

truXTRAC™ FFPE DNA microTUBE Kit – Column Purification (25)

Adaptive Focused Acoustics™ (AFA) -based DNA extraction and purification from Formalin-Fixed, Paraffin-Embedded Tissue using columns

Product PN 520136

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INTENDED USE

The truXTRAC FFPE DNA Kit is intended for use in molecular biology applications. This product is not intended for the diagnosis, prevention, or treatment of a disease.

INTRODUCTION

The truXTRAC FFPE DNA Kit is designed for the controlled and efficient extraction of DNA from Formalin Fixed, Paraffin Embedded (FFPE) tissue samples with the Covaris Adaptive Focused Acoustics (AFA™). AFA enables the active removal of paraffin from FFPE tissue samples in aqueous buffer, allowing simultaneous tissue rehydration. Compared to traditional passive, chemical-based methods of paraffin removal, this mechanical process is not as limited by the thickness of FFPE tissue sections. The AFA process enables the use of thicker sections, which can increase DNA yield and minimize the impact of increased DNA degradation at the exposed surfaces of a section. The truXTRAC process results in high yields of high-quality DNA well suited for analytical methods such as next-generation sequencing or qPCR.

This protocol is optimized for sections up to 25 µm in thickness and cores up to 1.2 mm in diameter.

Important Notes on FFPE Samples:

The yield of DNA from FFPE tissue blocks is highly variable. Factors such as fixation time, size and thickness of the sections, the ratio of tissue to wax, the type of tissue, and the age of the FFPE block are the main causes for this variability.

The quality of DNA isolated from FFPE samples is also highly variable. During the fixation process, DNA is cross-linked to proteins and other nucleic acid molecules to varying degrees. Incomplete reversal of this crosslinking may cause the isolated DNA to perform less well in downstream applications such as PCR and qPCR. In addition, the size of DNA fragments isolated from FFPE samples is generally smaller than that of DNA isolated from fresh or frozen tissues. This is particularly evident in older FFPE sample blocks or sample blocks stored at elevated temperatures.

Note for First Time Users:

Given the highly variable yield of DNA from FFPE tissue blocks, we recommend using FFPE blocks that have been well characterized for yield and quality for initial testing of the truXTRAC FFPE kit. Ideally, samples should be extracted immediately after sectioning.

Please contact Covaris at Application Support (ApplicationSupport@covarisinc.com) if you have any questions.

REVISION HISTORY

Part Number	Revision	Date	Description of change
010178	I	03/15	Update M220 Holder requirement and M220 settings for options B and C
010178	J	05/15	Change name of Heat Block microTUBE Adapter
010178	K	07/15	Clarify sample input requirements
010178	L	05/16	Update product name, add calibration procedure, clarify preparation procedure, move option A and B to Appendix
010178	M	1/18	Change LE220 rack and water level, add ME220 settings
010178	N	8/19	Removed tissuePICK, sectionPICK and related accessories.

KIT CONTENTS

Tissue SDS Buffer	10 ml
B1 Buffer	7.5 ml
B5 Buffer	7 ml
BW Buffer	15 ml
BE Buffer	7.5 ml
PB Buffer	1.5 ml
Proteinase K	30 mg
Purification Columns	25
Collection Tubes	50
microTUBE-130 AFA Fiber Pre-Slit Screw-Cap	25

SDS INFORMATION IS AVAILABLE AT <http://covarisinc.com/resources/safety-data-sheets/>

STORAGE

This kit should be stored at room temperature (18 – 25 °C) upon receipt.

After resuspension, Proteinase K should be stored at -20 °C.

SUPPLIED BY USERS

Covaris Instruments and Parts

Required parts						
Focused-ultrasonicator	LE220	E220 & E210	E220 evolution	S-Series	M220	ME220
Rack/ Holder	Rack 24 Place microTU BE Screw-Cap PN 500308	Rack 24 Place microTUBE Screw-Cap PN 500308	Rack E220e 4 Place microTUBE Screw Cap PN500432	Holder microTUBE Screw-Cap PN500339	Holder XTU PN500414 & Insert XTU PN500489 (*)	Rack 4-place microTUBE Screw-Cap PN500522
Intensifier	NA	PN500141	PN500141	NA	NA	Waveguide 4 Place PN500534
Accessories	Centrifuge and Heat Block microTUBE Adapter (PN500406)					

(*) Holder PN500358, although discontinued, can be used. This holder does not require an insert

Other supplies:

- Microcentrifuge with 11,000 x g capability
- Water bath, oven or dry block heater (e.g., Eppendorf ThermoMixer) for 1.5 or 2 mL tubes, capable of heating to 80°C.
- RNase A (DNase free) at 10 mg/ml (e.g., Thermo Scientific PN EN0531)
- Ethanol (>96%), molecular-biology grade (e.g., Thermo Scientific PN BP2818-100)
- 1.5 mL nonstick nuclease-free microfuge tubes (e.g., Life Technologies PN AM12450)

PROCEDURE WORKFLOW OVERVIEW

Three different options are possible with Covaris truXTRAC FFPE DNA Kit. The three options differ in the workflows for DNA extraction.

