

NANOBLOCK

Delivery Systems

NANOBLOCK Protein Delivery Kit

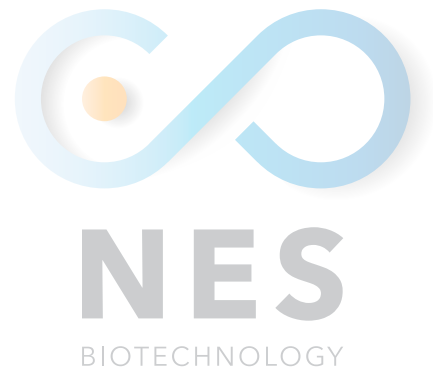


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NANOBLOCK DELIVERY SYSTEM

Introduction

The efficient delivery of bioactive molecules into mammalian cells is a central aspect of research involving cell biology and medical applications. NANOBLOCK Delivery System, gold nanoparticles (AuNP) functionalized to a universal cargo DNA, can be a convenient and efficient method of delivering nucleic acids, proteins, and antibodies to a wide variety of cells including primary cells and animals. In addition, this system can be achieved without causing significant cytotoxicity, even during long incubation periods (1–5 days). Such properties are advantageous in biological applications of bioactive molecules delivery systems, which require careful safety measures.



Overview – The Principle of NANOBLOCK Delivery Systems

Applications

NANOBLOCK delivery system is easily applicable to the delivery of various bioactive molecules into numerous cells in vitro and in vivo without cytotoxicity. This lego-like AuNP delivery system can be used universally for biological experiments including gene expression regulation, intracellular signaling, and transcriptional regulation.

NANOBLOCK Delivery System	Application	Conjugated Bioactive Molecule
NANOBLOCK-Nucleic Acid Delivery Kit	Transient gene knockdown	(ASO) : p53
		(shRNA) : p53, MCL1
	Alternative splicing modulation	(ASO) : MCL1, BCL6
	Sequestering of transcription factor	(DSO) : Estrogen response elements, p53 response elements
	Inhibition of miRNA activity	(AMO) : miR-29b, miR-21
	Inhibition of protein activity	(Aptamer) : beta-catenin, p50
	Synthesis of protein	(mRNA): BAX, dsRED, GFP
NANOBLOCK-Proteins Delivery Kit	Cancer research - intercellular signaling - cell cycle regulation - apoptosis - oncogenesis - transcriptional regulation	(Protein) : BIM, FOXL2, BCL-xL, EGF, RraAV1, Rnase III, AcrA, Enolase (Peptide) : TM-JM1/2, Lamin 406-567, Lamin 406-665
	Antimicrobial research	(AMP) : A3-APO, HPN3, HPA3P

- # ASO : antisense oligonucleotide
- # shRNA : small hairpin RNA
- # DSO : double-stranded oligonucleotide
- # AMO : anti-miRNA oligonucleotide
- # mRNA : messenger RNA
- # AMP : anti-microbial peptide

Example of bioactive molecules successfully applied to biological experiments with NANOBLOCK Delivery Systems

Cell Line	Cell Type	Culture Property	Species
HeLa	Cervical epithelial adenocarcinoma	Adherent	Human
293T	Embryonic kidney (epithelial)	Adherent	Human
K562	Chronic Myelogenous Leukemia (Lymphoblast-like)	Suspension	Human
LoVo	Colon epithelial adenocarcinoma	Adherent	Human
MCF-7	Breast epithelial adenocarcinoma (Mammary gland)	Adherent	Human
KGN	Ovarian granulosa cell (Solid Carcinomas)	Adherent	Human
HepG2	Hepatoblastoma (epithelial-like morphology)	Adherent	Human
A549	Lung epithelial adenocarcinoma	Adherent	Human
A431	Skin/epidermis (epithelial)	Adherent	Human
H1299	Lung epithelial carcinoma	Adherent	Human
COV434	Ovary (Polygonal & fusiform)	Adherent	Human
J1 mouse embryonic stem cell		Adherent	Mouse
Primary cells			
Cervical squamous carcinoma primary cell		Adherent	Human
primary granulosa cell		Adherent	Rat
Animals			
FvB mouse			
Sprague–Dawley rat			
BALB/c nu/nu immunodeficient mouse (Xenograft tumor model, LoVo cells)			
BALB/c nu/nu immunodeficient mouse (Xenograft tumor model, HeLa cells)			

