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# therascreen® PDGFRA RGQ PCR Kit Instructions for Use (Handbook)



Version 1



For In Vitro Diagnostic Use

For use with Rotor-Gene® Q MDx 5plex HRM instrument

For use with QIAamp® DSP DNA FFPE Tissue Kit





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## Intended Use

The QIAGEN therascreen® PDGFRA RGQ PCR Kit is a real-time qualitative in vitro diagnostic assay for the detection of the D842V somatic mutation in the *PDGFRA* gene using genomic DNA extracted from Gastrointestinal Stromal Tumour (GIST) patient's formalin-fixed, paraffinembedded (FFPE) tumour tissue.

The therascreen PDGFRA RGQ PCR Kit is intended to aid clinicians in identification of patients with GIST who may be eligible for treatment with AYVAKYT® (avaprinitib) based on a PDGFRA mutation detected result. FFPE tumour specimens are processed using the QIAamp® DSP DNA FFPE Tissue Kit for manual sample preparation and the Rotor-Gene® Q (RGQ) MDx 5plex HRM instrument for automated amplification and detection.

The therascreen PDGFRA RGQ PCR Kit is an in vitro diagnostic medical device.

## Intended User

This therascreen PDGFRA RGQ PCR Kit is intended to be used by trained personnel in a professional laboratory environment.

## Description and Principle

The therascreen PDGFRA RGQ PCR Kit comprises of one multiplex PCR amplification reaction to detect the point mutation of interest and a control region downstream of this in the same PDGFRA gene. The principal components of the kit are explained below.

#### PDGFRA reaction mix

The PDGFRA Reaction Mix contains primers and probes for both the PDGFRA D842V mutation and a control reaction.

#### **Mutation reaction**

Patient DNA is selectively amplified by qPCR using a forward and reverse primer and mutation specific labeled probe which is detected in the Texas-Red (Orange) channel of the Rotor-Gene Q MDx 5plex HRM instrument.

#### Control reaction

The Control reaction uses a forward and reverse primer and labeled probe which is detected in the Green (FAM) channel to amplify a short-conserved sequence of the *PDGFRA* gene. The Control reaction is used to determine if an appropriate level of amplifiable DNA is present in the sample and is a factor in the analytical calculations that determine mutation status.

The control target is used to both define a sample as mutant or wild type and as a DNA loading control. All samples will give a signal in the Green (FAM) channel in a predefined C<sub>T</sub> range if sufficient DNA has been added to the reaction tube.

#### Positive control

The positive control (tube PC) comprises a single plasmid representing both the mutant and control regions of the *PDGFRA* gene. Detection of both targets within acceptable ranges confirms the proper functioning of the reaction mix in the kit and overall assay setup.

#### **Negative control**

The no template control (tube NTC) contains nuclease- free water to be used for the "no template control" (NTC) reaction. The NTC serves as a negative control and assesses potential contamination during assay setup.

#### Sample diluent

The sample diluent (tube Dil.) contains nuclease-free water.

### Taq DNA Polymerase

Taq DNA polymerase is the enzyme for the polymerase chain reaction used by the therascreen PDGFRA RGQ PCR Kit.

## Summary and explanation

The *PDGFRA* gene codes for a cell surface tyrosine kinase receptor for members of the platelet-derived growth factor family. These growth factors are mitogens for cells of mesenchymal origin. The identity of the growth factor bound to a receptor monomer determines whether the functional receptor is a homodimer or a heterodimer, composed of both platelet-derived growth factor receptor alpha and beta polypeptides <sup>1-2</sup>. Studies suggest that this gene plays a role in organ development<sup>3</sup>, wound healing<sup>4</sup>, and tumour progression<sup>2,5</sup>. Mutations in this gene have been associated with idiopathic hypereosinophilic syndrome <sup>6-7</sup>, somatic and familial gastrointestinal stromal tumours<sup>8-9</sup>, and a variety of other cancers<sup>2,5</sup>.

The assay will contain the reagents necessary for the amplification and detection of the target DNA sequence using real-time qualitative PCR. It is designed to be used with the RGQ MDx 5plex HRM instrument in conjunction with Rotor-Gene AssayManager® (RGAM) software.

The assay will include a number of controls to be used as part of each assay run. A no template control (NTC) will be used to detect any contamination. A positive control plasmid containing the sequences of interest will be used to ensure the PCR run has performed within expected parameters.

The Rotor-Gene AssayManager (RGAM) MDx software generates a  $C_T$  value for the D842V and control sequence present in a sample. The  $C_T$  value difference ( $\Delta$   $C_T$ ) between the D842V  $C_T$  value and the control  $C_T$  value is used by the software to generate a mutation call ('D842V Mutation Detected' or 'No Mutation Detected' if the D842V mutation is present or absent).

## **Materials Provided**

#### Kit contents

| therascreen PDGFRA RGQ PCR<br>Kit                   |   |                        | (24)       |  |
|---|---|------------------------|------------|--|
| Catalog no.   |   |                        | 874911     |  |
| Number of reactions                                 |   |                        | 24         |  |
| Colour  | Identity                                    | Tube identification    | Volume     |  |
| Violet  | Reaction mix                                | PDGFRA Reaction<br>Mix | 720 μL     |  |
| Mint  | Taq DNA Poymerase                           | Таq                    | 1 x 85 µL  |  |
| Red   | Positive Control                            | PC                     | 40 µL      |  |
| Transparent   | Nuclease-Free water for No Template Control | NTC                    | 1 x 1.9 mL |  |
| Transparent   | Nuclease-Free water for Dilution            | Dil.                   | 1 x 1.9 mL |  |
| therascreen PDGFRA RGQ PCR Kit Instructions for Use |   |                        |            |  |

## Components of the kit

Reaction mixes are multiplex, containing fluorescently labeled reagents to detect targets and an internal control.

The principal components of the kit are explained in Table 1.

Table 1. Reagents supplied in the therascreen PDGFRA RGQ PCR Kit

| Reagent             | Active ingredients                               | Volume     |
|---------------------|--|------------|
| Reaction mix        | Control Unlabeled Primers, Control probe         | 1 x 720 μL |
|                     | labeled with Green (FAM) fluorophore,            |            |
|                     | Mutant Unlabeled Primers, Mutant Probe           |            |
|                     | labeled with Orange (Texas-Red) fluoro-          |            |
|                     | phore, Blocker Unlabeled Primer, Deoxy-          |            |
|                     | nucleotide Triphosphates, Magnesium              |            |
|                     | Chloride, Tris EDTA Buffer, PCR Buffer and       |            |
|                     | Nuclease-Free Water                              |            |
| Positive Control    | Single plasmid, Poly A RNA, Tris EDTA Buffer     | 1 x 40 µL  |
| Taq DNA Polymerase  | Taq Polymerase; 50% Glycerol/Nuclease-Free water | 1 x 85 µL  |
| No template control | Nuclease-Free Water                              | 1 x 1.9 mL |
| For sample dilution | Nuclease-Free Water                              | 1 x 1.9 mL |

### Platform and software

The therascreen PDGFRA RGQ PCR Kit is specifically designed for use with the Rotor-Gene Q MDx 5plex HRM instrument installed with Rotor-Gene AssayManager version 2.1 or later. Refer to the instrument user manual for information concerning the instrument.

The Rotor-Gene AssayManager software needs to be loaded with the therascreen\_PDGFRA\_ FFPE\_CE\_V1\_0\_1 Assay Profile for the cycle parameters specific to the *therascreen* PDGFRA RGQ PCR Kit.

The Rotor-Gene Q MDx 5plex HRM instrument must be maintained according to the requirements in the instrument user manual.

## Materials Required but not Provided

## Additional reagent

 QIAamp DSP DNA FFPE Tissue Kit (QIAGEN, cat. no. 60404; see "Procedure" on page 20)

### Consumables

- 01. mL Strip Tubes and Caps, for use with 72-well rotor (QIAGEN, cat. no. 981103 or 981106)
- · Nuclease-free, low DNA-binding microcentrifuge tubes for preparing master mixes
- Nuclease-free pipette tips with aerosol barriers

## Equipment

- Permanent marker
- Rotor-Gene Q MDx 5plex HRM instrument \* with 72-well rotor (QIAGEN, cat. no. 9002032)
- Rotor-Gene AssayManager version 2.1 or later
- therascreen\_PDGFRA\_FFPE\_CE\_V1\_0\_1
- Loading Block  $72 \times 0.1$  mL Tubes, aluminum block for manual reaction setup (QIAGEN, cat. no. 9018901)
- Dedicated pipettes (adjustable) for sample preparation

<sup>\*</sup>Prior to use, ensure that instruments have been checked and calibrated according to the manufacturer's recommendation.

- Dedicated pipettes (adjustable) for PCR master mix preparation
- Dedicated pipettes (adjustable) for dispensing of template DNA
- Benchtop centrifuge with rotor for 1.5 mL tubes
- Thermomixer, heated orbital incubator, heating block, or water bath capable of incubation at 56°C, 70°C, and 90°C

## Warnings and Precautions

For customers in the European Union, please be aware that you are required to report serious incidents that have occurred in relation to the device to the manufacturer and the competent authority of the Member State in which the user and/or the patient is established.

For in vitro diagnostic use.

For use only with Rotor-Gene Q MDx 5plex HRM instrument.

For use only with QIAamp DSP DNA FFPE Tissue Kit.

**Note**: For safety information regarding the Rotor-Gene Q MDx 5plex HRM instrument, see the user manual supplied with the instrument.

**Note**: For safety information regarding the QIAamp DSP DNA FFPE Tissue Kit (cat. no. 60404), see the QIAamp DSP DNA FFPE Tissue Kit Handbook at **www.qiagen.com**.

**Note**: The *therascreen* PDGFRA RGQ PCR Kit is to be used by trained personnel in a professional laboratory environment.

## Safety information

When working with chemicals, always wear a suitable lab coat, disposable gloves, and protective goggles. For more information, please consult the appropriate safety data sheets (SDSs). These are available online in convenient PDF format at <a href="https://www.qiagen.com/safety">www.qiagen.com/safety</a> where you can find, view, and print the SDS for each QIAGEN kit and kit component.