Option A: Shear DNA during extraction to a size suitable for next-generation sequencing library construction. Fragment size can be tuned between 200 and 400 bp.

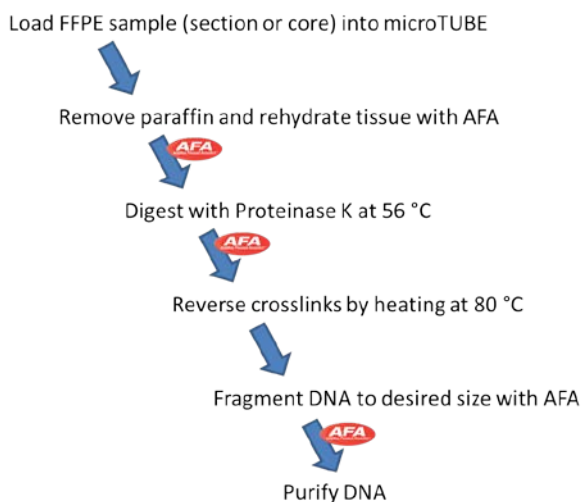
Option B: Extract ~2kb DNA fragments. This protocol is recommended for most analytical applications, including PCR. Note that actual DNA fragment size will depend of the quality of the starting material.

Option C: Extract large “genomic” DNA without any additional fragmentation. Actual DNA fragment size will depend on the quality of the starting material. For high-quality FFPE tissue blocks, we typically see an average fragment size of ≥ 8 kb.

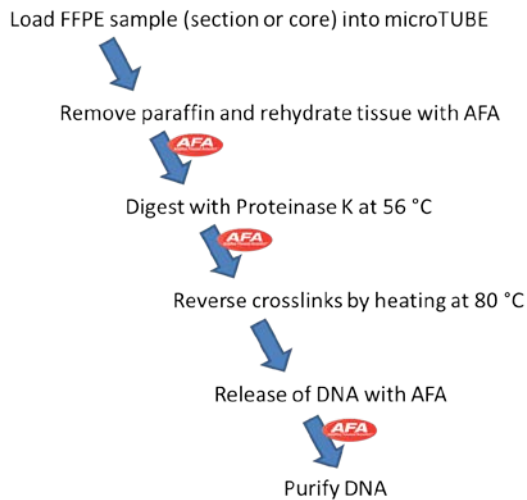
Only Option C is described in the main part of the protocol, please refer to Appendix A for Options A and B.

Please refer to Appendix B for examples of final DNA fragment size distribution.

OPTION A – EXTRACT AND FRAGMENT DNA (FOR NGS) (APPENDIX A)

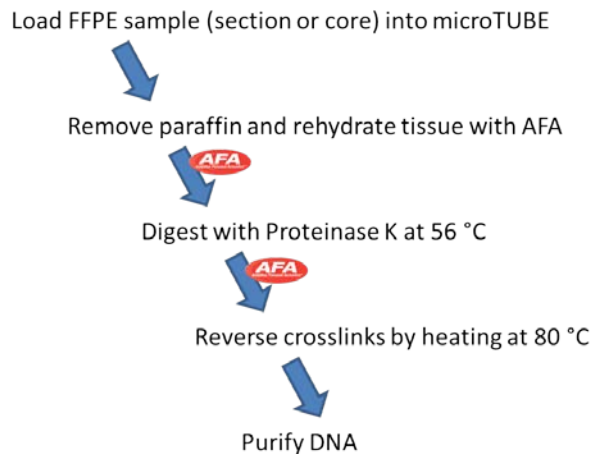


OPTION B - EXTRACT LARGE DNA FRAGMENTS (>2 KB)* WITH IMPROVED YIELD (APPENDIX A)



* Actual DNA fragment size will depend of the quality of the starting tissue block

OPTION C - EXTRACT “GENOMIC” DNA*



* Actual DNA fragment size will depend of the quality of the starting tissue block

1 - PREPARATION

FFPE Tissue Sample

1. Sample Input requirements

The truXTRAC process is highly efficient at removing paraffin even from relatively thick FFPE sections while simultaneously rehydrating the tissue. Use of thicker sections is often desirable, both for increased yield and that DNA or RNA in the exposed surfaces of a section tends to degrade quickly. **We recommend using sections between 15 and 25 μm thick, or cores of 1.2 mm.**

NOTE: Excess paraffin will adversely affect the yield and quality of DNA extracted from FFPE. We strongly advise trimming off any excess of paraffin before sectioning an FFPE tissue block, or after the section has been cut from the FFPE block. A ratio of 80% tissue to 20% paraffin or higher is ideal.

	FFPE Sections Mounted on slide		FFPE Sections "Scrolls" or "Curls"		FFPE Cores
Size (thickness or diameter)	4 to 10 μm	7 to 10 μm	7 to 15 μm	16 to 25 μm	< 1.2 mm diameter
Size (length)	NA		<10 mm (see Note A)		
Collection tool	Scalpel or razor blade	Scalpel or razor blade	NA		
Maximum number of samples Per Tube*	2 (200 mm^2 tissue for a 5 μm section)	2	2*	1*	1*

* Numbers represent trimmed sections only

NOTES

A. If the FFPE sample is longer than about 10 mm, cut it in half before loading.

WARNING: The total mass of FFPE sample processed per extraction should be between 2 to 5 mg. Lower amounts may result in insufficient yield and higher amounts may cause spin columns to become partially or fully clogged.

2. Tissue Fixation Requirements

The yield and quality of DNA extracted from FFPE tissue blocks is highly dependent on tissue collection and paraffin embedding procedures. For good yields of high quality DNA:

- Use a maximum fixation time of 24 hours
- Use Formalin solution, neutral buffered, 4%
- Fix tissue sample as quickly as possible after collection

Buffers

1. **Add ethanol to Buffer B5:** Add 28 ml of ethanol (>96%) to Buffer B5 concentrate and mark label on the cap. After preparation, Buffer B5 can then be stored for one year at room temperature.
2. **Resuspend Proteinase K:** Add 1.35 ml Buffer PB to the lyophilized Proteinase K vial. Proteinase K solution can be stored for 6 months at -20 °C. When re-suspended in the supplied Buffer PB, the solution should not freeze when stored at -20 °C.
3. **Check Buffer B1 and Tissue SDS Buffer:** A white precipitate may form during storage. Incubate the bottles at 50–70 °C before use to dissolve any precipitate.

Focused-ultrasonicator

- For S, E, or LE-Series Focused-ultrasonicators, set up the instrument as shown in Table below. Wait for the water to reach temperature and to degas.
- For the M220 Focused-ultrasonicators, put the Holder PN500414 and the Insert PN500489 (or the discontinued Holder PN500358 without insert) in place and fill the water bath until the water reaches the top of the holder. Allow system to reach temperature (20°C).
- For ME220 Focused-ultrasonicators, position the ME220 Waveguide 4 Place PN500534 into place in the water bath. Allow system to reach 20°C. Load samples into Rack 4 Place PN500522 and place into the rack holder.

For more detailed instructions on how to prepare your particular instrument please refer to your instrument's User Manual.

Focused-ultrasonicator setup

Instrument	Water level*	Chiller temp	Intensifier	Plate definition**	Holder or Rack
S-Series	15	18°C	NA	NA	PN500339
E220 & E210	10	18°C	Yes	500308 Rack 24 Place microTUBE Screw-Cap	PN500308
E220 evolution	10	18°C	Yes	500432 Rack E220e 4 Place microTUBE Screw Cap	PN500432
LE-Series	15	18°C	NA	500308 Rack 24 Place microTUBE Screw-Cap	PN500308

* Use RUN side of FILL/RUN water level label when transducer is submerged.

**If you do not see a plate definition on your system, please contact Covaris technical support at TechSupport@covarisinc.com

Heating Blocks, Water Baths, or Ovens

Preheat dry block heaters, water baths, or ovens to 56°C (or **T set₁** - see Appendix E) and 80°C (or **T set₂** - see Appendix E).

When using a dry block heater, microTUBEs need to be placed into microTUBE Adapters (PN500406) first.

It is important to confirm that the desired temperatures are actually reached. See Appendix E for instructions on how to calibrate your heating device. Subsequent steps will require 70°C.

