NUCLEIC ACID PURIFICATION PURE AND SIMPLE®

# **IONIC<sup>®</sup>** Purification System

Nucleic Acid Purification – Pure and Simple™

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IONIC



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## Why We Developed a New Approach to Nucleic Acid Purification

The commonly used bead- and column-based extraction technologies have followed the same fundamental workflow for over 20 years. This workflow uses ethanol, chaotropic salts, and other solutions to bind nucleic acid to a silica membrane or surface-labeled bead, which is then washed prior to the nucleic acid being stripped off the solid support into an elution buffer. During this typically laborious process, the nucleic acid is denatured, dehydrated, and fragmented. The eluate is also susceptible to contamination from wash buffers or beads.





## Isotachophoresis, a Superior Approach to Nucleic Acid Separation

Isotachophoresis (ITP) separates and concentrates charged molecules in solution solely based on their electrophoretic mobility. Biological samples are gently lysed and added to the Purigen Ionic<sup>®</sup> Fluidic Chip. An electric field is then applied to the chip and the nucleic acid is isolated in its natural, native form. The nucleic acid is not denatured or dehydrated, and there's no binding to, or stripping from, fixed surfaces. The result is a higher yield of pure nucleic acid that is less fragmented and free from bead or wash buffer contamination.



### Simple, Automated Charge-based Sample Prep in Solution

Nucleic A	Acid			
Cathodic Buffer	Separation Extraction Buffer Buffer	Anodic Buffer		
Current ap	<b>plied</b> $\rightarrow$ nucleic acid moves	and concentrates	Impurities	Nucleic Aci
>	Nucleic	)		
			1085950M	
Cathodic Buffer	Separation Extraction Buffer Buffer	Anodic Buffer		
	mple collected Nucleic	3	🔒 Impurity 🥚 Trailing Ions	🏹 Nucleic Acid 🛛 🙆 Leadin

## Ionic<sup>®</sup> Purification System

### **How it Works**

The lonic<sup>®</sup> Fluidic Chip is placed on the lonic system and separation buffers are loaded. The chip is then primed. Next, biological samples are added into the 8 sample wells and purification using isotachophoresis begins. By applying an electric field across the length of a chip microchannel, the lonic system separates and concentrates nucleic acid between buffers with higher and lower mobilities. Impurities fall behind the low mobility buffer and are separated from target nucleic acids. As target nucleic acids pass through the channel, an integrated sensor stops the current once nucleic acids reach the extraction well.



### The Next-Gen Sample Preparation System

The revolutionary Ionic Purification System requires no binding, stripping, or washing from fixed surfaces for higher yields, higher quality nucleic acids, and ultimately, better data for your research.

- No organic solvents
- No harsh, high-salt buffers
- No system programming
- No beads or repetitive washing
- No hands-on mixing, separation, sample transfers, or buffer exchanges
- No pumps, valves, or other moving parts

### **Rapid Purification of Precious Samples**

in just ONE hour



### **Simplified Nucleic Acid Preparation**

The lonic system is so different, its advantages are most readily understood in contrast with conventional nucleic acid extraction and purification methods:

- Higher nucleic acid yields
   No sample loss associated with binding nucleic acids to, or stripping from, fixed surfaces
- Simple workflows with fully automated separations No columns or beads and no repetitive washing
- Reduced nucleic acid fragmentation No harsh high-salt buffers or organic solvents

## Simplified FFPE Workflow Saves Time and Money

Purigen's FFPE protocols greatly simplify the processing of FFPE samples. For example, the lonic® FFPE to Pure DNA protocol reduces the hands-on time to less than 3 minutes per sample and enables working directly from scrolls. The protocol also eliminates the need for a separate paraffin removal step.



### **Column-based Kit Workflow**

### Flexibility for Working with Scrolls or Slides

The lonic system produces more DNA and RNA from FFPE samples without requiring the use of slides (slide use and microdissection is optional). The ability to obtain comparable nucleic acid yields when using scrolls (versus slide mounted FFPE slices) greatly simplifies the workflow when sample micro-dissection is not required. This allows projects to be completed faster and at a lower cost.



### **Superior Nucleic Acid Recoveries from FFPE Samples**

A vast majority of clinical samples used in oncology research are stored as FFPE tissues, which often contain degraded or fragmented nucleic acid. Conventional extraction methods are labor intensive and can further damage nucleic acid during the extraction and purification process. The lonic system simplifies and accelerates nucleic acid purification, resulting in higher yields of higher quality DNA.



