

# Monarch<sup>®</sup> Nucleic Acid Purification

**TECHNICAL GUIDE** 



delivered by VWr



•

*be* INSPIRED *drive* DISCOVERY *stay* GENUINE

# Make the right choice and migrate to Monarch

Monarch<sup>®</sup> Nucleic Acid Purification Kits are the perfect complement to many molecular biology workflows. Recover pure, intact DNA and RNA in minutes with our fast, user-friendly protocols and optimized buffer systems, and focus your time on the experiments that will drive your research forward. The Monarch nucleic acid purification portfolio can serve your needs, whether you are isolating nucleic acids from biological samples, cleaning up DNA and RNA from enzymatic reactions, extracting DNA fragments from gels, or purifying plasmids.

Monarch kits are all designed with sustainability in mind; whenever possible, kits and components are made with significantly less plastic and are packaged with responsiblysourced, recyclable packaging. Furthermore, plastic recovered during the manufacture of Monarch columns is used to manufacture other plastic-based NEB products.

#### **AVAILABLE KITS:**

#### Monarch Genomic DNA Purification Kit

- · Purify high-quality, genomic DNA from several sample types
- Achieve excellent DNA yields with fast, user friendly protocols

#### Monarch HMW DNA Extraction Kits

- Quickly and easily extract ultra-high molecular weight DNA
- Available for cells & blood as well as tissues, bacteria and other samples

#### Monarch Total RNA Miniprep Kit

- Extract total RNA from a variety of sample types with this all-in-one kit
- Validated with cells, blood, saliva, swabs, tissues, bacterial, plant and more

#### Monarch RNA Cleanup Kits

- Purify and concentrate RNA after enzymatic reactions and RNA synthesis
- Choose between 3 convenient binding capacities (10 μg, 50 μg, and 500 μg)

#### Monarch Plasmid Miniprep Kit

- · Easily purify plasmids from bacterial cultures
- · Monitor your progress with our convenient colored-buffer system

#### Monarch PCR & DNA Cleanup Kit

- Purify DNA in 5 minutes and elute in as little as 6 μl
- Unique column design eliminates buffer carryover, resulting in highly-pure DNA
- A modified protocol enables the purification of small DNA fragments and oligos

#### Monarch DNA Gel Extraction Kit

- Quickly extract highly-pure DNA from agarose gels with excellent yields
- Column design enables elution in as little as  $6 \ \mu$ l and prevents buffer carryover

#### TABLE OF CONTENTS

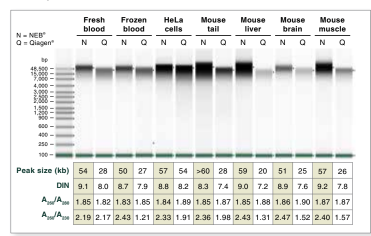
- 3 Monarch Genomic DNA Purification Kit
  - 4 Performance Data
  - 5 Compatibility with Next Generation Sequencing
  - 6 Sample Inputs and Expected Recovery
- 7 Monarch HMW DNA Extraction Kits
  - 8 Performance Data: Cells & Blood
  - 9 Performance Data: Tissue & Other Samples
  - 10 Suitability for Long Read Sequencing
- 11 Monarch Total RNA Miniprep Kit
  - 12 Performance Data
  - 13 Sample Inputs & Expected Recovery
- 14 Monarch Kits for RNA Cleanup
- 16 Monarch DNA Gel Extraction Kit
- 16 Monarch PCR & DNA Cleanup Kit (5 μg)
- 18 Monarch Plasmid Miniprep Kit
- 19 Troubleshooting Guides
  - 19 Genomic DNA Purification
  - 20 Total RNA Extraction & Purification
  - 21 RNA Cleanup
  - 22 DNA Cleanup & Plasmid Purification
- 23 Ordering Information



## Monarch Genomic DNA Purification Kit

The Monarch Genomic DNA Purification Kit is a comprehensive solution for cell lysis, RNA removal, and purification of intact genomic DNA (gDNA) from a wide variety of biological samples, including cultured cells, blood and mammalian tissues. Excellent results are achieved even with challenging samples like fatty (e.g., brain) and fibrous (e.g., muscle, mouse tail) tissues. Additionally, bacteria, yeast and insects can be processed with minor protocol modifications to enhance lysis in these tough-to-lyse samples. Protocols are also included to enable purification from clinically-relevant samples, such as saliva and cheek swabs, as well as rapid cleanup of previously extracted gDNA. Purified gDNA has high quality metrics and minimal residual RNA. The purified gDNA is suitable for downstream applications, such as endpoint PCR, qPCR and library prep for next generation sequencing (NGS).

The Monarch Genomic DNA Purification Kit provides excellent yields of higher quality, higher molecular weight DNA than the Qiagen<sup>®</sup> DNeasy<sup>®</sup> Blood & Tissue Kit



Agilent Technologies<sup>®</sup> 4200 TapeStation<sup>®</sup> Genomic DNA ScreenTape was used for analysis of gDNA purified from blood, cultured cells and tissue samples using the relevant protocols of the Monarch Genomic DNA Purification Kit and the Qiagen DNeasy Blood & Tissue Kit. gDNA was eluted in 100 µl and 1/100 of the eluates (~1 µl) was loaded on a Genomic DNA ScreenTape. Starting materials used: 100 µl fresh human whole blood, 100 µl frozen pig blood, 1 × 10<sup>6</sup> HeLa cells and 10 mg frozen tissue powder. Monarch-purified gDNA samples typically show peak sizes 50–70 kb and DINs of ~9. DNeasy-purified gDNA peak sizes are typically < 30 kb with DINs ~7–8. DNeasy kits produce lower yields and low  $A_{280/230}$  ratios for liver, brain, muscle and frozen blood.

#### ADVANTAGES

- · Use with a wide variety of sample types
- Achieve higher yields, especially with difficult tissue samples (e.g., brain and muscle)
- Effectively remove RNA (< 1% residual RNA) with optimized buffer chemistry and included RNase A
- Isolate longer DNA (peak size > 50 kb), which is suitable for long read sequencing platforms
- Save time with fast protocols, efficient lysis steps and minimal hands on time
- · Can also be used to clean up genomic DNA

#### SPECIFICATIONS

- · Recommended Input Amount: Varies by sample type. See page 6.
- Binding Capacity: 30 µg genomic DNA
- Genomic DNA Size: Peak size > 50 kb for most sample types; may be lower for saliva and buccal swabs
- Elution Volume: ≥ 35 µl (100 µl is recommended)
- Purity:  $A_{260/280} \ge 1.8$ ,  $A_{260/280} \ge 2.0$
- RNA Content: < 1% (with included RNase A treatment)
- Compatible Downstream Applications: endpoint PCR, qPCR, library preparation for NGS (including Oxford Nanopore Technologies<sup>®</sup> and PacBio<sup>®</sup>)



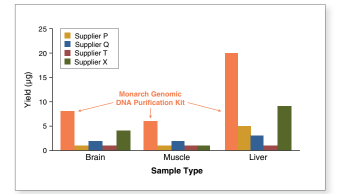
This kit yielded the highest purity DNA I have ever seen from a commercial spin column kit.



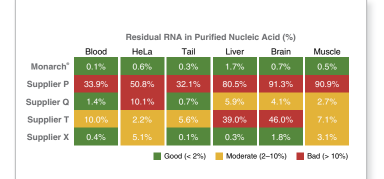
<sup>-</sup> Stephen, Lake Superior State University

### Purified DNA is High Yield, Highly Pure, Free from RNA and Ready for use in Downstream Applications

The Monarch Genomic DNA Purification Kit provides excellent yields for difficult tissue types



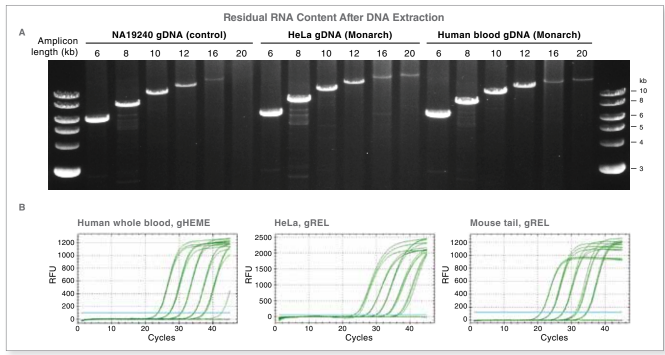
Duplicate 10 mg samples of RNAIater®-stabilized rat tissue were cut to small pieces and subsequently lysed and purified according to the protocols provided for each kit. Optional RNase A steps were included. Elution was carried out with 100 µl elution buffer provided in the respective kits. Yields displayed are averages of the duplicate samples, and represent the genomic DNA yield after correcting for the RNA content as determined by LC-MS. Results indicate that the Monarch Genomic DNA Purification Kit provides excellent yields for a wide range of tissues, which can be problematic for other commercial kits. DNA purified with the Monarch Genomic DNA Purification Kit has significantly lower residual RNA across all sample types



RNA content present in genomic DNA eluates from various kits was evaluated by LC-MS. All samples were processed in duplicate according to manufacturers' recommendations and were eluted in 100 µl. Starting materials used: 100 µl human blood, 1 x 10<sup>6</sup> HeLa cells, 10 mg of RNAlater-stabilized mouse (tail) and rat tissue samples (others). 1 µg of each sample was treated with the Nucleoside Digestion Mix (NEB #M0649) and subjected to LC-MS.

Values displayed are averages of duplicate measures and indicate the percentage of riboguanoside (rG) versus the total amount of ribo- and deoxyriboguanoside in the samples. Actual RNA content may be lower for all samples, since rG is more abundantly co-purified in silica preps than other RNA bases. The Monarch Genomic DNA Purification Kit consistently delivers residual RNA below 1%–2% levels, which is usually undetectable with most analysis methods and lower than what is seen for other commercial kits.

The Monarch Genomic DNA Purification Kit generates high quality genomic DNA suitable for sensitive applications like long range PCR and qPCR



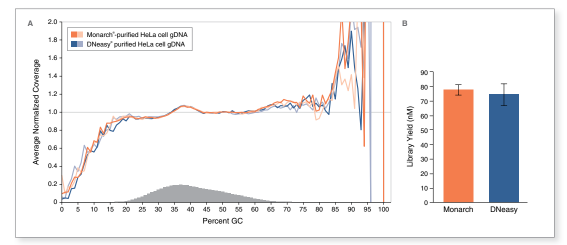
A. Amplification reactions were set up with primer pairs specific for 6, 8, 10, 12, 16, 20 kb amplicons from human DNA. LongAmp® Hot Start Taq 2X Master Mix was used and 25 ng template DNA was added to each sample. PCR reactions were carried out on an Applied Biosystems 2720 Thermal Cycler. Monarch-purified genomic DNA isolated from HeLa cells and human blood were compared to commercially available reference DNA from the human cell line NA19240 F11. 10 µl was loaded on a 1.5% agarose gel, using the 1 kb DNA Ladder as a marker. Results indicated DNA was of high-integrity and suitable for long range PCR.

B. Monarch-purified genomic DNA from human whole blood, HeLa cells and mouse tail was diluted to produce a five log range of input template concentrations. The results were generated using primers targeting gHEME (human whole blood) and gREL (HeLa, mouse tail) for qPCR assays with the Luna<sup>®</sup> Universal qPCR Master Mix and cycled on a BioRad<sup>®</sup> CFX Touch qPCR thermal cycler. Results indicated that DNA is highly pure and free from inhibitors, optimal for qPCR.

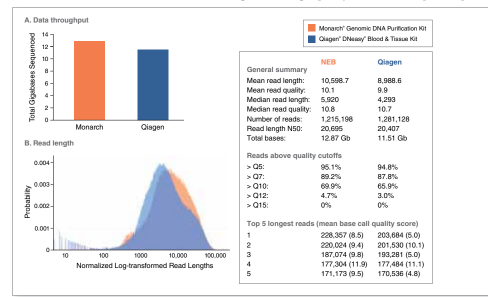
### An Outstanding Choice for Illumina® and Long Read Sequencing

The Monarch Genomic DNA Purification Kit is an excellent choice for DNA extraction upstream of library preparation for next generation sequencing. DNA purified with this kit is high quality and of high molecular weight (peak size > 50 kb), making it an outstanding choice for preparation of libraries for nanopore sequencing and other long read platforms. The kit is also optimized to selectively bind DNA, not RNA, and is also supplied with RNase A for (optional) removal of any residual RNA, allowing for purification of DNA with extremely low RNA contamination.

The Monarch Genomic DNA Purification Kit generates excellent input material for NGS library preparation with NEBNext<sup>®</sup> kits for Illumina



- A. Duplicate libraries were made from 100 ng HeLa cell gDNA purified with Monarch (orange) or Qiagen DNeasy Blood & Tissue Kit (blue) using the NEBNext Ultra<sup>™</sup> II FS DNA Library Prep Kit for Illumina. Libraries were sequenced on an Illumina MiSeq<sup>®</sup>. Reads were mapped using Bowtie 2.2.4 and GC coverage was calculated using Picard's CollectGCBiasMetrics (v1.117). Expected normalized coverage of 1.0 is indicated by the horizontal grey line, the number of 100 bp regions at each %GC is indicated by the vertical grey bars, and the colored lines represent the normalized coverage for each library. Monarch GC coverage matched Qiagen DNeasy results.
- B. High yield libraries are achieved from Monarch-purified gDNA. Library yields of the samples described above were assessed on an Agilent Technologies 2100 BioAnalyzer® using a High Sensitivity DNA Kit.