2 – DNA EXTRACTION FROM FFPE TISSUE: DE-PARAFFINIZATION, PROTEIN DIGESTION, AND DE-CROSSLINKING

Option C - Extract genomic DNA

Option C is designed for extraction of the largest possible DNA fragments from FFPE tissue. Note that actual DNA fragment size will depend of the quality of the starting material.

1. Open microTUBE Screw-Cap, add 100 µl Tissue SDS Buffer into microTUBE, and load FFPE tissue (section or core). Affix Screw-Cap back in place.

NOTE: if the FFPE tissue samples are loose or broken the samples may be added to the microTUBE prior to Tissue SDS Buffer addition to facilitate easier loading.

2. Process the sample using the settings provided in Table 1 to dissociate the paraffin and rehydrate the tissue. During the AFA process it is normal for the solution to turn milky white as the paraffin is emulsified. (Please see the example in Appendix C.)

Table 1 - Paraffin removal and tissue rehydration settings

System	Duty Factor	Peak Incident Power	Cycles per burst	Treatment Time	Temperature (Instrument)
S220 or E220	10%	175 Watts	200	300 sec	20 °C
S2 or E210	10%	5 (Intensity)	200	300 sec	20 °C
M220	20%	75 Watts	200	300 sec	20 °C
ME220	25%	75 Watts	1000	390 sec	20 °C
LE220	30%	450 Watts ⁽¹⁾	200	300 sec	20 °C

⁽¹⁾ As Covaris LE220 process multiple samples at a time, its PIP is distributed across microTUBES, and power received by individual microTUBE stays within the 200 W limit.

3. Open microTUBE Screw-Cap, add 20 µl of Proteinase K solution to the sample, and affix Screw-Cap back in place.

4. Process the sample using the settings provided in Table 2 to properly mix Proteinase K with the sample.

Table 2 - Proteinase K mixing settings

System	Duty Factor	Peak Incident Power	Cycles per burst	Time	Temperature (Instrument)
S220 or E220	10%	175 Watts	200	10 sec	20 °C
S2 or E210	10%	5 (Intensity)	200	10 sec	20 °C
M220	20%	75 Watts	200	10 sec	20 °C
ME220	25%	75 Watts	1000	10 sec	20 °C
LE220	30%	450 Watts ⁽¹⁾	200	10 Sec	20 °C

5. **Proteinase K digestion at 56°C** (or **T set₁** - see Appendix D): Incubate samples for 1 hour (sections $\leq 10 \mu\text{m}$) or overnight (sections $> 10 \mu\text{m}$ or cores) at 56°C for proteinase K digestion. If the digestion is incomplete after overnight incubation, add another 20 μl of Proteinase K solution, mix, and incubate for 1 more hour. **When using a dry block heater, microTUBEs need to be placed into microTUBE Adapters first.**
6. **Crosslink reversal at 80°C** (or **T set₂** - see Appendix D): Incubate samples for 1 hour at 80°C to reverse formaldehyde crosslinks. **When using a dry block heater, microTUBEs need to be placed into microTUBE Adapters first.**

NOTE: If you are using the same heating source for both the 56°C & 80°C incubations, the microTUBE should be stored at room temperature until the heating source reaches 80°C or **T set₂**

7. Proceed to Section 3 – DNA Purification

3 – DNA PURIFICATION

NOTE: Set heating source to 70°C and preheat the required volume of Buffer BE in a 1.5mL microfuge tube: (number of samples x 100 µl x 1.1)

1. Transfer the sample to a clean 1.5 ml microcentrifuge tube.
2. **Optional:** The sample can be treated with RNase A to remove RNA before DNA purification. Add 5µl of RNase A solution and incubate for 5 minutes at room temperature.
3. Add 140 µl Buffer B1 to your sample and vortex thoroughly.
4. Add 160 µl ethanol (>96%) to the sample and vortex thoroughly.
5. Centrifuge at 10,000 x g for 2 minutes at room temperature. After centrifugation, much of the paraffin will have formed a white layer, floating on top of the liquid.
6. Place a Purification Column into a provided Collection Tube.
7. While holding the sample tube at about the same angle as in the rotor, use a pipette to slowly recover the liquid layer, and transfer to the column. Transfer of a small amount of paraffin particles to the column is acceptable and will not interfere with the DNA purification.
8. Spin the assembly at 11,000 x g for 1 minute.
9. Discard the flow-through and place the Column back in the Collection Tube.
10. **1st wash:** Add 500 µl Buffer BW. Spin the assembly at 11,000 x g for 1 minute.
11. Discard the flow-through and place the Column back in the Collection Tube.
12. **2nd wash:** Add 600 µl Buffer B5. Spin the assembly at 11,000 x g for 1 minute.
13. Discard the flow-through and place the column in a new Collection Tube (provided).
14. **Dry column:** Spin the assembly at 11,000 x g for 1 minute.
15. **Elute DNA - 1st step:** Place the Purification Column into a new 1.5 ml microfuge tube (not provided) and add 50 µl pre-warmed Buffer BE (70 °C) to the center of the column. Incubate at room temperature for 3 minutes. Spin the assembly at 11,000 x g for 1 minute.
16. **Elute DNA – 2nd step:** Add a second aliquot of 50 µl pre-warmed Buffer BE. Incubate again at room temperature for 3 minutes. Spin the assembly at 11,000 x g for 1 minute.
17. **DNA is eluted in 100 µl Buffer BE (5mM Tris HCl pH 8.5).**

