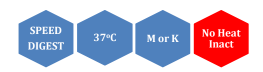
Pvu II



5' ···CAG▼CTG···3'
3' ···GTC▲GAC···5'

Pvull is a restriction enzyme purified from a recombinant *E.coli* strain.

<u>Catalogue No</u> 128-1, 500 U

128-2, 2500 U 128-3, 12500 U

Concentration 10-12u/µl and 40-

60u/µl*

Reagents supplied: 10x M and 10x K

buffer

Unit substrate: Lambda DNA.

Unit calculation assay conditions: 50 mM NaCl, 10 mM Tris-HCl (pH 7.9 @ 25° C), 10 mM MgCl₂, 1 mM dithiothreitol, 100 µg/ml bovine serum albumin and DNA. Incubate at 37° C.

Absence of contaminants: 100 units of Pvu II do not produce any unspecific cleavage products after 16 hrs incubation with 1 μg of Lambda DNA at 37°C. After 10-fold overdigestion with Pvu II greater than 95% of the DNA fragments can be ligated and recut with this enzyme.

Storage buffer: 50 mM KCl, 10 mM Tris-HCl (pH 7.4), 0.1 mM EDTA, 1 mM dithiothreitol, 200 μ g/ml bovine serum albumin and 50% glycerol. Store at -20°C.

Heat inactivation: No.

Methylation Sensitivity:

dam methylation: Not sensitive dcm methylation: Not sensitive CpG methylation: Not sensitive **Star activity:** Conditions of low ionic strength, high enzyme concentration, glycerol concentration >5% or pH >8.0 may result in star activity.

Percent Activity in MINOTECH Buffers

L	М	Н	SH	Α	K
25-50	100	100	25-50	50	100

General reaction mixture:

10U Pvull	1μΙ				
10x M or K buffer *	2μΙ				
DNA substrate	<1µg				
Sterile ultrapure water	Up to 20 μl				
Incubate for 15 min at 37°C					

^{*}In the case of M buffer we recommend the addition of BSA to a final concentration of $100 \mu g/ml$.

Frequency of Cutting

λ	Ad-2	Фх174	pUC18	M13mp18	pBR322
15	24	0	2	3	1



Lambda DNA 1.0 % agarose



^{*}Add an H to cat.# to order the high concentration