- All chemicals and biological materials are potentially hazardous. Specimens and samples
  are potentially infectious and must be treated as biohazardous materials.
- Discard sample and assay waste according to your local safety procedures.

#### **Precautions**

- The test is for use with formalin-fixed, paraffin-embedded GIST specimens.
- All chemicals and biological materials are potentially hazardous. Specimens and samples
  are potentially infectious and must be treated as biohazardous materials.
- Discard sample and assay waste according to your local safety procedures.
- Reagents for the therascreen PDGFRA RGQ PCR Kit are diluted optimally. Do not dilute reagents further as this may result in a loss of performance. Do not use reaction volumes (master mix plus sample) of less than 25 μL.
- All reagents supplied in the therascreen PDGFRA RGQ PCR Kit are intended to be used solely with other reagents supplied in the same therascreen PDGFRA RGQ PCR Kit. Do not substitute the reagents in the therascreen PDGFRA RGQ PCR Kit or between therascreen RGQ PCR Kits, as this may affect performance.
- Only use the Taq DNA polymerase (tube Taq) that is provided in the therascreen PDGFRA RGQ PCR Kit. Do not substitute with Taq DNA polymerase from other kits of the same or any other type or with Taq DNA polymerase from another supplier.
- Refer to the Rotor- Gene Q MDx 5plex HRM instrument user manual for additional warnings, precautions, and procedures.
- Do not use expired or incorrectly stored components.

**Note**: Use extreme caution to prevent contamination of the control and reaction mix reagents with the synthetic materials that are contained in the positive control reagent.

**Note**: Use extreme caution to prevent cross-contamination between samples. Cap the tubes promptly after addition of each sample.

**Note**: Thoroughly decontaminate the loading block before using it for preparation of assay master mixes. The use of DNAZap solutions followed by Distel and IPA wash is recommended. Loading block must be dry before use.

**Note**: Use individual, dedicated pipettes for setting up reaction mixes and adding positive control reagents.

**Note**: Perform preparation and dispensing of reaction mixes in an area separate from the one used for the addition of the positive control.

Note: Do not open the Rotor-Gene Q MDx 5plex HRM instrument until the run has finished.

**Note**: Do not open Rotor-Gene Q tubes after the run has finished. This is to prevent laboratory contamination with post-PCR products.

**Note**: Caution must be observed to ensure correct sample testing with emphasis on eliminating wrong sample entry, loading error, pipetting error, and ensuring the PCR strip tubes are placed into the appropriate positions of the 72-well rotor.

**Note**: Ensure kit reagents are thawed and mixed. Absence of sufficient mixing may affect performance.

#### Tag DNA Polymerase

Contains: Ethoxylated nonylphenol. Harmful to aquatic life with long lasting effects. Avoid release to the environment. Wear protective gloves/ protective clothing/ eye protection/ face protection. Dispose of contents/ container to an approved facility in accordance with local, regional, national, and international regulations.

### **Emergency information**

**CHEMTREC** 

Outside USA & Canada +1 703-527-3887

## Disposal

For disposal of chemicals, please consult the appropriate safety data sheets (SDSs). These are available online in convenient and compact PDF format at <a href="https://www.qiagen.com/safety">www.qiagen.com/safety</a>, where you can find, view, and print the SDS for each QIAGEN kit and kit component.

Specimens and samples are potentially infectious. Discard sample and assay waste according to your local safety procedures.

## Reagent Storage and Handling

Attention should be paid to the expiration dates and storage conditions printed on the box and labels of all components. Do not use expired or incorrectly stored components.

## Shipping conditions

The *therascreen* PDGFRA RGQ PCR Kit is shipped on dry ice and must be frozen on arrival. If any component of the *therascreen* PDGFRA RGQ PCR Kit is not frozen on arrival, the outer packaging has been opened during transit, or the shipment does not contain a packing note, handbooks, or the reagents, please contact one of the QIAGEN Technical Service Departments or local distributors (see back cover or visit **www.qiagen.com**).

## Storage conditions

The therascreen PDGFRA RGQ PCR Kit should be stored immediately upon receipt at -30 to  $-15^{\circ}$ C in a constant-temperature freezer and protected from light until the stated expiration date.

## In-use stability

Once opened, reagents can be stored in their original packaging at -30 to  $-15^{\circ}$ C for 24 months (or until the stated expiry date shown on the packaging if this is less than 24 months once the reagents have been opened for the first time). Repeated thawing and freezing should be avoided. Do not exceed a maximum of five freeze—thaw cycles.

The reagents must be thawed at room temperature  $(15-25^{\circ}C)$  for 1 hour (and up to a maximum of 4.5 hours) before use. Once the reagents are ready to use, the PCR reactions can be set up. The Rotor-Gene Q tubes, containing the master mix and the DNA sample, can be

loaded onto the Rotor-Gene Q MDx 5plex HRM immediately. The total time from start of PCR setup to the start of the run should not exceed 4.5 hours if stored at room temperature.

**Note**: Fluorescently labeled molecules included in the reaction mix reagents are light sensitive. Protect reaction mix reagents from light to avoid photo bleaching.

Reagents in the *therascreen* PDGFRA RGQ PCR Kit are diluted optimally, and no further purification or treatment is required prior to their use in analysis as directed by the *therascreen* PDGFRA RGQ PCR Kit Instructions for Use (this document).

Attention should be paid to the expiration dates and storage conditions printed on the box and labels of all components. Do not use expired or incorrectly stored components.

**Note**: Reagents required for extraction should be stored as described in the QIAamp DSP DNA FFPE Tissue Kit Handbook.

## Specimen Storage and Handling

The therascreen PDGFRA RGQ PCR Kit is for use with DNA samples extracted from GIST patient's formalin-fixed, paraffin-embedded (FFPE) tumour tissue. Tumours are heterogeneous in terms of both genotype and phenotype. Mutation-positive tumours can contain wild type DNA and similarly histology can show regions of non-tumour tissue. All tissue specimens should be treated as potentially hazardous. All samples should be treated as potentially infectious. Discard sample and assay waste according to your local safety procedures.

Store FFPE blocks and slides at room temperature.

Genomic DNA may be stored at  $2-8^{\circ}$ C for 1 week post-extraction, then at -30 to  $-15^{\circ}$ C for up to 8 weeks before use.

The workflow is described in Figure 1.

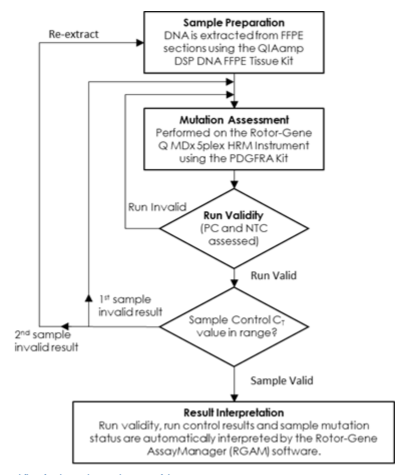


Figure 1. Workflow for the qualitative detection of the PDGFRA D842V Mutation.

Repeat PCR testing of an invalid sample should only be done once. If the required conditions are not met, samples should be re-extracted from fresh FFPE slides. In case of low slide availability, slides from a new specimen can be used.

## **Procedure**

### Protocol: DNA Extraction

### Important points before starting

- Use the QIAamp DSP DNA FFPE Tissue Kit (QIAGEN, cat. no. 60404) with the stipulations
  described below for purifying genomic DNA from samples prepared from FFPE GIST
  specimens.
- The therascreen PDGFRA RGQ PCR Kit has been developed using DNA extracted using the QIAamp DSP DNA FFPE Tissue Kit. Do not use any other DNA extraction product.
- Use dry scalpels. Do not perform this step in a laminar flow or fume hood.

### Handling reagents

- Label, handle, and store tumour specimens, blocks, slides, samples, and microcentrifuge tubes ready for extraction in a controlled fashion according to local procedures.
- Store FFPE blocks and slides at room temperature.

## Things to do before starting

 Using standard materials and methods, fix the tissue specimen in 10% neutral buffered formalin (NBF), and embed the tissue specimen in paraffin. Using a microtome, cut 5 µm serial sections from the paraffin block and mount them on glass slides.

#### Procedure

Carry out the DNA extraction according to instructions in the *QIAamp DSP DNA FFPE Tissue Kit Handbook* noting the following:

- Use a trained individual (e.g., a pathologist) to assess a Hematoxylin & Eosin (H&E)stained section to confirm that there is tumour present.
- The stained sections must not be used for DNA extraction.
- The QIAamp DSP DNA FFPE Tissue Kit must be used manually only. Do not use the automated protocol for the QIAcube® described in the QIAamp DSP DNA FFPE Tissue Kit Handbook.
- Use the number of slides and elution volumes as recommended in the FFPE tissue resections section of this handbook.
- Do not use the QIAGEN Deparaffinization Solution. Use only the xylene/ethanol method for deparaffinization described in the QIAamp DSP DNA FFPE Tissue Kit Handbook.
- Allow additional centrifugation (repeat Step 4 in QIAamp DSP DNA FFPE Tissue Kit Handbook) if tissue not pelleted after first centrifugation.
- Ensure that molecular biology grade ethanol is used for all required steps.
- Incubate open tube at 37°C for 10 minutes or until all residual ethanol has evaporated (Step 7 in QIAamp DSP DNA FFPE Tissue Kit Handbook).
- Do not use the RNase step described in the QIAamp DSP DNA FFPE Tissue Kit Handbook.

<sup>\*</sup>Do not use denatured alcohol, which contains other substances such as methanol or methylethylketone.

### Protocol: FFPE Tissue Resections

#### **Procedure**

- 1. Scrape the entire tissue area from 2 sections (5 μm) into labelled microcentrifuge tubes using a fresh scalpel for each specimen.
- 2. Proteinase K digestion must be performed for 1 hour.
- 3. For resected tissue specimens, purified genomic DNA must be eluted as a multi-step process.
  - a. Pipette 60 µL of Buffer ATE (provided in the QIAamp DSP DNA FFPE Tissue Kit) and incubate at room temperature (15–25°C) for 2.5 minutes.
  - b. Centrifuge at full speed for 1 minute.
  - c. Retaining the collection tube and flow-through, carefully open the lid of the QIAamp MinElute column again and apply a further 60  $\mu L$  Buffer ATE to the centre of the membrane.
  - d. Incubate at room temperature (15–25°C) for 2.5 minutes.
  - e. Centrifuge at full speed for 1 minute.
  - f. Discard the QIAamp MinElute column and store extracts at -30 to  $-15^{\circ}$ C as needed.

**Note**: Genomic DNA may be stored at  $2-8^{\circ}$ C for 1 week post-extraction, then at -30 to  $-15^{\circ}$ C for up to 8 weeks before use.

### Protocol: Detection of PDGFRA Mutation

## Important points before starting

- Up to 24 samples can be assessed over four runs using the Reaction Mix available in each
  kit. The optimal use is four runs with each run containing a maximum of six samples.
   Smaller sample batch sizes will mean that fewer samples can be tested with the kit.
- Do not vortex the Taq DNA polymerase (tube Taq) as this may inactivate the enzyme.
   However, once the Taq has been added to reaction mix to make the master mix, the Taq can be vortexed as part of the master mix.
- Pipette the Taq DNA polymerase by carefully placing the pipette tip just under the liquid surface to avoid the tip being coated in excess enzyme.

### Things to do before starting

- Ensure that the Rotor-Gene AssayManager version 2.1 software or later and the therascreen\_PDGFRA\_FFPE\_CE\_V1\_0\_1 Assay Profile is installed before first use of the Rotor-Gene Q MDx 5plex HRM instrument.
- Before each use, all reagents (except the Taq DNA polymerase (tube Taq)) and DNA samples need to be thawed completely for 1 hour at room temperature (15–25°C), mixed by vortexing for 3–5 seconds, and centrifuged briefly to collect the contents at the bottom of the tube.
- Ensure that the *Taq* DNA polymerase (tube *Taq*) is at room temperature (15–25°C) for 1 hour and mixed by inverting 10 times before each use. Centrifuge the tube briefly to collect the enzyme at the bottom of the tube.

#### Procedure

Completely thaw the reaction mix, nuclease-free water for no template control (tube NTC), positive control (tube PC), and DNA samples at room temperature (15–25°C) for 1 hour. Bring the Taq DNA polymerase to room temperature at the same time as other reagents. After 1 hour mix all reagents (except the Taq DNA polymerase) by vortexing each tube for 3–5 seconds to avoid localised concentration of salts. Mix the Taq DNA polymerase by inverting the tube 10 times. Centrifuge all reagents briefly to collect the contents at the bottom of the tube.

Important: Insufficient mixing of the reagents could produce invalid assay results.

- 2. Label the Lo-Bind tubes with appropriate assay contents.
- 3. Prepare sufficient master mix (reaction mix plus *Taq* DNA polymerase) for the DNA samples, 1 positive control (tube PC) reaction, and 1 nuclease-free water for no template control (tube NTC) reaction according to the volumes in Table 2.

**Note**: The master mix contains all the components needed for the PCR except the sample.

**Note**: When preparing the master mix, the required volume of reaction mix is added to the relevant tube first and the *Taq* DNA polymerase is added last.

Table 2. Preparation of assay master mix\*

| Volume of reaction mix | Volume of <i>Taq</i> DNA polymerase (tube <i>Taq</i> ) |  |  |  |
|------------------------|--|--|--|--|
| 19.8 µL × (n+2)        | 0.2 µL × (n+2)   |  |  |  |

<sup>\*</sup> n = number of samples. Two extra reactions are counted to ensure sufficient overage for the PCR setup and the controls.

4. Mix the tube containing the master mix by vortexing for 3–5 seconds followed by brief centrifugation.

5. Immediately after the master mix is ready, place the appropriate number of PCR 4-strip tubes (each strip has 4 tubes) in the loading block according to the example layout in Table 3 (adding more tubes if required, depending on the number of samples to be tested). Do not cap the tubes. Immediately add 20 µL master mix to each PCR strip tube.

**Note**: Leave the caps in the plastic bag container until required.

**Note**: Ensure that the loading block is appropriately decontaminated (see "Precautions" on page 13), and dry before use.

**Note**: Refer to Table 3 for the example run layout while setting up the reaction mixes.

Note: Change pipette tips between each template addition to avoid contamination.

Table 3. Run layout for detection of mutation

| Column 1 | Column 2 | Column 3 | Column 4 | Column 5 | Column 6 | Column 7 | Column 8 | Column 9 |
|----------|----------|----------|----------|----------|----------|----------|----------|----------|
| PC       | EMPTY    |
| NTC      | EMPTY    |
| Sample 1 | EMPTY    |
| Sample 2 | EMPTY    |
| Sample 3 | EMPTY    |
| Sample 4 | EMPTY    |
| Sample 5 | EMPTY    |
| Sample 6 | EMPTY    |

- 6. Immediately add 5  $\mu L$  of nuclease-free water for no template control (tube NTC) to the NTC tube and cap the tube.
- 7. Add 5  $\mu L$  of each DNA sample to the sample tubes and cap the tubes immediately after each sample to avoid sample to sample cross-contamination.
- 8. Add 5 µL of positive control (tube PC) to the PC tube and cap the tube.

**Important**: Run should be performed immediately after the run setup.

- 9. Using a permanent marker, mark the lids of the first tubes in the lowest numerical position in each PCR 4-strip tube to show the orientation to load the tubes into the 72-well rotor of the Rotor-Gene Q MDx 5plex HRM instrument.
- 10. Place all PCR 4-strip tubes into the appropriate positions of the 72-well rotor according to the run layout (Table 3). Take extra care to ensure that tubes will be transferred to the correct positions in the 72-well rotor (tube position in 72-well rotor should be the same as tube position in the Loading Block).

**Note**: All unused positions on the rotor must be filled with capped, empty tubes. This ensures that the thermal efficiency of the Rotor- Gene Q MDx 5plex HRM instrument is maintained.

- 11. Place the 72-well rotor into the Rotor-Gene Q MDx 5plex HRM instrument. Ensure that the locking ring (supplied with the Rotor-Gene Q MDx 5plex HRM instrument) is placed on top of the rotor to secure the tubes during the run.
- 12. The cycling parameters are shown in Table 4.

**Table 4. Cycling parameters** 

| Cycles | Temperature | Time       | Data acquisition                   |
|--------|-------------|------------|------------------------------------|
| 1      | 95°C        | 15 minutes | None                               |
| 40     | 95°C        | 30 seconds | None                               |
|        | 61°C        | 60 seconds | Green (FAM) and Orange (Texas-Red) |

13. Double-click the Rotor-Gene AssayManager version 2.1 (or later) icon desktop of the laptop connected to the RGQ instrument.



14. The Setup environment appears by default.

15. Click New Manual work list to create a new work list.

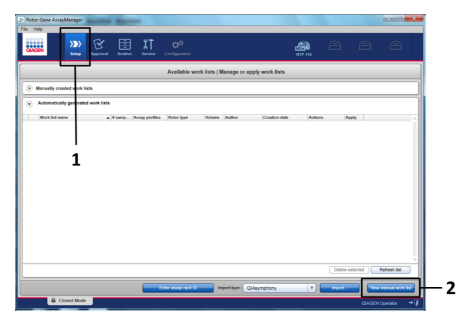


Figure 2. Setting up new manual work list. 1 – Select Setup tab, 2 – Click New manual work list.

16. Click the Assays tab (1) on the left-hand side of the main window. Click therascreen\_ PDGFRA\_FFPE\_CE\_V1\_0\_1 (2) from the Available assay profiles list. Then, click the blue arrow (3) to move the selected assay profile to the Selected assay profiles section. If the assay profile name is truncated, move the mouse pointer over the assay profile to see the full name (Figure 3).

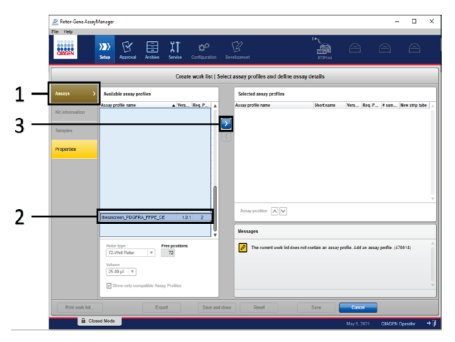


Figure 3. Setting up new manual work list – select assay profile name. 1 – Click Assays tab, 2 – Select therascreen\_ PDGFRA\_FFPE\_CE from the available assay profiles, 3 - Click the arrow.

17. In the Selected assay profiles section, enter the number of test samples to be tested excluding the number of run controls (1) (Figure 4)

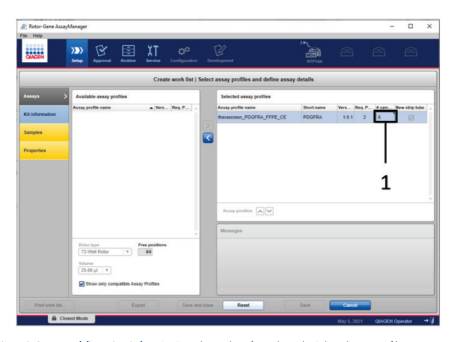
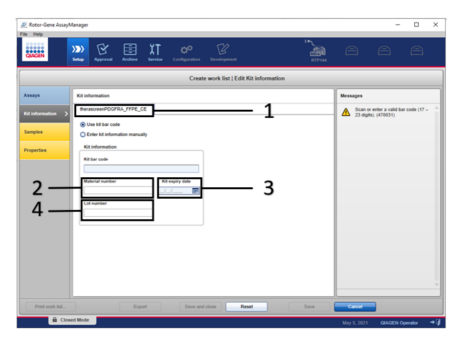


Figure 4. Create work list main window. 1 – Enter the number of samples in the Selected assay profiles section.

- 18. Click **Kit Information**. Select **Enter kit information manually** (1) and enter the following information:
  - Material number (2)
  - Kit expiry date (3)
  - Lot number (4), if available (Figure 5)



**Figure 5. Create work list main window.** In the Kit information tab, select Enter kit information manually, 1. Enter the Material number 2, Kit expiry date 3 and Lot number 4.

19. Click Samples (1) to enter sample information. Enter the sample names (2) manually.

**Important**: If using a previously generated work list, ensure that the correct sample names are entered. Sample names must be ≤40 characters in length. All sample names must be unique. Multiple entries of the same sample name will be highlighted, and the work list cannot be applied until resolved.

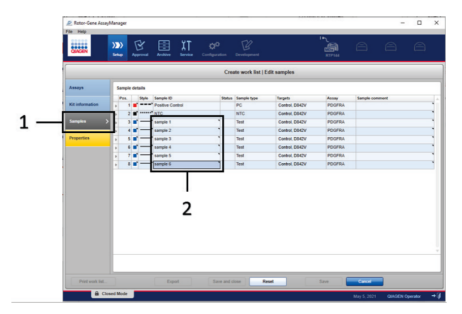


Figure 6. Create work list main window. 1 – "Samples" tab, 2 - Entering sample names. Sample names can also be entered by pasting from a notepad document.

20. Click **Properties** (1) and enter the name of the work list (2). After the work list name is entered, ensure that the **is editable** and **work list is complete** boxes are checked (3). Click **Apply**.

A new window will appear.

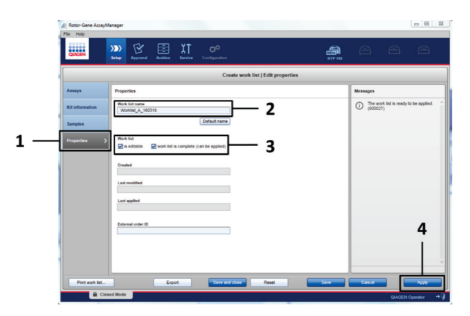


Figure 7. Create work list main window. 1 – Properties tab, 2 – Entering work list name, 3 – Select "is editable" and "work list is complete", 4 – Apply button.

21. Enter the experiment name in the **Experiment name** (1) field. Select a cycler from the Cycler selection list (only compatible cyclers will be available) (2) and ensure that the **Ring** attached box (3) is checked. After all steps have been performed, click **Start run** (4).

**Note**: A progress indicator will appear below the Rotor-Gene Q MDx 5plex HRM icon at the top of right of the screen to indicate that the run has started (Figure 8).

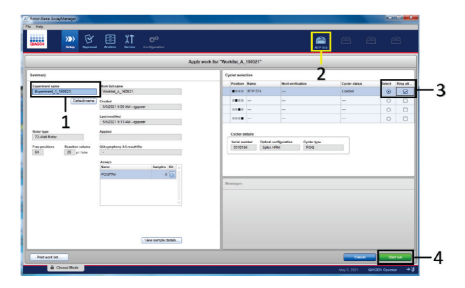


Figure 8. Applying work list and run start. 1 – Enter experiment name, 2 – Select the instrument, 3 – Ensure that Ring attached is selected, 4 – Start the run.

**Note**: The Cycler icon changes its appearance depending on the progress and result of the run. Full descriptions of these cycler icons can be found in the *Rotor-Gene AssayManager version 2.1 Core Application User Manual*.

Example cyclers are shown in Figure 9.





Figure 9. Cycler icons that may be displayed.

22. After the run is complete, click **Finish run** (Figure 10).

**Note**: During the run process, the amplification curves will be displayed and updated in real time. A progress indicator at the bottom left will show the remaining time.

**Important**: Do not close the window when the run is in progress as data loss may occur.

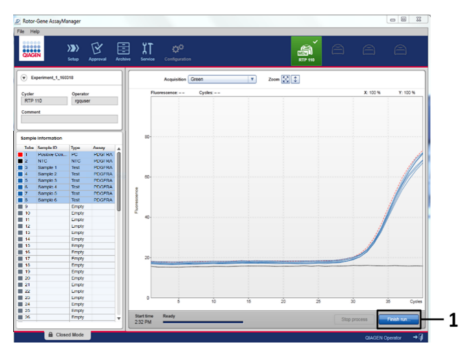


Figure 10. Finishing a run. 1 – Finish run button.

23. Click **Release and go to approval** (1) to enter the Approval tab and release the RGQ instrument. The RGQ icon at the top-right of the screen will change from green to blue indicating that the RGQ instrument is ready to perform another run (Figure 11). Regardless if a run is successful or not, the run must be released and approved.

**Note**: For a list of potential failures and error codes presented in RGAM, see the *Rotor-Gene AssayManager version 2.1 Core Application User Manual* and the *Rotor-Gene AssayManager version 2.1 Gamma Plug-in User Manual* (available from the QIAGEN website, visit **www.qiagen.com**).



Figure 11. Finish Run pop up window. 1 – Release and go to approval button.

24. Select the experiment in the Assay selection section of the Approval environment (1). Click Start approval (2) (Figure 12). If the software has been closed after the run has finished, it may be necessary to search for the experiment.

**Note**: To view a stored experiment, click the **Approval** tab (A) and search for the experiment using the search criteria in the Filter Options section (B). Click **Apply filter** (C) to search. Check the box corresponding to the experiment you want to approve (1) and click **Start approval** (2).

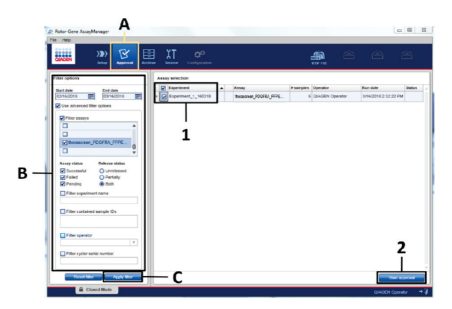


Figure 12. Starting the release process in the Approval Environment. 1 – Assay selected to approve, 2 – Start approval button, A – Approval tab, B – Filter options, C – Apply filter.

Raw Data, Processed data, Experiment information, Assay, and Audit trail information can be found in the Plots and information section (1). Assay results can be found in the Results section (Figure 13).

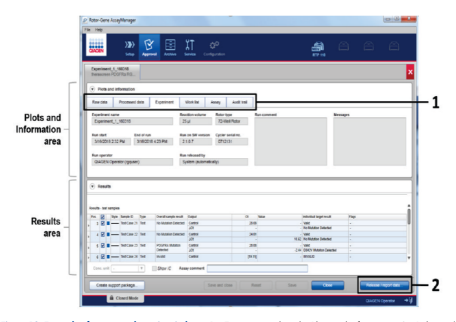


Figure 13. Example of assay results main windows. 1 – Experiment tab in the Plots and information. 2 – Release / report data button.

- If the Positive Control and No Template Control are within an acceptable range, the Sample Status column will report "Valid"; otherwise, an "Invalid" sample status will be reported.
- If either of the run controls fail, the run will be invalidated. All samples will be flagged as "ASSAY INVALID".
- See "Results" on page 42 for instructions about how to proceed.

**Note**: The Assay Profile for the *therascreen* PDGFRA RGQ PCR Kit contains all of the rules for automatic assay and sample analysis and result interpretation. The software will therefore assess the validity or invalidity of samples and controls automatically.

25. Click Release / report data. The Release / report data window will open (Figure 14).



Figure 14. Example of "Release / report data" window.

26. To save the experiment to the archive, create a LIMS output, and run report, click **OK**. The Status of release window will open (Figure 15). Run reports and LIMS exports will be saved in the default report directory. The default directory can be found in the Default data export directories in the Configuration tab.



Figure 15. Example of Status of release window.

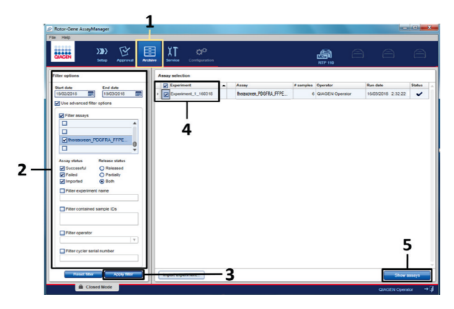
27. Click OK. The Assay is fully released dialog box will open (Figure 16).



Figure 16. Example of Assay is fully released window.

28. Click **OK**.

29. To view an experiment stored in the experiment archive, click **Archive** (1) and search for the experiment using the search criteria in the Filter Options section (2). Click **Apply filter** to search (3). Check the box next to the experiment you want to view (4) and click **Show** assays (5).



**Figure 17. Example of "Experiment Archive" main window.** 1 – Archive tab, 2 – Search options, 3 – Apply filter, 4 – Selecting experiment name, 5 – Show assays tab.

## Results

Based on predetermined analytical  $C_T$  values, the RGAM software qualitatively determines the mutation status of the DNA samples and reports which samples contain a mutation.

The run controls (PC and NTC) are assessed to ensure that acceptable  $C_T$  values are met, and the reactions have been performed successfully. If the sample Control  $C_T$  value is below the acceptable range, this means that the DNA input is too high, and the sample needs to be diluted as described in the Control  $C_T$  value section of this handbook.

All of these assessments are performed automatically and require no manual interpretation. The system automatically checks the run validity and sample validity criteria and will not report mutation status in the event of an invalid sample or invalid run.

Rotor-Gene AssayManager version 2.1 (or later) determines the result of a target by combining all relevant analysis results according to core analysis algorithms such as normalisation, and sample and assay rules defined in the corresponding assay profile.

The following results could be assigned to an individual sample:

- D842V Mutation Detected
- No Mutation Detected
- INVALID if one or more flags are assigned to the sample during analysis by Rotor-Gene AssayManager version 2.1 (or later) that are defined to set the target result to INVALID.

All possible flags corresponding to Rotor-Gene AssayManager version 2.1 Gamma Plug-in are listed *Rotor-Gene AssayManager version 2.1 Gamma Plug-in User Manual*. Additional flags which are specific to therascreen\_PDGFRA\_FFPE\_CE\_V1\_0\_1 Assay Profile are shown in Table 5.

**Note**: If an error has occurred during the run, the samples in the RGQ must be disposed of and not be retested.

Table 5. Software flags used in the PDGFRA Assay Profile

| Software flags             | Description  | Action to be taken   |
|----------------------------|--|--|
| PC_CONTROL_HIGH_CT         | PC control $C_T$ value is above the specification range              | Invalid Run – the run must be repeated   |
| PC_CONTROL_LOW_CT          | PC control C <sub>T</sub> value is below the specification range     | Invalid Run – the run must be repeated   |
| PC_CONTROL_NO_CT_ VALUE    | No $C_{\overline{1}}$ value detected for the PC in the control       | Invalid Run – the run must be repeated   |
| PC_D842V_HIGH_CT           | PC mutant C <sub>T</sub> value is above the specification range      | Invalid Run – the run must be repeated   |
| PC_D842V_LOW_CT            | PC mutant $C_T$ value is below the specification range               | Invalid Run – the run must be repeated   |
| PC_D842V_NO_CT_ VALUE      | No $C_{T}$ value detected for the PC in the mutant channel           | Invalid Run – the run must be repeated   |
| NTC_CONTROL_ UNEXPECTED_CT | A C <sub>T</sub> value has been detected in the NTC control channel  | Invalid Run – the run must be repeated   |
| NTC_D842V_UNEXPECTED_CT    | A C <sub>T</sub> value has been detected in the NTC mutant channel   | Invalid Run – the run must be repeated   |
| CONTROL_HIGH_CT            | Sample control C <sub>T</sub> value is above the specification range | Sample is invalid. Retest the sample once. If the same result is obtained after the retest then re-extract the sample and test the re-extracted sample. If the re-extracted sample is invalid for the same reason, then retest one more time. If the sample is still above specification the sample should be reported as indeterminate. |
| CONTROL_LOW_CT             | Sample control C <sub>T</sub> value is below the specification range | Sample is invalid. Sample is too concentrated and must be diluted. Follow instructions in the Control section of this handbook.  |

Table 5. Software flags used in the PDGFRA Assay Profile (continued)

| Software flags       | Description  | Action to be taken  |
|----------------------|--|---|
| CONTROL_NO_CT_VALUE  | No C <sub>T</sub> value detected for the sample in the control channel             | Sample is invalid. Retest the sample once. If the same result is obtained after the retest then re-extract the sample and test the re-extracted sample. If the re-extracted sample is invalid for the same reason, then retest one more time. If the sample still has no C <sub>T</sub> the sample should be reported as indeterminate.   |
| CURVE_SHAPE_ANOMALY  | A PCR curve anomaly has been detected in the control channel                       | If this flag is seen for a Run Control either with or without another PC and/or NTC flag; Invalid Run - the run must be repeated.  If this flag is seen for a sample; Sample is invalid. Retest the sample once. If the same result is obtained after the retest then re-extract the sample and test the re-extracted sample. If the re-extracted sample is invalid for the same reason, then retest one more time. If the sample still has no C <sub>T</sub> the sample should be reported as indeterminate. |
| OTHER_TARGET_INVALID | A PCR curve anomaly has been detected in the control channel or the mutant channel | If this flag is seen for a Run Control either with or without another PC and/or NTC flag; Invalid Run - the run must be repeated.  If this flag is seen for a sample; Sample is invalid. Retest the sample once. If the same result is obtained after the retest then re-extract the sample and test the re-extracted sample. If the re-extracted sample is invalid for the same reason, then retest one more time. If the sample still has no C <sub>T</sub> the sample should be reported as indeterminate. |

## Control C<sub>T</sub> value

Based on the  $C_T$  values obtained in the Control reaction, the system automatically checks the sample validity criteria and will not report 'D842V mutation detected' or 'mutation not detected' in the event of an invalid sample.

There are three possible flags for an invalid sample due to Control C<sub>T</sub> value:

### Control\_LOW\_CT

Sample is too concentrated and will overload the mutation assay. In order to obtain a valid sample result, the sample must be diluted. Samples should be diluted on the basis that diluting by half will increase the C<sub>T</sub> value by 1. Samples should be diluted using the water provided in the kit (Water for Dilution [Dil.]).

To calculate the required Control  $C_T$  shift  $(X_R)$  and estimate the dilution factor required (as detailed in Table 6):

$$X_R = 25 - X$$

where 25 is given as an example target Control  $C_T$  for the diluted sample and X is an actual Control  $C_T$  of sample to be diluted.

If X is not a whole number, round up to the next whole number, e.g., 2.1 is rounded up to 3.0. This value is  $X_R$ . Obtain the Dilution Factor required from Table 6.

Table 6. Calculation of dilution factor

| Xr | Dilution factor | Ratio of sample (µL) | Ratio of diluent (µL)) |
|----|-----------------|----------------------|------------------------|
| 1  | 2-fold          | 1                    | 1                      |
| 2  | 4-fold          | 1                    | 3                      |

Table 6. Calculation of dilution factor (continued)

| Xr | Dilution factor | Ratio of sample (µL) | Ratio of diluent (μL)) |
|----|-----------------|----------------------|------------------------|
| 3  | 8-fold          | 1                    | 7                      |
| 4  | 16-fold         | 1                    | 15                     |
| 5  | 32-fold         | 1                    | 31                     |
| 6  | 64-fold         | 1                    | 63                     |

### CONTROL HIGH CT

Quantity of DNA is not sufficient for mutation analysis. Retest the sample where sufficient DNA eluate is available (>5  $\mu$ L). If the quantity of DNA is still insufficient upon retest, re-extract from fresh FFPE sections. If not possible, the sample will be reported as indeterminate.

## CONTROL\_NO\_CT\_VALUE

Quantity of DNA is either not sufficient for detection or not present in the sample. Retest the sample where sufficient DNA eluate is available (>5  $\mu$ L). If the quantity of DNA is still insufficient upon retest, re-extract from fresh FFPE sections. If not possible, the sample will be reported as indeterminate.

## Contraindication

There are no known contraindications.

## Limitations

Samples with results reported as No Mutation Detected may harbour *PDGFRA* mutations, which were not detected by the *therascreen PDGFRA RGQ PCR Kit*.

Detection of mutations is dependent on sample integrity and the amount of amplifiable DNA present in the specimen. The procedure should be repeated if DNA in the sample indicates that the quantity is either not sufficient or too high for mutation analysis.

Determination of DNA input for the assay is based on DNA qualification and is performed using the Control Reaction  $C_T$  to assess the amount of amplifiable DNA. The amount of amplifiable DNA in a sample (represented by the  $C_T$  value of the Control Reaction) needs to be within the pre-defined Control Working Range (CWR) in order for the sample to proceed to mutation assessment. The acceptable measurable range of the Control assay is Control  $C_T$  20.85–30.99. The PDGFRA assay profile will automatically determine if the sample has a Control  $C_T$  within the acceptable range.

Macrodissection of FFPE specimens is not performed prior to extraction of DNA. The performance of the *therascreen PDGFRA RGQ PCR* Kit was validated with DNA extracted from FFPE GIST specimens with 2.0–99.0% tumour content.

The therascreen PDGFRA RGQ PCR Kit is used in a polymerase chain reaction (PCR) procedure. As with all PCR procedures, samples may become contaminated by external sources of DNA in the test environment and the DNA in the positive control. Use caution to avoid contamination of samples and kit reagents.

The therascreen PDGFRA RGQ PCR Kit is only for use with DNA samples extracted from formalin-fixed, paraffin-embedded GIST specimens. The therascreen PDGFRA RGQ PCR Kit is only for use with the QIAamp DSP DNA FFPE Tissue Kit (QIAGEN, cat. no. 60404) and users should follow the extraction protocol stipulations as stated in this handbook.

Only the Rotor- Gene Q MDx 5plex HRM instrument has been validated for use with the *therascreen* PDGFRA RGQ PCR Kit. No other thermal cycler with real-time optical detection can be used with this product.

There is no information if the *therascreen PDGFRA RGQ PCR* Kit shows cross-reactivity (results of D842V Mutation Detected) to additional *PDGFRA* mutations different to that included in the kit.

The therascreen PDGFRA RGQ PCR Kit is a qualitative test. The test is not for quantitative measurements of percent mutation.

Performance of the *therascreen PDGFRA RGQ PCR* Kit is unknown if microbial contamination is introduced during assay procedures.

Performance of the *therascreen PDGFRA RGQ PCR* Kit could be affected in the presence of haemoglobin and triglycerides in the test samples.

## Performance Characteristics

## Analytical performance

The specific performance characteristics of the *therascreen* PDGFRA RGQ PCR Kit were determined using formalin- fixed, paraffin- embedded (FFPE) specimens collected from Gastrointestinal Stromal Tumour (GIST) patients.

### Limit of Blank (LoB)

The Limit of Blank (LoB) is defined in CLSI guideline EP17-A2 as "the highest measurement result that is likely to be observed (with a stated probability) for a blank sample". For the therascreen PDGFRA RGQ PCR Kit, this is the data point that corresponds to the upper 95% percentile in the blank (wild type, i.e., D842V mutation negative) samples.

The LoB has been determined by measuring the levels of breakthrough for the *PDGFRA* D842V mutation reaction, where breakthrough is defined as the non-specific, low-level amplification of a DNA template. For the *therascreen* PDGFRA RGQ PCR Kit, this refers to non-specific binding of the mutant probe to the wild type DNA.

Based on this analysis, the LOB for the assay has been calculated as a  $\Delta$  C<sub>T</sub> of 9.854.

## $\Delta C_T$ cutoff value

The assay cutoff is a specific  $\Delta C_T$  value used to determine whether a sample is classed as positive or negative for the *PDGFRA* D842V mutation. Samples which generate  $\Delta C_T$  values at or below the cutoff are classified as D842V mutation positive (i.e., *PDGFRA* D842V mutation detected) and  $\Delta C_T$  values generated above the cutoff are classified as D842V mutation negative (i.e., no mutation detected). The false-negative and false-positive rates for the D842V assay were used to determine a cutoff value such that a result equal or less than the cutoff will

result in a D842V mutation detected classification. The cutoff for the *therascreen* PDGFRA RGQ PCR Kit is  $9.0~\Delta$  C<sub>T</sub>.

## Effect of DNA input on $\Delta C_T$ values (linearity)

The DNA input level is defined as the total quantity of amplifiable DNA in a sample as determined by the  $C_T$  value from the Control reaction. To demonstrate that the performance of the *therascreen* PDGFRA RGQ PCR Kit is consistent across the Control reaction working range (CWR) (20.85–30.99  $C_T$ ), a 9-level serial dilution of a D842V mutation positive sample with varying DNA input levels (with the lower levels being outside of the CWR) and 13.5% MAF (representing 1.5x LoD) was tested.

The evaluation was performed using one therascreen PDGFRA RGQ PCR Kit lot with six replicates tested per DNA input level. The data was analysed using regression analysis to determine the linear range. For the assay to be determined as linear across the DNA input range, there should be no change across the range in  $\Delta C_T$ , i.e., no statistically significant linear, quadratic, or cubic effect.

Parameter estimates for the tested dilutions were reported along with the corresponding standard errors, 95% confidence limits, and p-values, and the data showed no significant linear, quadratic, or cubic trends. A linear  $\Delta C_T$  range was observed across the DNA input range tested for the assay.

## Limit of Detection (LoD)

The LoD is defined as the lowest amount of the *PDGFRA* D842V mutation (in a background of wild type DNA) that is possible to detect 95% of the time. For the *PDGFRA* D842V assay, the LoD is reported as % mutant allele frequency (MAF).

Genomic DNA was extracted from D842V mutation positive clinical GIST FFPE specimens and was diluted into normalised DNA extracted from wild type clinical GIST FFPE specimens and serially diluted to % MAF levels above, at, and below the estimated LoD.

At least 60 replicates of each % MAF dilution in the series per DNA input level (medium and low DNA input) were tested using two *therascreen* PDGFRA RGQ PCR Kit. The LoD was determined using a Probit model and reported as the maximum observed LoD value across the two *therascreen* PDGFRA RGQ PCR Kit lots tested.

The LoD of the *therascreen* PDGFRA RGQ PCR Kit was determined at 9% MAF. The LoD was verified in a low level of background DNA and where the level of background DNA was not limiting.

## Cross-reactivity

The *therascreen* PDGFRA RGQ PCR Kit is comprised of one multiplex PCR amplification reaction to detect the D842V mutation at codon 842 in Exon 18 of the *PDGFRA* gene, and a control region downstream of this in the *PDGFRA* gene. There is no reaction that specifically measures the wild type sequence at codon 842 in Exon 18. The *therascreen* PDGFRA RGQ PCR Kit "No Mutation Detected" result is inferred by the absence of any positive mutation result.

## Analytical specificity

The level of potential cross-reactivity between primers, probes, and the blocker used within the *therascreen* PDGFRA RGQ PCR Kit and nonspecific targets within human and non-human genomes, and the level of potential heterodimer formation was investigated. An in silico analysis was performed to determine whether the primers, probes, and blocker used with the *therascreen* PDGFRA RGQ PCR Kit assays bind non-specifically within any genome, including

the human genome. An additional in silico analysis was performed to determine whether the oligonucleotides used within the multiplex assay bind non-specifically to each other.

The *in silico* analysis of oligo heterodimers indicated that there is a low prevalence of heterodimer formation. The primers and probes do not cross react with any targets not detected by the *therascreen PDGFRA RGQ PCR* Kit and therefore will not generate a false signal.

## Repeatability and reproducibility

The precision and reproducibility of the *therascreen PDGFRA RGQ PCR* Kit was investigated by testing DNA extracted from D842V mutation positive clinical GIST FFPE specimens and wild type clinical GIST FFPE specimens.

To demonstrate repeatability, D842V mutation positive samples at two mutation levels (1x LoD and 1.5x LoD) and two DNA input levels (medium ( $C_T$  26) and low ( $C_T$  29) DNA input), and wild type samples also at medium and low DNA input levels were tested at one site (located in the United Kingdom). The samples were tested in triplicate with four runs per day across multiple days, Rotor- Gene Q instruments, and operators and using one lot of *therascreen* PDGFRA RGQ PCR Kit to generate a total of 108 replicates per sample (Table 7).

Table 7. Assay repeatability\*

| Template                              | No. of correct calls | Percentage of correct calls (%) | Lower two-sided<br>95% confidence limit<br>(%) | Upper two-sided 95% confidence limit (%) |
|---------------------------------------|----------------------|---------------------------------|--|--|
| D842V 1.5x LoD<br>(C <sub>T</sub> 26) | 108/108              | 100.00                          | 96.64  | 100.00                                   |
| D842V 1.5x LoD<br>(C <sub>T</sub> 29) | 108/108              | 100.00                          | 96.64  | 100.00                                   |
| D842V 1x LoD<br>(C <sub>T</sub> 26)   | 108/108              | 100.00                          | 96.64  | 100.00                                   |
| D842V 1x LoD<br>(C <sub>T</sub> 29)   | 108/108              | 100.00                          | 96.64  | 100.00                                   |
| WT<br>(C <sub>T</sub> 26)             | 108/108              | 100.00                          | 96.64  | 100.00                                   |
| WT<br>(C <sub>T</sub> 29)             | 107/107              | 100.00                          | 96.61  | 100.00                                   |

<sup>\*</sup> Number of correct calls and two-sided 95% confidence limits for D842V mutation positive samples at 1x LoD and 1.5x LoD and wild type samples tested across one site.

To demonstrate reproducibility, two runs per day were performed per operator (two operators per site) by two additional sites (both located in the USA) over six days to give an additional 108 data points for each additional site. The same test panel was used for this study as for the repeatability study and one lot of the *therascreen PDGFRA RGQ PCR* Kit (the same lot for all three sites) was used. The proportion of correct mutation calls for each sample across all three sites was calculated and is reported in Table 8.

Table 8. Assay reproducibility\*

| Template                              | No. of correct calls | Percentage of correct calls (%) | Lower two-sided 95% confidence limit (%) | Upper two-sided 95% confidence limit (%) |
|---------------------------------------|----------------------|---------------------------------|--|--|
| D842V 1.5x LoD<br>(C <sub>T</sub> 26) | 321/321              | 100.00                          | 98.86                                    | 100.00                                   |
| D842V 1.5x LoD<br>(C <sub>T</sub> 29) | 323/324              | 99.69                           | 98.29                                    | 99.99                                    |
| D842V 1x LoD<br>(C <sub>T</sub> 26)   | 324/324              | 100.00                          | 98.87                                    | 100.00                                   |
| D842V 1x LoD<br>(C <sub>T</sub> 29)   | 324/324              | 100.00                          | 98.87                                    | 100.00                                   |
| WT<br>(C <sub>T</sub> 26)             | 324/324              | 100.00                          | 98.87                                    | 100.00                                   |
| WT<br>(C <sub>T</sub> 29)             | 323/323              | 100.00                          | 98.86                                    | 100.00                                   |

<sup>\*</sup> Number of correct calls and two-sided 95% confidence limits for D842V mutation positive samples at 1x LoD and wild type samples tested across all sites.

## Specimen handling (Extraction reproducibility)

This study assessed sample handling variability, specifically during the DNA extraction step using sections taken from four specimens; two wild type (i.e. D842V mutation negative) clinical GIST FFPE specimens and two D842V mutation positive clinical GIST FFPE specimens. Sections from the specimens were divided into three independent sets to be extracted with the QIAamp DSP DNA FFPE Tissue Kit at three independent laboratories. All specimens were blinded before extraction. Four extractions for each specimen were performed at each of the three sites, yielding a total of 12 data points per specimen. One QIAamp DSP DNA FFPE Tissue Kit lot was used for the extractions across all three sites. All DNA samples were then tested using the *therascreen* PDGFRA RGQ PCR Kit at one site.

The proportion of correct calls across all three sites was 100%, supporting the reproducibility and repeatability for the *therascreen PDGFRA RGQ PCR* Kit System at the pre-analytical step of DNA isolation

## Lot interchangeability

The therascreen PDGFRA RGQ PCR Kit System utilises the QIAamp DSP DNA FFPE Tissue Kit, for the isolation of gDNA, and the therascreen PDGFRA RGQ PCR Kit for the amplification of gDNA and detection of PDGFRA D842V mutation status. Lot interchangeability was demonstrated using three lots of the QIAamp DSP DNA FFPE Tissue Kit and three lots of the therascreen PDGFRA RGQ PCR Kit to test five wild type and five D842V mutation positive clinical samples.

Each sample was extracted with three different lots of QIAamp DSP DNA FFPE Tissue Kit. Two replicate extractions were carried out per QIAamp DSP DNA FFPE Tissue Kit to give a total of six extractions per sample. All extracts were tested with three different lots of the *therascreen* PDGFRA RGQ PCR Kit.

The overall percentage of correct calls across lots for all mutation positive and wild type samples was 100% (180/180).

## Cross-contamination/analytical carryover

The purpose of this study was to evaluate the *therascreen* PDGFRA RGQ PCR Kit for carryover when high *PDGFRA* D842V mutation positive samples are tested adjacent to *PDGFRA* D842V mutation negative samples. This study examined the entire *PDGFRA* Assay System from extraction to PCR amplification and investigated if carryover occurred between samples, extractions, and within or between runs.

Sections from one D842V mutation positive and three wild type clinical GIST FFPE specimens were divided into two independent sets, Set A and Set B. Two operators performed extractions

for each set of samples, following a pre-defined extraction matrix, to introduce the risk of cross-contamination in wild type samples. Nine extractions from the D842V mutation positive specimen and seven extractions from each of the three wild type specimens were extracted for each of the Set A and Set B sample sets.

The extracts were tested across five PCR runs following a run matrix designed to introduce the risk of sample cross-contamination. The five PCR runs were set up consecutively by the same operator using the same equipment and Rotor-Gene Q MDx 5plex HRM instrument, with no other runs set up using this instrument between these runs. Extracts were tested with one lot of the therascreen PDGFRA RGQ PCR Kit.

A total of 126 wild type replicates were tested, all of which generated the correct mutation status call ('no mutation detected', i.e., wild type sample). Therefore, the percentage of false-positive calls for the wild type samples was 0%, demonstrating no cross-contamination of the wild type samples by D842V mutation positive samples sharing the same DNA extraction and run setup procedure.

## Interfering substances

The effects of potential interfering substances introduced from the QIAamp DSP DNA FFPE Tissue Kit (exogenous substances) or from the sample itself (endogenous substances) on assay performance were measured by comparison of  $\Delta C_T$  between interferent-spiked and control-spiked extracts of D842V mutation positive DNA samples, and by comparison of the Control reaction  $C_T$  for wild type DNA samples and by comparison of mutation status calls.