3.5x More DNA from FFPE Samples | (80bp target)

FIGURE 1: Comparison of nucleic acid yields from replicate sections of 32 FFPE samples purified by either the lonic system or a commercially available column-based kit. The concentration of amplifiable DNA purified from each sample was determined with the Qiagen MRef Multicopy Reference Assay. For optimal performance from the columns, sections purified by this method were mounted onto slides prior to lysis. Sections purified by lonic system were processed as unmounted scrolls to demonstrate improved performance using a simpler workflow. The lonic system yield exceeds that of column-based extraction kit for 31 of 32 samples. Error bars indicate the 95% confidence interval for each data point.

Nucleic acid yields from scrolls using the lonic Purification System are on average 3.5x higher when compared to yields from slide-mounted slices using a column-based kit.



#### Higher Quality DNA vs. Column-based Kits

FIGURE 2: Samples were assessed with the QC Plex assay (Agilent) and the resulting amplicons were analyzed on the Agilent Technologies 4200 TapeStation. The resulting traces were scored using the DQC algorithm (Agilent). The results show better amplification from the samples purified with the lonic system. In addition, 14 of the samples purified by column resulted in no amplification. Amplification was observed from every sample purified by the lonic system.

### **Improved NGS Data Quality from FFPE Samples**

In addition to increased yields and a greatly simplified workflow, data quality is also improved. Data below shows purified DNA from FFPE samples analyzed using the Agilent SureMASTR Tumor Hotspot sequencing panel which includes 252 amplicons ranging in size from 128–245 bps. To highlight coverage differences related to the sample purification technology, all data was normalized to a reference sample to remove the effect of coverage differences introduced by differences in target amplicon amplification efficiencies. The reference data set was generated using the average results obtained from the purification of a high-quality sample using both lonic system and column-based techniques (as such the reference sample is not biased to either technology).



### More Uniform Coverage vs. Column-based Kits

FIGURE 3: Results shown on a chromosome level. Ionic system eluates show tighter clustering more centered around the zero line. This is indicative of more uniform sequencing coverage.

### Less Coverage Bias vs. Column-based Kits



Coverage Ratio vs. Length

Coverage Ratio vs. % GC Content

FIGURE 4: Results shown relative to amplicon length and amplicon GC content. Ionic system samples show superior uniformity for both amplicon length and GC content, with very little deviation from the expected coverage. Column-based purification shows a bias towards shorter amplicon lengths and lower GC content amplicons.

- Ionic system purification shows no bias towards amplicon length.
- lonic system purification shows no bias towards GC content.



### A Better Solution for RNA from FFPE Samples

The lonic Purification System provides for the automated purification of RNA from FFPE tissue samples with less hands-on time than conventional bead and column-based methods. To help scientists overcome the sample preparation bottleneck commonly associated with FFPE samples, the lonic system provides a simple workflow that co-purifies both mRNA and miRNA with higher yields versus column-based extraction kits.

2x Higher Yields of RNA from FFPE Samples



FIGURE 5: Replicate 10 µm sections from 17 FFPE tissue blocks were extracted and purified by both the lonic FFPE to Pure RNA Kit and a market-leading columnbased RNA extraction kit. The extracted and purified material from each kit was measured using a Qubit fluorometer and a Qubit High Sensitivity RNA assay. The average yield across the replicate samples processed by each method for each block is plotted as a bar graph.



#### **High Quality Sequencing Results**

FIGURE 6: Libraries were prepared from RNA extracted and purified from replicate samples using the lonic FFPE to Pure RNA Kit and a market-leading column-based RNA extraction kit following the AmpliSeq Illumina Immune Response Panel protocol. Libraries were then sequenced on an Illumina MiSeq sequencer. The mRNA expression results from each kit were normalized and compared by correlation analysis for each tissue type sampled. The correlation coefficient for each analysis indicates a strong correlation between the results of each kit for each tissue type sampled.

### **Get Higher miRNA Yields from FFPE Samples**

For researchers studying either the relationship between gene expression and microRNA expression or focusing purely on microRNA expression in FFPE tissue samples, the lonic system provides more miRNA than the market-leading column-based miRNA kit. More impressive, is that no additional steps are required. The lonic FFPE to Pure RNA kit produces both mRNA and more miRNA from FFPE samples in a single, simple workflow.



