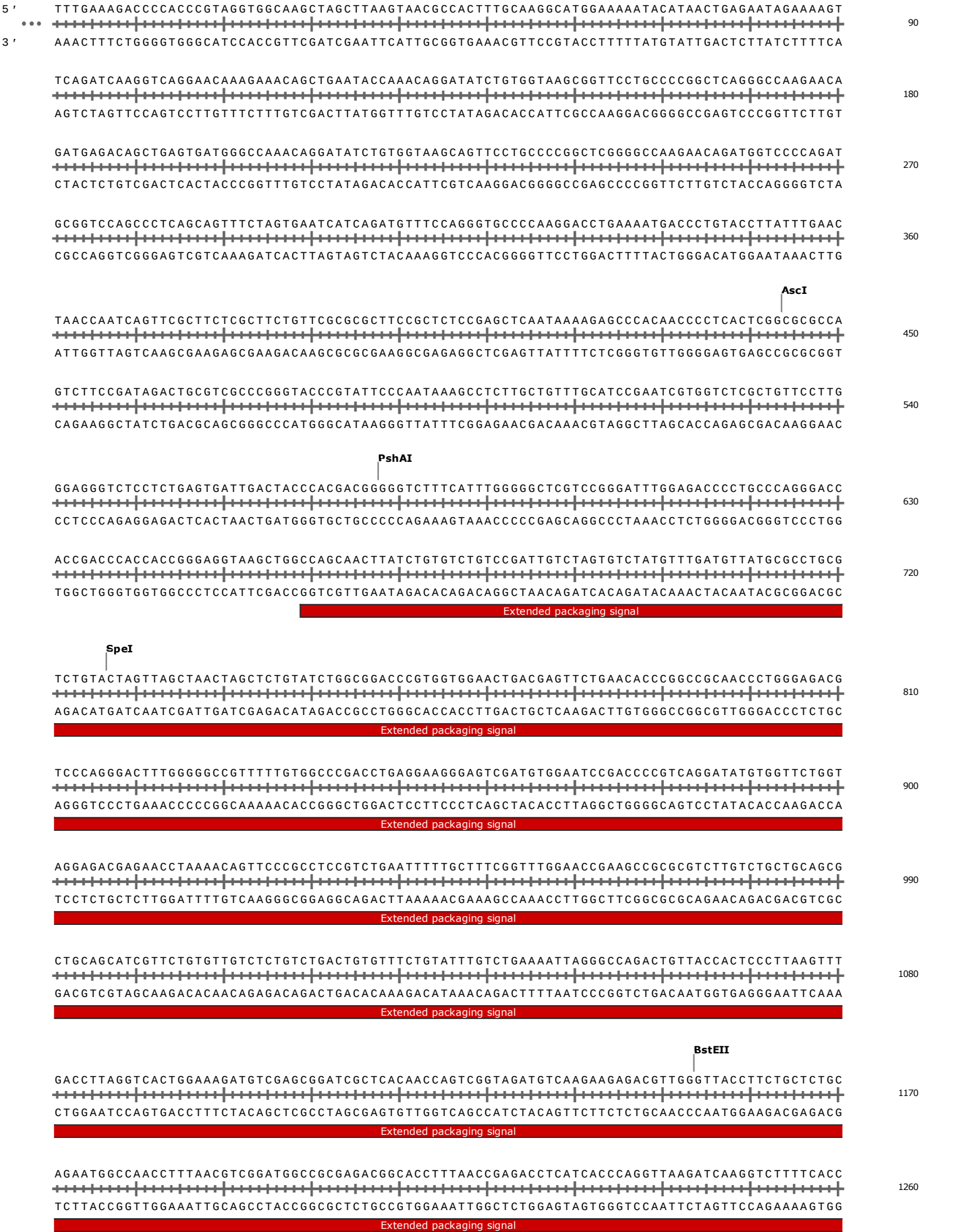
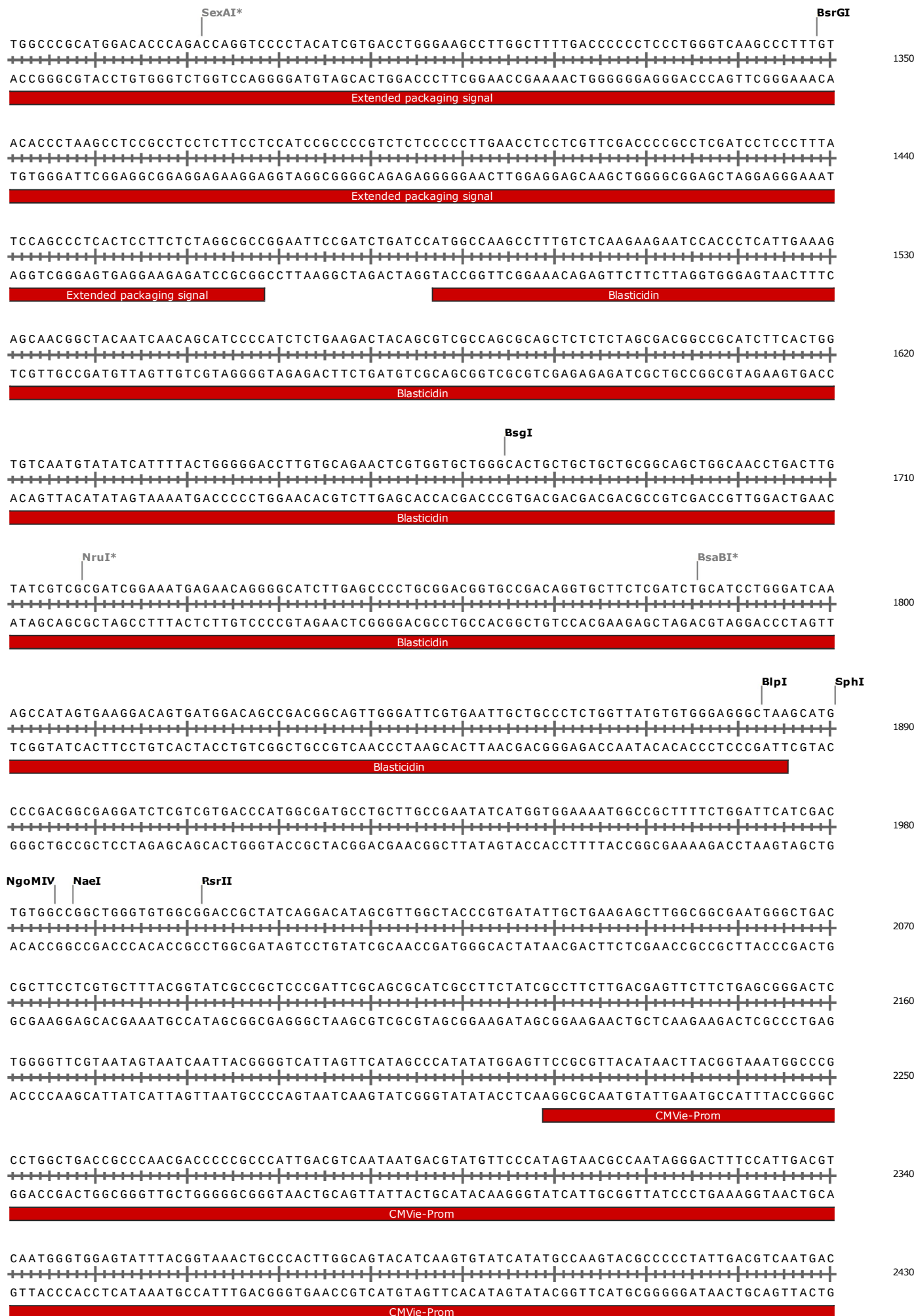






SnaBI

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 CMVie-Prom

2520

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 CMVie-Prom

2610

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 CMVie-Prom

2700

BglII XhoI
PaeR7I HindIII

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 CMVie-Prom

2790

NotI SalI StuI
ClaI BspDI

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2880

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2970

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3060

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3150

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3420

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3510

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3600

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3690

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3780

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3870

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3960

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4140

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4320

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4590

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4680

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4770

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4860

Amp

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4950

Amp

AseI

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5040

Amp

FspI

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 +-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+-----+
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5130

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5220

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5310

ScaI

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5400

XmnI

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5490

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5580

SspI

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5670

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5760

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5850

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5940

SacII

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5978

DNA Type: Synthetic DNA

Laboratory Host:

Bacterial Transformation Strain: Unspecified
Dam⁺ Dcm⁺ EcoKI⁺

Description: Ligation of Fragment 2 into Fragment 2

Created: Samstag, 10. Aug 2024

Last Modified: Samstag, 10. Aug 2024

Accession Number:

Code Number:

Sequence Author:

Comments:

References:

Embedded Files: