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Storage

Upon receipt of the kit, store at −20°C until use. At this temperature the reagents are stable for 6 months.

After first use, store all of reagents at 2-8°C and keep them protected from direct light. At this condition the reagents are stable for 1 month.
Notice to Purchaser

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Introduction

Mutector™ KRAS Codon 61 Mutation Analysis Reagents are designed to detect and differentiate the following 5 mutations occurring in codon 61 of the KRAS gene.

<table>
<thead>
<tr>
<th>Codon 61 mutations</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gln 61 His Q61H (CAA &gt;CAT)</td>
</tr>
<tr>
<td>Gln 61 Leu Q61L (CAA &gt;CTA)</td>
</tr>
<tr>
<td>Gln 61 Arg Q61R (CAA &gt;CGA)</td>
</tr>
<tr>
<td>Gln 61 Glu Q61E (CAA &gt;GAA)</td>
</tr>
<tr>
<td>Gln 61 His Q61H (CAA &gt;CAC)</td>
</tr>
</tbody>
</table>

The mutation detection is performed in a single tube. Each kit provides reagents enough for 32 reactions.

The assay's products are analyzed on an Applied Biosystems Genetic Analyzer using fragment analysis software.

The kit uses Shifted Termination Assay* (STA) technology to enrich the mutation signal and is able to accurately detect low-level somatic mutations.

* Shifted Termination Assay (STA)

Shifted Termination Assay is a proprietary technology that uses uniquely designed primers, mixtures of modified enzymes and specially synthesized nucleotides. STA technology extends primers by multiple bases to increase signal strength and fragment size, creating mutation peaks that are easily distinguished from wild type. The enriched mutation signals are then detected by fragment analysis. The STA technology can detect low-level mutations often missed by sequencing.
Overview of Mutector™ Assay

- **PCR Amplification**
  - 1.5 hours*
  - *Time varies by thermal cycler used

- **PCR Product Clean-up**
  - 30 min

- **STA reaction (Mutation detection)**
  - 40 min*
  - *Time varies by thermal cycler used

- **Sample Loading**
  - To Sequencer

- **Capillary Electrophoresis**
  - Fragment analysis
  - 25-40 min*
  - *Time varies depending on the type of sequencer
**Materials Provided:**

The Mutector™ KRAS Codon 61 Mutation Detection kit contains reagents enough for 32 tests.

<table>
<thead>
<tr>
<th>Reagents</th>
<th>Quantity</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Master Mix</td>
<td>650 μl</td>
<td>Master Mix Reagents for DNA amplification</td>
</tr>
<tr>
<td>KRAS 61 PCR Primers</td>
<td>50 μl</td>
<td>PCR primer mix for amplification of KRAS gene codon 61</td>
</tr>
<tr>
<td>C-UP1</td>
<td>20 μl</td>
<td>Enzyme 1 for cleanup of PCR products</td>
</tr>
<tr>
<td>C-UP2</td>
<td>20 μl</td>
<td>Enzyme 2 for cleanup of PCR products</td>
</tr>
<tr>
<td>C-UP Buffer</td>
<td>430 μl</td>
<td>Buffer for C-UP reaction</td>
</tr>
<tr>
<td>KRAS ST-61*</td>
<td>430 μl</td>
<td>Pre-mixed STA reagents for detection of KRAS codon 61 mutations</td>
</tr>
<tr>
<td>KRAS DP-61</td>
<td>80 μl</td>
<td>Pre-mixed detection primers for KRAS codon 61 mutations</td>
</tr>
<tr>
<td>KRAS CTL-61</td>
<td>60 μl</td>
<td>Mutation controls for KRAS codon 61</td>
</tr>
<tr>
<td>Loading Buffer*</td>
<td>1000 μl</td>
<td>Sample loading buffer with size standards</td>
</tr>
</tbody>
</table>

* **Light Sensitive:** Keep these reagents protected from direct light.
**Materials required:**

0.2 ml PCR tubes (8-well strip tube)

DS-32 Matrix Standard kit (Applied Biosystems Cat. No. 4345831). This kit is a one-time calibration to set up the correct spectral channels. This is required for all Mutector II assays.