The Monarch Genomic DNA Purification Kit generates high quality DNA for nanopore sequencing

HeLa cell genomic DNA was extracted using either the Monarch Genomic DNA Purification Kit or the Qiagen DNeasy Blood & Tissue Kit. One microgram of purified DNA was used to prepare Oxford Nanopore Technologies sequencing libraries following the 1D Ligation Sequencing Kit (SQK-LSK109) protocol without DNA fragmentation. Libraries were loaded on a GridION® (Flow cell R9.4.1) and the data was collected for 48 hrs. Libraries produced using the Monarch gDNA exceeded the Qiagen libraries on common sequencing metrics including: A. total sequencing data collected, B. read length. Data was generated using NanoComp (Bioinformatics, Volume 34, Issue 15, 1 August 2018, Pages 2666–2669).

### Guidelines for Choosing Sample Input Amounts When Using the Monarch Genomic DNA Purification Kit

Genomic DNA yield, purity and integrity vary immensely based on sample type, input amount and sample condition. Below, we have provided some empirical yield, purity, and DIN data from a wide variety of sample types as well as guidance on the maximal input amounts for each of those samples when using the Monarch Genomic DNA Purification Kit. It is very important not to overload the column and the buffer system when extracting and purifying gDNA, as DNA yields, purity, integrity, and length may suffer. Inputs that will result in ~100 ng of gDNA should be considered the minimum input amount for this kit (5  $\mu$ l whole blood, 1 x 10<sup>4</sup> cultured cells or 0.2 mg tissue). If using smaller amounts, the use of carrier RNA is recommended (see product manual for more details).

SAMPLE TYPE	RECOMMENDED	TYPICAL YIELD (µg)	DIN	MAXIMUM INPUT AMOUNT
TISSUE*				
Tail (mouse)	10 mg	12–20	8.5-9.5	25 mg
Ear (mouse)	10 mg	18–21	8.5-9.5	10 mg
Liver (mouse and rat)	10 mg	15–30	8.5–9.5	15 mg
Kidney (mouse)	10 mg	10–25	8.5–9.5	10 mg
Spleen (mouse)	10 mg	30–70	8.5–9.5	10 mg
Heart (mouse)	10 mg	9–10	8.5–9.5	25 mg
Lung (mouse)	10 mg	14–20	8.5–9.5	15 mg
Brain (mouse and rat)	10 mg	4–10	8.5–9.5	12 mg
Muscle (mouse and rat)	10 mg	4–7	8.5–9.5	25 mg
Muscle (deer)	10 mg	5	8.5–9.5	25 mg
BLOOD**				
Human (whole)	100 µl	2.5–4	8.5–9.5	100 µl
Mouse	100 µl	1–3	8.5–9.5	100 µl
Rabbit	100 µl	3–4	8.5–9.5	100 µl
Pig	100 µl	3.5–5	8.5–9.5	100 µl
Guinea pig	100 µl	3–8	8.5–9.5	100 µl
Cow	100 µl	2–3	8.5–9.5	100 µl
Horse	100 µl	4–7	8.5–9.5	100 µl
Dog	100 µl	2-4	8.5–9.5	100 µl
Chicken (nucleated)	10 µl	30–45	8.5–9.5	10 µl
CELLS				
HeLa	1 x 10 <sup>6</sup> cells	7–9	9.0–9.5	5 x 10 <sup>6</sup> cells
HEK293	1 x 10 <sup>6</sup> cells	7–9	9.0–9.5	5 x 10 <sup>6</sup> cells
NIH3T3	1 x 10 <sup>6</sup> cells	6–7.5	9.0–9.5	5 x 10 <sup>6</sup> cells
BACTERIA				
E. coli (Gram-negative)	2 x 10 <sup>9</sup> cells	6–10	8.5–9.0	2 x 10 <sup>9</sup> cells
Rhodobacter sp. (Gram-negative)	2 x 10 <sup>9</sup> cells	6–10	8.5–9.0	2 x 10 <sup>9</sup> cells
B. cereus (Gram-positive)	2 x 10 <sup>9</sup> cells	6–9	8.5–9.0	2 x 10 <sup>9</sup> cells
ARCHAEA				
T. kodakarensis	2 x 10 <sup>9</sup> cells	3–5	8.5–9.0	2 x 10 <sup>9</sup> cells
YEAST				
S. cerevisiae	5 x 10 <sup>7</sup> cells	0.5–0.6	8.5–9.0	5 x 10 <sup>7</sup> cells
SALIVA/BUCCAL CELLS***				
Saliva (human)	200 µl	2–3	7.0–8.0	500 µl
Buccal swab (human)	1 swab	5–7	6.0-7.0	1 swab

\* Tissue gDNA yields are shown for frozen tissue powder, frozen tissue pieces and RNAlater-stabilized tissue pieces. Though frozen tissue powder results in highly-intact gDNA, lower yields can be expected than when using frozen or RNAlater-stabilized tissue pieces. Residual nuclease activity in tissue pieces will cut the gDNA, resulting in a slightly smaller overall size; however, this gDNA is optimal for silica-based purification.

\*\* Human whole blood samples stabilized with various anticoagulants (e.g., EDTA, citrate and heparin) and various counter-ions were evaluated and results were comparable in all cases. Additionally, all indicated blood samples were tested both as fresh and frozen samples, yielding comparable results. Human samples were donated by healthy individuals; yields from unhealthy donors may differ.

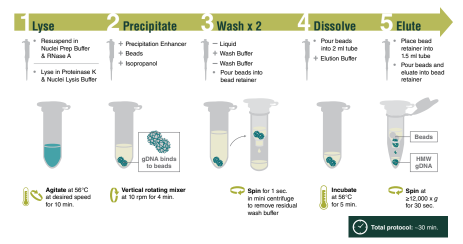
### Monarch HMW DNA Extraction Kits

The Monarch HMW DNA Extraction Kits provide a rapid and reliable process for extracting high molecular weight, intact genomic DNA from cultured cells, whole blood, various tissues, bacteria and other sample types (e.g., amphibian, insect). Utilizing a novel and optimized process that combines lysis with tunable fragment length generation, followed by precipitation of the extracted DNA onto the surface of large glass beads, the prep proceeds rapidly and efficiently. DNA size ranges from 50 kb into the Mb range, depending on the agitation speed used during lysis. Purified DNA is recovered in high yield with excellent purity, including nearly complete removal of RNA. For cells, the process time is only 30 minutes, while blood samples require erythrocyte lysis and are processed in approximately 60 minutes. Tissues and bacteria are processed in about 90 minutes, leading the market in speed. Purified HMW DNA is suitable for a variety of downstream applications including long read sequencing (Oxford Nanopore Technologies and Pacific Biosciences<sup>®</sup>), optical mapping, and linked-read genome assembly.

Glass beads used for HWM DNA Extraction



#### Workflow for cell samples





We've had great success with obtaining HMW DNA for long read sequencing from a variety of cell types, using less input and obtaining a comparable yield...It is straightforward and easy to use."

 Inswasti Cahyani & Matt Loose, DeepSeq, University of Nottingham

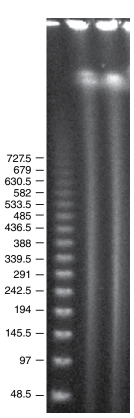
#### ADVANTAGES

- Fast workflow (cells: 30 min, blood: 60 min, tissue/bacteria: 90 min)
- Extract DNA into the megabase (Mb) range with cells, blood, soft organ tissues, and bacteria
- Tune DNA size based on agitation speed during lysis
- · Achieve best-in-class yields and purity
- · Consistently achieve reproducible results
- · Effectively remove RNA
- · Elute DNA easily and completely

#### SPECIFICATIONS

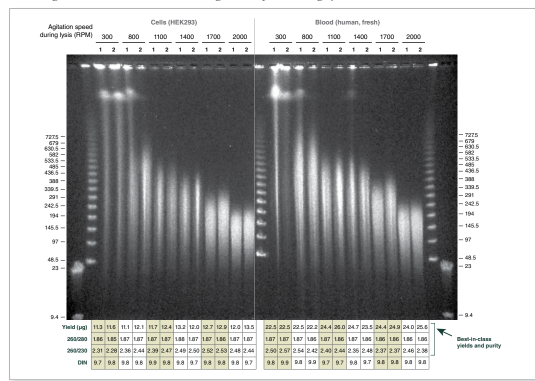
- Binding Mechanism: precipitation on glass beads
- Input Amount: Cells: 1 x  $10^5 1$  x  $10^7$  cells Blood: 100 µl - 2 ml Tissue: 2 - 25 mg Bacteria: 5 x  $10^8 - 5$  x  $10^9$  cells
- Genomic DNA Size: 50 kb up to several MB; dependant on agitation speed and sample quality/type
- Purity: OD  $_{\rm 260/280}$  typically 1.8 1.9 OD  $_{\rm 260/280}$  typically 2.1 2.5
- RNA Content: < 2%

#### HMW DNA from HEK293 cells



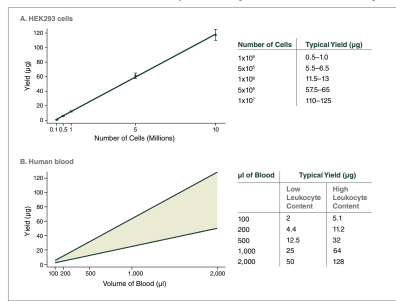
# Easily Extract Megabase-sized DNA from Cells and Blood with High Yields and Purity

DNA fragment size is tunable based on agitation speed during lysis



Preps were performed on duplicate aliquots of 1 x 10<sup>6</sup> HEK 293 cells and 500 µl fresh human blood. Samples were agitated at the indicated speed during the lysis step to control the fragmentation of the DNA. Equal amounts of DNA from the replicates (cells: 500 ng; blood: 650 ng) were resolved by PFGE (1% agarose gel, 6 V/cm, 13°C for 20 hours, switch times ramped from 0.5–94 seconds on a BioRad CHEF-DR III System). Yield anpurity ratios of the individual preps are shown in the accompanying tables. Lambda PFG Ladder and Lambda DNA-Hind III Digest were used as molecular weight standards. Yield, purity ratios and DINs of the individual preps are shown in the accompanying tables.

Linear correlation between DNA yield and input for cell and blood samples



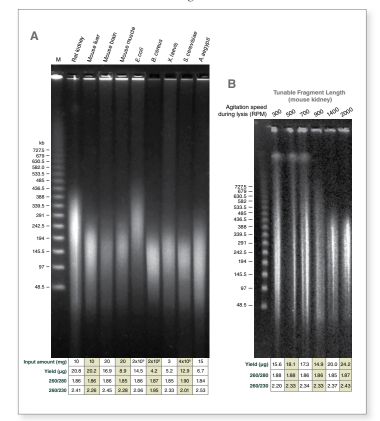
Summarized yield data for HMW DNA preps are shown carried out at 2,000 rpm during lysis, using HEK293 cultured cells and fresh human blood samples from different donors as input material in the corresponding protocols. The starting materials were diluted to 5 different concentrations to cover the entire recommended input range. Cell samples  $\leq 5 \times 10^5$  cells and blood samples < 500 µl were purified using the recommended volumes for low input samples. Obtained yields show a high degree of linearity over the displayed input range.

#### Validated Sample Types\* Cells & Blood Kit

Cells	Mammalian Blood**			
HEK293	Human			
HeLa	Mouse			
NIH3T3	Rat (fresh only)			
Jurkat	Rabbit			
K562 (suspension cells)	Pig			
HCT116	Horse			
A549	Cow			
U50s	Rhesus monkey			
HepG2	Goat			
NCI-460	Nucleated Blood**			
SK-N-SH	Chicken			
Aa23				
	Turkey			
*Fresh and frozen samples have been validated				
**Compatible with all common a	nticoagulants			

### Simply and Effectively Isolate HMW DNA from a Variety of Tissue Types, with Tunable Fragment Length

Successful extraction of HMW DNA from various tissue samples with tunable DNA size for soft organ tissues



**A.** HMW genomic DNA extracted from various samples using the Monarch HMW DNA Extraction Kit for Tissue (species and input amounts indicated in the figure). Preps were performed according to the kit instructions, with sample agitation at 2000 rpm during lysis. A modified workflow was used to process S. cerevisiae samples. **B.** HMW genomic DNA from mouse kidney (10 mg) was purified using the Monarch HMW DNA Extraction Kit for Tissue. Samples were agitated at the indicated speed during the lysis step to control the fragmentation of the DNA. 500 ng (**A**) or 300 ng (**B**) of purified DNA was resolved by PFGE. Yield and purity ratios of the individual preps are shown in the accompanying tables. Lambda PFG Ladder was used as a molecular weight standard.

#### TIPS FOR SUCCESSFUL HMW DNA EXTRACTION

- Pay close attention to input amounts and follow the protocol guidance for your recommended inputs. Lysis volumes may need to be reduced for optimal binding, especially when working with low input tissue samples.
- 2. For optimal results in the blood protocol, resuspend leukocyte pellets carefully and completely at each resuspension step.
- When working with low-input tissue samples, stop agitation in the thermal mixer after 15 minutes, and continue the incubation without shaking.
- Carry out the inversions exactly as directed to ensure the DNA binds completely to the beads. If inversions are done manually, do them slowly and gently, each inversion taking about 5-6 seconds.
- If using low agitation speeds for the highest molecular weight DNA, additional inversions during DNA binding will maximize yields.

#### Tissue workflow highlights

Following lysis and phase separation, use the same simple worflow for binding and elution as in the cell workflow





Phase separation step optimizes protein removal.