### **Total RNA Purification with Higher Yields of miRNA**

FIGURE 7: RNA from replicate sections of 8 FFPE sample blocks were purified using either the Ionic FFPE to Pure RNA Kit or a column-based miRNA extraction kit. The extracted and purified samples from each kit were analyzed by qPCR and the Applied Biosystems TaqMan Advanced miRNA Assays for miR-16 and miR-21. The concentration of the target miRNA represented in each sample was extrapolated and plotted against the tissue type of the source FFPE sample block. The Ionic system produced samples with a higher concentration of miRNA in all but one of the samples tested. For several samples the column-based miRNA extraction kit did not yield a detectable amount of miRNA.



#### Reproducible miRNA Expression Profiles

FIGURE 8: Samples from "Colon 1" of FIGURE 9 were analyzed for miRNA expression using the NanoString nCounter Human miRNA panel. The level of miRNA expression between replicate samples purified using the lonic system has a Pearson correlation of 0.98. The level of miRNA expression between replicate samples purified using the lonic system and the column-based miRNA kit has a Pearson correlation of 0.95. This analysis indicates a high reproducibility of miRNA expression across replicate samples purified using the lonic system that is comparable to that of the column-based miRNA extraction kit.



## Simultaneous Extraction of RNA and DNA from FFPE

The lonic® FFPE Complete Purification Kit is used with the lonic system to enable the automated purification of DNA and RNA, including microRNA from FFPE tissue samples. The kit provides a protocol, lonic® Fluidic Chips and reagents to enable the lonic system to automate DNA and RNA purification using an innovative isotachophoresis technology. Samples are prepared for purification on the lonic system using a simple lysis procedure that can be automated using a programmable thermomixer without any need for micro-dissection or de-paraffinization using harsh chemicals.

	IONIC	Manual Bead-based	Manual Column-based	
Lysis time	<b>1.5</b> hrs	Overnight	<b>1</b> hr	
RNA isolation	<b>2</b> hrs	<b>2</b> hrs	<b>2.5</b> hrs	
Lysis time	<b>1.7</b> hrs	<b>3</b> hrs	<b>3.5</b> hrs	
Total time	<b>5.2</b> hrs	<b>13</b> hrs	<b>6.5</b> hrs	
Total hands-on time	<b>1.5</b> hrs	6 hrs	<b>7</b> hrs	

### Comparison of Total Hands-on Time vs. Manual Methods

**TABLE 1:** In a study conducted by a third-party genomic services lab, this table shows 3 extraction methods that were used to compare the hands-on time and total time to extract and purify RNA and DNA from 8 samples. Replicate 10 µm sections of FFPE samples were extracted and purified using either the lonic system, a market-leading manual bead-based kit, or a manual column-based kit.

Six adjacent sections of a 10 µm thickness were harvested from 6 FFPE tissue blocks containing brain, breast, colon, or lung tissue. DNA and RNA were extracted and purified from 4 of the 6 sections using the published workflow for the lonic FFPE Complete Purification Kit. DNA and RNA were extracted and purified from the remaining sections using the published workflow for either a market-leading manual column-based kit or a market-leading manual bead-based kit.

The average estimated time to process 8 samples through the Ionic FFPE Complete Purification kit was 5 hours and 12 minutes with a hands-on time of 1 hour and 30 minutes (**TABLE 1**). This results in 11.25 minutes of hands-on time per sample to extract both DNA and RNA. The estimated time to process 8 samples through the column-based kit was 7 hours with most of that time being hands-on. This results in a hands-on time of 52.5 minutes per sample. Using a similar calculation, the hands-on time for the manual bead-based approach was 45 minutes per sample.

## 1.2x Improvement to RNA Yield with Comparable DNA Yield

The simplified workflow of the Ionic FFPE Complete Purification Kit provides simultaneous extraction and purification of FFPE samples without compromising yield.



FIGURE 9: Replicate 10 µm sections from 14 FFPE tissue blocks were extracted and purified by both the Ionic FFPE Complete Purification Kit and a market-leading column-based DNA and RNA extraction kit. The extracted and purified material from each kit was measured using a Qubit fluorometer with the Qubit RNA High Sensitivity assay. In comparison to the column-based kit, the average yield improvement across the sample set for RNA purified using the Ionic system was 1.2x.

### Improved DNA Yield with Optional Secondary Incubation



**FIGURE 10:** Replicate 10 µm sections from FFPE tissue blocks were extracted and purified by both the lonic FFPE Complete Purification Kit and a market-leading column-based DNA and RNA extraction kit. For the lonic kit, the incubation time prior to loading sample on the lonic system were varied as follows: 1) 2-hour incubation and 2) 9-hour (Overnight) incubation. The extracted and purified DNA from each kit was measured by qPCR using the Qiagen Multi-copy Reference Assay. In comparison to the column-based kit, the average yield improvement across the sample set for DNA purified using the lonic system was 2.7x with overnight incubation and 1.2x without.



