

nzytech



Your Partner for Life Sciences

NZYTech designs, manufactures and supplies key reagents, kits and master mixes for life science research and molecular diagnostics. Powered by in-house Research and Development, NZYTech engineers products to aid researchers and clinicians around the world - Serving more than 70 Countries, across 5 Continents.

From qPCR Master Mixes and cDNA Synthesis Kits to One-Step RT-qPCR Master Mixes, Key Components for qPCR assays, Bst DNA Polymerases and RNA/DNA Isolation Kits.

NZYTech is a global supplier-of-choice providing IVD kits for SARS-CoV-2 and Respiratory Viruses detection as well as more than 100 qPCR detection kits for Veterinary, Agro Pathogen, Food/Water and Human Pathogen industries searching for molecular research and diagnostic qPCR kits.

NZYTech has global database and supply of Glycobiology products. With more than 80 enzyme categories available, listing more than 1000 CAZymes to choose from. As well as analytical test kits for agro-food and fermentation industries.

Committed to the ever growing field of scientific research, our scientists are deeply engaged in the development of novel products every year. Dedicated to advancing science and being your trusted partner in Life Sciences.







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DNA & RNA purification

PLASMID DNA PURIFICATION GENOMIC DNA PURIFICATION RNA PURIFICATION

NUCLEIC ACIDS CLEAN-UP DNA & RNA PURIFICATION ENZYMES

DNA & RNA PURIFICATION

PLASMID DNA PURIFICATION

NZYTech provides kits for the rapid preparation of highly pure plasmid DNA from recombinant *Escherichia coli* strains at different scales (NZYMiniprep, NZYMidiprep and NZYMaxiprep). The resulting purified plasmid DNA is suitable for use in the most demanding molecular biology applications. Buffers for plasmid purification kits, columns and RNase A are also sold separately for your convenience. Our portfolio also includes kits designed for manual or automated large-scale purification of DNA plasmids from *E. coli* in 96-well format (NZYMiniprep 96-well plate), or for the fast purification of multiples DNA molecules simultaneously (NZYDNA Clean-up 96-well plate).

Plasmid DNA kits	Plasmid type	Typical yield	Scale/ Format
NZYMiniprep	high and low-copy number, BACs or cosmids	up to 45 µg	Mini/spin-column
NZYMiniprep 96-well plate	high and low-copy number	up to 15 µg	Mini/96-well plate
NZYSpeedy Miniprep	high and low-copy number, BACs or cosmids	up to 35 µg	Mini/spin-column
NZYMidiprep	high and low-copy number, BACs or cosmids	up to 150 µg	Midi/gravity flow column
NZYMaxiprep	high and low-copy number, BACs or cosmids	up to 1000 µg	Maxi/gravity flow column
NZYMaxiprep Endotoxin Free	high and low-copy number, BACs or cosmids	up to 500 µg	Maxi/gravity flow column

Miniprep

NZYMinipre	ep
MB01001	50 columns
MB01009	2 x 50 columns
MB01002	200 columns
MB01008	5 x 200 columns
	ep 96-well plate
MB20201	2 plates
MB20202	4 × 2 plates
NZYSpeedy	y Miniprep
MB21001	50 columns
MB21004	2 x 50 columns
MB21002	200 columns
MB21003	5 x 200 columns
-	pin Columns & Collection Tubes
MB18901	50 units
MB18902	4 x 50 units

NZYTech 2023

Procedure	Time	Quality grade	Application
Manual – centrifugation	< 25 min	Molecular Biology	Cloning, sequencing, PCR, enzymatic reactions, transfection
Manual or Automated – under vacuum or centrifugation	~ 45 min	Molecular Biology	Cloning, sequencing, PCR, enzymatic reactions, transfection
Manual – centrifugation	≤ 12 min	Molecular Biology	Cloning, sequencing, PCR, enzymatic reactions, transfection
Manual – centrifugation	~ 70 min	Transfection	Transfection, cloning, sequencing, PCR, enzymatic reactions
Manual – centrifugation	~ 70 min	Transfection	Transfection, cloning, sequencing, PCR, enzymatic reactions
Manual – centrifugation	~ 80 min	Endotoxin free	Transfection, cloning, sequencing, PCR, enzymatic reactions

Midiprep

NZYMidiprep	
MB05003	5 columns
MB05004	20 columns
MB05005	3 x 20 columns

NZYTech Plasmid Midi Columns			
MB19001	5 units		
MB19002	4 x 5 units		

NZYMidiprep, no columns

MB14101 20 preps

Maxiprep

NZYMaxiprep		NZYMaxipr	ep, no columns
MB05101	5 columns	MB14201	5 preps
MB05102	2 x 5 columns		
MB05103	5 x 5 columns		
NZYTech Pla	asmid Maxi Columns	NZYMaxipr	ep Endotoxin Free
NZYTech Pla MB20101	asmid Maxi Columns	NZYMaxipro MB39901	ep Endotoxin Free 5 columns

GENOMIC DNA PURIFICATION

NZY gDNA Isolation kits are spin column silica-based systems designed for the simple and rapid small-scale purification of genomic DNA from various sources. Resulting purified DNA has the highest quality and integrity and is suitable for use in most sensitive downstream applications.

Genomic DNA kits	Sample Material	Typical yield	Scale/Format	Time	Application
NZY Tissue gDNA Isolation kit	Animal tissue, cultured cells, bacte- rial cells, rodent tails, buccal swabs, paraffin embedded tissue and fecal material, whole blood*, serum, plasma, body fluids		Small-scale/ Spin-column	20 min ²	PCR, qPCR, genotyping, sequencing, enzimatic manipulations
NZY Plant/Fungi gDNA Isolation kit	Plant and fungal tissues	up to 30 µg	Small-scale/ Spin-column	30 min	PCR, qPCR, genotyping, sequencing, enzimatic manipulations
NZY Soil gDNA Isolation kit	Soil, sludge, sediment and stool samples	2-10 µg	Small-scale/ Spin-column	~ 15 min²	PCR, qPCR, genotyping, sequencing, enzimatic manipulations
NZY Microbial gDNA Isolation kit	Cell pellets of gram-positive and gram-negative bacteria	5-25 µg¹	Small-scale/ Spin-column	~ 15 min	PCR, qPCR, genotyping, sequencing, enzimatic manipulations

* Fresh, frozen or treated; ¹ From 30 mg wet weight cell pellet; ² Excluding lysis step.

NZY Tissue gDNA Isolation kit		NZY Plan	t/Fungi gDNA Isolation kit		
MB13502	50 columns	MB17701	MB17701 50 columns		
MB13503	200 columns	MB17702	MB17702 4 x 50 columns		
NZY Soil gDNA Isolation kit		NZY Micro	bial gDNA Isolation kit		
MB21802	50 columns	MB21702	50 columns		

Different sample sources (animal tissues, cultured cells, bacterial cells, mouse tails, yeast, stool, forensic and clinical samples) can be used for gDNA isolation with NZY Tissue gDNA Isolation kit

All protocols are optimized for different biological matrices to provide high quantity of pure genomic DNA

NZYTech recently discontinued NZY Blood gDNA Isolation kit (Cat. No. MB136) since the NZY Tissue gDNA Isolation kit serves for the same use

RNA PURIFICATION

NZYTech's RNA Isolation kits are spin column silica-based systems designed for the easy and fast purification of RNA at the highest integrity from a variety of sources, such as animal tissues, cultured cells and bacteria (NZY Total RNA Isolation kit), serum, plasma, saliva, nasal samples, blood and environmental samples (NZY Viral RNA Isolation kit). Recently, NZYTech developed the NZY Mag Viral RNA/DNA Isolation kit, IVD (MD04881/2), which is a magnetic bead technology-based nucleic acid purification kit designed to recover RNA and DNA from viral particles contained in transport medium from human respiratory swabs.

ZY Total RNA Isolation kit
B13402 50 columns
Nase Cleaner
1B16001 500 mL
uffer NVL
IB40801 100 mL

NUCLEIC ACIDS CLEAN-UP

NZYGelpure kit is designed for the purification of DNA (50 bp to 20 kb) from TAE/TBE agarose gels and for direct purification of enzymatic reactions. NZYGelpure purification kit utilizes a silica-gel based membrane which selectively adsorbs up to 20 µg of DNA fragments in the presence of specialized binding buffers. NZYDNA Clean-up in 96 well format is available for high-through-put protocols.

DNA Clean-Up

NZYGelpure		NZYDNA Clean-up 96-well plate	
MB01101	50 columns	MB20001	2 plates
MB01104	2 x 50 columns	MB20002	4 x 2 plates
MB01102	200 columns		
MB01103	5 x 200 columns		

NZYGelpure: includes a pH indicator, allowing to analyze the optimal pH for DNA binding

DNA & RNA PURIFICATION ENZYMES

NZY DNase I	
MB19901	200 U/vial

100 mg

500 mg

Proteinase K MB01901

MB01902

NZY RNAse A		
MB18701	100 mg	



end-point pcr

DNA POLYMERASES & OTHER PCR ENZYMES MASTER MIXES MULTIPLEX MIXES COMPONENTS/SUPPLEMENTS

END-POINT PCR

DNA POLYMERASES & OTHER PCR ENZYMES

NZYTech's DNA polymerases offer great performance expressed by high yields and extreme sensitivity. Each enzyme presents different features to cover a wide range of applications. Thus, we organized enzymes by applications allowing you to direct your selection to the enzymes that most suit your experiment. The Supreme version of each enzyme displays a hot-start-like activity for improved specificity and sensitivity.

	DNA polymerase	Features	Product length	Extension (sec/kb)	Proof reading	Sensitivity*	Product overhang	Cat. No.
ē	NZYTaq II	High yields with minimal optimization	≤ 6 kb	15-30	No	5 pg	3'-A	MB354
Routir	Supreme NZYTaq II	Hot-start-like version of NZYTaq II	≤ 6 kb	15-30	No	1 pg	3'-A	MB355
Power & Routine	Speedy NZYTaq	Powerful PCR in 30-60 min	≤ 6 kb	5-10#	No	50 pg	3'-A	MB403
P	Speedy Supreme NZYTaq	Hot-start-like PCR in 15-60 min	≤ 6 kb	5-10#	No	l pg	3'-A	MB390
	NZYProof	High fidelity proofreading enzyme	≤ 10 kb	60	Yes	5 ng	Blunt	MB146
ility	Supreme NZYProof	Hot-start-like version of NZYProof	≤ 10 kb	15-30	Yes	lng	Blunt	MB283
Fidelity	Speedy NZYProof	High fidelity PCR in 30-60 min	≤ 2 kb	5	Yes	10 ng	Blunt	MB404
	Speedy Supreme NZYProof	Fast & powerful fidelity PCR	≤ 2 kb	2	Yes	lng	Blunt	MB436
Long	NZYLong	Increased processivity for extended PCR	≤ 20 kb	60	No	5 ng	Mixed	MB003
2	Supreme NZYLong	Hot-start-like version of NZYLong	≤ 25 kb	60	No	lng	Mixed	MB331

*Measures the minimal quantity of human gDNA required to amplify a 1 kb template

* For DNA fragments lower than 2kb, use 5 sec/kb

Powerful PCR

Taq II [ONA polymerase	NZYTaq II w	vith 5x Gel Load Reaction Buffer
35401	500 U	MB36401	500 U
35402	1000 U	MB36402	1000 U
35403	2500 U	MB36403	2500 U
upreme N	ZYTaq II DNA polymerase	Supreme N	ZYTaq II with 5x Gel Load Buffe
IB35501	50011	MB36501	500 U
וטכככםו	500 U		5000
B35502	1000 U	MB36502	1000 U

Supreme NZYTaq II DNA polymerase: High sensitivity, reproducibility and performance

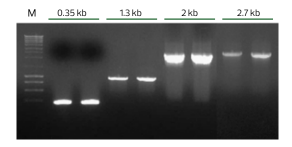
	Supreme NZYTaq II (batch A)				Supreme NZYTaq II (batch B)				Competitor enzyme				me						
М	1	2	3	4	5	NC	1	2	3	4	5	NC	1	2	3	4	5	NC	М
100	-	-	_				_	-	-	1			_	-					111 III