The exogenous substances present in the DNA extraction process tested were:

- Paraffin wax
- Xylene
- Ethanol

- Buffer ATL
- Proteinase K
- Buffer Al
- Buffer AW1
- Buffer AW2

D842V mutation positive and wild type samples to be spiked with exogenous interferents were first normalised to Control reaction  $C_T$  26.00 and Control reaction  $C_T$  29.00. The mutation positive samples were then diluted with wild type DNA sample (also normalised to the corresponding Control reaction  $C_T$ ) to give the % MAF representing 1.5x LoD.

Samples spiked with haemoglobin and triglycerides (endogenous interferents) during the extraction process were not normalised to Control reaction  $C_T$  26.00, or  $C_T$  29.00 nor diluted to 1.5x LoD prior to mutation assessment, but used immediately following extraction. This was to avoid removing any variability which may have been introduced by the interferent.

The study required the preparation of a test sample set and a blank sample set (Buffer ATE for exogenous substances, water for haemoglobin and Xylene for triglycerides). The test sample set included all D842V mutation positive samples and wild type samples spiked with an interferent. The blank sample set included D842V mutation positive samples and wild type samples spiked with an appropriate control substance. Samples tested with haemoglobin and triglycerides were spiked during the extraction process to reflect what would be introduced via the FFPE sample.

The test concentrations of haemoglobin and triglycerides and the estimated tissue volume used in the extraction process were based on CLSI guidelines (CLSI EP07-A2, Appendix D, 2005, Interference Testing in Clinical Chemistry; Approved Guideline). Both substances were tested at the highest (worst-case) and lowest feasible level of the interfering substance listed in the CLSI guidelines:

- Haemoglobin (2 mg/mL & 4 mg/mL)
- Triglycerides (3.7 µmol & 7.4 µmol)

Samples tested with potential exogenous interferents were spiked, following normalisation to Control reaction  $C_T$  26.00 and Control reaction  $C_T$  29.00 and dilution of the D842V mutation positive samples to 1.5x LoD, at a concentration representing the highest (worst-case) feasible level of the interfering substance carryover into a sample.

In total, eight replicates of each sample/interferent combination were tested with one therascreen PDGFRA RGQ PCR Kit lot.

All mutation status calls in the D842V mutation positive samples and wild type samples were as expected.

All exogenous substances tested had no impact on the *therascreen PDGFRA RGQ PCR Kit's* performance. The study also showed that haemoglobin had no impact on the *therascreen PDGFRA RGQ PCR Kit's* performance when testing D842V mutation positive samples.

There was, however, a statistically significant impact on the *therascreen* PDGFRA RGQ PCR Kit's performance observed between the spiked and control samples for wild type (WT) samples spiked with low and high concentrations of haemoglobin, D842V mutation positive (MT) samples spiked with the low concentration of triglycerides, and MT and WT samples spiked with the high concentration of triglycerides (Table 9).

Table 9. Summary of the impact of the endogenous substances

| Potential interferant (endogenous) | Concentration   | Sample type | Green channel/Delta C <sub>T</sub>                                    | Mutation status call |
|------------------------------------|-----------------|-------------|---|----------------------|
| Haemoglobin                        | 2 mg/mL (low)   | WT          | Higher Green channel C <sub>T</sub> value                             | No change            |
| Haemoglobin                        | 4 mg/mL (high)  | WT          | Higher Green channel C <sub>T</sub> value                             | No change            |
| Triglycerides                      | 3.7 µmoL (low)  | MT          | Lower delta C <sub>T</sub> value                                      | No change            |
| Triglycerides                      | 7.4 μmoL (high) | MT<br>WT    | Higher delta CT value<br>Higher Green channel C <sub>T</sub><br>value | No change            |

## Accuracy: Comparison to the analytical reference method

A study demonstrated the concordance in mutation detection of the *therascreen* PDGFRA RGQ PCR Kit relative to a validated next generation sequencing (NGS) method. In this study, a total of 166 FFPE samples from the BLU-285-1101 clinical trial were tested, 147 subjects received results from both methods, of which 40 were D842V positive by NGS. The Overall Percent Agreement (OPA), Positive Percent Agreement (PPA), and Negative Percent Agreement (NPA) between the *therascreen* PDGFRA RGQ PCR Kit and NGS (with NGS as the reference method) were 99.32%, 97.50%, and 100.00%, respectively.

Analytical accuracy was determined by comparing the result from the *therascreen* PDGFRA RGQ PCR Kit for both clinical trial samples and procured clinical samples against the NGS result. Of the 166 clinical trial samples and 51 procured clinical samples, 194 samples gave a valid result with both NGS and the *therascreen* PDGFRA RGQ PCR Kit. Seven clinical trial samples had a valid result with NGS but could not be evaluated using the PDGFRA Kit. The OPA, PPA, and NPA between the *therascreen* PDGFRA RGQ PCR Kit and NGS (with NGS as the reference method, and excluding PDGFRA Kit invalid results) were 99.48%, 98.44%, and 100.00%, respectively.

The OPA, PPA, and NPA were also evaluated including the seven invalid samples (samples without PDGFRA Kit results), with corresponding two-sided exact 95% CI. The results demonstrate point estimates of PPA, NPA, and OPA of 96.92%, 95.59% and 96.02%, respectively.

## Clinical performance

The *therascreen* PDGFRA RGQ PCR Kit is intended to be used as a companion diagnostic test to aid clinicians in identifying GIST patients who may be eligible for treatment with AYVAKYT (avapritinib) based on a *PDGFRA* D842V mutation detected result in GIST tumour tissue samples<sup>10</sup>.

## Clinical study BLU-285-1101

The Clinical Study BLU-285-1101 was an open-label, multicentre, international, Phase 1, first-in-human (FIH) study to evaluate the safety and efficacy of AYVAKYT (avapritinib) in adult patients with unresectable or metastatic GIST. The study was expanded to include patients with advanced GIST based on initial efficacy observed in dose escalation. A total of 237 patients, with or without a mutation in the *PDGFRA* gene, were enrolled across all parts of the study and 204 of those patients were treated with at least 1 dose of either 300 mg or 400 mg once daily (*quaque die* [QD]) of AYVAKYT (avapritinib) (33 patients were treated with a dose of <300 mg QD).

The primary endpoints were to determine maximum tolerated dose (MTD) and recommended Phase 2 dose (RP2D) of AYVAKYT (avapritinib), to determine overall safety and tolerability of avapritinib, and to determine the overall response rate (ORR) by modified response evaluation criteria in solid tumours (mRECIST) version 1.1 at the MTD/RP2D of avapritinib in patients with GIST:

- A D842V mutation in PDGFRA;
- Progressed following treatment with imatinib and at least 1 other kinase inhibitor, and who
  are not known to have a D842V mutation in PDGFRA; or
- Progressed or experienced intolerance to imatinib, including in the adjuvant setting, and who had not received additional kinase inhibitor therapy and did not have a known D842V mutation in PDGFRA.

The analysis of effectiveness of avapritinib was based on the Primary Efficacy endpoints of Overall Response Rate (ORR) in 38 Clinical Trial Assay (CTA) D842V positive, evaluable patients at the database cutoff.

Of these 38 patients, 31 patients in the 300/400 mg starting dose group had PDGFRA D842V mutations identified by the *therascreen* PDGFRA RGQ PCR Kit, 29 had confirmed complete response (CR) or partial response (PR) based on central radiology review for an ORR of 94% (95% CI: 78.6%, 99.2%). Best response for these 31 patients was CR in 2 patients, PR in 27 patients, and stable disease (SD) in 2 patients, for a clinical benefit rate (CBR) of 100%. Median duration of response (DOR) was not reached in the 300/400 mg dose group; 22 patients (76%) were censored at the time of the data cutoff based on FDA censoring rules with 64% and 51% estimated to be in response at 12 months and 18 months, respectively.

The BLU-285-1101 Study met its primary objective, demonstrating treatment with AYVAKYT (avapritinib) provides meaningful clinical benefit to *PDGFRA* D842V mutant GIST patients, as measured by ORR. ORR for patients with GIST harboring *PDGFRA* Exon 18 D842V mutations was 89% (with 95% CI 75%, 97%). Collectively, the efficacy and safety results from the BLU-285-1101 study demonstrate that AYVAKYT (avapritinib) has a favorable benefit-risk profile and is a useful treatment in patients with GIST who have *PDGFRA* D842V mutation.

## Concordance between CDx assay and CTA (Bridging study)

In the BLU-285-1101 study, the PDGFRA D842V mutation status for screening and enrolment of patients was determined by a CTA.

Concordance between the *therascreen PDGFRA RGQ PCR Kit (CDx)* and the CTA was assessed on DNA extracted from GIST samples from 154 patients in study BLU-285-1101.

Of the 154 samples, 147 gave a valid therascreen PDGFRA RGQ PCR Kit result (CDx evaluable patients). The estimated PPA, NPA, and OPA between the therascreen PDGFRA RGQ PCR Kit and the CTA (with CTA as the reference method) for CDx evaluable patients are detailed in Table 10. The result of 98.64% OPA met the study acceptance criteria of  $\geq$ 80%.

Table 10. Measures of agreement between CDx and CTA\*

| Measure of agreement          | Frequencies | Percent agreement (%) | Lower two-sided 95% confidence limit† (%) | Upper two-sided 95% confidence limit† (%) |
|-------------------------------|-------------|-----------------------|---|---|
| Overall Percent Agreement     | 145/147     | 98.64                 | 95.17                                     | 99.83                                     |
| Positive Percent Agreement    | 40/42       | 95.24                 | 83.84                                     | 99.42                                     |
| Negative Percent<br>Agreement | 105/105     | 100.00                | 96.55                                     | 100.00                                    |

<sup>\*</sup>CTA as reference method and CDx evaluable patients

Seven (7) subjects with invalid therascreen PDGFRA RGQ PCR Kit results were then included as discordant results in the analysis across all CDx testable patients. The estimated PPA, NPA, and OPA between the therascreen PDGFRA RGQ PCR Kit and the CTA (with CTA as reference method) for all CDx testable patients are detailed in Table 11. The result of 94.16% OPA met the study acceptance criteria of ≥80%.

