

CYCV17-CHANGYU pTOPO-Blunt Simple Kit

Contents	20 rxns (CYCV1701)	80 rxns (CYCV1702)
pTOPO-Blunt Simple Vector(30ng/μl)	20 μl	80 μl
1000bp Control (30ng/μl)	5 μl	5 μl
10 × Enhancer	20 μl	80 μl

Note: All reagents, when stored at -20 °C, are stable for 12 months

❖ Description

The Zero Background Blunt Simple Topoisomerase Cloning Kit is designed for fast cloning of blunt-ended DNA fragments up to 10 kb generated by high fidelity DNA polymerase such as KOD, Pfu and Phusion etc. DNA fragments obtained by restriction digestion or mechanical shearing can also be cloned after end polishing to become blunt ended. It utilizes DNA strand transfer activity of *Vaccinia* virus topoisomerase I. *Vaccinia* virus DNA topoisomerase I forms a 3'-phosphoryl intermediate with the plasmid vector containing cleavage recognition motif of 5'CCCTT ↓. Covalently bound topoisomerase I then transfers the incised vector DNA strand to the DNA fragment to be cloned with free 5'-OH termini. This transferring reaction is rapid and reproducible. The cloning vector pTOPO-Blunt Simple included in this kit is a high-copy-number plasmid engineered to tolerate mildly toxic genes. Regions flanking the cloning site of pTOPO-Blunt Simple vectors do not contain any common restriction sites which eliminates possible redundancy of restriction sites.

❖ Procedure

1. Set up the topoisomerase cloning reaction by mixing the reagents in the order shown.

DNA Fragment or 1μl 1000bp control	0.5-8μl
pTOPO-Blunt Simple Vector	1μl
10 × Enhancer	1μl
ddH2O	Xμl
Final Volume	10μl

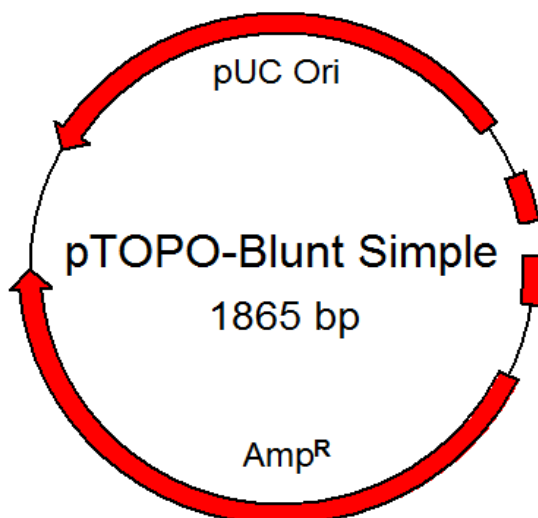
DNA fragments MUST NOT be 5'-phosphorylated. Blunt-ended DNA fragment is preferred. Highest cloning efficiency was observed from PCR fragments generated by high fidelity enzymes such as KOD, Phusion, Pfu etc. 10-150 ng of DNA

fragments ranging from 100 bp to 5000 bp have been tested to give satisfactory results. Excess amount of DNA inserts e.g. >300ng, will reduce cloning efficiency. Refer to the following table.

Fragment size (bp)	Optimal amount (ng)
100-1000	10-40
1000-2000	40-80
2000-5000	

2. Mix the reaction gently and incubate for 5 minutes at room temperature between 15-30°C. In any case, do not let the incubation go beyond 5 minutes. Extended incubation for larger inserts up to 5 kb is unnecessary and may introduce background.
3. Add 5 µl of the cloning reaction into 50 µl chemically competent E. coli and mix gently. Do not mix by pipetting up and down.
4. Incubate on ice for 2-30 minutes.
5. Follow the competent cell manufacturer's instructions to complete the transformation.

❖ **Map of pTOPO-Blunt Simple:**

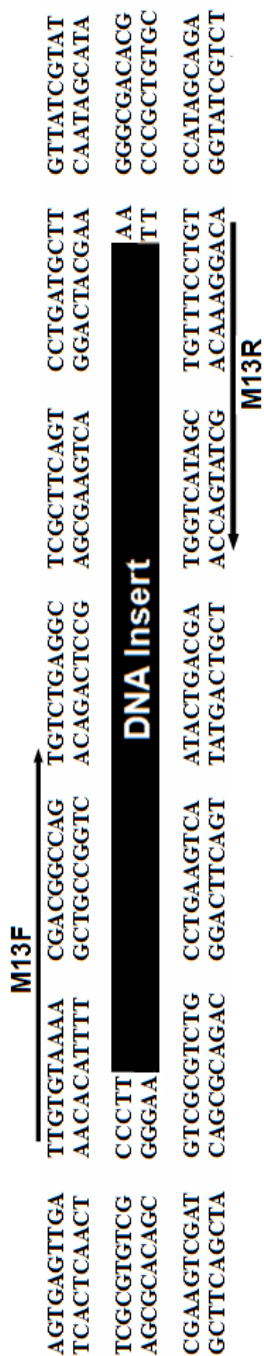


❖ **Sequencing Primer:**

M13F: TGTAACGACGGCCAGT

M13R: CAGGAAACAGCTATGACC

❖ Cloning site information:



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 Not for diagnostic or therapeutic use.