Amplification of a 1-kb fragment from human genomic DNA using a 1:5 dilution series ranging from 20 ng to 32 pg. 1) 20 ng; 2) 4 ng; 3) 0.8ng; 4) 0.16 ng; 5) 0.032 ng; NC) No template control M) NZYDNA Ladder III

Speedy NZYTaq DNA polymerase						
MB40301	125 U					
MB40302	500 U					

Speedy Supreme NZYTaq DNA polymerase						
MB39001	500 U					
MB39002	1000 U					
MB39003	2500 U					

Speedy NZYTaq DNA polymerase: Successful amplification of different-sized targets with limited extension time (5 sec/kb)



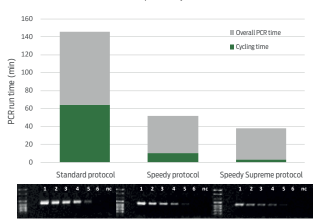
High Fidelity PCR

MB14601 125 U MB28301 125 U MB14602 500 U MB28302 500 U MB14603 1000 U MB28303 1000 U	NZYProof DI	NA polymerase	Supreme NZYProof DNA polymerase			
	MB14601	125 U	MB28301	125 U		
MB14603 1000 U MB28303 1000 U	MB14602	500 U	MB28302	500 U		
	MB14603	1000 U	MB28303	1000 U		

Speedy NZYProof DNA polymerase					
MB40401	125 U				
MB40402	500 U				

Speedy Supre	me NZYProof D	NA polymerase
IVIDZOJUJ	10000	

Speedy Supreme NZYProof DNA polyme								
MB43601	125 U							
MB43602	500 U							



Speedy Supreme NZYProof DNA polymerase: The best choice for ultra-rapid fidelity PCRs

Amplification of a 1-kb fragment from human genomic DNA using dilution series ranging from 80 ng to 0.625 ng. The PCR was conducted with **Speedy Supreme NZYProof DNA polymerase** (2.5 U) in a 50 μ L final volume in different cycling protocols (standard, speedy and speedy supreme).

The cycling times for each protocol are shown in blue, while the overall PCR times on a Thermal Cycler System with a ramp rate of 4°C/sec are shown in grey. PCR products analysis through agarose gel electrophoresis (1% v/v TAE) is presented below to the chart, using NZYDNA Ladder VIII (Cat. No. MB175); numbers represent the amount of DNA template used.

1) 80 ng; 2) 40 ng; 3) 20ng; 4) 10 ng; 5) 2.5 ng; 6) 0.625 ng; NC) No template control

Long PCR

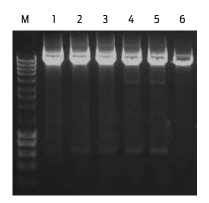
NZYLong DNA polymerase					
125 U					
500 U					
1000 U					

Supreme NZYLong DNA polymerase					
MB33101	125 U				
MB33102	500 U				
MB33103	1000 U				

NZYLong DNA polymerase: Successful amplification of DNA fragments from 0.5 to 15 kb

М	0.5	1	2.5	5	10	15	М	(kb)
					•	-		
			-					
		-					=	
	-							

Supreme NZYLong DNA polymerase: High sensitivity with large-sized fragments using different amounts of *E. coli* gDNA



1) 40 ng; 2) 20 ng; 3) 10 ng; 4) 5 ng; 5) 2.5 ng; 6) 1.25 ng ; M) NZYDNA Ladder III

		R Enzymes
NZY Uracil-DNA Glycos		IA Glycosylase
	MB44601	100 µL
	MB44602	3 x 100 µL

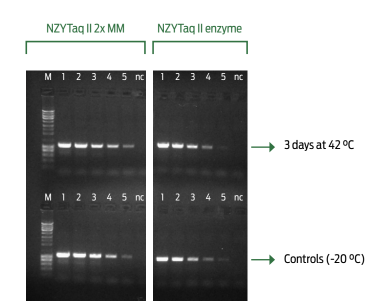


MASTER MIXES

NZYTech presents a variety of master mixes containing all PCR components (except primers and template) at optimal concentrations for efficient DNA amplification. Green Master Mixes allow amplification reactions to be directly loaded onto agarose gels. The mixes are formulated in 2x concentrated solutions.

DNA polymerase	Master Mix	Loading dye	Cat. No.
	NZYTaq II 2x Green Master Mix	Yes	MB358
NZYTaq II DNA polymerase	NZYTaq II 2x Colourless Master Mix	No	MB357
	Supreme NZYTaq II 2x Green Master Mix	Yes	MB360
Supreme NZYTaq II DNA polymerase	Supreme NZYTaq II 2x Colourless Master Mix	No	MB359
	Speedy NZYTaq 2x Green Master Mix	Yes	MB362
speedy NZYTaq DNA polymerase	Speedy NZYTaq 2x Colourless Master Mix	No	MB361
	Speedy Supreme NZYTaq 2x Green Master Mix	Yes	MB391
speedy Supreme NZYTaq DNA polymerase	Speedy Supreme NZYTaq 2x Colourless Master Mix	No	MB392
	NZYProof 2x Green Master Mix	Yes	MB287
IZYProof DNA polymerase	NZYProof 2x Colourless Master Mix	No	MB288
	Supreme NZYProof 2x Green Master Mix	Yes	MB285
Supreme NZYProof DNA polymerase	Supreme NZYProof 2x Colourless Master Mix	No	MB286
	NZYLong 2x Green Master Mix	Yes	MB139
IZYLong DNA polymerase	NZYLong 2x Colourless Master Mix	No	MB332
	Supreme NZYLong 2x Green Master Mix	Yes	MB333
Supreme NZYLong DNA polymerase	Supreme NZYLong 2x Colourless Master Mix	No	MB334

NZYTaq II DNA polymerase: Robust stability at high temperatures



Activity of NZYTaq II DNA polymerase (individual enzyme and master mix formats) after 3 days at high temperature storage. NZYTaq II DNA polymerase and NZYTaq II 2x Green Master Mix were stored during 3 days at 42 °C. Controls of the same batches were kept at -20 °C. Then, the activity was assayed in a PCR experiment to amplify a 1-kb fragment using as template a dilution series of human genomic DNA (from 20 ng to 32 pg); numbers represent the amount of DNA template used (1: 20 ng; 2: 4 ng; 3: 0.8ng; 4: 0.16 ng; 5: 0.032 ng; nc: negative control without DNA). M: NZYDNA Ladder III (Cat. No. MB044)

NZYTaq II-based master mixes

NZYTaq II 2x Green Master Mix	
35801 500 U (100 x 50 μL rxs)	
1000 U (200 x 50 µL rxs)	
3 5000 U (1000 x 50 µL rxs)	

Supreme NZYTaq II-based master mixes

Supreme NZYTaq II 2x Green Master Mix	
MB36001	500 U (100 x 50 µL rxs)
MB36002	1000 U (200 x 50 µL rxs)
MB36003	5000 U (1000 x 50 µL rxs)

Speedy Supreme NZYTaq-based master mixes

Speedy Supreme NZYTaq 2x Green MM	
MB39101	500 U (100 x 50 µL rxs)
9102	1000 U (200 x 50 µL rxs)
MB39103	5000 U (1000 x 50 µL rxs)

Speedy NZYTaq-based Master Mices

NZYProof-based master mixes

NZYProof 2x Green Master Mix	
MB28701	500 U (100 x 50 µL rxs)
28702	1000 U (200 x 50 µL rxs)
IB28703	5000 U (1000 x 50 µL rxs)

Supreme NZYProof-based master mixes

Supreme NZ	YProof 2x Green Master Mix
MB28501	500 U (100 x 50 µL rxs)
MB28502	1000 U (200 x 50 µL rxs)
MB28503	5000 U (1000 x 50 µL rxs)

Supreme NZY	Proof 2x Colourless Master Mix
MB28601	500 U (100 x 50 µL rxs)
MB28602	1000 U (200 x 50 µL rxs)
MB28603	5000 U (1000 x 50 µL rxs)

NZYLong 2x Green Master Mix:

Convenience of direct gel loading of different-sized PCR products in a high-throughput PCR experiment (96 amplifications)

Green Master Mixes offer convenience in colony PCR and high-throughput applications





NZYLong-based master mixes

NZYLong 2x	Green Master Mix
MB13902	500 U (100 x 50 µL rxs)
MB13903	1000 U (200 x 50 µL rxs)
MB13904	5000 U (1000 x 50 µL rxs)

NZYLong 2x C	olourless Master Mix
MB33201	500 U (100 x 50 µL rxs)
MB33202	1000 U (200 x 50 µL rxs)
MB33203	5000 U (1000 x 50 µL rxs)

Supreme NZYLong-based master mixes

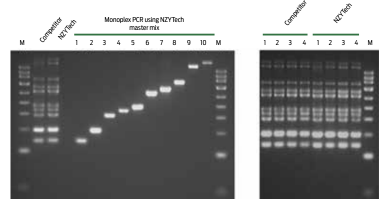
Supreme NZYLong 2x Green Master Mix		Supreme N	ZYLong 2x Colourless Master Mi
MB33301	500 U (100 x 50 µL rxs)	MB33401	500 U (100 x 50 µL rxs)
MB33302	1000 U (200 x 50 µL rxs)	MB33402	1000 U (200 x 50 µL rxs)
MB33303	5000 U (1000 x 50 µL rxs)	MB33403	5000 U (1000 x 50 µL rxs)

MULTIPLEX MIXES

NZYTech developed two PCR master mixes based on our non-proofreading and proofreading enzymes for the simultaneous amplification of multiple DNA fragments (up to 15 targets) in a single tube. The mixes are provided in 2x concentrated solutions, in green (direct-gel load) and colourless versions.

Multiplex F	PCR NZYTaq 2x Green Master Mix
MB33501	500 U (100 x 50 µL rxs)
MB33502	1000 U (200 x 50 µL rxs)
MB33503	5000 U (1000 x 50 µL rxs)
Multiplex F	PCR NZYProof 2x Green Master Mix
MB33701	500 U (100 x 50 µL rxs)
MB33702	1000 U (200 x 50 µL rxs)

Multiplex PCR NZYTaq 2x Green Master Mix: High performance in a multiplex PCR reaction amplifying 10 different fragments from human genomic DNA



1) 65 ng; 2) 30 ng; 3) 15 ng; 4) 7,5 ng

COMPONENTS/SUPPLEMENTS

dNTPs

dNTPs NZYSet		
MB08701	100 mM (4 x 0.25 mL)	

dNTPs NZYMix		
MB08601	25 mM each, 1 mL	
MB08602	25 mM each, 5 mL	
MB08603	10 mM each, 0.2 mL	
MB08604	10 mM each, 1 mL	
MB08605	10 mM each, 5 mL	

PCR Supplements

NZYTech supplements are available to optimize PCR conditions with NZYTaq II DNA polymerase.

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NZYTaq 5x Optimizer Solution		
MB06001	1 mL	
MB06002	3x1mL	

NZYTaq 2x GC-Enhancer Solution

MB14301	1 mL
MB14302	5x1mL

MOLECULAR BIOLOGY

real-time pcr

qPCR MASTER MIXES ONE-STEP RT-qPCR KITS ONE-STEP RT-qPCR MASTER MIXES qPCR COMPONENTS

anda

REAL-TIME PCR

qPCR MASTER MIXES

NZYTech has developed and optimized two of the most widely used real-time fluorescent PCR chemistries: the probe-detection technology (Probe Mixes) and the intercalating green dye chemistry (Green Mixes). NZYSupreme mixes are ultra-sensitive mixes developed with a dual hot-start mode and suitable for standard thermal cycling protocols. NZYSpeedy mixes present a higher performance for the faster real-time PCR protocols. Passive reference dye based on ROX[™] dye is used to normalize the fluorescent reporter signal in real-time PCR. NZYTech provides highly optimized mixes that are compatible with different thermocyclers available on the market. NZYTech recently increased its portolio of real-time qPCR products by introducing lyophilized master mixes, which are very stable formulations that allow an eco-friendly and cost-effective room temperature shipment.