Example of cells efficiently transported bioactive molecules with NANOBLOCK Delivery Systems

RELATED PRODUCTS

High performance transfection efficiency of proteins into the living cells

Catalog No.	Product	Quantity
NES001-01	NANOBLOCK Protein Delivery Kit	1 ml
NES001-02	NANOBLOCK Protein Delivery Kit Bulk	5 X 1 ml

Delivery of non-histag proteins are required additional customized aptamer construction. If you would like to order the customized products, please contact our Technical Support Department by email, info@nesbiotech.com.

High performance transfection efficiency of antibodies into the living cells

Catalog No.	Product	Quantity
NES002-01	NANOBLOCK Antibody Delivery Kit	1 ml
NES002-02	NANOBLOCK Antibody Delivery Kit Bulk	5 X 1 ml

High performance transfection efficiency of nucleic acids into the living cells

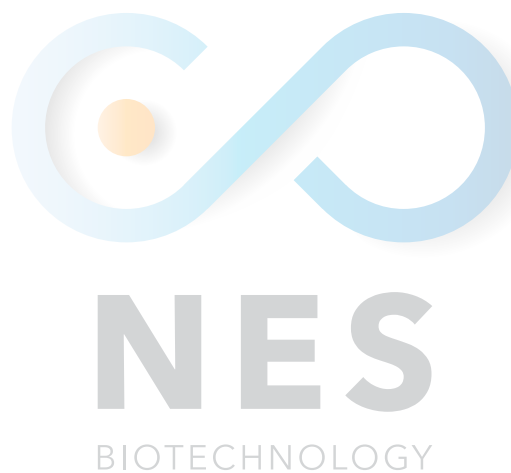
Catalog No.	Product	Quantity
NES003-01	NANOBLOCK Nucleic Acid Delivery Kit	1 ml
NES003-02	NANOBLOCK Nucleic Acid Delivery Kit Bulk	5 X 1 ml

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Technical Support



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NANOBLOCK

PROTEIN DELIVERY KIT

The Nanoblock-Protein Delivery Kit is a unique gold nanoparticle(AuNP)-based protein delivery system that can be used to deliver biologically active proteins or peptides into living cells.

The Nanoblock-Protein Delivery Kit is a simple, efficient, and versatile protein delivery system that allows any his-tagged recombinant protein to be loaded without additional modifications and delivered into mammalian living systems in a manner independent of their size, isoelectric point, and cellular localization. And this system could deliver various proteins into variety of cell types (on primary and stem cells, indicating that its use is not limited to fast-dividing cells) in vitro without showing cytotoxicity. Furthermore, this system was also effective for the local and systemic targeted delivery of proteins in vivo. Therefore, the Nanoblock-Protein Delivery Kit may be used in applications relevant to intercellular signaling, cell cycle regulation, apoptosis, oncogenesis and transcriptional regulation.

KIT CONTENTS

Nanoblock-Protein Delivery Kit (NES001-01)

Catalog No.	Reagent	Quantity	Storage/Stability
NES001-01-01	AuNP ^{His} Reagent (25 nM)	1 ml	4°C for 2 month
NES001-01-02	EGFP ^{His} -control protein	20 µl	4°C for 2 month
NES001-01-03	10x Binding buffer	1 ml	RT for 1 year
NES001-01-04	PBS buffer	1 ml	RT for 1 year
NES001-01-05	MgCl ₂ (0.1 M)	1 ml	RT for 1 year

*Additional Materials Required.