NOTE: To increase the concentration of the recovered DNA, perform the second elution with the eluate from the first elution: Reload the first eluate to the top of the column, incubate for 3 min at room temperature and re-spin the assembly at 11,000 x g for 1 minute collecting the sample in the same 1.5 ml microcentrifuge tube. The eluted DNA is now in 50 µl.

APPENDIX A – ALTERNATIVE PROTOCOLS FOR DNA EXTRACTION FROM FFPE

Note for ME220 Users: AFA Settings for the ME220 instrument have yet to be determined for Option A or Option B.

Option A - Extract and fragment DNA (for NGS)

This protocol allows direct fragmentation of DNA to a size suitable for Next Generation Sequencing library construction during the extraction process.

1. Open microTUBE Screw-Cap, add 100 µl Tissue SDS Buffer into microTUBE, and load FFPE tissue (section or core). Affix Screw-Cap back in place.

NOTE: if the FFPE tissue samples are loose or broken the samples may be added to the microTUBE prior to Tissue SDS Buffer addition to facilitate easier loading.

2. Process the sample using the settings provided in Table 1 to dissociate the paraffin and rehydrate the tissue. During the AFA process it is normal for the solution to turn milky white as the paraffin is emulsified. (Please see the example in Appendix C.)

Table 1 - Paraffin removal and tissue rehydration settings

System	Duty Factor	Peak Incident Power	Cycles per burst	Treatment Time	Temperature (Instrument)
S220 or E220	10%	175 Watts	200	300 sec	20 °C
S2 or E210	10%	5 (Intensity)	200	300 sec	20 °C
M220	20%	75 Watts	200	300 sec	20 °C
LE220	30%	450 Watts ⁽¹⁾	200	300 sec	20 °C

⁽¹⁾ As Covaris LE220 process multiple samples at a time, its PIP is distributed across microTUBEs, and power received by individual microTUBE stays within the 200 W limit.

3. Open microTUBE Screw-Cap, add 20 µl of Proteinase K solution to the sample, and affix Screw-Cap back in place.

4. Process the sample using the settings provided in Table 2 to properly mix Proteinase K with the sample.

Table 2 - Proteinase K mixing settings

System	Duty Factor	Peak Incident Power	Cycles per burst	Time	Temperature (Instrument)
S220 or E220	10%	175 Watts	200	10 sec	20 °C
S2 or E210	10%	5 (Intensity)	200	10 sec	20 °C
M220	20%	75 Watts	200	10 sec	20 °C
LE220	30%	450 Watts ⁽¹⁾	200	10 Sec	20 °C

5. **Proteinase K digestion at 56°C** (or T set₁ - see Appendix E): Incubate samples for 1 hour (sections <= 10 µm) or overnight (sections > 10 µm or cores) at 56°C for proteinase K digestion. If the digestion is incomplete after overnight incubation, add another 20 µl of Proteinase K solution, mix, and incubate for 1 more hour. **When using a dry block heater, microTUBEs need to be placed into microTUBE Adapters first.**
6. **Crosslink reversal at 80°C** (or T set₂ - see Appendix E): Incubate samples for 1 hour at 80°C to reverse formaldehyde crosslinks. **When using a dry block heater, microTUBEs need to be placed into microTUBE Adapters first.**

NOTE: If you are using the same heating source for both the 56°C & 80°C incubations, the microTUBE should be stored at room temperature until the heating source reaches 80°C or T set₂

7. DNA fragments size can be tuned to the desired average fragment size by using the settings in Table 3 below.

NOTE: If the target size is not achieved then the treatment time should be adjusted.