Choose the mix with the reference dye that is most appropriate for your instrument with the help of the qPCR Selection Guide below.

	ROX plus	ROX	(no ROX*)
Agilent			
AriaMX		8	
MX3000P™, MX3005P™, MX4000P™		8	
Applied Biosystems™			
7000/7300/7700	8		
7500/7500 FAST		8	
7900/7900HT/7900HT FAST	8		
QuantStudio™ 5, 6, 7, 12k Flex/ViiA7™		8	
StepOne™/StepOne™plus	8		
Bio-Rad®			
CFX Opus/CFX96™/CFX384™			8
Opticon™/Opticon™ 2			8
Fluidigm®			
BioMark™		8	
Illumina®			
Eco™			8
Qiagen			
Rotor-Gene™ 3000			8
Rotor-Gene™ 6000			8
Rotor-Gene™ Q			8
Roche			
Lightcycler® 96			8
Lightcycler® 480			8
Lightcycler® Nano			8

* For qPCR instruments that require ROX reference dye, it is possible to add ROX in a separate step, according to instructions provided in the respective Master Mix product brochure.

Green Master Mixes

NZYSupreme qPCR Green MM (2x), ROX plus		
MB44001	2 mL (200 x 20 µL rxs)	

MB44001	2 ML (200 X 20 µL IXS)
MB44002	5 mL (500 x 20 µL rxs)
MB44003	20 mL (2000 x 20 µL rxs)

MB44101	2 mL (200 x 20 µL rxs)
MB44102	5 mL (500 x 20 µL rxs)
MB44103	20 mL (2000 x 20 µL rxs)

NZYSupreme qPCR Green Master Mix (2x)		
MB41901	2 mL (200 x 20 µL rxs)	

MB41902	5 mL (500 x 20 µL rxs)
MB41903	20 mL (2000 x 20 µL rxs)

Hot-start like activity
Reproducibility
Highly sensitive

Also Availa	able:
NZYSpeedy	qPCR Green Master Mix (2x), ROX plus
MB22201	2 mL (200 x 20 µL rxs)
MB22202	5 mL (500 x 20 µL rxs)
MB22203	20 mL (2000 x 20 µL rxs)
NZYSpeedy	qPCR Green Master Mix (2x)
MB22401	2 mL (200 x 20 µL rxs)
11000/00	5 mL (500 x 20 µL rxs)
MB22402	JIIIE (JOO X 20 PE IXS)

Probe Master Mixes

NZYSupreme PCR Probe MM (2x), ROX plus NZYSupreme PCR Probe MM (2x), ROX MB43901 2 mL (200 x 20 µL rxs) MB43801 2 mL (200 x 20 µL rxs) MB43902 5 mL (500 x 20 µL rxs) MB43802 5 mL (500 x 20 µL rxs) MB43903 2 0 mL (2000 x 20 µL rxs) MB43803 2 0 mL (2000 x 20 µL rxs)

NZYSupreme qPCR Probe Master Mix (2x)		
MB41601	2 mL (200 x 20 µL rxs)	
MB41602	5 mL (500 x 20 µL rxs)	
MB41603	20 mL (2000 x 20 µL rxs)	

Lyo NZYSupreme qPCR Probe Master Mix (2x)

MB41702 For 1.5 mL (150 x 20 µL rxs)

NZYSupreme Multiplex qPCR Probe MM (2x))
---	---

MB45201	2 mL (200 x 20 µL rxs)
MB45202	5 mL (500 x 20 µL rxs)
MB45202	20 mL (2000 x 20 µL rxs)

MB45301 For 1.5 mL (150 x 20 µL rxs)

High efficiency	
High specificity	
Efficient multiplexing	

Also Availa	ble:
NZYSpeedy	qPCR Probe Master Mix (2x), ROX plus
MB22801	2 mL (200 x 20 µL rxs)
MB22802	5 mL (500 x 20 µL rxs)
MB22803	20 mL (2000 x 20 µL rxs)
NZYSpeedvo	gPCR Probe Master Mix (2x)
MB23001	2 mL (200 x 20 µL rxs)
MB23002	5 mL (500 x 20 µL rxs)
MB23003	20 mL (2000 x 20 µL rxs)

ONE-STEP RT-qPCR KITS

NZYTech provides One-step real-time PCR kits designed to directly amplify RNA samples on your real-time PCR instrument. These kits were developed to enable cDNA synthesis from input RNA followed by PCR amplification of the cDNA in the same reaction well, with no extra hands-on requirement or further reagent addition. This not only reduces the number of sample manipulations but also saves time. One-step kits are available for Probe and Green detection. Choose the One-step real-time PCR kit that most suits your experiment and that is most appropriate for your instrument through the analysis of the qPCR Selection Guide presented in page 22.

One-Step RT-qPCR Green Kits

Fast

MB34501

MB34502

🛕 One-step R	T-qPCR Green Kit (2x), ROX plus	\Lambda One-step R	T-qPCR Green Kit (2x)
MB34401	100 reactions	MB34601	100 reactions
MB34402	500 reactions	MB34602	500 reactions
⚠ One-step R	T-qPCR Green Kit (2x), ROX		

One-Step RT-qPCR Probe Kits

100 reactions

500 reactions

Fast

One-step RT-qPCR Probe Kit (2x), ROX plus

MB35001	100 reactions
MB35002	500 reactions

One-step RT-qPCR Probe Kit (2x), ROX

MB35101	100 reactions
MB35102	500 reactions

One-step RT-qPCR Probe Kit (2x)

,,, _,, _	
MB35201	100 reactions
MB35202	500 reactions

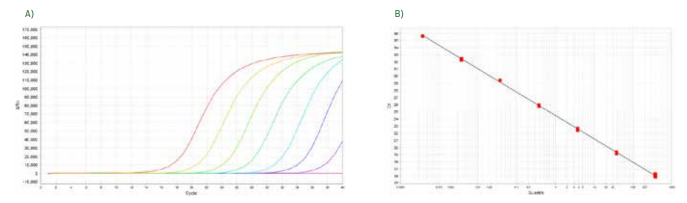
ONE-STEP RT-qPCR MASTER MIXES

NZYTech provides One-step real-time qPCR master mixes containing all required components (except primers/probe and template) to perform reverse transcription and qPCR amplification in a single-step. This offers great convenience and minimizes the risk of errors and contaminations. These master mixes were engineered with a dual hot-start enzyme control mechanism to provide the highest detection sensitivity. In addition, the latest developments in PCR enhancers were introduced. Lyophilized versions are also available as very stable room temperature options.

One-Step RT-qPCR Probe Master Mixes

NZYSupreme RT-qPCR Probe MM (2x)			me Multiplex RT-qPCR Probe MM (2x)
MB41401	2 mL (200 x 20µL rxs)	MB44201	2 mL (200 x 20 µL rxs)
MB41402	5 mL (500 x 20 µL rxs)	MB44202	5 mL (500 x 20 µL rxs)
MB41403	20 mL (2000 x 20 µL rxs)	MB44203	20 mL (2000 x 20 µL rxs)

NZYSupreme One-step RT-qPCR Probe Master Mix: High-performance across a wide dynamic range of very low RNA inputs (<0.5 pg)



A 10-fold serial dilution of total RNA from mouse liver (375 ng to 0.375 pg) was used as template for a one-step real-time RT-qPCR experiment to detect the rpl27 housekeeping gene. Panel A: evidence of the high performance and linearity when using NZYSupreme One-step RT-qPCR Probe Master Mix. Panel B: standard curve (slope: -3.28; efficiency: 101.7%).

Ultra-sensitive: Detect low-copy number targets (8 copies) Dual Hot-Start mode for Supreme versions Include RNase Inhibitor Efficient multiplexing Lyo formats stable at Room Temperature



Lyo One-step RT-qPCR Master Mixes

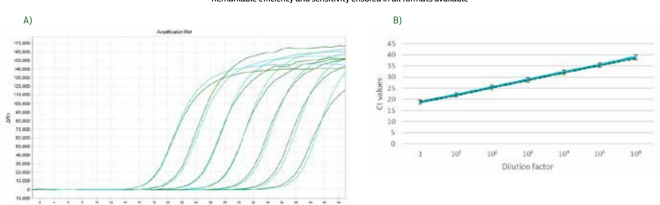
Lyo NZYSupreme RT-qPCR Probe MM (2x)

MB41501 For 1.5 mL (150 x 20 µL rxs)

Lyo NZYSupreme Multiplex RT-qPCR Probe MM (2x)

MB44301 For 1.5

For 1.5 mL (150 x 20 µL rxs)



NZYSupreme One-step RT-qPCR Probe Master Mix vs Lyo NZYSupreme One-step RT-qPCR Probe Master Mix: Remarkable efficiency and sensitivity ensured in all formats available

High sensitivity and linearity of the two formulations of NZYSupreme One-step RT-qPCR Probe Master Mix – liquid (with glycerol) and lyophilized (tested after rehydrated with the respective reconstitution buffer), across a wide range of input RNA (from 375 ng to 0.375 pg) to amplify the rpl27 mouse gene. Panel A: Amplification curves evidencing the high reproducibility between the liquid (blue curves) and lyophilized (green curves) formats. Panel B: Comparison of Ct variation across template dilutions.

qPCR COMPONENTS

DEPC-treated Water	
MB43701	5x1mL



RNA & cDNA

RNA SYNTHESIS

cDNA SYNTHESIS

RNases

RNA PROTECTION

RNA & cDNA

RNA SYNTHESIS

NZYTech offers convenient kits for the *in vitro* transcription of DNA into RNA using a T7 RNA polymerase that is highly specific for T7 phage promoters. The enzyme is also provided separately for your routine protocols.

NZY T7 RNA	A Synthesis kit	NZY T7 Hig	h Yield RNA Synthesis ki
MB35301	50 reactions	MB36301	50 reactions
T7 RNA poly	ymerase		
MB08001	10000 U (20 U/µL)		
MB08003	10000 U (200 U/µL)		

cDNA SYNTHESIS

Reverse transcriptases

NZY M-MuLV and NZY Reverse Transcriptases lack 3'-5' exonuclease activity and have no intrinsic RNase H activity. NZY Reverse Transcriptase is a thermostable (50-55 °C) and sensitive enzyme with fast synthesis capacity (30 min reaction). Supreme NZYReverse Transcriptase was recently developed to increase inhibitor resistance, range and thermostability.

NZY M-MuL	V Reverse Transcriptase	NZY Revers	se Transcriptase
AB08301	20000 U	MB12401	20000 U
IB08302	100000 U	MB12402	100000 U
yo NZY Re	verse Transcriptase		
B40901	100000 U		
			Standard Curve
	NZYReverse Transcrip	tase vs Competitor T	:
55	Anplife	ation Plot	
14 13			
11		AFF	NZYReverse Transcriptase
*		/////	Slope: -3.298; Eff%: 101.03
§ 7 *			Standard Curve
8 4	/		:
2	/		
,		Cycle	5
			Competitor T Slope: -3.346; Eff%: 99.02

NZY

NZYReverse Transcriptase vs Competition. Two-step RT-qPCR to detect the GAPDH gene from mouse total RNA (dilutions from 1 µg to 0.1 ng). After reverse transcription, the cDNA obtained was used in a qPCR assay with NZYSupreme qPCR Green Master Mix (2x) (Cat. No. MB419).

NZYTech 2023

cDNA kits

NZYTech offers convenient, reliable and cost-effective kits to generate high quality cDNA for different downstream applications, such as standard PCR, cDNA library construction, or two-step RT-PCR assays. cDNA kits contain all the components required to synthesize first-strand cDNA (except the template RNA), at optimal conditions. A mixture of NZY Ribonuclease Inhibitor and one of the NZYTech's reverse transcriptases is included in all kits. NZY First-Strand cDNA Synthesis kits provide more specific synthesis of cDNA, while NZY M-MuLV First-Strand Synthesis kits are a cost-effective alternative, also providing high yields of full-length cDNA at low reaction temperatures.