Sterile 1.5 ml microcentrifuge tube

PROTOCOL

QUICK GUIDE

Process	Description	Materials	96-well	24-well	6-well
1	 <p>The cells - Seed to be 70 ~ 80% confluent</p>	Number of the cells ($\times 10^4$)	1.5 ~ 0.5	10 ~ 5	50 ~ 25
2	 <p>  AuNP^{His} Reagent Pre-incubate at 80 °C for 5 min and cool down at RT for 5 min </p>	AuNP ^{His} Reagent (μ l)	2	10	40
3	 <p>  His-tagged protein solution - Prepare protein solution with PBS buffer and MgCl₂ </p>	MgCl ₂ (0.1 M)	1	5	20
		PBS buffer (μ l)	6	30	120
		40 μ M His-tagged protein (μ l)	1	5	20
4	 <p>  Conjugation [AuNP^{His} - (His)protein] - Mix AuNP^{His} reagent and His-tagged protein solution - Incubate for 10-15 min at RT </p>				
5	 <p>Transfection - Add conjugated solution to the cells</p>	Conjugated solution (μ l)	10	50	200

DETAILED INSTRUCTION

Step 1. Cells Preparation

Protocol

- 1) The day before protein delivery experiment, seed/split the cells at appropriate density on appropriate culture vessel for your experiment. We suggest cell number to seed in table 1.
- 2) The suitable cell density will depend on the growth rate and the condition of the cells. Cells should not be more than 70-80% confluent at the time of experiment.

Experimental consideration

Table 1: Suggested number of cells to seed.

Culture vessel	Number of adherent cells	Number of suspension cells	Cell overlay volume
96 well	0.5 – 1.5 X 10 ⁴	0.5 – 1 X 10 ⁵	90 µL
24 well	0.5 – 1 X 10 ⁵	0.5 – 5 X 10 ⁵	450 µL
6 well	2.5 – 5 X 10 ⁵	5 – 20 X 10 ⁵	1.8 mL

Step 2. AuNP^{His} / Protein Conjugation

Protocol

- 1) Prepare AuNP^{His} reagent in microtube.
 - Pre-incubate the AuNP^{His} reagent at 80 °C for 5 min
 - Cool down at RT for 5 min to refold the secondary structure of aptamer.
- 2) Prepare His-tagged protein solution.
 - Adjust His-tagged protein concentration to 40 µM.
 - Make protein solution with PBS buffer and MgCl₂ (0.1 M).
[Note] High concentration of protein can lead to protein precipitation. We recommend that final concentration of protein should NOT be more than 100 µM.
- 3) Construct conjugates of AuNP^{His} and His-tagged protein.
 - Add AuNP^{His} reagent into protein solution and gently tapping.
 - Incubate 10-15 min at room temperature.

* The amounts of reagents depending on culture vessel are in table 2.

Experimental consideration

Table 2: Suggested amount of protein and AuNP^{His}

Culture vessel	40 μ M Protein	MgCl ₂ (0.1 M)	PBS buffer	AuNP ^{His}	Total Volume
96 well	1 μ L	1 μ L	6 μ L	2 μ L	10 μ L
24 well	5 μ L	5 μ L	30 μ L	10 μ L	50 μ L
6 well	20 μ L	20 μ L	120 μ L	40 μ L	200 μ L

* Because proteins differ one from another, reflecting a variety of physical properties, conjugation efficiency of proteins with AuNP^{His} reagent are variable. So, users need to determine optimum conditions to deliver your protein. If your sample is aggregate, add 10X pH Buffer instead of PBS Buffer.

[Note] The presence of NaCl (>300 mM) or MgCl₂ (>10 mM) precipitate the AuNPs. If NaCl and MgCl₂ are present as high concentration in your protein sample, we recommend removing it before proceeding with the delivery assay.

* Any impurities, contaminants present with your protein might affect the delivery efficiency. Also, additives such as detergents, glycerol, sodium azide may inhibit the delivery. Consequently, we suggest using a recombinant protein as pure as possible. Otherwise, it can be removed by dialysis.

Step 3. Transfection

Protocol

- 1) Disperse onto the cells growing in their regular culture medium (with serum)
- 2) Incubate the cells at 37°C in a CO₂ incubator under standard conditions until the evaluation of the protein delivery efficiency (1-48 h). Incubation time will depend on your experimental purpose (for example , incubate 1~4 h to detect delivered protein, and incubate over 24 h to determine cell toxicity of delivered protein)

Experimental consideration

* AuNP^{His} reagent can be used onto cells in serum-free media. In this case, replace the complete culture medium by serum-free medium. This procedure can be more efficient to deliver certain proteins in some cells. After 3-4 h, add some serum-containing medium if further incubation time is necessary.