### A Better Solution for DNA Purification from Tissue Samples

The lonic® Tissue to Pure DNA Kit provides automated purification of DNA from fresh frozen tissue samples with less hands-on time than conventional bead and column-based methods.



### Comparison of Ionic Tissue to Pure DNA Kit to 3 Column-based Methods

FIGURE 11: DNA extraction was performed on 8 different tissue types with four different extraction methods on a total of 632 specimens. Each tissue type/condition has an n-value ranging from 6 to 47 individual specimens. Pancreas, brain, breast, and heart specimens were derived from one donor and kidney, liver, lung, spleen were from two donors. Error Bars represent the 95% CI on the mean DNA yield (µg) per mg of tissue from replicate specimens (1-10 mg).



### **Linearity of Yield Across Tissue Input Amounts**

**FIGURE 12:** DNA yields were normalized to the lonic kit yield for 1 mg on a by-tissue basis. The dotted line represents a theoretical linear extraction efficiency up to 10 mg tissue (assuming optimal recovery by lonic kit at 1 mg specimens). The lonic kit tracked the theoretical linear extraction efficiency more closely than the column-based methods.

## Maximize Yields For WBCs, PBMCs, and Cultured/Sorted Cells

The Ionic<sup>®</sup> Cells to Pure DNA Kit supports a wide range of cell types including white blood cells (WBC) or peripheral blood mononuclear cells (PBMC) isolated from blood as well as cultured or sorted cells. The standard input range is 50k to 5 million cells. Customized protocols for as few as 10 cells are available upon request.



### Higher Yields for Blood-based, Cultured, and Sorted Cells

FIGURE 13: Peripheral Blood Mononuclear Cells (PBMCs) were isolated via Ficoll gradient, White Blood Cells (WBCs) were pelleted from lysed whole blood, and GM24385 cells were pelleted from culture media. Extractions were performed for each cell type at amounts ranging from 1-5 million cells then quantified via Qubit assay.



### Consistent Length Profiles with Average Length Above 20k bp

PURIGEN

# **Purigen Products**

Instrument		Configuration	ı	Part No.
	Ionic <sup>®</sup> Purification System Includes: Power cord, installation, warranty	Standard		44001
P.H. GED	<ul> <li>Warranty Information</li> <li>12 months coverage</li> <li>Initial response within 8 business hours</li> <li>On-site response within 3 business days</li> <li>Includes parts and materials</li> <li>Parts, materials</li> <li>On-site labor</li> </ul>			
Kits		Configuration	1	Part No.
	Ionic® FFPE to Pure DNA Kit Includes: Fluidic chips, room temp reagents, -20°C reagents	6-Chip Kit	Data and	33006
parini ja n	Ionic® FFPE to Pure RNA Kit Includes: Fluidic chips, room temp reagents, -20°C reagents	6-Chip Kit		33010
	Ionic® FFPE Complete Purification Kit Includes: Fluidic chips, room temp reagents, -20°C reagents	6-Chip Kit		33015
	Ionic® Cells to Pure DNA Includes: Fluidic chips, room temp reagents, -20°C reagents	6-Chip Kit		33005
partie (s)	Ionic® Tissue to Pure DNA Kit Includes: Fluidic chips, room temp reagents, -20°C reagents	6-Chip Kit		33019
Service Contracts	Includes		Term	Part No.

Service Contracts	Includes	Term	Part No.
Purigen Comprehensive Service Plan	<ul> <li>1 planned maintenance visit</li> <li>Initial response within 8 business hours</li> <li>On-site response within 3 business days</li> <li>Minimum 90-day warranty on replacement parts</li> <li>Parts, materials</li> <li>On-site labor, per diem charges</li> <li>Phone and email support</li> <li>Remote support sessions</li> <li>Software upgrades</li> </ul>	12 months	44900
Purigen Planned Maintenance-only Plan	<ul> <li>1 planned maintenance visit</li> <li>Initial response within 12 business hours</li> <li>On-site response within 5 business days</li> <li>90-day warranty on all replacement parts</li> <li>10% discount on Parts and Materials</li> <li>10% discount on on-site labor, per diem charges</li> <li>Phone and email support</li> <li>Remote support sessions</li> <li>Software upgrades</li> </ul>	12 months	44901
Time and Materials Service	Hourly service work	N/A	44902



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