

**Quick to assay  
protocol handbook**

for use with

**prepiT<sup>®</sup>•Q2A**

**DNAGENOTEK**



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Please visit our website at [www.dnagenotek.com](http://www.dnagenotek.com) for a full page version of each protocol and any additional languages.

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Some DNA Genotek products may not be available in all geographic regions.

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




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### Label legend:

	In vitro diagnostic medical device
	Catalog number
	Storage instructions
	Manufacturer
	Lot number

**Patent ([www.dnagenotek.com/legalnotices](http://www.dnagenotek.com/legalnotices))**

# prepIT®•Q2A product overview

## For preparation of genomic DNA from ORAcollect® and Oragene®

prepIT®•Q2A will enable a rapid, liquid-based removal of inhibitors found in saliva samples collected with ORAcollect® and Oragene® devices.

The following step-by-step protocols describes how to prepare genomic DNA for direct input into downstream applications:

## For preparation of genomic DNA from 100 µL of sample from ORAcollect (OCD-100, OCD-100A, OCR-100, OC-175) in 96-well plate format

### prepIT•Q2A reagents included

- Reagent AG (ref: PT-QAG-96 or PT-QAG-384)
- Reagent ST (ref: PT-QST-96 or PT-QST-384)

### Equipment and reagents required, not provided

- Heating block for a 96-well plate at 75°C
- 96-well PCR plate
- Dilution reagent: 10mM Tris (pH 7.5 – 8.0), nuclease-free water or similar (as required)
- Proteinase K (> 30 mAU/mg activity) (**required for OCD-100A preparation only**)

#### Proteinase K stock preparation:

Prepare a 24 mg/mL stock solution by dissolving lyophilized Proteinase K in nuclease-free water. Store in aliquots at -20°C.

1 mL stock solution is sufficient for 500 sample preparations.

- 96-well storage plate

### Warning and precautions

**Precaution:** Use Reagent ST in a well-ventilated area. Keep container closed when not in use. See MSDS at [www.dnagenotek.com](http://www.dnagenotek.com)

### Product use limitations

Use prepIT•Q2A only as directed in this product handbook. This protocol is intended to be performed by an automated liquid handler. Consult your DNA Genotek representative for a manual version of this protocol.

## Procedure

Purification steps	Notes
1. Transfer a 100 $\mu$ L aliquot of each ORAcollect sample to a 96-well plate.	
2. <b>This step applies to OCD-100A ONLY.</b> Proceed to Step 3 if preparing OCR-100, OC-175 or OCD-100 samples.  Add 2 $\mu$ L of a 24 mg/mL Proteinase K (PK) suspension. Mix by pipetting 5 $\times$ with volume set at 80 $\mu$ L.	<ul style="list-style-type: none"> <li>• See Proteinase K stock preparation instructions on page 3.</li> </ul>
3. Heat the plate at 75°C for 20 minutes.	<ul style="list-style-type: none"> <li>• <b>This heat treatment is essential.</b> Failure to adhere to these parameters will negatively impact performance on downstream assay.</li> <li>• Samples can remain unsealed during heating.</li> </ul>
4. Add 10 $\mu$ L of Reagent AG.	
5. Add 20 $\mu$ L of Reagent ST and mix thoroughly by pipetting 12 $\times$ with volume set at 100 $\mu$ L.	<ul style="list-style-type: none"> <li>• Use in a well ventilated area, keep bottle closed when not in use. Reagent has a noticeable aroma.</li> <li>• Thorough mixing is required to ensure sufficient removal of impurities from the sample.</li> </ul>
6. Incubate the samples undisturbed at room temperature for 15 minutes.	<ul style="list-style-type: none"> <li>• A phase separation will occur in this step. The upper phase contains DNA.</li> <li>• This step may alternatively be performed manually by centrifuge at 2,500 <math>\times</math> g for 2 minutes.</li> </ul>
7. Transfer 25 $\mu$ L of the upper phase to a 96-well storage plate.	<ul style="list-style-type: none"> <li>• Be careful not to disturb the bottom phase as it contains impurities.</li> <li>• DNA is fully prepared at this point.</li> </ul>

Purification steps	Notes
8. Proceed directly to assay.	<ul style="list-style-type: none"> <li>• A dilution may be required for optimal assay performance. See suggested dilution reagents in the equipment and reagents section on page 3.</li> <li>• Samples are not suitable for DNA purity assessment by spectrophotometry due to reagent interference.</li> <li>• If DNA quantification is desired, quantification should be performed by a fluorescent assay, such as with PicoGreen® or SYBR® Green I.</li> </ul>
9. Prepared DNA can be stored at 4°C for up to 1 week or at -20°C for long-term storage.	<ul style="list-style-type: none"> <li>• Ensure tube or plate are properly sealed to prevent evaporation.</li> </ul>