**Equipment required:**

**Thermal Cycler:** Any type of thermal cycler with a 0.2 ml tube block is acceptable for performing the assay.

**Sequencer:**

Applied Biosystems Genetic Analyzer

<table>
<thead>
<tr>
<th>Instrument</th>
<th>Data Collection</th>
<th>Data Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genetic analyzer 3100</td>
<td>Data Collection Software v 3.0 or v 3.1</td>
<td>GeneMapper® Software v 4.0 or v 4.1</td>
</tr>
<tr>
<td>Genetic analyzer 3700</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Genetic analyzer 3130</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Genetic analyzer 3500</td>
<td>3500 Data Collection Software v 1.0</td>
<td>GeneMapper® Software v 4.1</td>
</tr>
</tbody>
</table>
DNA Sample Preparation:

Reagents for DNA preparation are not provided with the kit.

Paraffin (FFPE) and fresh or frozen tissue samples

TrimGen has developed the WaxFree DNA extraction kit especially for FFPE samples. The kit uses special resins that bind and remove PCR inhibitors in the tissue extracts, leaving all DNA or RNA fragments in the extract. This method recovers more DNA in comparison with other extraction methods. The kit has been validated in many laboratories using a variety of FFPE samples as well as fresh and frozen tissue samples. WaxFree’s simple procedure and high DNA yield ensures a PCR amplification success rate of > 95%.

Product information:
WaxFree™ DNA for 50 samples (Cat. WF-50)
WaxFree™ DNA for 100 samples (Cat. WF-100)

DNA concentration:
When using a column or bead DNA extraction method, adjust the final concentration of extracted DNA to **20-80 ng /µl**

When using TrimGen’s WaxFree DNA kit, follow the user manual to perform PCR reaction.

Sequencer setup:

First time users should set up the analysis program for the ABI sequencer (one time setup). After setup, the program can apply to all Mutector™ tests for data analysis.

GeneMapper® Analysis

**Step I. GeneMapper® Setup**
www.trimgen.com/docs/PartI-GeneMapper-Setup.pdf

**Step II. Data Collection® Software Setup**
www.trimgen.com/docs/PartII-Data-Collection-Setup.pdf

**Step III. Data Analysis Using GeneMapper®**
www.trimgen.com/docs/PartIII-Data-Analysis-GeneMapper.pdf
**Important**

**Spectral calibration is required before running the test**

The sequencer needs to be calibrated with the DS-32 calibration kit (Applied Biosystems cat No. 4345831). This is a one-time calibration to set up spectral channels to collect the test results. Refer to the DS-32 Matrix standards kit to prepare the DS-32 matrix standards. Run a Matrix Standard Set DS-32 (5FAM, JOE, NED, ROX) to perform a spectral calibration.

**Thermal Cycling Programs:**

<table>
<thead>
<tr>
<th><strong>Program 1</strong> (PCR)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1 cycle</td>
<td>94°C 5 min</td>
</tr>
<tr>
<td>35 cycles</td>
<td>94°C 30 sec</td>
</tr>
<tr>
<td></td>
<td>52°C 30 sec</td>
</tr>
<tr>
<td></td>
<td>72°C 30 sec</td>
</tr>
<tr>
<td>1 cycle</td>
<td>72°C 5 min</td>
</tr>
<tr>
<td></td>
<td>Hold at 4°C</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Program 2</strong> (Clean-up)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>37°C 25 min</td>
</tr>
<tr>
<td></td>
<td>95°C 5 min</td>
</tr>
<tr>
<td></td>
<td>Hold at 4°C</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th><strong>Program 3</strong> (EM reaction)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>1 cycle</td>
<td>94°C 4 min</td>
</tr>
<tr>
<td>20 cycles</td>
<td>94°C 20 sec</td>
</tr>
<tr>
<td></td>
<td>60°C 30 sec</td>
</tr>
<tr>
<td></td>
<td>70°C 20 sec</td>
</tr>
<tr>
<td></td>
<td>Hold at 4°C</td>
</tr>
</tbody>
</table>
**Mutector™ Assay Protocol:**

**A. PCR Amplification**

Thaw all reagents and keep on ice. Spin down the reagents before use.