Features	NZY First-Strand cDNA Synthesis Kits	NZY M-MuLV First-Strand cDNA Synthesis Kits
Reverse Transcriptase	NZY Reverse Transcriptase	NZY M-MuLV Reverse Transcriptase
Product length	up to 7 kb	up to 7 kb
Optimal reaction temperature	50 °C	37 ºC
Reaction time	30 min	50 min
Sensitivity	10 pg – 5 µg total RNA	10 pg – 5 µg total RNA
Amplification of GC-rich templates	yes	-
Amplification of secondary structure-rich templates	yes	-
Available as kits with separate oligos	yes	yes

NZY First-S	trand cDNA Synthesis kit	NZY cDNA S	Synthesis kit, sep. oligos
MB12501	50 reactions	MB17001	50 reactions
MB12502	250 reactions	MB17002	250 reactions
\Lambda NZY M-MuL	V First-Strand cDNA Synthesis kit	NZY M-MuL	V cDNA Synthesis kit, sep oligos
MB17201	50 reactions	MB17301	50 reactions
MB17202	250 reactions	MB17302	250 reactions
мв40001 RNA prir		Denderste	
	, primer mix	 Random he	
MB12801	27 μg (100 μL)	MB12901	25 μg (500 μL)
) ₁₈ primer: hybridizes to the poly(v used when cDNA is used for clon		



Random hexamers: priming ssDNA or RNA for extension by DNA polymerase or Reverse transcriptases; during cDNA synthesis, random hexamer will perform random priming throughout the entire length of the RNA to generate a cDNA pool containing various lengths of cDNA

RNases

NZY RNase H	(E. coli)
MB08501	250 U
MB08502	1250 U

 NZY RNase A

 MB18701
 100 mg

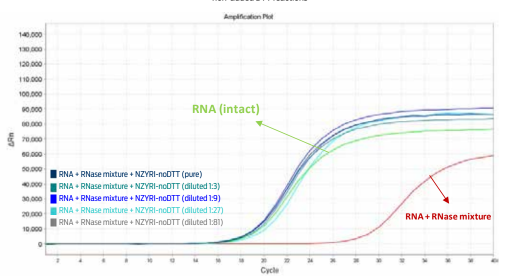
RNA PROTECTION

NZYTech Ribonuclease Inhibitors are active againts RNases (E. C. 3.1) of the pancreatic type (RNase A, B and C) and are useful in applications where eukaryotic RNase contamination is a potential problem. For reactions that do not tolerate higher levels of DTT, please use the oxidation-resistant NZY Ribonuclease Inhibitor (no DTT). RNase Cleaner is a reagent that completely removes RNase contamination surfaces and plasticware, allowing to maintain a RNAse-free working area.

ZY Ribonu	uclease Inhibitor	Lyo NZY Ri	bonuclease Inhibitor
MB08401	2500 U	MB41102	20000 U
MB08402	5 x 2500 U	MB41103	5 x 20000 U
NZY Ribonu	uclease Inhibitor (no DTT)	Lyo NZY Ri	bonuclease Inhibitor (no DTT)
	uclease Inhibitor (no DTT) 2500 U	Lyo NZY Ri MB41202	bonuclease Inhibitor (no DTT) 100000 U
MB41001	2500 U	MB41202	100000 U

RNase Cleane

MB16001 500 mL



NZYRibonuclease Inhibitor (no DTT): Efficient protection of RNA against RNases in non-added DTT reactions

The ability of NZYRibonuclease Inhibitor (no DTT) to inhibit ribonuclease (RNase) activity was tested by pre-mixing 125 ng of mouse RNA with different amounts of the inhibitor enzyme (1:3 dilution series) and a cocktail of RNases from serum origin. No DTT was added in the reactions, which were incubated at 37 °C for 1 hour. A negative control (without NZYRibonuclease Inhibitor (no DTT)) and a positive control (only RNA not exposed to the RNases mixture) were introduced. The integrity of RNA was judged through a real-time one-step RT-qPCR experiment. Complete preservation of RNA integrity is observed in the presence of NZY Ribonuclease Inhibitor (no DTT) (in all dilutions tested), as measured by the successful amplification of the desired target in the real-time RT-PCR assay (the signal overlaps to that emitted by the positive control – intact RNA).

MOLECULAR BIOLOGY

restriction enzymes

CONVENTIONAL

SPEEDY

RESTRICTION ENZYMES

NZYTech offers a vast portfolio of restriction enzymes used in recombinant DNA technology. We present two different types of enzymes: Conventional and Fast Digestion (Speedy). Each conventional restriction enzyme is provided with a specific reaction buffer in which the enzyme is 100% active. For double digestions, we recommend to use the 10x NZYBuffer U (sold separately). Speedy restriction enzymes are a new generation of DNA modifying enzymes that were developed for rapid DNA digestion (digestion periods range from 5-15 min) and are 100% active in the 10x NZYSpeedy Buffers Colourless or Orange.

CONVENTIONAL

Ascl	MB23101	500 U	Hpal GTT‡AAC	MB07101	500 U
GG1CGCGCC	MB23102	2500 U		MB07102	2500 U
BglII	MB06501	1000 U	Mbol	MB24101	1000 U
A↓GATCT	MB06502	5000 U	↓GATC	MB24102	5000 U
Ddel	MB23601	500 U	Mlui	MB24301	1000 U
C↓TNAG	MB23602	2500 U	A↓CGCGT	MB24302	5000 U
Dpnl	MB07801	100 U	Ncol	MB06601	500 U
G(mA)↓TC	MB07802	1000 U	C↓CATGG	MB06602	2500 U
Dpnll	MB23301	1000 U	Pstl CTGCA↓G	MB07301	4000 U
↓GATC	MB23302	5000 U		MB07302	20000 U
EcoRI	MB06701	5000 U	Sali	MB07701	2000 U
G↓AATTC	MB06702	25000 U	G↓TCGAC	MB07702	10000 U
HindIII	MB07001	5000 U	Taql	MB23501	1000 U
A↓AGCTT	MB07002	25000 U	T↓CGA	MB23502	5000 U
Hinfl	MB23901	500 U	Xhol	MB07401	2000 U
G↓ANTC	MB23902	2500 U	C↓TCGAG	MB07402	10000 U

NZYTech 2023

SPEEDY

s	Speedy Ascl GG↓CGCGCC	MB23201	50 reactions	
G		MB23202	250 reactions	
s	peedy BglII	MB09301	100 reactions	
	A↓GATCT	MB09302	500 reactions	
S	Speedy Dpnll	MB23401	100 reactions	
	↓GATC	MB23402	500 reactions	
S	Speedy EcoRI	MB09501	500 reactions	
	G↓AATTC	MB09502	2500 reactions	
Sp	eedy HindIII	MB09701	500 reactions	
	A↓AGCTT	MB09702	2500 reactions	
S	peedy Hinfl	MB24001	50 reactions	
	G↓ANTC	MB24002	250 reactions	
S	peedy Hpal	MB09801	50 reactions	
	GTT↓AAC	MB09802	250 reactions	

Speedy Mbol ↓GATC	MB24201	100 reactions	
	MB24202	500 reactions	
Speedy Mlui	MB24401	100 reactions	
A ↓CGCGT	MB24402	500 reactions	
Speedy Ncol	MB10001	50 reactions	
C↓CATGG	MB10002	250 reactions	
Speedy Pstl	MB10301	400 reactions	
CTGCA↓G	MB10302	2000 reactions	
Speedy Sall	MB10401	200 reactions	
G↓TCGAC	MB10402	1000 reactions	
Speedy Xhol	MB10701	200 reactions	
C↓TCGAG	MB10702	1000 reactions	

Buffers

10x NZYBuffe	rU
11001	

MB11001 500 μL MB11002 1000 μL



DNA & RNA modifying enzymes

LIGASES POLYMERASES NUCLEASES OTHER MODIFYING ENZYMES

DNA & RNA MODIFYING ENZYMES

NZYTech offers a variety of enzymes optimized for the development of different molecular biology protocols, including nucleases, polymerases, ligases, DNA binding enzymes and others.

LIGASES **T4 DNA Ligase** Speedy Ligase MB00703 500 U MB13001 50 ligations MB00704 2500 U ∕∿ DNA Ligase (E. coli) T7 DNA Ligase MB42401 200 U MB42501 100000 U Taq DNA Ligase **T4 ssRNA Ligase** MB42601 2000 U MB42701 1000 U T4 dsRNA Ligase 150 U MB42801

NUCLEASES

A Endonuclease V (<i>E. coli</i>)	T7 Endonuclease I
MB21301 250 U	MB21201 250 U
UltraPrecise T7 Endonuclease I	Nt.BbvCl, Nicking Endonuclease
MB34001 50 reactions	MB09401 1000 U
MB34002 200 reactions	MB09402 5000 U
	\wedge
NZY DNase I	Exonuclease I <i>(E. coli)</i>
MB19901 200 U/vial	MB42901 3000 U
Exonuclease III (E. coli)	Endonuclease IV (Tth)
MB43001 5000 U	MB43101 500 U
<u>A</u>	
Exonuclease VII (E. coli)	T5 Exonuclease
MB43201 200 U	MB43301 1000 U

OTHER MODIFYING ENZYMES

MutS (E	. coli)	Topoisomerase I (<i>E. coli</i>)
MB21101	50 µg	MB43501 100 U
MB21102	250 µg	
\wedge		
dam Me	thyltransferase	🛕 Alkaline Phosphatase (E. coli)
MB43401	500 U	MB01801 200 U
	nucleotide Kinase (T4 PNK)	
MB0080	500 U	
NEW NZY Ur	acil-DNA Glycosylase	NZY Thermolabile Uracil-DNA Glycosylase
MB4460	1 100 µL	MB44501 100 µL
MB4460	2 3 x 100 µL	MB44502 3 x 100 µL





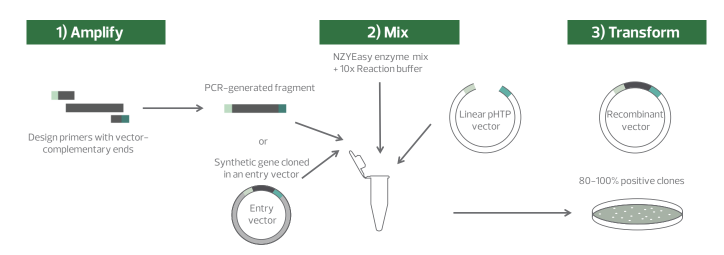
DNA cloning and mutagenesis kits

EASY CLONING & EXPRESSION STANDARD CLONING DNA MUTAGENESIS

DNA CLONING KITS

EASY CLONING & EXPRESSION

The NZYEasy Cloning & Expression System was designed to allow directional cloning of PCR-generated fragments or synthetic genes previously cloned in pUC-vectors into a linearized pHTP vector in a single reaction mediated by NZYEasy enzyme mix. The system allows achieving high cloning efficiencies and does not require the use of DNA ligases. In addition, no further treatment (e.g. restriction digestion, phosphorylation or blunt-end polishing) of the inserts is required.