## For preparation of genomic DNA from 100 $\mu$ L of sample from Oragene (OGX-XXX) in 96 deepwell plate format

### prepIT•Q2A reagents included

- Reagent AG (ref: PT-QAG-96 or PT-QAG-384)
- Reagent ST (ref: PT-QST-96 or PT-QST-384)

### Equipment and reagents required, not provided

- Air or water incubator at 50°C
- 96 deepwell heating block at 75°C
- 96 deepwell plates (e.g., Abgene™ 1.2 mL round bottom, AB-0564)
- Vortexer at 1,300 rpm
- Dilution reagent: 10mM Tris (pH 7.5 – 8.0), nuclease-free water or similar (as required)
- 96-well storage plate

### Warning and precautions

**Precaution:** Use Reagent ST in a well-ventilated area. Keep container closed when not in use. See MSDS at [www.dnagenotek.com](http://www.dnagenotek.com)

### Product use limitations

Use prepIT•Q2A only as directed in this product handbook. This protocol is intended to be performed by an automated liquid handler. Consult your DNA Genotek representative for a manual version of this protocol.

### Procedure

Purification steps	Notes
1. Mix the sample by inverting the capped tube 5 $\times$ .	• This is to ensure that viscous samples are properly mixed.

Purification steps	Notes
<p>2. Incubate the sample at 50°C in a water incubator for a minimum of 1 hour or in an air incubator for a minimum of 2 hours.</p>	<ul style="list-style-type: none"> <li>• This heat-treatment step is essential to ensure that DNA is adequately released and that nucleases are permanently inactivated.</li> <li>• The entire sample must be incubated in the original collection tube before aliquoting to ensure sample homogeneity.</li> <li>• This incubation step may be performed at any time after sample is collected and before it is purified.</li> <li>• The sample may be incubated at 50°C overnight if it is more convenient.</li> <li>• A longer time is required in an air incubator because temperature equilibration is slower than in a water incubator.</li> </ul>
<p>3. Transfer a 100 µL aliquot of each Oragene sample to a 96 deepwell plate.</p>	
<p>4. Heat the 96 deepwell plate at 75°C for 10 minutes.</p>	<ul style="list-style-type: none"> <li>• <b>This heat treatment is essential.</b> Failure to adhere to these parameters will negatively impact performance on downstream assay.</li> <li>• Samples can remain unsealed during heating.</li> </ul>
<p>5. Add 10 µL of Reagent AG.</p>	

Purification steps	Notes
6. Add 20 $\mu$ L of Reagent ST and mix thoroughly by vortexing for 2 minutes at 1,300 rpm.	<ul style="list-style-type: none"> <li>• Use in a well ventilated area, keep bottle closed when not in use. Reagent has a noticeable aroma.</li> <li>• Thorough mixing is required to ensure sufficient removal of impurities from the sample.</li> <li>• Vortexing can be performed unsealed with the 96 deepwell plate, however, vortexing above 1,300 rpm is not recommended to avoid spillage across wells.</li> <li>• If sealed plates are preferred, take caution not to get reagent on the surface-edge of the plate-well. This will result in insufficient plate sealing and cross-contamination while vortexing.</li> </ul>
7. Incubate the samples undisturbed at room temperature for 15 minutes.	<ul style="list-style-type: none"> <li>• A phase separation will occur in this step. The upper phase contains DNA.</li> <li>• This step may alternatively be performed manually by centrifuge at <math>2,500 \times g</math> for 2 minutes.</li> </ul>
8. Transfer 25 $\mu$ L of the upper phase to a 96-well storage plate.	<ul style="list-style-type: none"> <li>• Be careful not to disturb the bottom phase as it contains impurities.</li> <li>• DNA is fully prepared at this point.</li> </ul>
9. Proceed directly to assay.	<ul style="list-style-type: none"> <li>• A dilution may be required for optimal assay performance. See suggested dilution reagents in the equipment and reagents section on page 6.</li> <li>• Samples are not suitable for DNA purity assessment by spectrophotometry due to reagent interference.</li> <li>• If DNA quantification is desired, quantification should be performed by a fluorescent assay, such as with PicoGreen or SYBR Green I.</li> </ul>
10. Prepared DNA can be stored at 4°C for up to 1 week or at -20°C for long-term storage.	<ul style="list-style-type: none"> <li>• Ensure plate is properly sealed to prevent evaporation.</li> </ul>