Table 3 - DNA Shearing settings

E- and S-Series Focused-ultrasonicator			
Targeted fragment size	200 bp	300 bp	400 bp
Treatment Time	300 sec	110 sec	80 sec
PIP (S220 and E220)	175 W	175 W	175 W
Intensity (S2 and E210)	5	5	5
Duty Factor	10%	10%	10%
Cycle per Burst	200	200	200
Temperature	20 °C	20 °C	20 °C

M220 Focused-ultrasonicator			
Targeted fragment size	200 bp	300 bp	400 bp
Treatment Time	450 sec	200 sec	120 sec
PIP	75 W	75 W	75 W
Duty Factor	20%	20%	20%
Cycle per Burst	200	200	200
Temperature	20 °C	20 °C	20 °C

LE220 Focused-ultrasonicator			
Targeted fragment size	200 bp	300 bp	400 bp
Treatment Time	300 sec	150 sec	80 sec
PIP	450 W ⁽¹⁾	450 W ⁽¹⁾	450 W ⁽¹⁾
Duty Factor	30%	30%	30%
Cycle per Burst	200	200	200
Temperature	20 °C	20 °C	20 °C

8. Proceed to Section 3 – DNA Purification.

Option B - Extract large DNA fragments (>2 kb)* with improved yield

Option B enhances the release of DNA from the tissue while preserving a fragment size of >2kb. Note that actual DNA fragment size will depend of the quality of the starting material.

1. Open microTUBE Screw-Cap, add 100 µl Tissue SDS Buffer into microTUBE, and load FFPE tissue (section or core). Affix Screw-Cap back in place.

NOTE: if the FFPE tissue samples are loose or broken the samples may be added to the microTUBE prior to Tissue SDS Buffer addition to facilitate easier loading.

2. Process the sample using the settings provided in Table 4 to dissociate the paraffin and rehydrate the tissue. During the AFA process it is normal for the solution to turn milky white as the paraffin is emulsified. (Please see the example in Appendix C.)

Table 4 - Paraffin removal and tissue rehydration settings

System	Duty Factor	Peak Incident Power	Cycles per burst	Treatment Time	Temperature (Instrument)
S220 or E220	10%	175 Watts	200	300 sec	20 °C
S2 or E210	10%	5 (Intensity)	200	300 sec	20 °C
M220	20%	75 Watts	200	300 sec	20 °C
LE220	30%	450 Watts ⁽¹⁾	200	300 sec	20 °C

⁽¹⁾ As Covaris LE220 process multiple samples at a time, its PIP is distributed across microTUBES, and power received by individual microTUBE stays within the 200 W limit.

3. Open microTUBE Screw-Cap, add 20 µl of Proteinase K solution to the sample, and affix Screw-Cap back in place.
4. Process the sample using the settings provided in Table 5 to properly mix Proteinase K with the sample.

Table 5 - Proteinase K mixing settings

System	Duty Factor	Peak Incident Power	Cycles per burst	Time	Temperature (Instrument)
S220 or E220	10%	175 Watts	200	10 sec	20 °C
S2 or E210	10%	5 (Intensity)	200	10 sec	20 °C
M220	20%	75 Watts	200	10 sec	20 °C
LE220	30%	450 Watts ⁽¹⁾	200	10 Sec	20 °C

5. **Proteinase K digestion at 56°C** (or **T set₁** - see Appendix E): Incubate samples for 1 hour (sections <= 10 µm) or overnight (sections > 10 µm or cores) at 56°C for proteinase K digestion. If the digestion is incomplete after overnight incubation, add another 20 µl of Proteinase K solution, mix, and incubate for 1 more hour. **When using a dry block heater, microTUBEs need to be placed into microTUBE Adapters first.**

6. **Crosslink reversal at 80°C** (or **T set₂** - see Appendix E): Incubate samples for 1 hour at 80°C to reverse formaldehyde crosslinks. **When using a dry block heater, microTUBEs need to be placed into microTUBE Adapters first.**

NOTE: If you are using the same heating source for both the 56°C & 80°C incubations, the microTUBE should be stored at room temperature until the heating source reaches 80°C or **T set₂**