NZYTech provides ready-to-use pHTP vectors in separate kits. The portfolio of pHTP prokaryotic expression vectors, which includes a different range of fusion tags, offers the possibility to quickly assay levels of expression and solubility of the desired protein in multiple expression vectors simultaneously. Choose the pHTP vectors that most suits your experiments on the table below:

Vector	Features	Comments	Kit Cat. No.
pHTP0	<i>lac</i> promoter and <i>lacZ_a</i> reporter	High-copy number cloning vector for the NZYEasy Cloning & Expres- sion System; allows blue/white screening during cloning (if adding IPTG and X-GAL)	MB281
pHTP1	N- and/or C-terminal 6xHis	Standard vector for high-level protein expression in E. coli	Kit I, MB282
pHTP2	N- and/or C-terminal 6xHis; N-terminal LLDsbC	Leader Less DsbC promotes cytoplasmic isomerization of disulfide bonds	Kit II, MB319
pHTP3	N- and/or C-terminal 6xHis; N-terminal mutDsbC	Inactive DsbC promotes cytoplasmic solubilization without isomerization of disulfide bonds	Kit III, MB320
pHTP4	N- and/or C-terminal 6xHis; N-terminal DsbC	DsbC promotes periplasmic isomerization of disulfide bonds	Kit IV, MB321
pHTP7	N- and/or C-terminal 6xHis; N-terminal DsbA	DsbA promotes periplasmic formation of disulfide bonds	Kit VII, MB322
pHTP8	N- and/or C-terminal 6xHis; N-terminal Trx	Trx enhances solubility of tagged proteins	Kit VIII, MB323
pHTP9	N- and/or C-terminal 6xHis; N-terminal GFP	GFP is a reporter molecule that allows monitoring protein localization	Kit IX, MB324
pHTP10	N- and/or C-terminal 6xHis; N-terminal NusA	NusA enhances solubility of tagged proteins	Kit X, MB325
pHTP11	N- and/or C-terminal 6xHis; N-terminal GST	GST enables glutathione-based affinity purification of tagged pro- teins while enhancing protein solubility	Kit XI, MB326
pHTP13	N- and/or C-terminal 6xHis; N-terminal GB1	GB1 enhances solubility of tagged proteins	Kit XIII, MB327
pHTP14	N- and/or C-terminal 6xHis; N-terminal KSI	KSI enhances solubility of tagged proteins	Kit XIV, MB328
pHTP16	N- and/or C-terminal 6xHis; N-terminal CpA	CpA enhances solubility of tagged proteins	Kit XVI, M3290
pHTP17	N- and/or C-terminal 6xHis; N-terminal CpB	CpB enhances solubility of tagged proteins	Kit XVII, MB330

PCR-generated fragments can be cloned into the pHTPO cloning vector (included in the NZYEasy Cloning kit) or, alternatively, into one of the various kanamycin-resistant pHTP expression vectors (included in the different NZYEasy Cloning & Expression kits) without the need to go through the tedious and laborious intermediate stages.

DNA Cloning

The NZYEasy Cloning kit was designed for time-saving and cost-effective DNA cloning. It includes the pHTPO vector (pUC-derivative) that allows blue/white screening for positive bacterial colonies (if adding IPTG and X-GAL).

\wedge	NZYEasy Cloning kit		
	MB28101	8 reactions	
	MB28103	96 reactions	

DNA Cloning & Expression

NZYEasy Cloning & Expression kits include different expression vectors that use the T7/*lac* promoter for regulated high-level protein expression in *E. coli* strains containing the λ DE3 lysogen, such as BL21(DE3).

ning & Expression kit I	$\mathbf{\Lambda}$	NZYEasy Clo	ning & Expression kit II
8 reactions		MB31901	8 reactions
96 reactions		MB31903	96 reactions
ning & Expression kit III		NZYEasy Clo	ning & Expression kit IV
8 reactions		MB32101	8 reactions
96 reactions		MB32103	96 reactions
ning & Expression kit VII		NZYEasy Clo	ning & Expression kit VIII
8 reactions		MB32301	8 reactions
96 reactions		MB32303	96 reactions
	96 reactions ning & Expression kit III 8 reactions 96 reactions ning & Expression kit VII 8 reactions	8 reactions 96 reactions ning & Expression kit III 8 reactions 96 reactions ning & Expression kit VII 8 reactions	8 reactions MB31901 96 reactions MB31903 ning & Expression kit III MB31903 8 reactions MB32101 96 reactions MB32103 ning & Expression kit VII MB32103 8 reactions MB32103

NZYEasy Cloning & Expression kit IX

MB324018 reactionsMB3240396 reactions

A NZYEasy Cloning & Expression kit X

MB32501	8 reactions
MB32503	96 reactions

NZYEasy Cloning & Expression kit XIII

8 reactions

96 reactions

MB32701

MB32703

NZYEasy Cloning & Expression kit XI

MB326018 reactionsMB3260396 reactions

NZYEasy Cloning & Expression kit XIV

MB32801	8 reactions
MB32803	96 reactions

NZYEasy Cloning & Expression kit XVI

MB32901 8 reactions MB32903 96 reactions

A NZYEasy Cloning & Expression kit XVII

MB33001	8 reactions
MB33003	96 reactions

STANDARD CLONING

NZYTech's DNA cloning kits are optimized to provide high-efficiency cloning based on easy protocols with no requirement for time-consuming restriction digests. To clone PCR-amplified fragments take into account the type of DNA polymerase that was used to generate the DNA fragment and choose the appropriate cloning kit: NZY-A PCR cloning kits are designed for cloning of DNA fragments amplified using non-proofreading polymerases, while NZY-blunt PCR cloning kit is designed to clone blunt-end PCR products amplified by proofreading enzymes. For a faster DNA cloning, the speedy version of NZY-A PCR cloning kit (NZY-A Speedy PCR cloning kit) should be chosen.

Blunt-end

NZY-blunt PCR cloning kit was designed to allow the direct cloning of PCR products with blunt-ends which result from amplifications using proofreading DNA polymerases such as NZYProof DNA polymerase (MB146).

NZY-blunt PCR cloning kit			
MB12101	24 ligations + competent cells		
MB12102	24 ligations		

A-overhang

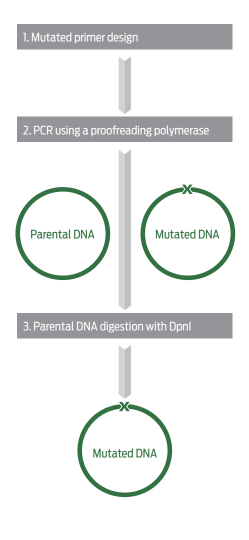
NZY-A PCR cloning kits were designed to clone PCR products produced by non-proofreading DNA polymerases such as NZYTaq II DNA polymerase (MB354). They take advantage of the terminal transferase activity of these polymerases which adds a single 3'-A overhang to each end of the PCR product. For a faster DNA cloning, NZYTech provides the NZY-A Speedy PCR cloning kit which allows direct cloning of PCR products with 3'-A overhangs in only 5 minutes at room temperature. Blunt-ended PCR fragments generated by amplification with proofreading polymerases can also be cloned using NZY-A PCR cloning kits after conducting an A-tailing procedure.

NZY-A PCR	cloning kit	NZ	Y-A Speed	y PCR cloning kit
MB05301	24 ligations + competent cells	MB	313701	24 ligations + competent cells
MB05302	24 ligations	MB	813702	24 ligations
	ency cloning: >95% positive clon e colony screening	es	U	
	cloning with NZY-A Speedy PCR	cloning kit		
L				1

DNA MUTAGENESIS

Site-Directed Mutagenesis can help you in a range of applications allowing to edit the desired DNA sequence by incorporation of single or multiple point mutations in any type of plasmid DNA. NZYTech's Mutagenesis kits provide simple and highly efficient methods to generate point mutations and delete or insert single (or multiple) nucleotides in plasmid DNA using PCR. NZYSupreme Mutagenesis kit was recently developed to reduce labor time and increase the efficiency of DNA editing.

Mutagenesis kits contain a proofreading DNA polymerase for PCR amplification of dsDNA plasmid to be mutated. NZYSupreme Mutagenesis kit includes Supreme NZYProof DNA polymerase, an engineered highly accurate, fast and sensitive variant of NZYProof DNA polymerase (the DNA polymerase present in NZYMutagenesis kit), formulated in a 2x concentrated master mix solution. Both Supreme NZYProof and NZYProof DNA polymerases ensure high fidelity for the exponential PCR amplification, thus reducing the unwanted secondary mutations and enabling amplification of large plasmids up to 15 kb. In addition, mutagenesis system requires the provision of two synthetic oligonucleotide primers containing the desired mutation. Mutated primers can be designed following standard guidelines (overlapping primers) or using an improved methodology (non-overlapping primers) recommended on NZYSupreme Mutagenesis kit. The mutagenesis protocol includes only three main steps:



1. Mutated primer design

Primers should have between 25 and 45 bases in length, with a melting temperature (T_m) of \ge 78 °C; a GC content of 40% and should terminate in one or more C or G bases;

2. PCR amplification

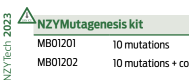
Extension of the oligonucleotide primers with a proofreading DNA polymerase generates a mutated plasmid containing staggered nicks;

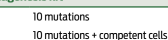
3. Digestion with Dpnl

Digestion of PCR product with DpnI endonuclease for elimination of the parental methylated and hemimethylated DNA template and selection of the mutation-containing synthetic DNA (not methylated).

NZYTech provides convenient versions of the kit, which include highly efficient competent cells for mutaded plasmid recovering.

	NZYMutagenesis kit	NZYSupreme Mutagenesis kit
Primer design:	Standard (overlapping primers)	Improved (non-overlapping primers)
DNA polymerase:	NZYProof DNA polymerase	Supreme NZYProof 2x Colourless Master Mix
PCR time:	2h 30	<1h 30 min
Digestion time:	1 hour (at 37 ºC)	5-15 min (at 37 ºC)





NZYSupreme Mutagenesis kit

MB44701 10 mutations MB44702 10 mutations + competent cells



MOLECULAR BIOLOGY

COMPETENT CELLS

CELL PREPARATION & GROWTH

COMPETENT CELLS & MEDIA

COMPETENT CELLS

Efficient DNA transformation of *Escherichia coli* competent cells is essential for successful cloning and protein expression applications. NZYTech offers competent *E. coli* host strains for high-efficiency transformation.

MB00401	20 transformations	MB00501	20 transformations
MB00402	40 transformations	MB00502	40 transformations
Routine cl	te colony screening oning : fhuA2fj(argF-lacZ)U169 phoA g	glnV44Ψ80 fj(lacZ)M15 gyrA9	GorecA1 relA1
	of pUC19 te colony screening ty plasmid preparation ion of cDNA libraries/gene banks		l
Constructi	-	hi -1 recA1 gyrA96 relA1 lac[F´p	proA+B+ lac-

NZYTech 2023

CELL PREPARATION & GROWTH

Competent cells preparation

NZYCompetent Cells Preparation Buffer is designed for the preparation of super competent *Escherichia coli* cells. The method is compatible with the classical heat shock transformation procedure and the transformation efficiencies are typically on the order of 10^8 - 10^9 transformants/µg plasmid DNA with the most common *E. coli* strains.

NZYCompetent Cells Preparation Buffer MB12001 100 mL

Culture media

NZYTech offers a selection of high quality culture media for a wide variety of applications. Our specific formulations were extensively tested for use in cloning, plasmid DNA preparation and protein expression.

902 903	500 g 1000 g	MB38901	500 g
B02903	1000 g		
	1000 8		
LB Broth (gra	nulated)	SOC Broth	
MB02802	500 g	MB28001	500 g
MB02803	1000 g		
Tryptone		Sodium Chlo	oride (NaCl)
IB16701	1000 g	MB15901	1000 g
LB Agar		Yeast Extra	ct, Micro Granulated
MB11802	1000 g	MB16401	1000 g

LB Broth (powder)			
MB14501	500 g		
MB14502	1000 g		

Auto-Induction media

NZY Auto-Induction LB medium (powder) is an innovative culture medium developed for growing *Escherichia coli* to high cell densities, while obtaining high-levels of recombinant protein expression when using IPTG-inducible bacterial expression systems.

