



NeoGeneStar
100 Randolph Road, Suite 2B
Somerset, NJ 08873
Tel: (732) 421-4567
Fax (908) 756-4483
www.NeoGeneStar.com

NeoGeneStar™ Cell Free DNA Purification Kit for 1ml Samples

For purification of cell-free DNA from 1ml of plasma, serum, CSF or urine

For Research Use Only.

Not for human or animal therapeutic or diagnostic use.



Binding Characteristics and Sample Volume

The superparamagnetic particles bind DNA molecules from ~30 bases to > 10,000 bases. The 1ml size NeoGeneStar™ circulating cell-free DNA Kit has been optimized for sample volumes of up to 1ml.

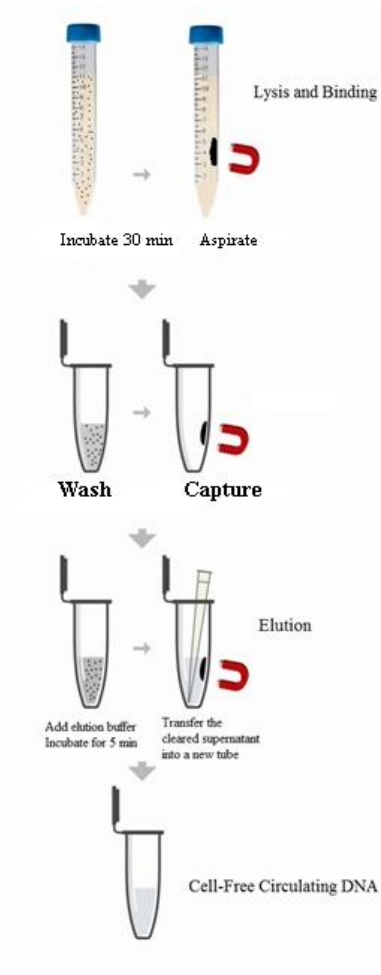
Catalog No	Sample Volume and Quantity	Pretreatment Buffer (20x)	NGS Protease (25X)	RNA Carrier	LYS ¹ Tubes	NGS™ Beads	Wash Buffer ²	Elution Buffer
NeoGeneStar™1ml-25-WPR	1ml x 25 preps	1.25ml	1ml	125µl	25	0.75 ml	40ml	2..5ml
NeoGeneStar™1ml-50-WPR	1ml x 50 preps	2.5ml	2ml	250µl	50	1.5ml	80ml	5.0ml
NeoGeneStar™1ml-100-WPR	1ml x 100 preps	5ml	4ml	500µl	100	3.0ml	160ml	10.0ml

¹LYS tubes contain chaotropic salts, which are irritants. Please wear gloves and handle with appropriate laboratory safety measures.

²Absolute ethanol must be added at 1:1 ratio prior to use for the Wash Buffer.

Procedure of the NeoGeneStar™ Circulating cfDNA Kit

For 1ml sample, add 50µl Pretreatment Buffer,
40µl NGS Protease, and 5µl RNA carrier to 1ml
sample in a 1.5 ml microcentrifuge tube,
incubate 30 minutes at 55-60°C.
Add the entire pretreated plasma slowly
into the LYS tube, dissolve at room
temperature, then add 0.18ml isopropanol
and 30µl NGS™ Beads and mix well
Incubate for 30 minutes by vortexing or
inverting
Wash 2 times with Wash Buffer
Wash 2 times with 80% Ethanol
Air dry 5-10 minutes
Elute





Note Regarding Wash Buffer

Wash Buffer must be diluted with an equal amount of absolute ethanol and stored at room temperature prior to use. In the event that the Wash Buffer is chilled, a precipitate may form; it would then be necessary to warm the Wash Buffer to fully dissolve the components prior to use.

Materials Needed That Are Not Supplied:

- Low DNA binding microcentrifuge tubes
- Normal Saline (0.9% w/v NaCl) solution (for volume adjustment of samples less than 1ml)
- Magnetic separation devices (for microcentrifuge tubes)
- Absolute ethanol and isopropanol
- Tube shaker / vortexer for microcentrifuge tubes.