Microtube pestle and grinding tube included for optimal sample homogenization.

Tissue input recommendations

Use of beads simplifies washing and elution.

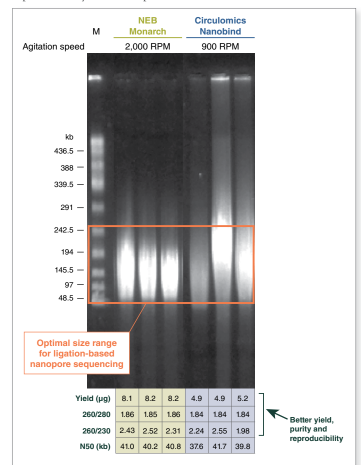
		RECOMMENDED INPUT (INPUT RANGE), mg	YIELD (µg) FOR RECOMMENDED INPUT (YIELD PER mg)
Mammalian Tissue			
Mouse brain	Fresh	15 (2**-20)	12–21
	Frozen	15 (2**–20)	15–21 (1–1.5)
Mouse liver	Fresh (w/NaCl)	10 (2–15)	7
	Frozen (w/NaCI)	10 (2–15)	17–19 (1.7–1.9)
	Fresh*	10 (2–15)	20
	Frozen*	10 (2–15)	27-31 (2.7-3.1)
Mouse muscle	Fresh	20 (2-25)	8–9
	Frozen	20 (2-25)	12-16 (0.6-0.8)
Mouse kidney	Fresh	10 (2-15)	23–34
	Frozen	10 (2–15)	32-41 (3.2-4.1)
Mouse tail	Frozen	20 (2-25)	20 (1.8–2.1)
Mouse ear punch	Fresh	10 (2-15)	15-16 (1.5-1.6)
Rat kidney	Frozen	10 (2–15)	20–25
Bacteria			
<i>E. coli</i> (Gram-negative)	Frozen	1 x 10 <sup>9</sup> cells (5 x 10 <sup>8</sup> - 5 x 10 <sup>9</sup> cells)	8–9
<i>B. cereus</i> (Gram-positive)	Frozen	2 x 10 <sup>8</sup> cells (2 - 4 x 10 <sup>8</sup> cells)	4–5
<i>M. luteus</i> (gram-positive)	Frozen	1 x 10 <sup>8</sup> cells	2.0
Amphibian			
X. laevis	Fresh	3–4	5
Yeast			·
S. cerevisiae	Fresh	3.8 x 108 cells	15
Insect			
A. aegypti	Frozen	15	6
*Standard protocol withou	t recommended NaCL	treatment	

\*Standard protocol without recommended NaCl treatment.

\*\* If working with input amounts < 5 mg, refer to the product manual for guidance on reducing buffer volumes.

### Reproducibly High Yields and Purity - Great for Long Read Sequencing

Monarch Kits extract HMW DNA with superior yields, purity and reproducibility when compared to Circulomics<sup>®</sup> Nanobind<sup>®</sup>



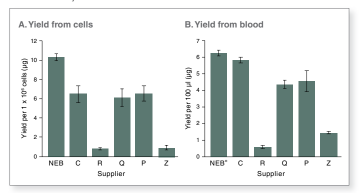
Genomic DNA was purified from 2 x 10<sup>6</sup> HEK293 cells and 0.7 x 10<sup>6</sup> K562 cells with the Monarch Kit and the Circulomics Nanobind CBB Big DNA Kit, according to manufacturers' recommendations. 400 ng of HMW genomic DNA was separated on a 0.75% gel using a Pippin pulse gel system (Sage Science) at the 5–430 kb program. M = Lambda PFG Ladder. NEB Monarch and Circulomics Nanobind samples were barcoded and analyzed on the same Oxford Nanopore Technologies flow cell.

Excellent performance in Oxford Nanopore Technologies Sequencing

	HEK293	HUMAN BLOOD	MOUSE KIDNEY
Mean read length	21338.9	21522.6	27120.7
Mean read quality	12.8	13.4	13
Median read length	10388	10130	23150
Median read quality	13.2	13.9	13.5
Number of reads	377687	538090	164000
Read length N50	45432	46542	44631
Total bases	8059414490 (8.1 Gb)	11581090785 (11.6 Gb)	4447789727 (4.4 Gb)

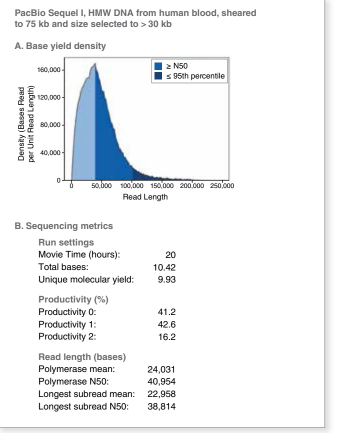
DNA used for the sequencing libraries was extracted from HEK293 cells (1x10<sup>6</sup> cells), human blood (500 µl) and mouse kidney (10 mg fresh, homogenized with rotor stator) using the Monarch kit, without further size selection. Libraries were prepared using the NEBNext Companion Module for Oxford Nanopore Technologies Ligation Sequencing and sequencing was performed on a GridION Mk1 (LSK109 kit, FLO-MIN106D flow cell) for up to 48 hours, or shorter if no more data was generated by the flow cell. No additional treatment of the flow cell (e.g., flushing) was employed. Read lengths are indicated in bases.

DNA yields from cell and blood preps using various commercially available kits



HMW DNA was isolated from 1 x 10° HEK293 cells (A) and fresh human blood (B) with kits from New England Biolabs (N), Circulomics [Nanobind CBB Big DNA Kit (C)], RevoluGer® [Fire Monkey (R)], QIAGEN [MagAttract HMW DNA Kit (Q)], Promega® [Wizard HMW DNA Extraction Kit (P)] and Zymd® Research [Quick-DNA HMW MagBead Kit (Z)]. Blood input volumes were used as specified in manufacturers' protocols (N: 500 µl, C: 200 µl, R: 500 µl, Q: 200 µl, P: 300 µl, Z: 200 µl). Yields for the blood samples were normalized for 100 µl blood.

#### Excellent performance in PacBio sequencing

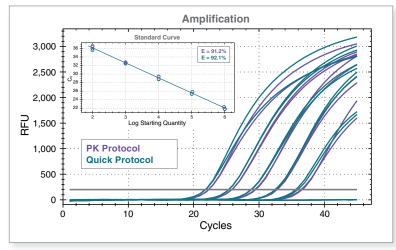


HMW DNA was extracted from human blood with the Monarch HMW DNA Extraction Kit for Cells & Blood with agitation speed of 2000 rpm during lysis. DNA was sheared to 75 kb with a Megaruptor, and SMRTbells were constructed with the SMRTbell Express Template Prep Kit 2.0 and size-selected to a minimum of 30 kb with a Blue Pippin. CLR reads were obtained on a PacBio Sequel I using 12 pM on-plate loading concentration and a 20 hour movie with no pre-extension.

### Monarch Total RNA Miniprep Kit

The Monarch Total RNA Miniprep Kit is a comprehensive solution for sample preservation, cell lysis, gDNA removal, and purification of total RNA from a wide variety of biological samples, including cultured cells, blood, and mammalian tissues. Additionally, tough-to-lyse samples, such as bacteria, yeast, and plant, can be processed with additional steps that enhance lysis. Total RNA, including viral RNA, can also be extracted from clinically-relevant samples like saliva, buccal swabs and nasopharyngeal swabs. Cleanup of enzymatic reactions or purification of RNA from TRIzol® -extracted samples is also possible using this kit. Purified RNA has high quality metrics, including  $A_{260/280}$  and  $A_{260/230}$  ratios  $\geq$  1.8, high RIN scores, and minimal residual gDNA. Captured RNA ranges in size from full-length rRNAs down to intact miRNAs. Additionally, differential binding conditions allow selective capture or exclusion of the sub-200 nucleotide RNA pool that includes miRNA, 5S rRNA, and tRNA. Purified RNA is suitable for downstream applications, such as RT-qPCR, cDNA synthesis, RNA-seq, Northern blot analysis, etc.

The Monarch Total RNA Miniprep Kit successfully purifies synthetic SARS-CoV-2 viral RNA from saliva samples



The Monarch Total RNA Miniprep Kit Proteinase K and Quick Protocols were used to isolate total RNA from saliva samples containing 10-fold serial dilutions of synthetic SARS-CoV-2 N-gene RNA. Purified RNA was eluted in 100 µl nuclease-free water to yield 50 to 500,000 copies of viral RNA/µl. Using the Luna Universal Probe One-Step RT-qPCR Kit, titers as low as 50 copies (the lowest input tested) were detected and linear, quantitative recovery of the SARS-CoV-2 N-gene was observed over a 5-Log range.

#### ADVANTAGES

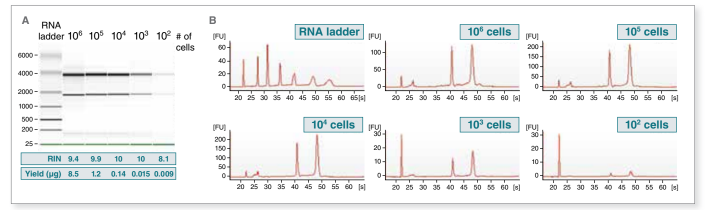
- Use with a wide variety of sample types, including clinically-relevant ones like swabs (buccal/NP) and saliva
- Purify RNA of all sizes, including miRNA & small RNAs  $\geq 20$  nucleotides
- Includes DNase I, gDNA removal columns, Proteinase K, and a stabilization reagent
- RNA Extraction from some samples is automatable on the QIAcube<sup>®</sup> and KingFisher Flex platforms
- · Save money with value pricing for an all-in-one kit

#### SPECIFICATIONS

- Binding Capacity: 100 µg RNA
- RNA Size: ≥ 20 nt
- Purity:  $A_{260/280}$  and  $A_{260/230}$  usually  $\geq 1.8$
- Input Amount: up to 107 cells or 50 mg tissue\*
- Elution Volume: 30–100 µl
- Yield: varies depending on sample type
- Compatible downsteam applications: RNA Library prep for NGS, RT-PCR, RT-qPCR, Northern blots
- \*See page 13 for more details and other sample types

The Monarch Total RNA Miniprep offers better yield and quality; easy protocol and good and consistent results among samples.

- Gisele, Mississippi State University

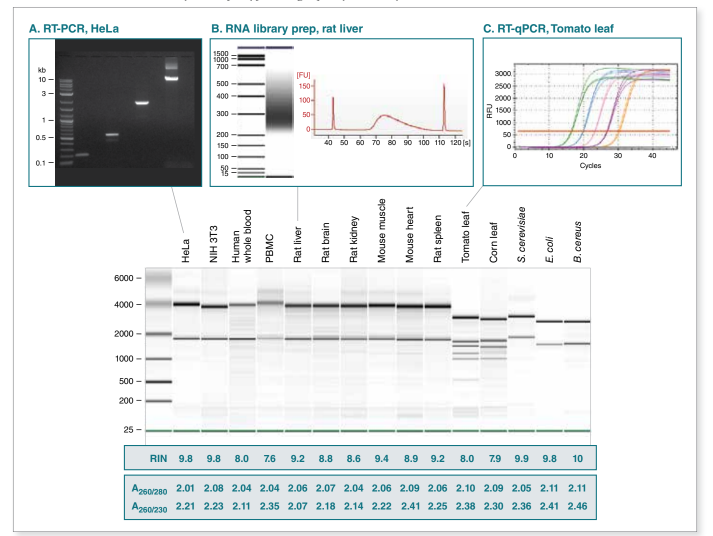


The Monarch Total RNA Miniprep Kit can generate high quality RNA from as few as 100 HeLa cells

Total RNA was isolated using the Monarch Total RNA Miniprep Kit from varying amounts of HeLa cells over 5 orders of magnitude and eluted in 100 µl of nuclease-free water. Samples were analyzed on a Bioanalyzer Pico chip, with RIN values and total yields shown below the lanes (A). Electropherograms are included as a reference (B).

### Monarch-purified RNA is High-quality and Compatible with a Wide Variety of Downstream Applications

RNA extracted from a wide variety of sample types is high quality and ready for downstream use



Total RNA from a broad array of sample types was purified using the Monarch Total RNA Miniprep Kit. Aliquots were run on an Agilent Bioanalyzer 2100 using the Nano 6000 RNA chip (S. cerevisiae RNA was run using a plant Nano assay). RIN values and O.D. ratios confirm the overall integrity and purity of the RNA. To demonstrate compatibility with downstream applications, samples were subsequently used for RT-PCR (+/– RT) (A) for detection of 4 different RNA species using Protoscript<sup>®</sup> II Reverse Transcriptase/LongAmp Taq DNA Polymerase, NGS library prep (B) using NEBNext Ultra Directional RNA Library Prep Kit and RT-qPCR (c) using Luna One-Step RT-qPCR Reagents.

### TIPS FOR SUCCESSFUL RNA EXTRACTIONS

- Prevent RNase Activity: Nucleases in your sample will degrade RNA, so inhibiting their activity is essential. Process samples
  quickly after harvest, use preservation reagents, and always ensure you are working in nuclease-free working environments.
- Inactivate RNases after harvesting your sample: Nucleases in your sample will lead to degradation, so inactivating them
  is essential. Process samples quickly, or use preservation reagents, and always ensure nuclease-free working environments.
- Do not exceed recommended input amounts: Buffer volumes are optimized for the recommended input amounts. Exceeding these can result in inefficient lysis and can also clog the column. See page 13.
- 4. Ensure samples are properly homogenized/disrupted: Samples should be disrupted and homogenized completely to release all RNA.
- For sensitive applications, ensure proper gDNA removal: gDNA is removed by the gDNA removal column and subsequent on-column DNase I treatment. Off-column DNase I treatment can also be employed.