NZY Auto-Induction LB medium (powder)*		
MB17901	100 g	
MB17903	1000 g	

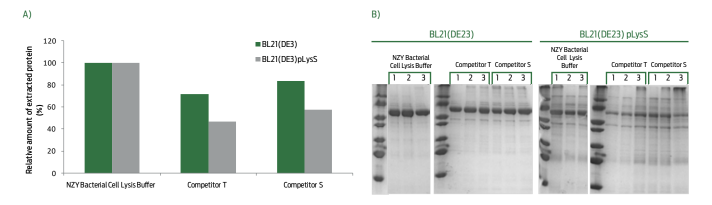
* Product not available for use or sale in the United States

No need for IPTG induction No need to monitor cell growth Ideal for High-Throughput (HTP) methods High protein yields

Cell disruption

NZY Bacterial Cell Lysis Buffer is an innovative product for the gentle disruption of *Escherichia coli* cell wall that generates a homogeneous cell-free extract. It provides a rapid and cost effective alternative to mechanical methods such as French Press or sonication for releasing recombinant and native proteins. This extraction reagent is a Tris buffered formulation (pH 7.5) with Lysozyme and DNase I provided separately.





Comparing the efficiency of *Escherichia coli* cell lysis for protein extraction using NZY Bacterial Cell Lysis Buffer and two similar competitor products. **Cells from two strains of** *E. coli* BL21(DE3) and BL21(DE3 pLysS) were harvested from 5 mL of cultured media and lysed (in triplicates) using three different protein extraction chemicals: NZY Bacterial Cell Lysis Buffer, Competitor T and Competitor S.

A) Levels of extracted protein obtained were evaluated. B) The recombinant proteins (in triplicates) were purified through IMAC and separated through SDS-PAGE.

MOLECULAR BIOLOGY

5

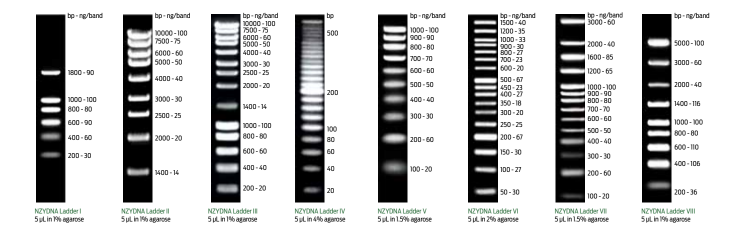
ladders & markers

DNA LADDERS PROTEIN MARKERS Life Science Research 2023

LADDERS & MARKERS

DNA LADDERS

NZYTech offers a ready-to-use set of DNA ladders suitable for standard gel electrophoresis applications. Ladders can be used to quantify the acid nucleic concentration based on band intensity and/or to accurately determine the molecular weight of the nucleic acids molecules.



NZYDNA Ladder I		
MB04101	90 µg (200 lanes)	
MB04102	225 µg (500 lanes)	

NZYDNA La	dder II
MB04301	83 µg (200 lanes)
MB04302	207 µg (500 lanes)

NZYDNA Lao	dder III	NZYDNA Lac	lder IV
MB04401	143 µg (200 lanes)	MB05801	10 µg (50 lanes)
MB04402	357 µg (500 lanes)	MB05802	30 µg (150 lanes)

NZYDNA Ladder V

MB06101 120 µg (200 lanes) MB06102 300 µg (500 lanes)

NZYDNA Lado	ler VI	

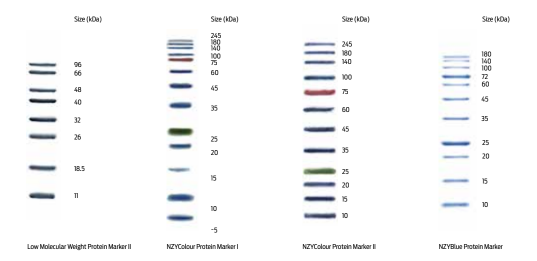
100 µg (200 lanes) MB08901 MB08902 250 µg (500 lanes)

NZYDNA Ladder VII		NZYDNA Ladder VIII	
MB14701	170 µg (200 lanes)	MB17501	150 µg (200 lanes)
MB14702	425 µg (500 lanes)	MB17502	375 µg (500 lanes)

PROTEIN MARKERS

NZYTech provides different ready to-use protein markers containing unstained or stained proteins. All protein markers are mixtures of highly purified proteins of known molecular weight developed for assessing proteins relative molecular weights.

nL (125 lanes)
nL (500 lanes)
ker
ACI
nL (125 lanes)
nL (500 lanes)
ine (500 tarie





loading & staining

LOADING

STAINING

LOADING & STAINING

LOADING

For DNA

 6x NZYDNA loading dye

 MB13101
 5 x 1 mL

For Proteins

 5x SDS-PAGE sample loading buffer

 MBI1701
 5 x 1 mL

STAINING

For DNA

NZYTech offers a DNA stain that can be used as a safer alternative to the traditional ethidium bromide (EtBr) and presenting equal sensitivity. GreenSafe Premium can be either incorporated in the agarose gels or used post-running.

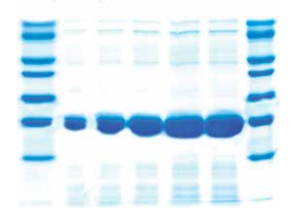
GreenSafe Premium		
MB13201	1 mL	

For Proteins

As an alternative to the traditional Coomassie Blue staining for detecting proteins in SDS-PAGE, NZYTech offers BlueSafe, a very sensitive and safe single-step protein stain.

BlueSafe	
MB15201	1000 mL

BlueSafe: High contrast and sensitivity





agaroses & buffers

AGAROSES

READY-TO-USE BUFFERS

AGAROSES & BUFFERS

AGAROSES

NZYTech offers two different agaroses that cover the most used applications in the laboratory.

	Analytical separation ≥ 0.5 kb	Preparative electrophoresis	DNA typing	Blotting
Electrophoresis grade	8		Ø	8
Ultrapure grade	8	8	Ø	8

Agarose (Agarose (electrophoresis grade)	
MB02702	100 g	
MB02703	500 g	
MB02704	1000 g	

Agarose (ul	trapure grade)	
MB05201	100 g	
MB05202	500 g	

READY-TO-USE BUFFERS

PBS (10x) pH 7.4 (liqui	d)
-------------------------	----

1000 mL

MB25201

PBS	(10x)	pH 7.4	(powder)

MB18201 1 pouch (1 x 10 L)

TAE Buffer (5	iOx) pH 8.3 (liquid)
MB20001	1000 ml

MB20901 1000 mL

TBE Buffer (10x) (liquid)

MB27701 1000 mL



antibiotics & biochemicals

ANTIBIOTICS BIOCHEMICALS

ANTIBIOTICS & BIOCHEMICALS

ANTIBIOTICS

Ampicillin (sodium salt)			
MB02101	5 g		
MB02102	25 g		
MB02103	50 g		

Chloramphenicol		
MB02402	25 g	

5g

5 g

MB16501

Gentamicin (sulphate)

MB16601

Kanamycin Monosulphate

5 g

MB02001

BIOCHEMICALS

D(+)-Gluce	ose Anhydrous	NaCl	
MB16801	1000 g	MB15901	1000 g

Glycerol	
MB16101	1000 mL

IPTG

MB02602 5 g MB02603 25 g

DEPC-treated Water

MB43701 5 x 1 mL

X-Gal		
MB02501	lg	
MB02502	5 g	

Water for Molecular Biology

MB11101 1000 mL

MOLECULAR BIOLOGY

protein electrophoresis & blotting

PROTEIN QUANTIFICATION PROTEIN ELECTROPHORESIS & TRANSFER WESTERN BLOTTING

PROTEIN ELECTROPHORESIS & **BLOTTING**

PROTEIN QUANTIFICATION

NZY Bradfo	ord reagent
MB19801	500 ml

PROTEIN ELECTROPHORESIS & TRANSFER

Acrylamide/b	is-Acrylamide (29:1 Solution)	
MB04501	500 mL	
Acrylamide/b	is-Acrylamide (37.5:1 Solution)	
MB15601	500 mL	
DTT		
MB03101	5g	
SDS Solution	20%	
MB11601	500 mL	
Tris base		
MB01601	1000 g	
TEMED		

MB03501 25 mL

Tris-Glycine-SDS Buffer (liquid)

MB19501 1000 mL

Tris-Glycine Buffer (liquid)MB194011000 mL

Ammonium Persulphate (APS)MB03403100 g

Glycine MB01401 1000 g

 SDS Powder

 MB18101
 500 g

WESTERN BLOTTING

BSA (Bovine Serum Albumin, Fraction V)					
MB04601	10 g				
MB04602	100 g				
MB04603	1000 g				

NZY Standard ECL

MB40101 250 mL

Milk, Dry Powder

MB26001 100 g

NZY Advanced ECL

MB40201 250 mL





PDZ DOMAINS

PDZ domains are structural modules of 80 to 90 amino acids found in signaling proteins of bacteria, yeast, plants, viruses and animals (Vicentelli et al. 2015, Nature Methods 12, 787-93). PDZ is an acronym combining the first letters of three proteins, which were first discovered to contain these domains: Post synaptic density protein (PSD95), Drosophila disclarge tumor suppressor (Dlg1), and Zonula occludens-1 protein (zo-1). PDZ domains play an essential role in a number of cellular processes by facilitating protein scaffolding and assembly of protein complexes. Protein complex formation between PDZ target molecules can lead to a number of signaling and regulatory cascades that may either promote or inhibit the activation of certain proteins. NZYTech offers a library of representative recombinant PDZ domains of the human proteome which includes 120 different constructs. For more information, please visit NZYTech website at www.nzytech.com.

VENOM PEPTIDES

Animal venoms are highly complex molecular cocktails containing a wide range of biologically active peptides that target, with high selectivity and efficacy, a variety of membrane receptors. These peptides and their biological activities possess important pharmacological, therapeutic and biotechnological values. Venom peptides are highly stable molecules displaying formidable affinity and selectivity while presenting low immunogenicity, making them attractive candidates for the development of novel therapeutics.

Venom peptides generally contain between 20 to 120 residues and include up to eight disulfide bonds that are critical for both biological activity and stability.

In recent years, considerable emphasis is being put on the discovery of novel molecules with therapeutic interest and venom peptides appear as a source of potential drugs. Unfortunately, the use of venom peptides as therapeutic or biotechnological molecules is still hampered by the difficulty to produce native and active proteins in sufficient amounts. In order to increase the availability of these molecules for academic or pharmaceutical partners, NZYTech is proud to offer a library of representative animal recombinant Venom Peptides. Our recombinant Venom Peptides are subjected to a variety of highly stringent quality control protocols to reach high purity levels. For more information, please visit NZYTech website at www.nzytech.com.



labware

PIPETTES & TIPS PCR TUBES, PLATES & STRIPS ADHESIVE SEALS



At NZYTech we believe your PCR samples are too precious for low quality consumables. Please test NZYTech throughly optimized and proved labware and take your PCR amplifications to the next level. Regarding PCR Plates and Strips compatibility please refer to www.nzytech.com

PIPETTES & TIPS

Tips 0.5-10 µ	JL (bulk)		Tips 0.5-10 µ	JL (rack sterile)
LW00101	1000 units	-	LW00102	96x10
Tips 20-200	μL (bulk)		Tips 20-200	µL (rack sterile)
LW00203	1000 units	-	LW00204	96x10
Tips 100-100)0 µL (bulk)		Tips 100-100	00 µL (rack sterile)
LW00301	1000 units	-	LW00302	100x10
Micropipette	25			
(single and mu	ltichannel; individual and sets)	-		
	BES, PLATES & STI	RIPS		
	idual PCR tubes with flat caps		-	mL PCR tubes w/ optically caps
LW01301	1000 tubes		LW03801	120 strips
Strips of 8 fl	at optical caps		Strip of 8 x 0).2 mL tubes flat strip caps
LW03201	300 strips	-	LW01201	125 strips + caps
Low P. 96 we	ll semi sk. PCR plate, Roche, white		96 Well non-	-skirted PCR plate, white
LW04101	50 plates		LW02601	50 plates
96 Well non	-skirted PCR plate		96 Well sem	i skirted PCR plate, white
LW00901	50 plates	-	LW03601	50 plates
96 Well sem	i skirted PCR plate		96 Well skirt	ted PCR plate, white
LW00801	50 plates		LW02501	50 plates