A negative control (water) is recommended to run with samples each time.

**A.1. Prepare PCR Reaction Mix:**

| Master Mix = 18 x ( _________ + 2*) x 1.1** = __________ μl |
|---------------|-----------------|
| # of Samples  |                 |

<table>
<thead>
<tr>
<th>KRAS 61</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCR Primers = 1 x ( _________ + <em>2) x 1.1</em>* = ________ μl</td>
</tr>
<tr>
<td># of Samples</td>
</tr>
</tbody>
</table>

* For negative and positive sample controls.

** For pipetting error.

Transfer entire volume of the reagents to one tube and gently mix (avoid bubble) the contents. This is the PCR Reaction Mix.

**A.2.** Collect 0.2 ml PCR strip tubes and label the tubes as follows:

Sample 1, 2, 3 ......

Neg: Negative Control

Pos: Positive Control

**A.3.** Transfer 19 μl of **PCR Reaction Mix** into all of the tubes.

**A.4.** Add 1 μl of nuclease-free water to the “**Neg**” tube.

**A.5.** Add 1 μl of **KRAS Codon 61 Positive Control** to the “**Pos**” tube.
A.6. Add 1-2 μl* of sample DNA (20-80 ng/μl) to each sample tube. When using TrimGen WaxFree kit for paraffin sample DNA extraction, add 0.5-1 μl* final extract to each sample tube.

*Add too much sample may cause an inhibition of PCR reaction.*

A.7. Place the PCR tubes in a thermal cycler and run Program 1.

<table>
<thead>
<tr>
<th>Program 1</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 cycle</td>
</tr>
<tr>
<td>94°C 5 min</td>
</tr>
<tr>
<td>35 cycles</td>
</tr>
<tr>
<td>94°C 30 sec</td>
</tr>
<tr>
<td>52°C 30 sec</td>
</tr>
<tr>
<td>72°C 30 sec</td>
</tr>
<tr>
<td>1 cycle</td>
</tr>
<tr>
<td>72°C 5 min</td>
</tr>
<tr>
<td>Hold at 4°C</td>
</tr>
</tbody>
</table>

Optional: The PCR products can be verified by agarose gel electrophoresis (5 μl loading). The correct band size is 120 bp.

**STOP**

The procedure can be temporarily stopped after **Program 1**. The PCR products can be stored at 4°C for 2-3 days.

*During the PCR amplification process, prepare steps B1-B2.*
B. PCR Products Clean Up

B.1. Prepare C-UP Mix:

\[
\begin{align*}
\text{C-Buffer} &= 10 \mu L \times (\text{# of PCR tubes}) \times 1.1^{**} = \text{________} \mu L \\
\text{C-UP1} &= 0.5 \mu L \times (\text{# of PCR tubes}) \times 1.1^{**} = \text{________} \mu L \\
\text{C-UP2} &= 0.5 \mu L \times (\text{# of PCR tubes}) \times 1.1^{**} = \text{________} \mu L \\
\end{align*}
\]

Mix the reagents and spin down

** For pipetting error

B.2. Collect 0.2 ml strip tubes, one tube for each PCR reaction. Label the tubes the same way as the PCR tubes.

B.3. Add 11 \(\mu l\) of C-UP Mix to each new tube.

B.4. Transfer 6 \(\mu l\) of PCR products to each tube (the remaining PCR products can be stored at \(-20^\circ C\) for re-test).

B.5. Mix the contents and spin all tubes.

B.6. Incubate the tubes in a thermal cycler using Program 2.

Program 2

37\(^{\circ}C\) for 25 min
95\(^{\circ}C\) for 5 min
Hold at 4\(^{\circ}C\)

During the clean-up incubation, prepare steps C1-C4.
C. STA Reaction (Mutation Detection)

C.1. Collect one 2 ml tube and label with “ST”. Mix the ST reagent and detection primers to make the pre-mixed ST Mix.