# Sample Inputs & Expected Recovery for the Monarch Total RNA Miniprep Kit

RNA yield, purity and integrity vary immensely based on sample type, input amount and sample condition. Empirical yield and RIN data from a wide variety of sample types are provided below, as well as guidance on the maximum input amounts. It is important NOT to overload the column when extracting and purifying RNA, as yields, purity and integrity may suffer.

SAMPLE TYPE (1	)	INPUT	AVERAGE YIELD (µg)	OBSERVED Rin	MAXIMUM Starting material
CULTURED MAN	IMALIAN CELLS				
HeLa		1 x 10 <sup>6</sup> cells	12–15	9–10	1 x 107 cells
HEK 293		1 x 10 <sup>6</sup> cells	12–14	9–10	1 x 10 <sup>7</sup> cells
NIH3T3		1 x 10 <sup>6</sup> cells	8–12	9–10	1 x 10 <sup>7</sup> cells
MAMMALIAN BL	-00D <sup>(2)</sup>				
Human	Fresh	200 µl	0.5–1.0	7–8	3 ml
	Frozen	200 µl	0.5–1.0	7–8	3 ml
	Stabilized	200 µl	0.5–1.0	7–8	3 ml
Rat	Frozen	100 µl	5.6	9	1 ml*
BLOOD CELLS				1	
PBMC (isolated fro	om 5 ml whole blood)	5 ml	3	7	1 x 10 <sup>7</sup> cells
TISSUE			·		· ·
Rat liver	Frozen pulverized	10 mg	25	8–9	20 mg
	Stabilized solid	10 mg	50-60	8–9	20 mg
Rat spleen (stabiliz	ed solid with bead homogenizer)	10 mg	40–50	9	20 mg
Rat kidney (frozen	pulverized)	10 mg	7–10	9	50 mg
Rat brain	Frozen pulverized	10 mg	2–3	8–9	50 mg
	Stabilized solid	10 mg	0.5–1.5	8–9	50 mg
	Stabilized solid with bead homogenizer	10 mg	5–8	8–9	50 mg
Rat muscle (frozen	pulverized)	10 mg	2–3	8–9	50 mg
Mouse muscle	Frozen pulverized	10 mg	3	8–9	50 mg
	Powder with bead homogenizer	10 mg	5	7–8	50 mg
	Stabilized solid with bead homogenizer	10 mg	8–10	9	50 mg
Mouse heart (stabi	lized solid with bead homogenizer)	10 mg	5–6	8–9	50 mg
YEAST				1	
S. cerevisiae	Frozen with bead homogenizer	1 x 107 cells	50	9–10**	5 x 10 <sup>7</sup> cells
	Fresh with Zymolyase	1 x 10 <sup>7</sup> cells	60	9**	5 x 10 <sup>7</sup> cells
BACTERIA				1	
E. coli	Frozen	1 x 10 <sup>9</sup> cells	5	10	1 x 10 <sup>9</sup> cells
	Frozen with bead homogenizer	1 x 10 <sup>9</sup> cells	10	10	1 x 10 <sup>9</sup> cells
	Frozen with lysozyme	1 x 10 <sup>9</sup> cells	70	10	1 x 10 <sup>9</sup> cells
B. cereus	Frozen with lysozyme	1 x 10 <sup>8</sup> cells	20–30	9	1 x 10 <sup>9</sup> cells
	Frozen with bead homogenizer	1 x 10 <sup>8</sup> cells	8	9–10	1 x 10 <sup>9</sup> cells
PLANT					
Corn leaf (frozen p	ulverized with bead homogenizer)	100 mg	45	8	100 mg
	pulverized with bead homogenizer)	100 mg	30	8	100 mg

(1) RNA for other samples including drosophila, zebrafish embryos/larvae, plasma, serum, saliva, buccal swabs and nucleated blood have been successfully purified with this kit; protocols are available in the product manual online.

(2) Protocol for nucleated blood (e.g., birds, reptiles) is also available.

\* Mouse blood also has a maximum input of 1 ml.

\*\* S. cerevisiae total RNA was run on an Agilent Nano 6000 Chip using plant assay.

## Monarch Kits for RNA Cleanup

The Monarch RNA Cleanup Kits provide a fast and simple silica column-based solution for cleanup and concentration of RNA after enzymatic reactions (including *in vitro* transcription (IVT), DNase I and Proteinase K treatment, capping, tailing and labeling) as well as after RNA isolation (e.g., TRIzol extraction). These kits can also be used to extract RNA from cells, saliva and buccal/nasopharyngeal swabs. Kits are available in three different binding capacities:  $10 \mu g$ ,  $50 \mu g$  and  $500 \mu g$ , each containing unique columns designed to prevent buffer retention and ensure no carryover of contaminants.

#### Monarch Kit Specifications:

MONARCH RNA Cleanup kit	(10 µg)	(50 µg)	(500 µg)
Binding Capacity	10 µg	50 µg	500 µg
RNA Size Range	≥ 25	nt ( $\ge$ 15 nt with modified p	rotocol)
Typical Recovery		70-100%	
Elution Volume	6–20 µl	20–50 µl	50—100 µl
Purity		.8	
Protocol Time	5 minutes of spin and incubation time		10–15 minutes of spin and incubation time
Common Downstream Applications	RT-PCR, RNA library prep for NGS, small RNA library prep for NGS, RNA labeling	RT-PCR, RNA library prep for NGS, formation of RNP complexes for genome editing, microinjection, RNA labeling, transfection	RT-PCR, RNA library prep for NGS, RNA labeling, RNAi, microinjection, transfection

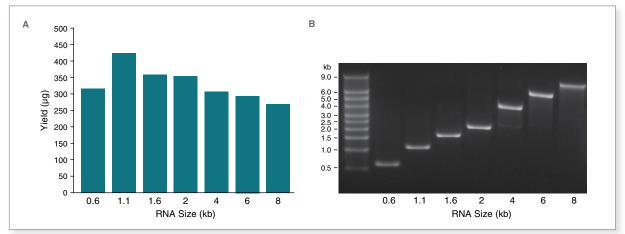
#### ADVANTAGES

- Isolate highly pure RNA ( $A_{260/280}$  and  $A_{260/230} \ge 1.8$ ) in minutes
- Clean up RNA with simple protocol utilizing a single wash buffer
- Elute in as little as 6 µl or 20 µl
- Bind up to 500 µg of RNA
- Adjust cutoff size down to 15 nt with a slight protocol modification
- Can be used for RNA extraction from some samples; extraction from saliva can be automated on the QIAcube and KingFisher Flex platforms

#### APPLICATIONS

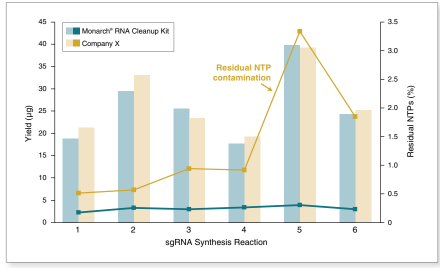
- Cleanup & concentration after enzymatic reactions (e.g., DNase I and Proteinase K treatment)
- Cleanup after RNA synthesis (IVT and sgRNA synthesis)
- Cleanup & concentration of previously-purified RNA (e.g., after TRIzol extraction)
- RNA extraction from cells, saliva and swabs (buccal/NP)
- RNA Gel Extraction

The Monarch RNA Cleanup Kit (500  $\mu$ g) is suitable for cleaning up large quantities (> 250  $\mu$ g) of RNA from *in vitro* transcription reactions



A. RNA transcripts of varying sizes (0.6-8 kb) were synthesized using the HiScribe<sup>™</sup> T7 Quick High Yield RNA Synthesis Kit. 40 µl of each in vitro transcription (IVT) reaction was cleaned up using the Monarch RNA Cleanup Kit (500 µg). RNA yields were calculated from the resulting A<sub>260</sub>, measured using a Nanodrop spectrophotometer and ranged from 268–425 µg of RNA per IVT reaction.

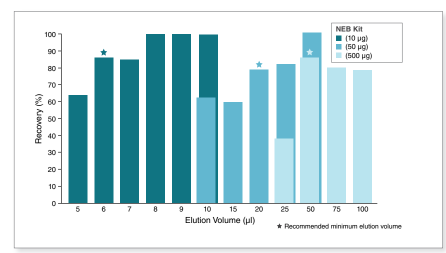
B. RNA integrity (200 ng/lane) was assessed on a 1% agarose-TBE gel stained with SYBR® Gold.



The Monarch RNA Cleanup Kit (50  $\mu g)$  produces sgRNA yields consistent with other competitor RNA cleanup kits and with lower residual NTP contamination

Six different sgRNA synthesis reactions from the EnGer® sgRNA Synthesis Kit, S. pyogenes were cleaned up using either the Monarch RNA Cleanup Kit (50  $\mu$ g) or a competitor kit (according to manufacturer's recommendations) and eluted in 50  $\mu$ l nuclease-free water. sgRNA yield was calculated from the resulting A<sub>200</sub> measured using a Trinean DropSense 16. The Monarch RNA Cleanup Kit produced sgRNA yields consistent with other commercially available RNA cleanup kits.

Following cleanup, residual nucleotides (NTPs) were measured by LC-MS and are reported as percent area NTPs (rATP+rCTP+rGTP+rUTP)/percent area sgRNA). The NEB Monarch RNA Cleanup Kit consistently outperforms other commercially available RNA cleanup kits in the removal of residual NTPs from sgRNA synthesis reactions.



Recovery of RNA from Monarch RNA Cleanup Kits with varying elution volumes

10, 50 or 500  $\mu$ g of RNA (16S and 23S Ribosomal Standard from E. coli, Sigma) was purified using a Monarch RNA Cleanup Kit (10  $\mu$ g) (50  $\mu$ g) (500  $\mu$ g). Nuclease-free water was used to elute the RNA. The percent recovery of the RNA was calculated from the resulting  $A_{260}$  as measured using a Trinean DropSense 16. ~80% of RNA can be efficiently recovered in 6  $\mu$ l from the Monarch RNA Cleanup Kit (10  $\mu$ g), 20  $\mu$ l from the Monarch RNA Cleanup Kit (50  $\mu$ g), and 50  $\mu$ l from the Monarch RNA Cleanup Kit (500  $\mu$ g).

We recently switched to NEB for our spin column kits, and I just wanted to say that I am so impressed with the thoughtfulness of the packaging—how the protocol is stored, the tiny bags that the columns are in, and the quality of the box itself, which we will keep to store items after we finish the kit. We also like that you can customize the elution volume. Cheers to your team for doing your part and designing a great product!

## Monarch Kits for your DNA Cleanup and Gel Extraction Needs

Monarch DNA cleanup kits rapidly and reliably purify up to 5  $\mu$ g of concentrated, high-quality DNA. These kits utilize a bind/wash/ elute workflow with minimal incubation and spin times. The columns provided with each kit ensure zero buffer retention and no carryover of contaminants, enabling elution of sample in volumes as low as 6  $\mu$ l. Monarch Buffers have been optimized, and do not require monitoring of pH. Eluted DNA is ready for use in restriction digests, DNA sequencing, ligation and other enzymatic manipulations.

### Monarch DNA Gel Extraction Kit

The Monarch DNA Gel Extraction Kit can be used to quickly purify DNA from agarose gels. Unlike other kits, there is no need to add isopropanol to the melted agarose prior to loading on the column, saving you a step. Enjoy high yields and minimal hands on time.

# Monarch PCR & DNA Cleanup Kit (5 µg)

The Monarch PCR & DNA Cleanup Kit (5  $\mu$ g) can be used to purify DNA from a variety of enzymatic reactions, such as PCR, restriction digestion, ligation and reverse transcription. The DNA Wash Buffer provided ensures enzymes, short primers ( $\leq 25$  nt), detergents and other low-molecular weight reaction components (e.g., nucleotides, DMSO, betaine) are removed. A simple protocol modification also enables purification of small DNA and oligonucleotides.



Optimized Monarch column design

Many purification columns are built with a retaining ring to hold the membrane in place, but this can trap buffer. Monarch columns' silica matrix is held in place without the use of a retaining ring, eliminating buffer retention and ensuring worry-free purification.

> This is a great, easy-to-use, small footprint kit... it was great to elute in such a small volume while feeling confident that the elution buffer managed to get to all of the surface area of the membrane.