96 Well skirted PCR plate

LW00701 50 plates

ADHESIVE SEALS

General adhesive plate seals

LW01701 100 sheets

qPCR adhesive clear plate seals

LW02101 100 sheets

Adhesive PCR plate seals

LW01801 100 sheets

Life Science Research 2023

LABWARE COMPATIBILITY

	Skirt type	Skirted 96		Half-Skirted 96			Non-Skirted		Strips	
	Number of wells								8	
	Profile	L	ow	Sta	Standard Lo		Sta	ndard	200µL	100µL
	Colour type	Clear	Clear White		Clear White		Clear	White	Clear	
	Catalogue Number	LW00701	LW02501	LW00801	LW03601	LW04101	LW00901	LW02601	LW03801	LW03901
ABI® / Life	Technologies/ Thermo Fisher Scientif	ic	•		÷	•			•	•
	GeneAmp® 2400								8	
	GeneAmp® 2700, 2720, 9600			8	8		8	8		
	GeneAmp® 9700			8	8		8	8	8	
	GeneAmp® 9800 FAST Block									8
hermal	MultiBlock System, MBS	8	8							8
yclers	PCR Sprint						8	8	8	8
	ProFlex 96 well			8	8		8	8		
	Simpliamp						8	8	8	
	Veriti 0.1 ml (96 well block) FAST									8
	Veriti 0.2 ml (96 well block)			8	8		8	8	8	
	7000, 7300, 7500, 7700, 7900			8	8				8	
PCR	QuantStudio™ 3, 5, 6, 7, 12K, Dx			8	8					
yclers	StepOne, StepOne Plus™								<u>.</u>	
	ViiA7™			8	8		8	8	8	
	310, 3100, 3130, 3130 XL Genetic Analyser			8	8		8	8		
equencers	3500, 3500 XL Genetic Analyser			8	8		8	8	8	
	3700, 3730, 3730XL DNA Analyser			8	8		8	8		
.gilent (St	ratagene)	·	•	•	÷	·	•	·	:	:
hermal	Surecycler 8800 96 well						8	8		
yclers	Gradient Cycler	8	8		•		8	8	•	
	AriaMx	8	8				8	8		
PCR	Mx3000P TM						8	8		
yclers	Mx3005P™	8	8				8	8		
	Mx4000™			8	8		8	8	<u>.</u>	
Biometra	: 								•	
	Flexcycler296			8	8		8	8	8	
	T1 Thermocycler, Tgradient			8	8		8	8	8	8
	T3 Thermocycler				-				8	
Thermal Cyclers	Tone, Tadvanced (96), TProfessional Gradient/XL			8	8		8	8	8	
	TRIO, Tpersonal				-	-	8	8	8	<u>.</u>
	Trobot 96			8	8					<u>.</u>
	SpeedCycler2			8	8		8	8	8	•••••
PCR	qTOWER3/G, touch	8	8	8	8		8	8	8	
qPCR Cyclers	Toptical Thermalcycler			8	8		8	8	8	

	Skirt type	Skirted 96		Half-Skirted 96			Non-Skirted		Strips 8	
	Number of wells									
	Profile	Low		Star	Standard Low		Standard		200µL 100µL	
	Colour type	Clear	White	Clear	W	nite	Clear	White	CI	ear
	Catalogue Number	LW00701	LW02501	LWOOBOI	LW03601	LW04101	LOGOONJ	LW02601	LW03801	LW03901
Bioer Tech	inologies		÷		÷	÷	÷	•	;	į.
hermal	GeneTouch			8	8		8	8	8	
lyclers	GeneQ			<u>.</u>			8	8	8	
BIO-RAD	•		·	:	:	÷	·	1	:	
	C1000 Touch	8	8							8
	Genecycler						8	8	8	
	iCycler™, MyCycler™			8	8		8	8	8	
hermal	\$1000	8	8							8
yclers	T100			8	8		8	8	8	8
	Mini Gradient						8	8	8	8
	PTC100™ (96 well block only)	8	8	8	8		8	8	8	8
	DNA Engine™, DNA Dyad™	8	8	8	8		8	8	8	8
	CFX Opus, CFX96 Touch/Touch Deep Well, CFX connect	8	8							
	iCycler™ IQ, IQ™ 4, IQ™ 5, MyiQ™			8	8		8	8	8	
IPCR Syclers	Chromo4™	8	8				8	8		
	MiniOpticon									
	Opticon™, Opticon2™, DNA Engine	8	8							
orbett Res	earch					. <u> </u>				
hermal Syclers	(Qiagen) Palm Cycler	8	8							8
IPCR Syclers	Rotor-Gene series									
PPENDOR	F		·	·	·	÷	·	·	·	ĺ
	MasterCycler® Pro, Pro S	8	8	8	8		8	8	8	
hermal	MasterCycler® nexus, gradient, eco, gradient eco	8	8	8	8		8	8	8	
yclers	MasterCycler® nexus X2, GX2, GX2e, X2e						8	8	8	
	MasterCycler® nexus SX1, GSX1	8	8	8	8		8	8	8	
IPCR Cyclers	Mastercycler™ ep realplex	8	8	8	8		8	8	8	
EQLAB (V	: WR)		:	:	:	; 	:	:	:	
hermal	peqSTAR 96	8	8				8	8	8	8
yclers	peqSTAR 2X		••••••				8	8	8	
OCHE	·		·	·	·	·		·	·	
	LC96/LC480					8				
	Nano									8





MOLECULAR SERVICES

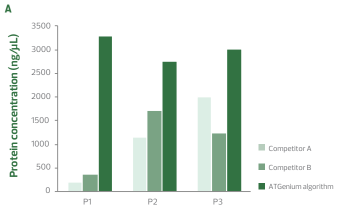
Life Science

GENE SYNTHESIS

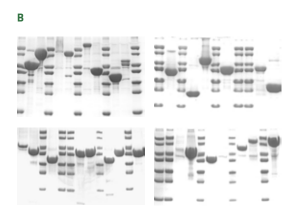
Gene synthesis has become an important tool in many fields of recombinant DNA technology. Whether you have limited cloning experience or simply want to save time, NZYTech's Gene Synthesis service helps you to quickly move your ideas from the planning stage to the laboratory. A highly motivated team is eager to provide you with a very efficient service affording considerable savings of time and money. Either if you select the custom or High-Throughput (HTP) Gene Synthesis service you will be offered a premium service that will deliver high quality and downstream efficacy.



NZYTech's scientists have also developed a highly robust gene design optimization software - **ATGenium** codon optimization algorithm - to maximize protein expression. Our proprietary algorithm comprehensively optimizes critical factors in transcription, translation and co-translational protein folding to deliver highest levels of expression in any host system. This service is offered at no extra cost for all gene synthesis services if required.



A. Levels of protein expression of three different eukaryotic proteins in *Escherichia coli* BL21 (DE3). The genes encoding these proteins were optimized by three algorithms for codons optimization, by two different competitors (light green) and using the ATGenium codon optimization algorithm developed by NZYTech (dark green).



B. High-throughput recombinant protein expression of CAZymes (Carbohydrate Active en-Zymes). Genes encoding these proteins were optimized using ATGenium, cloned into pHTP1 expression vector and overexpressed in *E. coli* BL21 (DE3). High protein expression yields were obtained.

CUSTOM GENE SYNTHESIS

NZYTech's Custom Gene Synthesis service provides in vitro chemical synthesis, cloning and 100% sequence verification of any desired DNA sequence.

Sub-cloning into NZYTech's pTHP vectors is available upon request.

Gene Synthesis	
Gene Synthesis (DNA fragments <500 bp)	GS00307
Gene Synthesis (DNA fragments 501-1000 bp)	GS00308
Gene Synthesis (DNA fragments 1001-2000 bp)	GS00309
Gene Synthesis (DNA fragments 2001-5000 bp)	GS00310
Gene Synthesis (DNA fragments > than 5001 bp)	GS00311
Gene sub-cloning	

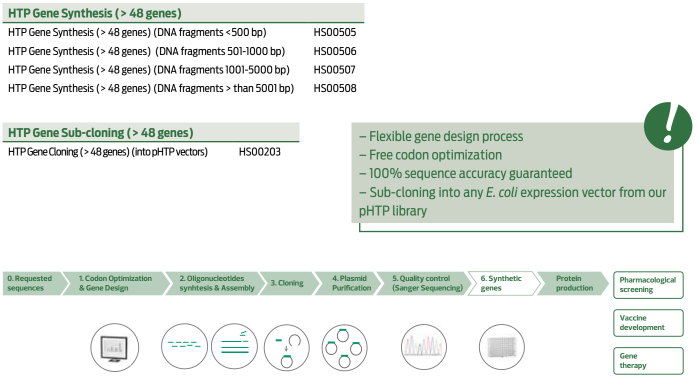
Gene sub-cloning into pHTP vectors

Flexible, custom gene design Free codon optimization - ATGenium Sequences 100% verified

HIGH-THROUGHPUT GENE SYNTHESIS

MS00209

NZYTech's R&D team has developed a High-Throughput (HTP) Gene Synthesis platform to efficiently generate hundreds to thousands of synthetic genes. We offer a high quality HTP Gene Synthesis service for DNA fragments below 4 kb. HTP Gene Synthesis will allow to tackle difficult research projects such as: determination of structure-function relationships in different proteins, creation of novel antibody libraries, screening of gene variant libraries or optimization of protein function and expression. Our codon optimization technology - ATGenium algorithm - allows obtaining the highest possible levels of gene expression in different expressions hosts while retaining high levels of protein solubility.



HTP Primer designer tool optimization algorithm

ATGenium codon

Multi-sequencing analysis tool

73

Molecular

engineering

GENE FRAGMENTS

Gene Fragments	
300-400 bp (price per fragment)	GF00108
401-500 bp (price per fragment)	GF00109
501-750 bp (price per fragment)	GF00110
751-1000 bp (price per fragment)	GF00111
1001-1250 bp (price per fragment)	GF00112
1251-1500 bp (price per fragment)	GF00113
1501-1800 bp (price per fragment)	GF00114

PEPTIDES

PEPTIDE SYNTHESIS

NZYTech offers a high-quality peptide synthesis service designed to ensure our customer's complete satisfaction. Every peptide is subjected to a variety of performance control steps to guarantee the highest quality. Peptides may be provided crude, desalted, or with 75% to 98% purity. In addition, NZYTech peptides can be subjected to a variety of chemical modifications upon request.

Lyophilized peptide delivered with the required sequence, purity and quantity Several modifications available: phosphorylation, methylation, acetylation, amidation, fluorescence/dye labeling Additional discounts for large-scale peptide synthesis orders



GLYCOBIOLOGY

Life Science

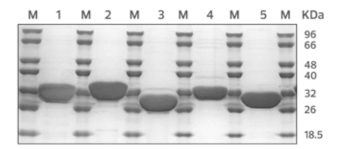
NZYTECH CAZYMES: HIGHLY PURE, STABLE AND OPTIMIZED RESEARCH TOOLS FOR SCIENCE AND INDUSTRY

1. A LARGE AND DIVERSE PORTFOLIO

NZYTech R&D department, individually or in collaboration with the CAZy community, is building a large and diverse portfolio to foster science and industrial progress. All proteins have a recombinant origin and are produced and purified from *Pichia pastoris* or *Escherichia coli*. In this catalogue you will find more than 1000 CAZymes, covering >200 CAZy families (www.cazy.com) and >150 EC number activities (www.enzyme.expasy.org).

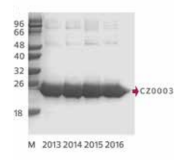
2. ULTRA-PURE FOR A VARIETY OF APPLICATIONS

NZYTech Cazymes are purified using a comprehensive list of chromatography protocols to achieve the highest level of protein purity. As an example, the figure below provides an SDS-PAGE analysis of Glycobiology CZ0604, CZ0580, CZ0621, CZ0633 and CZ0641. Purification level of all Glycobiology listed in this catalogue may be found at NZYTech website (www.nzytech.com).



