

NAUTIAGENE

Product Information

NautiaZ Culture Cell DNA Extraction Mini Kit

(100/300 prep)

Cat. No. : NGCZ-S100/NGCZ-S300

Sample : 10⁷ Culture Cell

Yield : Up to 50 µg

NGCZ-S100	NautiaZ Culture Cell DNA Extraction Mini Kit (100 prep)
NGCZ-S300	NautiaZ Culture Cell DNA Extraction Mini Kit (300 prep)
NGBZ-S100	NautiaZ Blood DNA Extraction Mini Kit (100 prep)
NGBZ-S300	NautiaZ Blood DNA Extraction Mini Kit (300 prep)
NGBAZ-S100	NautiaZ Bacteria/Fungi DNA Extraction Mini Kit (100 prep)
NGBAZ-S300	NautiaZ Bacteria/Fungi DNA Extraction Mini Kit (300 prep)
NGPZ-S100	NautiaZ Plant DNA Extraction Mini Kit (100 prep)
NGPZ-S300	NautiaZ Plant DNA Extraction Mini Kit (300 prep)
NGTZ-S100	NautiaZ Tissue DNA Extraction Mini Kit (100 prep)
NGTZ-S300	NautiaZ Tissue DNA Extraction Mini Kit (300 prep)
NGST-S100	NautiaZ Stool/Soil DNA Extraction Mini Kit (100 prep)
NGST-S300	NautiaZ Stool/Soil DNA Extraction Mini Kit (300 prep)

NGCF-S100	NautiaZ Cell-Free DNA Extraction Mini Kit (100 prep)
NGPK-S100	NautiaZ Whole Blood DNA Extraction Mini Kit (100 prep)

Contents

	NGCZ-S100T	NGCZ-S100	NGCZ-S300
N1 Buffer	2 ml	20 ml	20 ml
N2 Buffer	1.5 ml	35 ml	95 ml
N3 Buffer	2 ml	45 ml	125 ml
W1 Buffer	2 ml	45 ml	125 ml
W2 Buffer	300 ul x2	15 ml	25 ml x2
Elution Buffer	1 ml	10 ml	30 ml
GZ Column	4 pcs	100 pcs	300 pcs
Collection Tube	4 pcs	100 pcs	300 pcs
User Manual	1	1	1

*Add 1.2 ml x2 / 60 ml / 100 ml x2 ethanol (96-100%) to W2 Buffer prior to the initial use.

Buffer Preparation

- Add ethanol (96-100%) to the Wash Solution prior to first use

	NGCZ-S100T	NGCZ-S100	NGCZ-S300
W2 Buffer ethanol (96 ~ 100%)	300 ul x2 1.2 ml x2	15 ml 60 ml	25 ml x2 100 ml x2

Important Notes

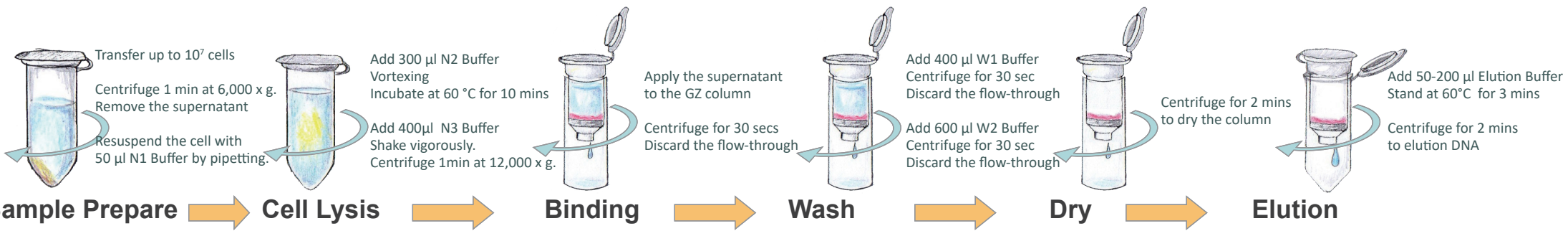
1. Buffers provided in this system contain irritants. Wear gloves and lab coat when handling these buffers.
2. Add ethanol (96- 100 %) to W2 Buffer when first open.
3. Prepare dry baths or water baths before the operation.
4. Resolve any precipitate by warming at 37°C.

Description

The NautiaZ Culture Cell DNA Extraction Mini Kit is designed for rapid extraction of pure genomic DNA from Bacteria and fungus cell. Efficiently remove cellular debris and inhibitors, this kit using column-type tube in purification process through three simple steps of binding, washing and then elution for the safe and convenient extraction of high-purity genomic DNA. The entire process can be completed in less than 1 hour without phenol/chloroform, and the final product can be used in PCR or other downstream experiments.

Storage

Store at room temperature.



PURIFICATION PROTOCOLS

For Culture Cell

STEP	PROCEDURE
1 Sample prepare	Transfer up to 10^7 cultured mammalian cells to a microcentrifuge tube (<i>not provided</i>). Centrifuge 1 min at $6,000 \times g$. Remove the supernatant completely. Resuspend the cell with $50 \mu\text{l}$ N1 Buffer by pipetting.
2 Cell Lysis	Add $300 \mu\text{l}$ N2 Buffer to the sample and mix thoroughly by vortexing. Incubate at 60°C for 10 mins until the sample lysate is clear. Invert the tube every 3 minutes during incubation.
Optional Step: RNase	<i>If RNA-free genomic DNA is required, perform this optional step.</i> Add $5 \mu\text{l}$ of RNase (10 mg/ml) to sample lysate and mix by vortexing. Incubate at room temperature for 5 mins.
3 Protein Removal	Add $400 \mu\text{l}$ N3 Buffer to the sample and shake vigorously. Centrifuge 1 min at $12,000 \times g$.
4 DNA Binding	Place a GZ column in Collection Tube. Transfer the supernatant to GZ column and centrifuge 30 seconds at $14,000 \times g$. Discard the flow-through and place GZ Column back in the Collection tube.

5-1 Wash	Add $400 \mu\text{l}$ W1 Buffer to GZ Column . Centrifuge at $14,000 \times g$ for 30 seconds. Discard the flow-through and place GZ Column back in the Collection tube.
5-2 Wash	Add $600 \mu\text{l}$ W2 Buffer (ethanol added) to GZ Column . Centrifuge at $14,000 \times g$ for 30 seconds. Discard the flow-through and place GZ Column back in the Collection tube.
6 Dry	Centrifuge at $14,000 \times g$ for 2 minutes to dry the column.
7 Elution	Place GZ Column to a clean 1.5 ml microcentrifuge tube (<i>not provided</i>). Add $50\text{-}200 \mu\text{l}$ of preheated Elution Buffer (70°C) or ddH_2O ($\text{pH}7.5\text{-}8.0$) into the center of the column matrix.
	Stand at 60°C for 3 minutes. Centrifuge at $14,000 \times g$ for 2 minutes to elute purified DNA.
8 Pure DNA	Store the DNA fragment at 4°C or -20°C .