Appendix

1 QUALITY CONTROLS

2 TROUBLESHOOTING

1. Low delivery efficiency

Check point	Suggestion
Protein purity	Make sure that the recombinant protein is highly pure and devoid of additives such as NaCl, MgCl ₂ BSA or detergents.
AuNP ^{His} reagent amount	Optimize the quantity of AuNP ^{His} reagent as described in the table 2.
Protein amount	Optimize the AuNP ^{His} / protein ratio. We recommend using at least 20 times as much protein as gold.
Cell density	A non-optimal cell density at the time of protein delivery can lead to insufficient uptake. The optimal confluence should range from 70 to 80%.
Cell condition	Cells that have been in culture for a long time (> 8 weeks) may become resistant to the delivery. Use freshly thawed cells that have been passaged at least once. Cells should be healthy and assay during their exponential growing phase. The presence of contaminants (mycoplasma, fungi) alters considerably the delivery efficiency.
Cell culture medium composition	For some cells, protein delivery efficiency can be increased without serum or under reduced serum condition. Thus, assay these cells in serum-free medium during the first 4h of incubation.
Incubation time	The optimal time range between delivery and assay varies with cells, type of protein, kinetics of biological function, etc. The delivery efficiency can be monitored after 4 to 96h. EGFP can be used to quantitatively monitored delivery kinetics.
Old AuNP ^{His} / protein complexes	AuNP ^{His} reagent / protein complexes must be freshly prepared every time. Complexes prepared and stored for more than 1 hour can be aggregated. Depending on the protein, reduce this time to avoid the aggregation which may occur during the complex formation.
Positive control	Ensure that your experiment is properly set up and includes a positive control. The EGFP provided in the kit can be used as positive control for delivery efficiency.
AuNP ^{His} reagent storage	Delivery efficiency can slowly decrease if AuNP ^{His} reagent is kept more than one week at room temperature.

2. Cellular toxicity

Check point	Suggestion
Concentration of AuNP ^{His} / protein too high	Decrease the amount of AuNP ^{His} / protein complexes added to the cells by lowering the protein amount or the AuNP ^{His} reagent. Complexes aggregation can cause some toxicity; prepare them freshly and adjust the ratio as outlined previously.
Unhealthy cells	Check cells for contamination Use new batch of cells Ensure culture medium condition (pH, type of medium used, contamination etc) Cells are too confluent or cell density is too low Verify equipments and materials
Protein is cytotoxic	Use suitable controls such as cells alone, AuNP ^{His} reagent alone or mock delivery (with positive EGFP provided).
Incubation time	Reduce the incubation time of complexes with the cells. Delivery medium can be replaced by fresh medium after 3 to 24 h if necessary.
Protein quality	Use high quality protein as impurities could lead to cell death.
Key protein delivered	If the protein delivered impacts cell survival this can lead to cell death, for instance as demonstrated with the recombinant caspase-3. In this way, the cell death is induced by the proteases.

3 EXAMPLE PROTOCOL AND RESULT

Delivery of a EGFP using a 24-well plate

1. Seed 0.5 to 1×10^5 cells per well in a 24-well plate or on a cover slip. Let grow overnight.
2. Pre-incubate the AuNP^{His} reagent at 80 °C for 5 min and cool down at RT for 5 min.
3. Mix 5 µl of EGFP^{His}, 5 µl of MgCl₂ (0.1 M) and PBS buffer up to 40 µl.
4. Add 10 µl of pre-incubated AuNP^{His} reagent into protein solution, mix by gently tapping.
5. Incubate 10-15 min at room temperature.
6. Distribute mixture 5 onto the cells.
7. Incubate cells in a 5% CO₂ incubator at 37°C for 1-24 hours.
8. After the incubation, wash the cells twice with PBS and proceed with the appropriate assay.

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THANK YOU FOR CHOOSING

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