3. REDUCED BATCH-TO-BATCH VARIATION AND LONG STORAGE STABILITY

NZYTech devotes significant efforts to minimize batch to batch variation and improve protein stability. In the Figure below 4 batches of Xylanase 11A from *Clostridium thermocellum* (CZ0003), produced in 2013, 2014, 2015 and 2016, were tested in March 2018 for molecular integrity and protein purity. The data reveals that there is minimal variation in molecular integrity up to 5 years after production (Panel B). Thus, it is important to follow the storage guidelines provided.



4. NZYTECH CAZYMES ARE THROUGHLY CHARACTERIZED AND THEIR BIOCHEMICAL PROPERTIES DESCRIBED IN LITERATURE

Although NZYTech provides CAZymes as highly pure recombinant proteins and not functionally assessed, the majority of NZYTech CAZymes have been thoroughly characterized and their enzymatic properties is described in the literature. The reference of the paper describing such properties is provided in the product manual which allows you to select the one most suited for your experiments.

Glycoside Hydrolases

ACETYLGALACTOSAMINIDASES	(5 entries)	PHOSPHO-β-GLUCOSIDASES	(16 entries)
ACETYLGLUCOSAMINIDASES	(19 entries)	GLUCURONIDASES	(14 entries)
AGARASES	(17 entries)	GLUCURONOXYLANASES	(4 entries)
AMYLASES	(15 entries)	HEXOSAMINIDASES	(13 entries)
AMYLOMALTASES	(7 entries)	HYALURONIDASES	(2 entries)
ARABINANASES	(8 entries)	INULASES	(2 entries)
ARABINOFURANOSIDASES	(35 entries)	LAMINARINASES	(16 entries)
ARABINOPYRANOSIDASES	(1 entries)	LEVANASES	(4 entries)
ARABINOXYLANASES	(2 entries)	LEVANSUCRASES	(3 entries)
CARRAGEENASES	(2 entries)	LICHENASES	(11 entries)
CELLOBIOHYDROLASES	(10 entries)	LYSOZYMES	(4 entries)
CELLODEXTRINASES	(2 entries)	α-MANNANASES	(3 entries)
CELLULASES	(79 entries)	β-MANNANASES	(25 entries)
CHITINASES	(7 entries)	α-MANNOSIDASES	(22 entries)
CHITOSANASES	(3 entries)	β-MANNOSIDASES	(6 entries)
DEXTRANASES	(4 entries)	MANNOSYLGLUCOSE	(2 entries)
FRUCTANASES	(3 entries)	PHOSPHORYLASES	
FRUCTOFURANOSIDASES	(4 entries)	OLIGOSACCHARIDE REDUCING	(2 entries)
FRUCTANASES	(3 entries)	-END XYLANASES	
FRUTCTOFURANOSIDADES	(4 entries)	PEPTIDOGLYCAN LYTIC EXOTRANSGLYCOSYLASES	(4 entries)
FUCOSIDASES	(16 entries)	POLYGALACTURONASES	(7 entries)
GALACTANASES	(6 entries)	PORPHYRANASES	(4 entries)
α -GALACTOSIDASES	(15 entries)	PULLULANASES	(3 entries)
β-GALACTOSIDASES	(21 entries)	RHAMNOGALACTURONASES	(16 entries)
$\textbf{PHOSPHO-}\beta\textbf{-}\textbf{GALACTOSIDASES}$	(4 entries)	SIALIDASES	(6 entries)
β-D-GALACTOSYL-1,4-L-RHAM-	(2 entries)	TREHALASES	(13 entries)
NOSE PHOSPHORYLASES		∆-4,5-UNSATURATED	(7 entries)
GALACTOSYL-N-ACETYLHEXOSA- MINE PHOSPHORYLASES	(2 entries)	β-GLUCURONYL HYDROLASES	(7 entities)
GALACTURONIDASES	(6 entries)	XYLANASES	(43 entries)
GLUCANSUCRASES	(5 entries)	XYLOGLUCANASES	(7 entries)
GLUCOSAMINIDASES	(3 entries)	XYLOSIDASES	(17 entries)
α-GLUCOSIDASES	(13 entries)	OTHER ACTIVITIES	(26 entries)
β-GLUCOSIDASES	(25 entries)		

Carbohydrate esterases

ACETYL XYLAN ESTERASES	(18 entries)
ACETYLGLUCOSAMINE DEACETYLASES	(8 entries)
DIACETYLCHITOBIOSE DEACETYLASES	(1 entries)
FERULOYL ESTERASES	(96 entries)
GLUCURONYL ESTERASES	(7 entries)
PECTIN ACETYL ESTERASES	(4 entries)
PECTIN METHYLESTERASES	(4 entries)

Polysaccharide lyases

ALGINATE LYASES	(14 entries)
CHONDROITIN LYASES	(4 entries)
HEPARIN LYASES	(6 entries)
HYALURONATE LYASES	(1 entries)
OLIGOGALACTURONATE LYASES	(2 entries)
PECTATE LYASES	(28 entries)
PECTIN LYASES	(2 entries)
POLY-a-GULURONATE LYASES	(2 entries)
RHAMNOGALACTURONAN LYASES	(8 entries)
ULVAN LYASES	(8 entries)
XANTHAN LYASES	(1 entries)
RHAMNOGLUCURONATE LYASES	(1 entries)

Auxiliar activities

LACCASES	(3 entries)
LYTIC POLYSACCHARIDE	(5 entries)
MONOOXYGENASES	

Carbohydrate-binding modules

CBMs	(103 entries)
GFP-CBM	(57 entries)
ZZ-CBM	(2 entries)

Mini-cellulosomes & other enzymes

MINI-CELLULOSOMES	(3 entries)
CELLOBIOSE DEHYDROGENASE	(1 entries)
XYLOSE ISOMERASE	(1 entries)



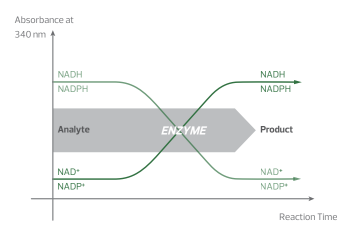
ANALYTICAL PRODUCTS

PRINCIPLES & FEATURES

UV TESTS

NZYTech test kits are based on enzymatic reactions and performed using spectrophotometric methods.

The principle of the enzymatic tests is based on the NAD(P)⁺/NAD(P)H system and uses highly pure enzymes engineered to display a premium performance. The enzymes used in these analytical kits produce or consume NAD(P)H, which strongly absorbs the UV radiation at 340 nm (extinction coefficient of 6300 M⁻¹cm⁻¹).



COLORIMETRIC TESTS

The principle of the enzymatic tests based on a chromogenic reaction is the formation of a coloured compound, which absorbs at the visible region of the spectrum. The coloured compound results from the interaction between the product of a first enzymatic reaction and a chromogenic compound. In this case, the concentration of the analyte must be determined by using a standard curve.

- Endpoint analysis
- Easy to use, simple protocols
- Rapid analysis
- Accuracy and precision
- Safe to the operator
- Standards included

REQUIRED MATERIAL

- Spectrophotometer
- \cdot Micropipettes set with disposable plastic tips to accurately dispense volumes from 20 µL to 1000 µL
- Cuvettes
- Basic filtering or other simple sample treatment device

APPLICATIONS

Nome	Kitsize	Range	Detection Limit	Food Industry	Feed Industry	Wine Industry	Fermen- tation Industry	Dairy Industry	Biofuel	Catalogue No
Acetaldehyde, UV Method	50 tests	0.25-200 mg/L	0.176 mg/L			3	⊗	8		AK00051
Acetic Acid, UV Method	53 tests	0.15-200 mg/L	0.14 mg/L	8	8	∞	⊗	8	8	AK00081
Ammonia, UV Method	96 tests	10-70 mg/L	0.07 mg/L	⊗	8	⊗	⊗			AK00091
L-Arginine/Urea/Ammonia, UV Method	50 tests each	50-400 mg/L L-arginine 20-140 mg/L urea 10-70 mg/L ammonia	0.37 mg/L L-arginine 0.13 mg/L urea 0.07 mg/L ammonia			8	8			АКООІЛ
Ethanol, UV Method	60 tests	0.12-120 mg/L	0.093 mg/L	8		⊗	⊗	8	⊗	AK00061
D-Fructose/D-Glucose, UV Method	IIO tests of each*	2-800 mg/L	0.66 mg/L	3	8	⊗		8	8	AK00041
D-Glucose HK, UV Method	110 tests	2-800 mg/L	0.66 mg/L	3			ᢒ	3		AKOOO31
D-Glucose (God-POD), Colorimetric Method	660 tests	100-1000 mg/L	100 mg/L	⊗			⊗	8		AK00161
L-Gutamine/Ammonia, UV Method	50 tests of each	10-400 mg/L L-glutamine 10-70 mg/L ammonia	0.54 mg/L L-glutamine 0.07 mg/L ammonia	⊗			⊗			AKOOTIT
L-Lactic Acid, UV Method	50 tests of each	0.30-300 mg/L	0.30 mg/L	8	8	8	⊗	8	8	AKOOI31
D-/L-Lactic Acid, UV Method	50 tests of each	0.30-300 mg/L	0.30 mg/L	3	3	ᢒ	3	8	3	AK00141
D-Malic Acid, UV Method	100 tests	0.25-400 mg/L	0.26 mg/L	8		⊗				AK00021
L-Malic Acid, UV Method	58 tests	0.25-300 mg/L	0.25 mg/L	3		3	⊗			AKOOOII
L-Malic Acid, Colorimetric Method	5x10 tests	8-800 mg/L	8 mg/L	8		8	⊗			AK00191
Sucrose/D-Fructose/D-Glucose, UV Method 100 tests of each	100 tests of each	20-800 mg/L	1.40 mg/L	3	3	3	3	8	⊗	AK00201
Sulfite, UV Method	30 tests	0.25-300 mg/L	0.25 mg/L	8		⊗	8			АКОООЛ
Urea/Ammonia, UV Method	50 tests each	1.5-140 mg/L urea 10-70 mg/L ammonia	0.13 mg/L urea 0.07 mg/L ammonia	⊗	⊗	3		⊗		AKOOIOI

analytical enzymes



L-ARGINI



ANALYTICAL PRODUCTS

ANALYTICAL ENZYMES

Enzyme	EC	Catalogue No	Pack size
Acetyl-CoA synthetase	6.2.1.1	AE00081	250 U
D-Alanine aminotransferase	2.6.1.21	AE00141	2500 U
Alcohol dehydrogenase	1.1.1.1	AE00131	1000 U
Aldehyde reductase YqhD	1.1.1.21	AE00021	2.0 mg
Arginase	3.5.3.1	AE00211	1950 U
Aspartate aminotransferase	2.6.1.1	AE00061	5000 U
Citrate synthase	2.3.3.1	AE00041	2500 U
Diaphorase	1.8.1.4	AE00231	1000 U
Glucokinase	2.7.1.2	AE00171	1400 U
Glucose-6-phosphate dehydrogenase , NADP+ dependent	1.1.1.49	AE00111	5000 U
Glucose-6-phosphate dehydrogenase , NAD⁺ dependent	1.1.1.49	AE00161	5000 U
Glucose-6-phosphate isomerase	5.3.1.9	AE00101	5000 U
Glutamate dehydrogenase	1.4.1.4	AE00051	3300 U
Glutaminase	3.5.1.2	AE00071	2500 U
Glutathione reductase	1.8.1.7	AE00221	500 U
nvertase	3.2.1.26	AE00241	100 kU
.actaldehyde dehydrogenase	1.2.1.22	AE00031	4.0 mg
D-Lactate dehydrogenase	1.1.1.28	AE00121	22 kU
D-Malate dehydrogenase	1.1.1.83	AE00151	200 U
Malate dehydrogenase	1.1.1.37	AE00091	50 kU
NADH peroxidase	1.11.1	AE00201	500 U
Sulfite oxidase molybdenum centre domain	1.8.3.1	AE00011	2.25 U
Jronate dehydrogenase	1.1.1.203	AE00191	24 kU

TERMS & CONDITIONS



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