<sup>†</sup> Clopper-Pearson (Exact) Binomial<sup>11</sup>

Table 11. Measures of agreement between CDx and CTA\*

| Measure of agreement          | Frequencies | Percent agreement | Lower two-sided 95% confidence limit† | Upper two-sided 95% confidence limit† |
|-------------------------------|-------------|-------------------|---------------------------------------|---------------------------------------|
| Overall Percent Agreement     | 145/154     | 94.16%            | 89.20%                                | 97.29%                                |
| Positive Percent Agreement    | 40/43       | 93.02%            | 80.94%                                | 98.54%                                |
| Negative Percent<br>Agreement | 105/111     | 94.59%            | 88.61%                                | 97.99%                                |

<sup>\*</sup>CTA as reference method and CDx evaluable patients

Effectiveness of treatment with AYVAKYT (avapritinib), when used with the *therascreen* PDGFRA RGQ PCR Kit, is shown in data collected in the BLU-285-1101 study of AYVAKYT (avapritinib), reanalysed in the bridging study. The *therascreen* PDGFRA RGQ PCR Kit demonstrated overall agreement of 98.64% (95% CI: 95.17, 99.83) for CDx evaluable patients. Treatment with AYVAKYT (avapritinib) provides meaningful clinical benefit to *PDGFRA* D842V mutant GIST patients, as measured by ORR. The ORR for PDGFRA D842V mutation positive patients when using the *therascreen* PDGFRA RGQ PCR Kit was 0.94 (95% CI: 0.79,0.99), corresponding to 94%.

<sup>†</sup> Clopper-Pearson (Exact) Binomial 11

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# Troubleshooting Guide

This troubleshooting guide may be helpful in solving any problems that may arise. For more information, see also the Frequently Asked Questions page at our Technical Support Center: www.qiagen.com/FAQ/FAQList.aspx (for contact information, visit www.qiagen.com).

#### **Comments and suggestions**

#### Positive Control

#### 'PC\_CONTROL\_NO\_CT\_VALUE' or 'PC\_D842V\_NO\_CT\_VALUE' flags

a. Incorrect configuration of the PCR

- Check your pipetting scheme and repeat the PCR.
- The storage conditions for one or more kit components did not comply with the instructions given in "Reagent Storage and Handling" on page 16
- Check the storage conditions (see kit label) of the reagents and use a new kit, if necessary.

#### 'PC\_CONTROL\_HIGH\_CT', 'PC\_D842V\_HIGH\_CT', 'PC\_CONTROL\_LOW\_CT', or 'PC\_D842V\_LOW\_CT' flags

a. Error during preparation of the PCR

Repeat the PCR ensuring accurate pipetting.

#### No Template Control (NTC)

#### 'NTC\_CONTROL\_UNEXPECTED\_CT' or 'NTC\_D842V\_UNEXPECTED\_CT' flags

a. Contamination occurred during preparation of the  $\ensuremath{\mathsf{PCR}}$ 

Repeat the PCR with new reagents. If possible, close the PCR tubes directly after addition of the sample to be tested. Ensure that the workspace and instruments are decontaminated at regular intervals.

#### Sample

#### 'CONTROL\_LOW\_CT' flag

a. Sample is too concentrated

Dilute sample to increase C<sub>T</sub> value. Samples should be diluted using the water provided in the kit (Water for Dilution [Dil.]).

#### Comments and suggestions

#### 'CONTROL\_HIGH\_CT' flag

a. Insufficient starting DNA template present in sample

Retest one more time. If the system shows the same flag a second time, re-extract the DNA using slides from the same tissue specimen and repeat the assay. If after re-extraction the system shows the same flag for the sample, retest for a second time. If the flag occurs again, the sample is not suitable for use. It must be recorded as "indeterminate" and no further testing performed.

#### 'CONTROL\_NO\_CT\_VALUE' flag

a. No amplifiable DNA template present in sample

Retest one more time. If the system shows the same flag a second time, re-extract the DNA using slides from the same tissue specimen and repeat the assay. If after re-extraction the system shows the same flag for the sample, retest for a second time. If the flag occurs again, the sample is not suitable for use. It must be recorded as "indeterminate" and no further testing performed.

#### Run control or sample

#### 'CURVE\_SHAPE\_ANOMALY' or 'OTHER\_TARGET\_INVALID' flags

a. Error during preparation of the PCR

#### Run Control

If either of these flags are seen for a Run Control with another flag for a Run Control, then follow the comments and suggestions for the other flag. If either of these flags are seen for a Run Control with no other flag, repeat the run ensuring accurate mixing and pipetting.

#### Sample

If either of these flags are seen for a sample, retest the sample ensuring accurate mixing and pipetting.

# **Symbols**

The following symbols appear in the instructions for use or on the packaging and labeling:

| Symbol      | Symbol definition   |
|-------------|---|
| <n></n>     | Contains reagents sufficient for <n> reactions</n>  |
| $\subseteq$ | Use by  |
| CE          | This product fulfills the requirements of the European Regulation $2017/746$ for in vitro diagnostic medical devices. |
| IVD         | In vitro diagnostic medical device  |
| REF         | Catalog number  |
| LOT         | Lot number  |
| MAT         | Material number (i.e., component labeling)  |
| COMP        | Component   |
| CONT        | Contains  |
| GTIN        | Global Trade Item Number  |

#### Symbol

#### Symbol definition



Number

Rn

R is for revision of the Instructions for Use and n is the revision number



Temperature limitation



Manufacturer



Consult instructions for use



Protect from light



Warning/Caution

## **Contact Information**

For technical assistance and more information, please see our Technical Support Centre at www.qiagen.com/Support, call 00800-22-44-6000, or contact one of the QIAGEN Technical Service Departments or local distributors (see back cover or visit www.qiagen.com).

# Ordering Information

| Product                                       | Contents  | Cat. no. |
|---|---|----------|
| therascreen PDGFRA RGQ PCR Kit                | For 24 reactions: Reaction mix, Positive Control, No-Template Control, Dilution Water, Taq DNA Polymerase   | 874911   |
| Rotor-Gene Q therascreen PDGFRA Assay Package | Software protocol package for use with the<br>therascreen PDGFRA RGQ PCR Kit and<br>Rotor-Gene Q MDx 5plex HRM instrument<br>Compatible with Rotor-Gene AssayManager  | 9027016  |
|   | version 2.1 or later  |          |
| Rotor-Gene Q MDx 5plex HRM and accessories    |   |          |
| Rotor-Gene Q MDx 5plex HRM Platform           | Real-time PCR cycler and High Resolution<br>Melt analyzer with 5 channels (green, yel-<br>low, orange, red, crimson) plus HRM chan-<br>nel, laptop computer, software, and<br>accessories.  | 9002032  |
| Rotor-Gene Q MDx 5plex HRM System             | Real-time PCR cycler and High Resolution<br>Melt analyzer with 5 channels (green, yel-<br>low, orange, red, crimson) plus HRM chan-<br>nel, laptop computer, software, accessories,<br>1-year warranty on parts and labor, install-<br>ation and training | 9002033  |
| Loading Block 72 x 0.1 mL Tubes               | Aluminum block for manual reaction setup with a single-channel pipette in $72 \times 0.1 \text{ mL}$ tubes  | 9018901  |
| Strip Tubes and Caps, 0.1 mL (250)            | 250 strips of 4 tubes and caps for 1000 reactions   | 981103   |
| Strip Tubes and Caps, 0.1 mL (250)            | $10 \times 250$ strips of 4 tubes and caps for 10000 reactions  | 981106   |
| PCR Tubes, 0.2 mL (1000)                      | 1000 thin-walled tubes for 1000 reactions of 20–50 $\mu\text{L}$  | 981005   |

| Product                             | Contents   | Cat. no. |
|-------------------------------------|--|----------|
| 72-Well Rotor                       | For holding Strip Tubes and Caps, 0.1 mL, with reaction volumes of 10 to 50 $\mu$ L; requires Locking Ring 72-Well Rotor                       | 9018903  |
| Locking Ring 72-Well Rotor          | For locking Strip Tubes and Caps, 0.1 mL in the 72-Well Rotor  | 9018904  |
|                                     | RELATED PRODUCTS   |          |
| QIAamp DSP DNA FFPE Tissue Kit (50) | For 50 DNA preps; QIAamp MinElute®<br>QIAamp DSP DNA FFPE Tissue Kit (50)<br>Columns, Proteinase K, Buffers, and Col-lec-<br>tion Tubes (2 mL) | 60404    |

For up-to-date licensing information and product-specific disclaimers, see the respective QIAGEN kit Instructions for Use. QIAGEN kit Instructions for Use are available at **www.qiagen.com** or can be requested from QIAGEN Technical Services or your local distributor.

# **Document Revision History**

| Revision        | Description  |
|-----------------|--|
| R1, March 2022  | Initial release  |
| R2, April 2024  | Updated "In-use stability" on page 16 and the number of freeze-thaw cycles   |
|                 | Update to software flags in Table 5:   |
|                 | <ul> <li>'MUTANT' replaced with 'D842V' in the flags</li> </ul>  |
|                 | <ul> <li>Added two additional flags: 'CURVE_SHAPE_<br/>ANOMALY' and 'OTHER_TARGET_INVALID'.</li> </ul>   |
|                 | Mutation call 'Mutation Detected' has been updated to 'D842V Mutation Detected'.   |
|                 | In "Warnings and Precautions" on page 12, 'reaction mix' has been updated to 'master mix' in the sentence ' Do not use reaction volumes (master mix plus sample) of less than 25 µL. |
|                 | Clarified that kit reagents should be thawed and mixed to ensure performance of kit is not affected.   |
|                 | Updated the section  |
|                 | Updated to clarify patient numbers   |
|                 | Some words have been updated from US English spelling to British English spelling.   |
| R3, August 2025 | Added Intended Use and Intended User sections  |
|                 | Corrected catalog number for Rotor-Gene Q therascreen PDGFRA Assay Package   |

#### Limited License Agreement for therascreen® PDGFRA RGQ PCR Kit

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