

CERTIFICATE OF ANALYSIS **Enzyme Properties:** 1 µl of Turbo Tru9 I cuts 1 µg of DNA in 1x SE-Buffer ROSE in 10 min (assayed on Lambda and plasmid DNA). A short time of DNA digestion requires high quality purification of DNA sample (PCR fragments should be

purified after amplification). Please note that supercoiled plasmid DNA and PCR fragments may have varying rates of cleavage and sometimes need more time to be completely digested.

Standard protocol of Turbo reaction: 20 µl of the reaction volume: 10x SE-Buffer ROSE - 2 µl $-0.2-1 \mu q$ Nuclease-free water - to 20 µl + 1 µl of Turbo Tru9 I Mix by pipette tip carefully. Incubate at 65°C for 10 min.

(10 min) DNA digestion in universal (ROSE) SE-Buffer. Source: An E.coli strain that carries the cloned Tru9 I gene from Thermus ruber 9. Supplied in: 10 mM Tris-HCl (pH 7.5), 50 mM NaCl, 0.1 mM EDTA, 1 mM DTT, 200 µg/ml BSA, 50% glycerol.

Description: Turbo Tru9 I is used for short time

endonuclease for 3 hours. Reagents Supplied with Enzyme:

most Restriction Endonucleases. ROSE Buffer is perfect for DNA cleavage with SE Turbo Restriction Endonucleases and for double digestion. **Heat Inactivation:** Enzyme is inactivated by incubation at 80°C for 20 min.

1x SE-Buffer ROSE, Incubate at 65°C.

Reaction Original SibEnzyme (ROSE) Buffer is a

specially designed universal reaction buffer for the

Reaction Conditions:

Applications:

-Fast preparation of vectors for cloning

Quality Control Assays

-Fast DNA analysis

Ligation: After digestion with 1 µl of Turbo Tru9 I,

with high-activity T4 DNA Ligase and recut.

approximately 95% of the DNA fragments can be ligated

Oligonucleotide Assay: No detectable degradation of a

single-stranded and double-stranded oligonucleotide

was observed after incubation with 1 µl of restriction

10X SE-Buffer ROSE.

-Double digestion