Purification Protocol:

1. For 1ml sample, add 50µl Pretreatment buffer, 40µl of NGS Protease, and 5µl RNA carrier in a 1.5 ml microcentrifuge tube, mix well and incubate at 55-60°C for 30 minutes.
Please note: If the sample volume is less than the designed kit processing volume, add the appropriate volume of 0.9% sodium chloride solution (normal saline) to bring the volume to the specified sample volume. (ie. if the sample volume is 0.5ml, add 0.5ml of 0.9% sodium chloride solution to result in a 1ml volume for processing sample).
 2. Add the entire pretreated sample to the 1ml LYS tube and mix thoroughly at room temperature, ensure that the reagents are fully dissolved, then add 0.18ml isopropanol, mix.
 3. **Resuspend the NGS™ Beads by vortexing for 1 minute**, then add 30µl of NGS™ Beads, vortex or invert for 30 minutes on a vortexer or 360° rotator. Proper shaking will result in visible foam above the liquid layer.
 4. Centrifuge briefly to reduce the foam, then place the 2ml tube on a magnetic stand for at least 2 minutes, the solution will clear. With the sample tube on the magnetic stand, carefully aspirate the cleared supernatant without aspirating the NGS™ Beads. Vacuum aspiration is convenient but not necessary.
 5. Remove the sample tube from the magnetic stand, add 1ml of Wash Buffer (diluted 1:1 with absolute ethanol) into the tube and carefully rinse the NGS™ Beads and transfer to a 1.5ml low DNA binding microcentrifuge tube, add 0.4ml diluted Wash Buffer to rinse the 2ml tube again and transfer to the 1.5 low DNA binding microcentrifuge tube.
 6. Place the microcentrifuge tube on a magnetic stand for at least 1 minute or until the solution clears, turn the tube over a few times while on the magnetic stand to remove the NGS™ Beads trapped on the lid. With the sample tube on the magnetic stand, carefully
-



aspirate the cleared supernatant without aspirating the NGS™ Beads.

7. Remove the microcentrifuge tube from the magnetic stand and add 1ml of Wash Buffer (diluted 1:1 with absolute ethanol) into the tube and resuspend the NGS™ Beads by vortexing for 30 seconds.
8. Place the sample tube onto the magnetic stand for at least 1 minute or until the solution clears, turn the tube over a few times while on the magnetic stand to remove the NGS™ Beads trapped on the lid. With the sample tube on the magnetic stand, carefully aspirate the cleared supernatant without aspirating the NGS™ Beads.
9. Remove the microcentrifuge tube from the magnetic stand and add 1ml of 80% ethanol to the tube. Completely resuspend the NGS™ Beads by vortexing for 30 seconds.
10. Place the microcentrifuge tube on the magnetic stand for at least 1 minute or until the solution clears, turn the tube over a few times while on the magnetic stand to remove the NGS™ Beads trapped on the lid. With the sample tube on the magnetic stand, carefully aspirate the cleared supernatant without aspirating the NGS™ Beads.
11. Wash the NGS™ Beads with 1ml of 80% ethanol again by repeating steps 9 and 10.
12. Pulse down and aspirate again to remove as much of the liquid as possible. Keeping the tube on the magnet, air-dry the NGS™ Beads at room temperature for 10 minutes.
13. Add 50µl-100µl Elution Buffer or elution buffer of your choice and resuspend the NGS™ Beads by pipetting up and down, then incubate 10 minutes at room temperature.
14. Place sample tubes on a magnetic stand, the solution will clear in about 1 minute. Transfer the cleared supernatant into a low DNA binding microcentrifuge tube. This is the purified cfDNA. Freeze the eluted cfDNA until you are ready for your downstream analysis.

Note: For certain applications, such as digital (microdroplet) PCR, it may be advantageous to briefly microcentrifuge to remove any NGS™ Beads that may be present in the eluate.

The information in this guide is subject to change without notice.

US Patent: **10619152** & **10844369**

DISCLAIMER

NeoGeneStar LLC and/or affiliate(s) disclaim all warranties with respect to this document, expressed or implied. Including, but not limited to, those of merchantability, fitness for a particular purpose, or non-infringement to the extent allowed by law. In no event shall NeoGeneStar and/or its affiliate(s) be liable whether in contract, tort, warranty, or under any statute or on any other basis for special, incidental, indirect, punitive, multiple or consequential damages in connection with or arising from this document, including but not limited to the use thereof.

Notice to Purchaser: Limited use label License: Research Use Only.

The purchase of this product conveys to the purchaser the limited, non-transferable right to use the purchased amount of the product only to perform internal research for the sole benefit of the purchaser. No right to resell this product or any of its components is conveyed expressly, by implication, or by estoppel.

This product is for internal research purposes only and is not for use in commercial applications of any kind, including, without limitation, quality control and commercial services such as reporting the results of purchaser's activities for a fee or other form of consideration. For information on obtaining additional rights, please contact NeoGeneStar LLC at 100 Randolph Road, Suite 2B, Somerset, NJ, 08873. Tel: (732) 421-4567