#### ADVANTAGES

- Elute in as little as 6 µl
- Prevent buffer retention and salt carryover with optimized column design
- Purify oligos and other small DNA fragments with simple protocol modification
- · Save time with fast, user-friendly protocols
- Designed with sustainability in mind
- With protocol modification, DNA ≥ 15 bp (dsDNA) or ≥ 18 nt (ssDNA) can be purified with Monarch PCR & DNA Cleanup Kit (5 µg)

#### SPECIFICATIONS

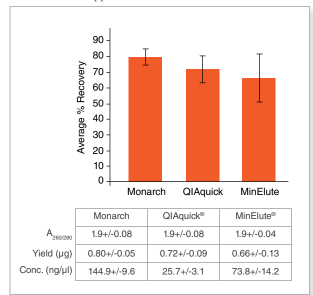
- Binding Capacity: up to 5 µg
- DNA Size Range: ~50 bp to 25 kb With protocol modification, oligos  $\geq$  15 bp (dsDNA) or  $\geq$  18 nt (ssDNA) can be purified with Monarch PCR & DNA Cleanup Kit (5 µg)
- Typical Recovery: DNA (50 bp to 10 kb): 70–90% DNA (11–25 kb): 50–70% ssDNA ≥ 18 nt and dsDNA ≥ 15 bp: 70–85% [Monarch PCR & DNA Cleanup Kit (5 μg) only]
- Elution Volume:  $\geq 6 \ \mu l$
- **Purity:** A<sub>260/280</sub> ≥ 1.8
- Protocol Time: Gel Extraction: 10 min of spin and incubation time PCR & DNA Cleanup: 5 min of spin and incubation time
- Compatible Downstream Applications: ligation, restriction digestion, labeling and other enzymatic manipulations, library construction and DNA sequencing

# TIPS FOR SUCCESSFUL

- 1. Use the smallest possible agarose plug: The less agarose in solution, the more efficient the extraction will be. More agarose means more melting time and more buffer needed to dissolve it (introducing more salts which can co-elute with your sample). If the plug is greater than 160 mg, the volume of agarose plus buffer will exceed that of the column reservoir (800 µl), and will require that your sample be loaded onto the column in two steps.
- Minimize exposure to UV light: Excise the gel slice as quickly as possible, as exposure to UV light damages DNA. As long as the excision is done quickly, damage will be negligible.
- Melt the agarose completely: If the agarose is not completely melted, DNA remains trapped inside and cannot be extracted properly.
- 4. Heat the elution buffer for large DNA fragments: Large DNA binds more tightly; heating the elution buffer helps to more efficiently release the DNA from the column matrix.

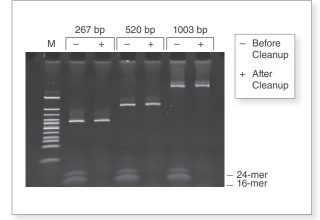
- Michelle, Central Michigan University

DNA purified from the Monarch DNA Gel Extraction Kit is recovered with similar efficiency and purity as the leading supplier, but is more highly concentrated, facilitating its use in downstream applications



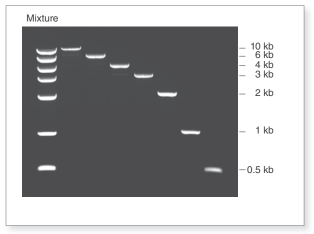
One microgram aliquots of a 3 kb fragment were resolved on a 1% w/v agarose gel, excised, and processed with different kits using manufacturer-specified minimum elution volumes. Values reported are the concentration and purity data determined by Nanodrop<sup>™</sup> readings, as well as recovery calculations based on the eluted DNA concentration and recovered volume.

Monarch PCR & DNA Cleanup Kit (5 µg) removes low molecular weight primers from dsDNA samples



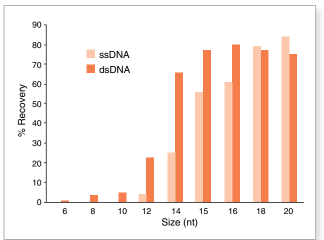
Three independent amplicons (267 bp, 520 bp, 1003 bp) were spiked with two oligonucleotides (16-mer, 24-mer) to a final concentration of 1  $\mu$ M. Half of each mix was purified with the Monarch PCR & DNA Cleanup Kit (5  $\mu$ g) following the included protocol. Equivalent fractions of the original mixture and the eluted material were resolved on a 20% TBE acrylamide gel at 100V for one hour and stained with SYBR Green II.

Monarch DNA Gel Extraction Kit reproducibly recovers DNA over a broad range of molecular weights



A mixture of 7 DNA fragments ranging from 10 kb to 0.5 kb was prepared and one-half of the mixture was resolved on a 1% w/v agarose gel. Each fragment was manually excised from the agarose gel and processed using the Monarch DNA Gel Extraction Kit. The entire elution of each fragment was resolved on a new gel with the remainder of the original mixture for comparison.

With simple protocol modification, the Monarch PCR & DNA Cleanup Kit can efficiently recover ssDNA and dsDNA oligonucleotides, expanding the utility of the kit

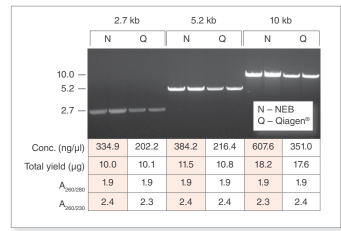


Synthesized ssDNA and dsDNA oligonucleotides (1 µg in 50 µl H<sub>2</sub>O) of varying lengths (6-20 nt) were purified using the oligonucleotide cleanup protocol and the Monarch PCR & DNA Cleanup Kit, and were eluted in 50 µl water. The average percent recovery (n-3) of the oligonucleotides was calculated from the resulting  $A_{260}$  as measured using a Trinean® DropSense<sup>™</sup> 16. Use of the Oligonucleotide Cleanup Protocol and the Monarch PCR & DNA Cleanup Kit results in the efficient removal of small oligonucleotides (6-12 nt) and > 70% recovery and cleanup of oligonucleotides ≥ 15 bp (dsDNA) or ≥ 18 nt (ssDNA).

# Monarch Plasmid Miniprep Kit

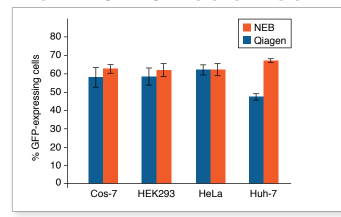
The Monarch Plasmid Miniprep Kit is a rapid and reliable method for the purification of high quality plasmid DNA. This method employs standard cell resuspension, alkaline lysis, and neutralization steps, with the additional benefit of color indicators at certain steps to easily monitor completion. Unique wash buffers ensure salts, proteins, RNA and other cellular components are removed, allowing low-volume elution of concentrated, highly pure DNA. Protocols are fast and userfriendly. Elution in as little as 30  $\mu$ l provides concentrated DNA for use in downstream applications, such as restriction digests, DNA sequencing, PCR and other enzymatic manipulations.

Monarch Plasmid Miniprep Kits consistently produce more concentrated plasmid DNA with equivalent yield, purity and functionality as compared to the leading supplier



Preps were performed according to recommended protocols using 1.5 ml aliquots of the same overnight culture. One microliter of each prep was digested with HindIII-HF® to linearize the vector and the digests were resolved on a 1% w/v agarose gel.

Plasmid DNA purified using the Monarch Plasmid Miniprep Kit produces transfection efficiencies equivalent to or better than plasmid DNA purified using the Qiagen QIAprep® Spin Miniprep Kit



Plasmid DNA encoding constitutively expressed GFP (pEGFP-C2) was prepared using either Monarch Plasmid Miniprep Kit or Qiagen QIAprep Spin Miniprep Kit. Four different cell lines (Cos-7, HEK293, HeLa, and Huh-7) were grown to 80-90% confluence and transfected with 100 ng of each plasmid, in complex with 0.3 µL Lipofectamine 2000, and 10 µL Opti-MEM. Five replicates for each cell type were performed using both DNA preps. GFP expressing cells were counted by flow cytometry 48 hrs post-transfection with a minimum of 2000 events collected per well. Average percentage of cells expressing GFP from all replicates is graphed and used as a measure of transfection efficiency.

#### ADVANTAGES

- Elute in low volumes
- Prevent buffer retention and salt carryover with optimized column design
- · Reduce hands on time with faster protocols and less spin time
- · Monitor completion of certain steps using colored buffer system
- · No need to add RNase before starting
- · Easily label columns using tab and frosted surfaces

#### **SPECIFICATIONS**

- Culture Volume: 1-5 ml, not to exceed 15 0.D. units
- Binding Capacity: up to 20 µg
- · Plasmid Size: up to 25 kb
- Typical Recovery: up to 20 µg, yield depends on plasmid copy number, host strain, culture volume, and growth conditions
- Elution Volume: ≥ 30 µl
- Purity:  $A_{260/280}$  and  $A_{260/230} \ge 1.8$
- Protocol Time: 91/2-121/2 minutes of spin and incubation time
- Compatible Downstream Applications: restriction digestion and other enzymatic manipulations, transformation, transfection of robust cells, DNA sequencing, PCR, labeling, cell-free protein synthesis, etc.

### <sup>)</sup> TIPS FOR SUCCESSFUL MINIPREPS

- Don't use too many cells (culture should not exceed 15 0.D. units): Using the optimal amount of cells increases lysis efficiency and ensures that excess cell debris does not clog the column.
- Lyse cells completely: In order to release all plasmid DNA, ALL of the cells need to be lysed. Resuspend cells completely, and incubate for the recommended time.
- Don't vortex cells after lysis: Vortexing can cause shearing of host chromosomal DNA, resulting in gDNA contamination.
- Allow the RNase to do its job: Do not skip or reduce the incubation with RNase (which is included in the neutralization buffer), otherwise you may observe RNA contamination.
- Don't skip any washes: Proper washes ensure the removal of cell debris, endotoxins and salts.
- Heat the elution buffer for large plasmids: Large DNA binds more tightly; heating the elution buffer helps to more efficiently release the DNA from the column matrix.

# Troubleshooting Guide for Genomic DNA Purification

PROBLEM	CAUSE	SOLUTION
LOW YIELD		
Cells	Frozen cell pellet was thawed and/or resuspended too abruptly	<ul> <li>Thaw cell pellets slowly on ice and flick tube several times to release the pellet from bottom of tube. Use cold PBS, and resuspend gently by pipetting up and down 5–10 times until pellet is dissolved</li> </ul>
	Cell Lysis Buffer was added concurrently with enzymes	Add Proteinase K and RNase A to sample and mix well before adding the Cell Lysis Buffer
	Blood was thawed, allowing for DNase activity	* Keep blood samples frozen and add Proteinase K, RNase A and Blood Lysis Buffer directly to the frozen samples
Blood	Blood sample is too old	* Fresh (unfrozen) whole blood should not be older than 1 week. Older samples will show progressive DNA degradation and loss of yield.
	Formation of hemoglobin precipitates	<ul> <li>In blood from species with high hemoglobin content, (e.g., guinea pig) insoluble hemoglobin complexes may accumulate and clog the membrane. Reduce Proteinase K lysis time from 5 to 3 minutes.</li> </ul>
	Tissue pieces are too large	• Cut starting material to the smallest possible pieces or grind with liquid nitrogen. In large tissue pieces, nucleases will destroy the DNA before the Proteinase K can lyse the tissue.
	Membrane is clogged with tissue fibers	<ul> <li>Proteinase K digestion of fibrous tissues (e.g., muscle, heart, skin, ear clips), brain tissue and all RNAlater-stabilized tissues leads to the release of small indigestible protein fibers, which block the binding sites of the silica membrane. To remove fibers, centrifuge lysate at maximum speed for 3 minutes, as indicated in the protocol. For ear clips and brain tissue, use no more than 12–15 mg input material.</li> </ul>
Tissue	Sample was not stored properly	<ul> <li>Samples stored for long periods of time at room temperature, 4°C or -20°C, will show degradation and loss of gDNA. Flash freeze tissue samples with liquid nitrogen or dry ice and store at -80°C. Alternatively, use stabilizing reagents to protect the gDNA.</li> </ul>
	Genomic DNA was degraded (common in DNase-rich tissues)	* Organ tissues (e.g., pancreas, intestine, kidney, liver) contain significant amounts of nucleases. Store properly to prevent DNA degradation. Keep on ice during sample preparation. Refer to the protocol for the recommended amount of starting material and Proteinase K to use.
	Column is overloaded with DNA	* Some organ tissues (e.g., spleen, kidney, liver) are extremely rich in genomic DNA. Using inputs larger than recommended will result in the formation of tangled, long-fragment gDNA that cannot be eluted from the silica membrane. Reduce the amount of input material.
	Incorrect amount of Proteinase K added	• Most samples are digested with 10 µl Proteinase K, but for brain, kidney and ear clips, use 3 µl.
DNA DEGRAD	ATION	
	Tissue samples were not stored properly	<ul> <li>Samples stored for long periods of time at room temperature, 4°C or -20°C, will show degradation and loss of gDNA. Flash freeze tissue samples with liquid nitrogen or dry ice and store at -80°C. Alternatively, use stabilizing reagents to protect the gDNA.</li> </ul>
Tissue	Tissue pieces are too large	<ul> <li>Cut starting material to the smallest possible pieces or grind with liquid nitrogen. In large tissue pieces, nucleases will degrade the DNA before Proteinase K can lyse the tissue.</li> </ul>
	High DNase content of soft organ tissue	<ul> <li>Organ tissues (e.g., pancreas, intestine, kidney, liver) contain significant amounts of nucleases. Store properly to prevent DNA degradation.</li> <li>Keep on ice during sample preparation. Refer to the protocol for the recommended amount of starting material and Proteinase K to use.</li> </ul>
	Blood sample is too old	• Fresh (unfrozen) whole blood should not be older than 1 week. Older samples will show progressive DNA degradation and loss of yield.
Blood	Blood was thawed, allowing for DNase activity	• Keep frozen blood samples frozen and add enzymes and lysis buffer directly to the frozen samples
SALT CONTAN	INATION	
	Guanidine thiocyanate salt	<ul> <li>When transferring the lysate/binding buffer mix, avoid touching the upper column area with the pipet tip and always pipet carefully onto the silica membrane.</li> </ul>
	from the binding buffer	* Avoid transferring any foam that may have been present in the lysate; foam can enter into the cap area of the spin column.
	was carried over into the eluate	<ul> <li>Close the caps gently to avoid splashing the mixture into the upper column area and move the samples with care in and out of the centrifuge.</li> <li>If salt contamination is a concern, invert the columns a few times (or vortex briefly) with oDNA Wash Buffer as indicated in the protocol.</li> </ul>
PROTEIN CON	ΤΑΜΙΝΑΤΙΩΝ	• It sait containination is a concern, invert the columnis a lew times (or vortex brieny) with going wash builer as indicated in the protocol.
FROTEIN GON	Incomplete digestion	<ul> <li>Cut samples to the smallest possible pieces. Incubate sample in the lysis buffer for an extra 30 minutes to 3 hours to degrade any remaining protein complexes.</li> </ul>
Tissue	Membrane is clogged with tissue fibers	<ul> <li>Proteinase K digestion of fibrous tissues (e.g., muscle, heart, skin, ear clips), brain tissue and all RNAlater-stabilized tissues leads to the release of small, indigestible protein fibers, which block the binding sites of the silica membrane. To remove fibers, centrifuge the lysate at maximum speed for 3 minutes as indicated in the protocol. For ear clips and brain tissue, use no more than 12–15 mg input material.</li> </ul>
	High hemoglobin content	Some blood samples (e.g., horse) are rich in hemoglobin, evidenced by their dark red color. Extend lysis time by 3–5 minutes for best purity results.
Blood	Formation of hemoglobin precipitates	<ul> <li>In blood from species with high hemoglobin content, (e.g., guinea pig) insoluble hemoglobin complexes may accumulate and clog the membrane. Reduce Proteinase K lysis time from 5 to 3 minutes.</li> </ul>
RNA CONTAM	INATION	
Tissue	Too much input material	<ul> <li>DNA-rich tissues (e.g., spleen, liver and kidney) will become very viscous during lysis and may inhibit RNase A activity. Do not use more than the recommended input amount.</li> </ul>
	Lysis time was insufficient	Extend lysis time by 30 minutes to 3 hours after the tissue piece has completely dissolved
ITSSUE DIGES	STION TAKES TOO LONG	• Out tissue pisses to the smallest percible size or grind with liquid altragen before statise typic
	Tissue pieces too large	Cut tissue pieces to the smallest possible size or grind with liquid nitrogen before starting lysis     Vortex to release pieces from the tube bottom, and immediately after adding Proteinase K and Tissue Lysis Buffer
	Tissue pieces are stuck to bottom of tube	
TISSUE LYSAT	Too much starting material	Use recommended input amount
HOOL LIGH	Formation of indigestible fibers	<ul> <li>Proteinase K digestion of fibrous tissues (e.g., muscle, heart, skin, ear clips), brain tissue and all RNAlater-stabilized tissues leads to the release of small indigestible protein fibers, which block the binding sites of the silica membrane. To remove fibers, centrifuge lysate at maximum speed for 3 minutes, as indicated in the protocol. For ear clips and brain tissue, use no more than 12–15 mg input material.</li> </ul>
RATIO A <sub>260</sub> /A <sub>23</sub>	> 2.5	
260 23	Slight variations in EDTA concentration in eluates	<ul> <li>EDTA in elution buffer may complex with cations like Mg<sup>2+</sup> and Ca<sup>2+</sup> samples present in genomic DNA, which may lead to higher than usual A<sub>260</sub>/A<sub>230</sub> ratio. In some cases, this ratio exceeds a value of 3.0 and is consistent with highly pure samples. In these cases, the elevated value does not have any negative effect on downstream applications.</li> </ul>