The pre-mixed ST Mix can be prepared using the following formula:

<table>
<thead>
<tr>
<th>Pre-mixed ST Mix</th>
</tr>
</thead>
<tbody>
<tr>
<td>KRAS ST-61 = 11 x (_________ + 1*) x 1.1** = _______ μL</td>
</tr>
<tr>
<td># of C-UP samples</td>
</tr>
<tr>
<td>KRAS DP-61 = 2 x (_________ + 1*) x 1.1** = _______ μL</td>
</tr>
<tr>
<td># of C-UP samples</td>
</tr>
</tbody>
</table>

*One extra tube for mutant controls (KRAS CTL-61)
** Adjustment for pipetting error.

Add reagents to the “ST” tube and mix gently.

C.2. Collect 0.2 mL strip tubes, one tube for each C-UP treated sample. Add an extra tube for mutant controls (KRAS CTL-61) and label the tubes as follows:

```
Sample 1, 2, 3 ....
CTL   Neg    Pos
1 2 3 4 5
```

Extra tube for mutant controls

The KRAS CTL-61 must be run each time.

C.3. Transfer 13 μl of ST Mix (from step C.1) into each tube.

C.4. Add 5 μl each of C-up treated controls and samples to their corresponding tube.

C.5. Add 2 μl of KRAS CTL-61 to the “CTL” tube.

C.6. Mix the contents and spin all tubes.

C.7. Place the tubes into a thermal cycler and perform ST reaction using Program 3.
During the STA reaction, prepare step D1-D3.

### D. Sample Loading

**D.1.** Add 15 µl of the **Loading buffer** to each well of a sequencer adapter plate.

**D.2.** Transfer 5 µl of the **ST products** into each well and remove any bubbles in the well.

**D.3.** Load the plate to sequencer and run the pre-set Data Collection Program (ref. page 8).
E. Data Analysis

E.1. Open the analysis software GeneMapper or GeneScan.

E.2. Follow the instructions to add the data for analysis.

The instructions are provided online:
GeneMapper: [www.trimgen.com/docs/PartIII-Data-Analysis-GeneMapper.pdf](http://www.trimgen.com/docs/PartIII-Data-Analysis-GeneMapper.pdf)

E.3. Confirm results of KRAS CTL-61 (mutant controls)

In the sample plot window (shows graphic data), find the results for the CTL-K61. The CTL shows 6 peaks. All peaks are located between 32-42 on the X-axis, zoom in on the X-axis to 25 (2\textsuperscript{nd} size marker) - 80 (6\textsuperscript{th} size marker). The peak size of KRAS CTL-61 is used as the standard for sample analysis.

### Result for KRAS CTL-61

<table>
<thead>
<tr>
<th>Peak #</th>
<th>Peak Color</th>
<th>Peak Size*</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Red</td>
<td>36.83</td>
<td>Mutation Q61H  (CAA &gt;CAT)</td>
</tr>
<tr>
<td>2</td>
<td>Blue</td>
<td>37.48</td>
<td>Mutation Q61R  (CAA &gt;CGA)</td>
</tr>
<tr>
<td>3</td>
<td>Black</td>
<td>37.94</td>
<td>Mutation Q61H  (CAA &gt;CAC)</td>
</tr>
<tr>
<td>4</td>
<td>Red</td>
<td>38.20</td>
<td>Mutation Q61L  (CAA &gt;CTA)</td>
</tr>
<tr>
<td>5</td>
<td>Blue</td>
<td>38.40</td>
<td>Mutation Q61E  (CAA &gt;GAA)</td>
</tr>
<tr>
<td>6</td>
<td>Black</td>
<td>39.51</td>
<td>Wild type</td>
</tr>
</tbody>
</table>

*Peak size may vary slightly depending on instrument, polymer type and the length of capillary.
E.4. Sample analysis

Zoom in on the X-axis to 25 (2nd size marker) – 80 (6th size marker). The wild type peak is observed in every sample. If the peak is not observed, it indicates that the DNA amplification failed (see troubleshooting section F.4.) or the sample is 100% mutant, such as mutant cell lines. If sample contains mutation(s), the mutation(s) will show as an additional peak(s). **Compare the peak size and color with the KRAS CTL-61** panel. The peak size may be slightly shifted due to migration differences between capillary tubes (Compare the wild type peak of the sample with the wild type peak of KRAS CTL-61 to identify the migration shift). Any peak that does not match with the mutant controls will not be considered (see trouble shooting F.6.).