7. Process the sample using the settings in Table 6 to release the DNA with AFA.

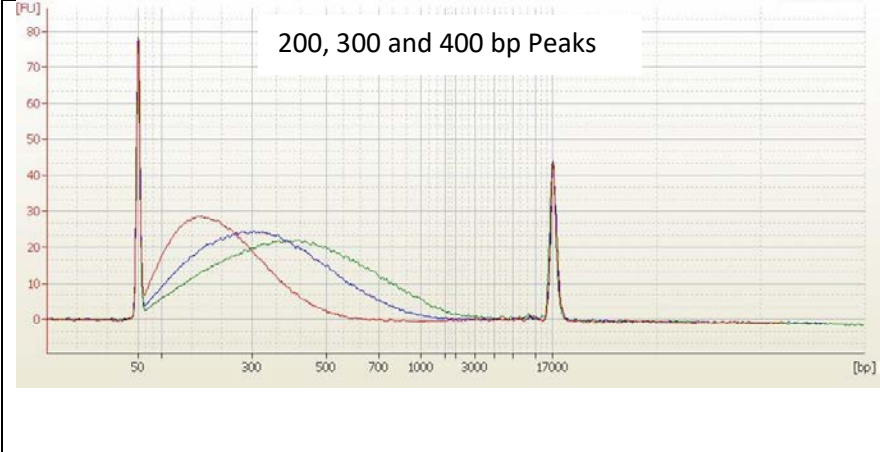
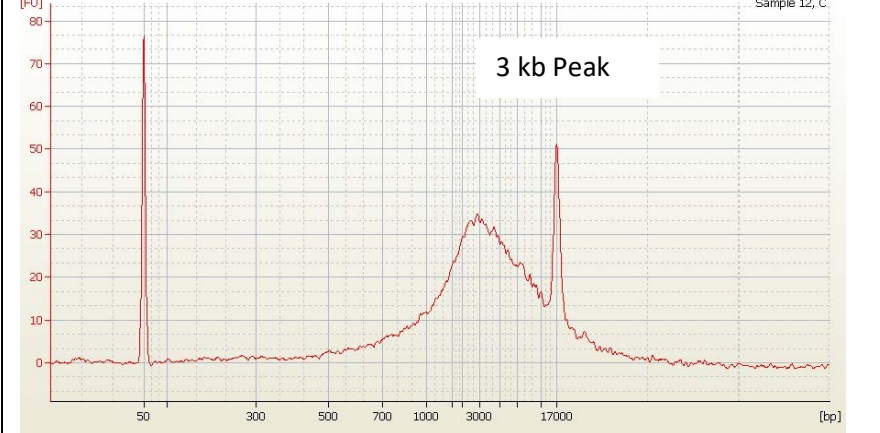
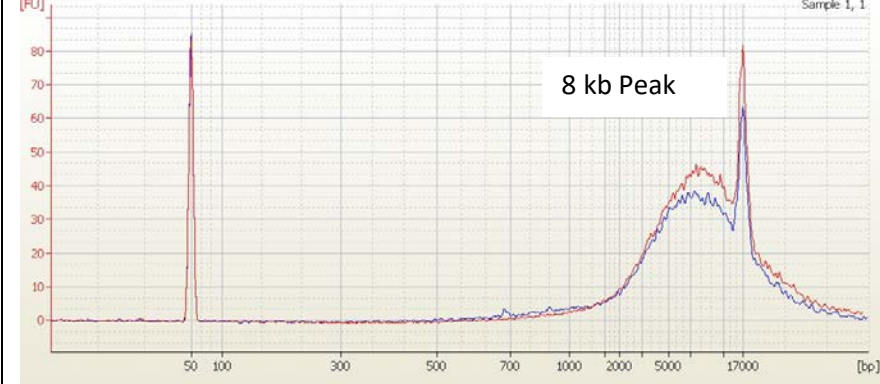
Table 6 – DNA release with AFA

System	Duty Factor	Peak Incident Power	Cycles per burst	Time	Temperature (Instrument)
S220 or E220	10%	105 Watts	200	10 sec	20 °C
S2 or E210	10%	3 (Intensity)	200	10 sec	20 °C
M220	20%	75 Watts	200	10 sec	20 °C
LE220	30%	300 Watts ⁽¹⁾	200	10 Sec	20 °C

8. Proceed to Section 3 – DNA Purification.

APPENDIX B – EXAMPLES OF DNA FRAGMENTS SIZE DISTRIBUTION

In these examples, DNA has been extracted from 10 µm sections from the same kidney tissue block with the Covaris FFPE kit following the 3 available options. The Bioanalyzer electropherograms below represent the fragment size distribution after purification. The size of the non-fragmented, genomic DNA (Option C) depends of the previous storage condition of the tissue block, including how it has been fixed and paraffin embedded.