# Troubleshooting Guide for Total RNA Extraction & Purification

PROBLEM	CAUSE	SOLUTION
Clogged column	Insufficient sample disruption or homogenization	<ul> <li>Increase time of sample digestion or homogenization</li> <li>Centrifuge sample after Proteinase K digestion or homogenization to pellet debris and use only supernatant for next steps</li> <li>Use larger volume of DNA/RNA Protection Reagent and/or RNA Lysis Buffer for sample disruption and homog- enization. See sample-specific protocols in the product manual.</li> </ul>
	Too much sample	<ul> <li>Reduce amount of starting material to match kit specifications to ensure buffer amounts are sufficient and column is not overloaded. See Guidelines for Choosing Sample Input Amounts on page 13.</li> </ul>
	Incomplete elution	<ul> <li>After addition of Nuclease-free Water to column matrix, incubate 5-10 min at room temperature and then centrifuge to elute</li> <li>Perform a second elution (note: this will dilute sample)</li> </ul>
	Sample is degraded	<ul> <li>Store input sample at -80°C prior to use</li> <li>Use Monarch DNA/RNA Protection Reagent to maintain RNA integrity during storage</li> </ul>
Low RNA yield	Insufficient disruption or homogenization	<ul> <li>Increase time of sample digestion or homogenization</li> <li>Centrifuge sample after Proteinase K digestion or homogenization to pellet debris and use only supernatant for next steps</li> <li>Use larger volume of DNA/RNA Reagent and/or RNA Lysis Buffer for sample disruption and homogenization. See sample specific protocol in the product manual.</li> <li>For Proteinase K treated samples, doubling Proteinase K (from 5% to 10%) may lead to an increase in RNA yield</li> </ul>
	Too much sample	<ul> <li>Reduce amount of starting material to match kit specifications to ensure buffer amounts are sufficient and column is not overloaded. See Guidelines for Choosing Sample Input Amounts on page 13.</li> </ul>
	Starting material not handled/stored properly	<ul> <li>Store input sample at -80°C prior to use. Degradation of RNA may occur if sample is not flash frozen or protected by a preservation reagent. Use Monarch DNA/RNA Protection Reagent to maintain RNA integrity during storage.</li> </ul>
RNA degradation	Deviation from the stated protocol may expose RNA to unwanted RNase activities	• Refer to the General Guidelines for Working with RNA in the product manual
	RNase contamination of eluted materials or kit buffers may have occurred	<ul> <li>See General Guidelines for Working with RNA in the product manual for advice on reducing risks of contamination</li> </ul>
	Low $\rm A_{\rm 260/280}$ values indicate residual protein in the purified sample	• Ensure the Proteinase K step was utilized for the recommended time. Ensure samples have no debris prior to addition of ethanol and loading onto RNA Purification Column.
Low OD ratios	Low A <sub>260/230</sub> values indicate residual guanidine salts have been carried over during elution	<ul> <li>Ensure wash steps are carried out prior to eluting sample. Use care to ensure the tip of the column does not contact the flow-through after the final wash. If unsure, please repeat centrifugation. When reusing collection tubes, blot rim of tube on a Kimwipe prior to reattachment to the column to remove any residual wash buffer.</li> </ul>
DNA contamination	Genomic DNA not removed by column	<ul> <li>Perform optional on-column DNase I treatment to remove unwanted gDNA from lysed sample</li> <li>Perform in-tube/off-column DNase I treatment to remove gDNA</li> </ul>
	Too much sample	<ul> <li>Reduce amount of starting material to match kit specifications to ensure buffer amounts are sufficient and column is not overloaded. See Guidelines for Choosing Sample Input Amounts on page 13.</li> </ul>
Low performance of RNA in downstream steps	Salt and/or ethanol carryover has occurred	<ul> <li>Use care to ensure the tip of the RNA Purification Column does not contact the flow-through after the final wash. If unsure, please repeat centrifugation.</li> <li>Be sure to spin the RNA Purification Column for 2 minutes following the final wash with RNA Wash Buffer</li> <li>When reusing collection tubes, blot rim of tube on a Kimwipe prior to reattachment to the column to remove any residual wash buffer</li> <li>Add additional wash step and/or extend spin time for final wash</li> </ul>
Unusual spectrophoto- metric readings	RNA concentration is too low for spectro- photometric analysis	<ul> <li>For more concentrated RNA, elute with 30 µl of nuclease-free water</li> <li>Increase amount of starting material (within kit specifications). See Guidelines for Choosing Sample Input Amounts on page 13.</li> </ul>
	Silica fines in eluate	<ul> <li>Re-spin eluted samples and pipet aliquot from the top of the liquid to ensure the A<sub>260/230</sub> is unaffected by possible elution of silica particles</li> </ul>

# Troubleshooting Guide for RNA Cleanup

PROBLEM	CAUSE	SOLUTION
	Reagents added incorrectly	<ul> <li>Check protocol to ensure correct buffer reconstitution, order of addition for buffers and ethanol, and proper handling of column flow-through and eluents</li> </ul>
	Insufficient mixing of reagents	<ul> <li>Ensure the ethanol is thoroughly mixed with mixture of RNA sample and RNA Binding Buffer before applying sample to the purification column</li> </ul>
Low RNA yield	Incomplete elution during prep	<ul> <li>Ensure the nuclease-free water used for elution is delivered directly to the center of the column so that the matrix is completely covered and elution is efficient. Larger elution volumes and longer incubation times can increase yield of RNA off the column at the cost of dilution of the sample and increased processing times. For typical RNA samples, the recommended elution volumes and incubation times should be sufficient. Additionally, multiple rounds of elution can be employed to increase the amount of RNA eluted, at the expense of dilution of the sample. The first elution can be used to elute a second time to maximize recovery and minimize sample dilution.</li> </ul>
	Complex secondary structure affects binding and elution of smaller RNAs (< 45 nt)	• Diluting your sample with 2 volumes of ethanol instead of one volume in Step 2.
	Poor extraction from Agarose gel	<ul> <li>Be sure to incubate the sample between 37–55°C after addition of both the RNA Cleanup Binding Buffer and the ethanol. Additionally, incubate the column with nuclease-free water at 65°C for 5 minutes prior to spinning to elute the RNA.</li> </ul>
Purified RNA is degraded	RNase contamination	<ul> <li>Wear gloves and use disposable RNase-free tips (not provided) and collection tubes during the procedure</li> <li>Keep all kit components tightly sealed when not in use</li> </ul>
	Improper storage of RNA	• Use RNA immediately in downstream applications or store at -70°C
		Ensure wash steps are carried out prior to eluting sample
Low OD ratios	Residual guanidine salts have been carried over during elution	<ul> <li>Use care to ensure the tip of the column does not contact the flow-through. If unsure, please repeat centrifugation</li> </ul>
		<ul> <li>When reusing collection tubes, blot rim of tube on a Kimwipe prior to reattachment to the column to remove any residual wash buffer</li> </ul>
Low performance of RNA in downstream steps	Salt and/or ethanol carry-over	• Use care to ensure that the tip of the column does not come into contact with the flow-through. If in doubt, re-spin for 1 minute to ensure traces of salt and ethanol are not carried over in the eluted RNA
DNA contamination	DNA present in sample	<ul> <li>Incubate RNA sample with DNase I and clean up RNA using the Monarch RNA Cleanup Protocol</li> </ul>