**Example of assay results**

Sample: FFPE sample, one section (1 x 0.5 cm, 10 µm). DNA extraction: WaxFree DNA kit. 1 µL extract was used for assay.
F. Troubleshooting

F.1. “Color leak-through”

When the sample DNA concentration is too high, the ST reaction generates a strong fluorescent signal >5,000 rfu. Fluorescence spillover will occur. For example, the black peak of the wild type signal may be observed in the red and/or blue channels. This color spillover is caused by limitation of the instrument. The “leak-through” peak will have the exact same peak size as the original peak. Because the mutation peaks have different peak size, leak-through will not affect data analysis.

F.2. The peak signal is too high

The assay is set at a condition to detect mutations in a small sample, such as DNA extracted from fine needle aspiration (FNA) sample. For regular FFPE sample, the assay signal may be too high to analyze (peak height >8000 rfu, cannot see the top of the peak or the peak is highlighted with pink color). **Diluting the final STA product with de-ionized water can efficiently reduce the signal and optimize the peak height.** Do not dilute the assay reagents, it will cause improper enzymatic reaction and generate a miss call. Each laboratory has different PCR instrument(s), the signal intensity may vary among the laboratories, first time users should define the dilution factor (1-20 times dilution). Once the dilution factor is determined, the assay will have consistent results.

F.3. Graphic data will not automatically show

- Check the raw data. If the signals from the sample and size standards are too low, the capillary tube may be blocked by a bubble. The sample needs to be re-loaded. When adding a sample to the loading plate, carefully add the sample to avoid bubbles.
- The ST products will compete with the size standard DNA to enter the capillary tube. If the sample signal is too strong and the size standard is too low, the software cannot detect the size standard correctly and the program will not show the graphic data. **Diluting the final ST product with de-ionized water and reloading the sample will easily resolve this problem.**
- The size standard may be miscalculated. Check the size standard and manually correct the size standard (see the
sequencer’s instruction manual). Reanalyze the data after correction of the size standard.

F.4. No wild type peak

The wild type peak is an internal control for sample DNA amplification; this peak should show in all samples. If the peak is not observed, it indicates that the PCR amplification failed. The possible causes could be poor DNA quality, low DNA concentration and/or existence of PCR inhibitors in the DNA sample (see page 8 for DNA sample preparation section).

F.5. Background noise

Normally, the background of the assay is low. When the peak signal is too strong (over 8000 rfu and highlighted with pink color), background noise may pull-up as peak. To resolve this issue, simply dilute the final ST product with de-ionized water and re-load the sample.

F.6. A peak that does not match with any peak in Mutant Controls (CTL)

If such peaks is detected, please contact our tech support for further analysis. In some circumstances, when the sample DNA concentration is too low or the PCR did not amplify DNA properly - an unusual peak will appear in a very different position (most of them are far from the wild type peak). Any peaks outside of the data interpretation zone (25-80 on x-axis) are not considered for analysis.

F.7. Mutation peak cut-off

For some samples, a small peak may be observed in one of the mutation positions. To verify the peak, you need to confirm the signal strength of the wild type peak. If the wild type peak is too high (cannot see the top of the peak and the peak is highlighted with pink color), your ST reaction is too strong and the small peak may be “pull up” from background noise. Follow F.2. to dilute the final product of the ST reaction with de-ionized water. After dilution, reload the sample. If you can see the top of the wild type peak, use the following calculation to identify the small peak:

\[ \text{Ratio} = \frac{\text{Area of mutant peak}}{\text{Area of wild type peak}} \]
If the ratio is larger than 0.06, the peak is determined to be a mutation peak (the ratio does not represent the percentage of the mutation present in the sample). Otherwise, the peak is a background pull-up and does not indicate the presence of a mutation in the sample.

F.8 “Bumper peak”

For some samples, there are peaks that show as a “bumper” (see figure below). Most of these peaks are background pull-up. The causes for the bumper peaks are over loading of the ST product. Refer to F.2. in the Troubleshooting to dilute the final ST product.

![Diagram showing wild type and mutant peaks]