	<p>Option A Extract and fragment DNA (for NGS)</p> <p>Subsequently to extraction, DNA is sheared to a size suitable for Next Gen Sequencing library construction. In this example, one sample has been sheared to 200 bp, one to 300 bp and one to 400 bp.</p>
	<p>Option B Extract large DNA fragments (>2 kb)*</p> <p>AFA energy is used to release the DNA from the tissue. During this process the DNA is also sheared into fragments larger than 2 kb.</p> <p>*Final size will depend of the quality of the starting tissue.</p>
	<p>Option C Extract genomic DNA*</p> <p>DNA size will be the largest possible and will depend on the quality of the starting tissue.</p> <p>*Final size will depend of the quality of the starting tissue</p>

APPENDIX C – PARAFFIN EMULSIFICATION WITH AFA ENERGY

Paraffin is emulsified in microTUBE Screw-Cap using a Covaris S220 Focused-ultrasonicator. Sample before (left side) and after (right side) processing. Sample was a 10 μm kidney tissue section.



APPENDIX D – TROUBLESHOOTING GUIDE

Issue	Cause	Solution	Comments / Suggestions
Low yield of DNA	Low tissue to wax ratio in FFPE section.	Repeat the procedure using additional sections until desired yield is achieved.	In your initial use of the truXTRAC FFPE kit, use FFPE blocks that have been well characterized for yield and quality.
	Insufficient tissue input	Increase FFPE tissue section thickness or use more sections up to 5mg total weight.	
	Proteinase K stored above recommended temperature or expired.	Repeat the procedure using fresh Proteinase K.	Always store proteinase K solution at -20°C.
No DNA	Ethanol not added to buffer B5.	Repeat the procedure with fresh samples and ensure ethanol is added to buffer B5.	
	Step 4 of option A, B or C has been omitted.	Repeat the procedure	Step 4 mixes the Proteinase K with the sample. As PB Buffer contains Glycerol, it falls at the bottom of the microTUBE and won't be in contact with the sample without this mixing step
DNA concentration is too low	Elution volume is too high.	Repeat procedure using lower elution volume (50 µl minimum volume is required). Concentrate samples using ethanol precipitation or other means.	
DNA does not perform well in downstream applications such as qPCR	DNA in FFPE sample blocks is severely cross-linked or degraded.	Design amplicons to be as small as possible (<100 bp).	DNA isolated using Covaris AFA technology is of the highest possible quality. Some FFPE sample blocks may be too degraded or cross-linked for some applications.
DNA fragments size too large when following Option A	Too much emulsified paraffin in the sample	Trim any excess paraffin from tissue blocks before proceeding with protocol. We recommend running a time course at step 7 and to increase the treatment time by 30 seconds steps.	Too much emulsified paraffin absorbs some of the acoustic energy and will adversely affect DNA Shearing efficiency.

APPENDIX E – HEATING SOURCE CALIBRATION PROCEDURE

1. If using a dry block heater, place the Covaris Heat Block microTUBE Adapters into the heating block.
2. Add water to one of the Heat Block microTUBE Adapters and insert a glass thermometer or place a glass thermometer into the water bath or the oven.
3. Set the heating source temperature to 56°C.
4. Wait for the heating source to reach the set point.
5. Check temperature displayed by the thermometer (**Tth**).
6. If **Tth** is between 55°C and 57°C (setpoint +/- 1 °C), use 56°C for **Tset₁**.
7. Otherwise, use the formula below to obtain **T set₁**:

$$\mathbf{T\ set_1\ (^{\circ}C) = 120^{\circ}C - Tth}$$

8. Repeat steps 3-7 with an initial set point of 80°C to obtain **T set₂**:

$$\mathbf{T\ set_2\ (^{\circ}C) = 160^{\circ}C - Tth}$$

Additional Notes

1. Covered by US Patent 9,080,167
2. Other patents pending
3. Best Practices for determining the yield and purity of isolated DNA:
 - To determine DNA yield with the highest level of accuracy, a fluorometric assay such as Qubit™ (Life Technologies) should be used.
 - In addition, spectrophotometric analysis of DNA for A260/280 and A260/230 ratios will determine if protein or peptide/salt contamination is present in the sample.
4. Tissue Blocks were obtained from: Theresa Kokkat, PhD and Diane McGarvey, Cooperative Human Tissue Network (CHTN), Eastern Division, University of Pennsylvania, USA
5. See following link: http://covarisinc.com/wp-content/uploads/pn_010178.pdf for updates to this document.
6. The treatment settings listed in this document are recommended guidelines. Actual results may vary depending on the tissue type, mass, and previous handling of FFPE samples.