# Troubleshooting Guide for DNA Cleanup & Plasmid Purification

No DNA purified         Reader Reamand Miningrey Rei         Relines added incomacily Plaunial lass during caluture growth caluture growth         In the contest squeece Plaunial lass during caluture growth         Ensure strong calutogical and concentration was used to maintain solecitor) and calutary growth           Morarch DNA Gie Extraction RA (Fig.)         Morarch DNA Gie Extraction RA (Fig.)         Partial added to wash buffer         Plann added to wash buffer         Plann added to wash buffer           Fig.)         Morarch DNA Gie Extraction RA (Fig.)         Interruption added to wash buffer         Plann added to wash buffer         Plann added to wash buffer           Fig.)         Fig.)         Interruption added to wash buffer         Plann added to wash buffer         Plann added to wash buffer           Fig.)         Fig.)         Interruption added to wash buffer         Plann added to wash buffer         Plann added to wash buffer           Fig.)         Fig.)         Fig.)         Plann added to wash buffer         Plann added to wash buffer           Fig.)         Fig.)         Fig.)         Fig.)         Plann added to wash buffer         Plann added to wash buffer           Fig.)	PROBLEM	PRODUCT	POSSIBLE CAUSE	SOLUTION
No DMA purified         Income to MMA call Education Kask and guidantial (scaling location)         Judies gradual           Moves DVIA Call Education Kask         Head In the added to weah buffer         Feature the proper amount of standard burdare buffer (SR) – calces and call charge transition of standard burdare buffer (SR) – calces and charge transition of estimated to deep feature in SR) – calces and charge transition of estimated to deep feature in SR) – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition of estimated and the adverted SD – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition of a consider buffer (SR) – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition of the standard burdare buffer (SR) – calces and charge transition and concentration are standare stan		Monarch Plasmid Miniprep Kit	Buffers added incorrectly	
Image:	No DNA purified		Plasmid loss during culture growth	Ensure proper antibiotic and concentration was used to maintain selection during culture growth
Knowski PCR & DNA Cleanay Ril (sp.)         Knowski PCR & DNA Cleanay Ril (sp.) <thknowski &="" cleanay="" dna="" pcr="" ril<br="">(sp.)         Knowski PC</thknowski>		Monarch DNA Gel Extraction Kit		
Lev DIA yield               isolate high and high			<ul> <li>Ethanol not added to wash buffer</li> </ul>	Ensure the proper amount of ethanol was added to Monarch DNA Wash Buffer
Lew DAA yield         Pearate loss during culue growth         Compare analysis of the server and server analysis of the server analysis of th			Incomplete lysis	<ul> <li>Pellet must be completely resuspended before addition of Plasmid Lysis Buffer (B2) – color should change from light to dark pink</li> <li>Avoid using too many cells; this can overload the column. If culture volume is larger than recommended, scale up buffers B1-B3 or consider using two columns.</li> </ul>
Lew DIA yield         Interact Plasmid Miniprep K1         Inclusion ground (under growh)         Inclusion growh)           Lew DIA yield         Inclusion growh         - Inclusion growh)         - Inclusion growh)         - Inclusion growh)           Lew DIA yield         Incomplete neutralization         - Increase yield incomplete neutralization         - Increase yield incomplete neutralization           Lew DIA yield         Incomplete neutralization         - Increase yield incomplete neutralization         - Increase yield incomplete neutralization           Monarch DIA Gel Extraction K1         Gel discoved abox 60°C         - Discove grow grow grow grow grow grow grow grow			Too many cells used	* The amount of cells used should not exceed 15 0.D. units
Low DNA yield <ul> <li> <ul> <li> <ul> <li> <li></li></li></ul></li></ul></li></ul>			Plasmid loss during culture growth	* Ensure proper antibiotic and concentration was used to maintain selection during culture growth
Lev DIA yield         Lysis to close during jub/wi         (-12-16 hours).           Low DIA yield         Incomplete neutralization         • hover tube several times unit close changes to a unitom yellow color           Low DIA yield         Particular biller directly to center of column         • Calver tube several times unit close changes to a unitom yellow color           Low DIA yield         Particular biller directly to center of column         • Calver tube several times unit close changes to a unitom yellow color           Morarch DIA GE Extraction Kitt         Buffers added incorrectly         • Several column         • Several column           Gel size not fully dissolved above 60°C         Disosve pel size in specified range (37-55°C). Higher temperatures can deter discolum and interfere with binding. Include to the content and interfere with binding. Include to the content and the context of the column         • Calver tiluino Buffer to 50°C and extend include to the content and the context of the column.           Morarch DRA & DIA Cleanup Kitt         Gel dissolved above 60°C         Dissolve pel size in specified range (37-55°C). Higher temperatures can deter discolumn.           Morarch PCR & DIA Cleanup Kitt         Gel dissolved above 60°C         Dissolve pel size in specified range (37-55°C). Higher temperatures can deter discolumn.           Morarch PCR & DIA Cleanup Kitt         Buffers added incorrectly         • Calver tiluino Buffer inclation times can increase yield incortectly and the context ode           Lew DIA quality         Parelicion of DIA - 100		Monarch Plasmid Miniprep Kit	Low-copy plasmid selected	* Increase amount of cells processed and scale buffers accordingly
Low DNA yield         • Deliver Elution Buffer directly to center of column           Low DNA yield         • Deliver Elution Outpaties and longer incubation films can increase yield           Monarch DNA Gel Extraction Kitt         Buffers added incorrectly         • Be surf buffers films be been reconstituted correctly and that reagents have builds in the corr and temperature.           Gel sile on thully dissolved         • • • • • • • • • • • • • • • • • • •			Lysis of cells during growth	
Incomplete elution         -Larger elution variums and longer incutation times can increase yield -fielding of plants by the the burn Buffer is SPC and eater incubation time is 5 minutes           Low DNA yield         Applet is added incorrectly         -B gusten buffers by been reconstituted correctly and that respents have be obtained in the correct order         -B gusten buffers by been reconstituted correctly and that respents have be obtained for Dissolving Buffer for proper time and interpretatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in specified range (37-55°C). Higher temperatures can dena Dissolve gel slice in solution can a			Incomplete neutralization	• Invert tube several times until color changes to a uniform yellow color
Low DNA yreid         Example in the constraint of t			Incomplete elution	Larger elution volumes and longer incubation times can increase yield     For elution of plasmids > 10 kb, heat the DNA Elution Buffer to 50°C and extend
Low DNA quality         Monarch DNA Gel Extraction Kit         Gel since from using discover dialoge 60°C         Discover gel silce in specified range (37-55°C). Higher temperatures can dera DNA           honarch DNA Gel Extraction Kit         Gel dissolved above 60°C         - Discover gel silce in specified range (37-55°C). Higher temperatures can dera DNA           honarch PRA Se DNA Cleanup Kit         Incomplete elution during preparation         - Peierer fultion of Unance and Ionger incubation times can increase yield           honarch PCR & DNA Cleanup Kit         Buffers added incorrectly         - Peierer Elution Buffer directly to center of column           honarch PCR & DNA Cleanup Kit         For elution of Unance and Ionger incubation times can increase yield           honarch PRR & DNA Cleanup Kit         For elution during preparation         - Peierer Elution Buffer directly to center of column           honarch PRR & DNA Cleanup Kit         For elution during preparation         - Peierer Elution Buffer directly to center of column           honarch Plasmid Miniprep Kit         For elution during preparation         - Peierer Elution Numerature te plasmid         - Peierer Elution Numerature te plasmid           Monarch Plasmid Miniprep Kit         Plasmid degradation         - Peierer elution vibring Barler (21) typis Buffer (22) to two minutes, as NaOH in the Duffer consource of the neuralization to finates           Monarch Plasmid Miniprep Kit         Improper storage         - Peierer elution Vibris end elution storage elemenad <tr< td=""><td>Low DNA yield</td><td></td><td>Buffers added incorrectly</td><td>Be sure that buffers have been reconstituted correctly and that reagents have been     added in the correct order</td></tr<>	Low DNA yield		Buffers added incorrectly	Be sure that buffers have been reconstituted correctly and that reagents have been     added in the correct order
Monarch DNA Gel Extraction Kit:         Bein dissibility abude 60 C         DNA         DNA         DNA         DNA           Incomplete elution during preparation         incomplete elution during or preparation         incarger elution volumes and longer incubation times can increase yield         - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend incubation times can increase yield           Monarch PCR & DNA Cleanup Kit         Buffers added incorrectly         - Be sure that buffers have been reconstituted correctly and that reagents have be added in the correct order           Monarch PCR & DNA Cleanup Kit         Incomplete elution during preparation         - Deliver Elution Buffer directly to center of column           - Law DNA quality         Monarch PCR & DNA Cleanup Kit         Incomplete elution during preparation         - Deliver Elution buffer directly to center of column           - Law DNA quality         Monarch PCR & DNA Cleanup Kit         Plasmid degradation         - Deliver Elution buffer directly to center of column           - Law DNA quality         Monarch Plasmid Miniprep Kit         Plasmid degradation         - Deliver Elution adhing investor mixing after cell tysis to avoid shearing of host cell chromosomal DNA. Do not vortex.           - Low DNA quality         Monarch Plasmid Miniprep Kit         Plasmid degradation         - Use careful investor mixing after cell tysis to avoid shearing of host cell chromosomal DNA. Do not vortex.           - Low DNA quality         Monarch Plasmid Miniprep Kit		Monarch DNA Gel Extraction Kit	Gel slice not fully dissolved	<ul> <li>Undissolved agarose may clog the column and interfere with binding. Incubate in Monarch Gel Dissolving Buffer for proper time and temperature.</li> </ul>
Image: section of the secting of the secting of the sectio			Gel dissolved above 60°C	Dissolve gel slice in specified range (37-55°C). Higher temperatures can denature DNA
Low DNA quality         Buffers added incorrectly         Be sure that buffers have been reconstituted correctly and that reagents have be added in the correct order           Low DNA quality         Incomplete elution during preparation              - Deliver Elution Numers and longer includation times can increase yield - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend includation - Elution Voltane and Plasmid UNI persons and DNA Du not vortex. - RNA contamination - Incubate sample in neutralization buffer for the full 2 minutes. For eell culture - for those sont corne in contact with flow through - Elute DNA in DNA Elution Buffer or nuclease-free water, and store at -20°C. Do not store in solutions containing magnesium. - Elution Vai i			Incomplete elution during preparation	<ul> <li>Larger elution volumes and longer incubation times can increase yield</li> <li>For elution of DNA &gt; 10 kb, heat the Elution Buffer to 50°C and extend incubation time to 5 minutes</li> </ul>
Monarch PCR & DNA Cleanup Nt (5 µg)         Monarch PCR & DNA Cleanup Nt (5 µg)         Incomplete elution during preparation         Larger elution volumes and longer incubation times can increase yield For elution of DNA > 10 kb, heat the Elution Buffer to 50°C and extend incubation ime to 5 minutes           Low DNA quality         Monarch Plasmid Miniprep Kit         Plasmid degradation         * Be cautious of strains with high levels of endogenous endonuclease (e.g., HB1 and JM 100 series)           Low DNA quality         Monarch Plasmid Miniprep Kit         Plasmid is denatured         * Limit incubation with Plasmid Lysis Buffer (B2) to two minutes, as NaOH in the buffer can denature the plasmid           Monarch Plasmid Miniprep Kit         Improper storage         * Limit incubation with Plasmid Lysis Buffer (B2) to two minutes. For cell culture volumes > 3 ml, increase the spin after meutralization to 5 minutes.           Monarch Plasmid Miniprep Kit         Ehanol has been carried over         * Centrifuge final wash for 1 minute to ensure complete removal * Ensure column tip does not come in contact with flow through           Low DNA purity         Monarch Plasmid Miniprep Kit         Guanidine carryover from Gel Dissolving Buffer         * Use smaller agarcse plugs, which will require less Gel Dissolving Buffer           Low DNA purity         Monarch Plasmid Miniprep Kit         Guanidine carried over         * Avoid strains with high amounts of endogenous carbohydrate (e.g., HB101 and JM 100 series). Be sure to follow protocol and include Plasmi Wash Buffer 1 step.			Buffers added incorrectly	• Be sure that buffers have been reconstituted correctly and that reagents have been
Low DNA quality       Monarch Plasmid Miniprep Kit       Plasmid is denatured          - Limit incubation with Plasmid Lysis Buffer (B2) to two minutes, as NaOH in the         buffer can denature the plasmid         Monarch Plasmid Miniprep Kit       gDNA contamination          - Use careful inversion mixing after cell lysis to avoid shearing of host cell         chromosomal DNA. Do not vortex.         RNA contamination          - Incubate sample in neutralization buffer for the full 2 minutes. For cell culture         volumes > 3 ml, increase the spin after neutralization to 5 minutes.         - Improper storage         Improper storage          - Elute DNA in DNA Elution Buffer or nuclease-free water, and store at -20°C. Do         not store in solutions containing magnesium.         Low DNA purity          - Monarch Plasmid Miniprep Kit         - Excessive salt in sample         - Excessive carbohydrate has been carried over         - Sensure column tip does not come in contact with flow through         - Excessive salt in sample         - Avoid strains with high amounts of endogenous carbohydrate         (e.g., HB101 and JM 100 series). Be sure to follow protocol and include Plasm         Wash Buffer 1 step.          Low DNA Gel Extraction Kit           Guanidine carried over         - Centrifuge final wash for 1 minute to ensure complete removal         - Ensure column tip does not come in contact with flow through         - Centrifuge final wash for 1 minute to ensure complete removal         - Sensure column tip does not come in contact with flow through         - Centrifuge final wash for 1 minute to ensure complete removal         - Centrifuge final wash for 1 minute to ensure complete removal         - Centrifuge fin			Incomplete elution during preparation	<ul> <li>Larger elution volumes and longer incubation times can increase yield</li> <li>For elution of DNA &gt; 10 kb, heat the Elution Buffer to 50°C and extend incubation time to 5 minutes</li> </ul>
Low DNA quality       Monarch Plasmid Miniprep Kit       gDNA contamination       * Use careful inversion mixing after cell lysis to avoid shearing of host cell chromosomal DNA. Do not vortex.         RNA contamination       * Incubate sample in neutralization buffer or the full 2 minutes. For cell culture volumes > 3 ml, increase the spin after neutralization to 5 minutes.         Improper storage       * Elute DNA in DNA Elution Buffer or nuclease-free water, and store at -20°C. Do not store in solutions containing magnesium.         Monarch Plasmid Miniprep Kit       Ethanol has been carried over       * Centrifuge final wash for 1 minute to ensure complete removal Excessive salt in sample         Low DNA purity       Monarch DNA Gel Extraction Kitt       Guanidine carryover from Gel Dissolving Buffer       * Vuse smaller agarose plugs, which will require less Gel Dissolving Buffer         Monarch DNA Gel Extraction Kitt       Ethanol has been carried over       * Use smaller agarose plugs, which will require less Gel Dissolving Buffer			Plasmid degradation	* Be cautious of strains with high levels of endogenous endonuclease (e.g., HB101 and JM 100 series)
Low DNA quarty       goina containination       chromosomal DNA. Do not vortex.         RNA contamination       incubate sample in neutralization buffer for the full 2 minutes. For cell culture volumes > 3 ml, increase the spin after neutralization to 5 minutes.         Improper storage       improper storage       eLute DNA in DNA Elution Buffer or nuclease-free water, and store at -20°C. Do not store in solutions containing magnesium.         Monarch Plasmid Miniprep Kit       Ethanol has been carried over       • Centrifuge final wash for 1 minute to ensure complete removal         Excessive salt in sample       • Use both plasmid wash buffers and do not skip wash steps         Excessive carbohydrate has been carried over       • Avoid strains with high amounts of endogenous carbohydrate (e.g., HB101 and JM 100 series). Be sure to follow protocol and include Plasmi Wash Buffer 1 step.         Low DNA purity       Guanidine carryover from Gel Dissolving Buffer       • Use smaller agarose plugs, which will require less Gel Dissolving Buffer         Monarch DNA Gel Extraction Kit       Ethanol has been carried over       • Centrifuge final wash for 1 minute to ensure complete removal         Ethanol has been carried over       • Use smaller agarose plugs, which will require less Gel Dissolving Buffer			Plasmid is denatured	<ul> <li>Limit incubation with Plasmid Lysis Buffer (B2) to two minutes, as NaOH in the buffer can denature the plasmid</li> </ul>
Low DNA purity       Monarch DNA Gel Extraction Kit       Guanidine carried over       • Centrifuge final wash for 1 minute to ensure complete removal • Ensure column tip does not come in contact with flow through         Improper storage       • Centrifuge final wash for 1 minute to ensure complete removal • Ensure column tip does not come in contact with flow through         Improper storage       • Centrifuge final wash for 1 minute to ensure complete removal • Ensure column tip does not come in contact with flow through         Improper storage       • Centrifuge final wash for 1 minute to ensure complete removal • Ensure column tip does not come in contact with flow through         Improper storage       • Centrifuge final wash for 1 minute to ensure complete removal • Ensure column tip does not come in contact with flow through	Low DNA quality	Monarch Plasmid Miniprep Kit	gDNA contamination	
Improper storage       Interproper storage       Interproper storage       Interproper storage         Improper storage       Improper storage       Interproper storage       Interproper storage       Interproper storage         Monarch Plasmid Miniprep Kit       Ethanol has been carried over       Improper storage       Improper storage       Improper storage         Excessive salt in sample       Improper storage       Improper storage       Improper storage       Improper storage         Excessive salt in sample       Improper storage       Improper storage       Improper storage       Improper storage         Excessive carbohydrate has been carried over       Improper storage       Improper storage       Improper storage         Monarch DNA Gel Extraction Kit       Guanidine carryover from Gel Dissolving Buffer       Improper storage       Improper storage         Monarch DNA Gel Extraction Kit       Ethanol has been carried over       Improper storage       Improper storage         Monarch DNA Gel Extraction Kit       Ethanol has been carried over       Improper storage       Improper storage         Monarch DNA Gel Extraction Kit       Ethanol has been carried over       Improper storage       Improper storage         Monarch DNA Gel Extraction Kit       Ethanol has been carried over       Improper storage       Improper storage         Monarch DNA Gel Extraction Kit<			RNA contamination	<ul> <li>Incubate sample in neutralization buffer for the full 2 minutes. For cell culture volumes &gt; 3 ml, increase the spin after neutralization to 5 minutes.</li> </ul>
Low DNA purity       Monarch DNA Gel Extraction Kit       Ethanol has been carried over       • Ensure column tip does not come in contact with flow through         Low DNA purity       Monarch DNA Gel Extraction Kit       Guanidine carryover from Gel Dissolving Buffer       • Use smaller agarose plugs, which will require less Gel Dissolving Buffer         Monarch DNA Gel Extraction Kit       Ethanol has been carried over       • Centrifuge final wash for 1 minute to ensure complete removal			Improper storage	<ul> <li>Elute DNA in DNA Elution Buffer or nuclease-free water, and store at -20°C. Do not store in solutions containing magnesium.</li> </ul>
Low DNA purity       Excessive safe in sample <ul> <li>Cose both plasmid wash bulkes and do not skip wash skips</li> <li>Avoid strains with high amounts of endogenous carbohydrate</li> <li>(e.g., HB101 and JM 100 series). Be sure to follow protocol and include Plasm Wash Buffer 1 step.</li> </ul> Monarch DNA Gel Extraction Kit <ul> <li>Guanidine carried over</li> <li>Centrifuge final wash for 1 minute to ensure complete removal</li> <li>Ensure column tip does not come in contact with flow through</li> </ul>			Ethanol has been carried over	
Low DNA purity         Excessive carbohydrate has been carried over         (e.g., HB101 and JM 100 series). Be sure to follow protocol and include Plasm Wash Buffer 1 step.           Nonarch DNA Gel Extraction Kit         Guanidine carryover from Gel Dissolving Buffer         • Use smaller agarose plugs, which will require less Gel Dissolving Buffer           • Centrifuge final wash for 1 minute to ensure complete removal         • Centrifuge final wash for 1 minute to ensure complete removal		Monarch Plasmid Miniprep Kit	Excessive salt in sample	Use both plasmid wash buffers and do not skip wash steps
Low DNA purity         Guanidine carryover from Gel Dissolving Buffer         • Use smaller agarose plugs, which will require less Gel Dissolving Buffer           Monarch DNA Gel Extraction Kit         Ethanol has been carried over         • Centrifuge final wash for 1 minute to ensure complete removal • Ensure column tip does not come in contact with flow through	Low DNA purity		Excessive carbohydrate has been carried over	(e.g., HB101 and JM 100 series). Be sure to follow protocol and include Plasmid
Monarch DNA Gel Extraction Kit Ethanol has been carried over • Centrifuge final wash for 1 minute to ensure complete removal • Ensure column tip does not come in contact with flow through			Guanidine carryover from Gel Dissolving Buffer	· · ·
	. ,	Monarch DNA Gel Extraction Kit	Ethanol has been carried over	
			Trace amounts of salts have been carried over	
Monarch PCR & DNA Cleanup Kit Ethanol has been carried over * Centrifuge final wash for 1 minute to ensure complete removal • Ensure column tip does not come in contact with flow through				* Centrifuge final wash for 1 minute to ensure complete removal
(5 µg) Trace amounts of salts have been carried over  • Ensure column tip does not come in contact with new tube			Trace amounts of salts have been carried over	

### Ordering Information

PRODUCTS FOR GENOMIC DNA EXTRACTION & PURIFICATION	VWR CAT. NO.	SIZE
Monarch Genomic DNA Purification Kit	76339-772	50 preps
	76339-770	150 preps
Monarch HMW DNA Extraction Kit for Cells & Blood	76470-754	5 preps
INDIDICITIENT DINA EXILACION NILTOL CENS & DIODU	76470-756	50 preps
Monarch HMW DNA Extraction Kit for Tissue	76470-750	5 preps
	76470-752	50 preps
COLUMNS, PLASTICS AND BEADS AVAILABLE SEPARATELY		
Monarch gDNA Purification Columns	76339-786	100 columns and tubes
Monarch Collection Tubes II	103529-160	100 tubes
Monarch DNA Capture Beads	76470-762	200 beads
Monarch Bead Retainers	76470-764	100 retainers
Monarch Pestle Set	76470-758	100 sets
Monarch 2 ml Tubes	76470-760	100 tubes
BUFFERS & REAGENTS AVAILABLE SEPARATELY		
Monarch gDNA Tissue Lysis Buffer	76339-774	34 ml
Monarch gDNA Cell Lysis Buffer	76339-776	20 ml
Monarch gDNA Blood Lysis Buffer	76339-778	20 ml
Monarch gDNA Binding Buffer	76339-780	65 ml
Monarch gDNA Wash Buffer	76339-782	60 ml
Monarch gDNA Elution Buffer	76339-784	34 ml
Monarch RNase A	76339-788	1 ml
Proteinase K, Molecular Biology	101076-584	2 ml
Monarch gDNA Nuclei Prep & Lysis Buffer Pack	76470-766	1 pack
Monarch RBC Lysis Buffer	76470-768	160 ml
Monarch HMW gDNA Tissue Lysis Buffer	76470-722	62 ml
Monarch Protein Separation Solution	76470-774	36 ml
Monarch Precipitation Enhancer	76470-776	10 ml
Monarch gDNA Elution Buffer II	76470-770	24 ml

PRODUCTS FOR RNA PURIFICATION	VWR CAT. NO.	SIZE
Monarch Total RNA Miniprep Kit	103529-148	50 preps
Monarch RNA Cleanup Kit (10 µg)	76307-360	10 preps
Nonaich nive cleanup kit (To µg)	76307-460	100 preps
Monarch RNA Cleanup Kit (50 µg)	76299-636	10 preps
	76299-650	100 preps
Monarch RNA Cleanup Kit (500 µg)	76299-654	10 preps
	76299-652	100 preps
COLUMNS AVAILABLE SEPARATELY		
Monarch RNA Purification Columns	103529-146	100 columns and tubes
Monarch gDNA Removal Columns	103529-158	100 columns and tubes
Monarch Collection Tubes II	103529-160	100 tubes
Monarch RNA Cleanup Columns (10 µg)	76307-362	100 columns and tubes
Monarch RNA Cleanup Columns (50 µg)	76299-642	100 columns and tubes
Monarch RNA Cleanup Columns (500 µg)	76299-656	100 columns and tubes
BUFFERS & REAGENTS AVAILABLE SEPARATELY		
Monarch DNA/RNA Protection Reagent	103529-150	56 ml
Monarch RNA Lysis Buffer	103529-152	100 ml
Monarch Total RNA Miniprep Enzyme Pack (contains DNase I, Prot K, and associated buffers)	103529-162	1 pack
Monarch RNA Priming Buffer	103529-154	56 ml
Monarch RNA Wash Buffer	103529-156	50 ml
Monarch RNA Cleanup Binding Buffer	76299-638	80 ml
Monarch RNA Cleanup Wash Buffer	76299-640	40 ml
Nuclease-free Water	103307-246	25 ml
ויענודמסטריווטל אימוטו	103307-278	100 ml

### Ordering Information (continued)

PRODUCTS FOR DNA CLEANUP & Plasmid Purification	VWR CAT. NO.	SIZE
Monarch DNA Gel Extraction Kit	102971-670	50 preps
	102971-668	250 preps
Monarch PCR & DNA Cleanup Kit (5 µg)	102971-674	50 preps
	102971-672	250 preps
Monarch Plasmid Miniprep Kit	102971-698	50 preps
	102971-696	250 preps
COLUMNS AVAILABLE SEPARATELY		
Monarch DNA Cleanup Columns (5 µg)	102971-676	100 columns and tubes
Monarch Plasmid Miniprep Columns	102971-666	100 columns and tubes
BUFFERS AVAILABLE SEPARATELY		
Monarch DNA Cleanup Binding Buffer	102971-680	235 ml
Monarch DNA Wash Buffer	102971-682	25 ml
Monarch DNA Elution Buffer	102971-688	25 ml
Monarch Gel Dissolving Buffer	102971-678	235 ml
Monarch Plasmid Lysis Buffer (B2)	102971-692	2 x 27 ml
Monarch Plasmid Neutralization Buffer (B3)	102971-694	110 ml
Monarch Plasmid Resuspension Buffer (B1)	102971-690	55 ml
Monarch Plasmid Wash Buffer 1	102971-684	2 x 27 ml
Monarch Plasmid Wash Buffer 2	102971-686	30 ml

ACCESSORIES	VWR CAT. NO.	SIZE
Monarch Microfuge Tube EcoRack	76326-060	2 racks
Monarch Collection Tubes II	103529-160	100 tubes
Monarch Pestle Set	76470-758	100 sets
Monarch Bead Retainers	76470-764	100 retainers
Monarch 2 ml Tubes	76470-760	100 tubes

#### Monarch Microfuge Tube EcoRack

The Monarch Microfuge Tube EcoRack is a bench-top tube rack made from plastic recovered during the manufacture of Monarch nucleic acid purification columns. Plastic that would otherwise be discarded during the injection molding process is recovered and re-molded into this useful lab accessory that can hold up to 48 tubes each side. One side can accommodate tubes 1.5-2 mls and the other can accommodate 0.5 ml tubes.





One or more of these products are covered by patents, trademarks and/or copyrights owned or controlled by New England Biolabs, Inc. The use of these products may require you to obtain additional third party intellectual property rights for certain applications.

Your purchase, acceptance, and/or payment of and for NEB's products is pursuant to NEB's Terms of Sale. NEB does not agree to and is not bound by ar other terms or conditions, unless those terms and conditions have been expressly agreed to in writing by a duly authorized officer of NEB.

AGILENT® and BIOANALYZER® are registered trademarks of Agilent Technologies, Inc. MISEO® and ILLUMINA® are registered trademarks of Illumina, Inc TRIZOL® is a registered trademark of Invitrogen. NANODROP® is a trademark of Thermo Fisher Scientific. OIAGEN®, DNEASY®, QIAPREP®, QIACUBE®, OIAQUICK® and MINELUTE® are registered trademarks of Diagen GmbH. TRINEN® and DROPESINS® are registered trademarks of Tinean NV. BIORAD® is a registered trademark of BioRad Laboratories. RNALATER® is a registered trademark of Ambion, Inc. GRIDION® and OXFORD NANOPORE TECHNOLOGIES® is a registered trademark of Oxford Nanopore Technologies Limited Corporation. PACBIO® is a registered trademark of Pacific Biosciences of California, Inc. CIRCULOMICS AND NANOBIND® are registered trademarks of Circulonics Inc. REVOLUGEN® is a registered trademark RevoluCen Limited Company. PROMEGA® is a registered trademark of Promega Corporation. ZYMO RESEARCH® is a registered trademark avolucen Limited Company. PROMEGA® is a registered trademark of Promega Corporation. ZYMO RESEARCH® is a registered trademark of Zymo Research Corp.

© Copyright 2020, New England Biolabs, Inc.; all rights reserved.

avantor delivered by VWP

#### VWR.COM

Prices, product, and/or services details are current when published and subject to change without notice. I Certain products or services may be limited by federal, state, provincial, or local regulations. I VWR, part of Avantor, makes no claims or warranties concerning sustainable/green products any claims concerning sustainable/green products are the sole claims of the manufacturer and not those of VWR International, LLC ant/or Avantor, Inc. or affiliates. All prices are in US dollars unless otherwise noted. Offers valid in US, vold where prohibited by law or company policy, while supplies last. I trademarks are owned by Avantor, Inc. or its affiliates, unless otherwise noted. I Visit vencom to view our privacy policy, trademark owners, and additional disclaimers. © 2021 Avantor, Inc. All rights reserved.

0421 Lit. No. 140035W