### diatech pharmacogenetics



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The "Easy<sup>®</sup> KRAS" Kit detects mutations of the KRAS codons 12, 13, 59, 61, 117 and 146 by Real-Time PCR.

For in vitro diagnostic use

RT001

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Diatech Pharmacogenetics srl a Socio Unico Società soggetta all'attività di direzione e coordinamento di Diatech srl con sede in Jesi Via Ignazio Silone, 1 b - 60035 Jesi (AN) Italy C.F./P.Iva/R.Imprese di Ancona n. 02483840423 Tel. +39 0731 213243 Fax +39 0731 213239 info@diatechpharmacogenetics.com www.diatechpharmacogenetics.com Changes made since the previous version 2015/02:

- Substitution of KRAS pos ctrl II with Easy KRAS pos ctrl.
  Update of section "Materials required but not provided" Genomic DNA extraction.
  Validation of the kit on ABI 7500.

For further details contact the technical support of the Diatech Pharmacogenetics (support@diatechpharmacogenetics.com).

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#### INTENDED USE

The *in vitro* diagnostic "**Easy**<sup>®</sup> **KRAS**" kit is intended for the qualitative detection by Real-Time PCR of *KRAS* somatic mutations in the genomic DNA isolated from fresh, frozen or formalin fixed paraffin-embedded (FFPE) tumor tissue.

The "Easy<sup>®</sup> KRAS" Kit is validated on the following instruments:

- CFX96 Bio-Rad (software v. 3.1) .
- ABI 7300- Applied Biosystems (software v. 1.4.1)
- .
- ABI 7500 Applied Biosystems (software v.2.0.5) Stratagene Mx3000P, Mx3005P Agilent Technologies (software v. 4.10 Build 389) .
- .
- Rotor-Gene Q Qiagen (software v. 1.7 Build 87) Rotor-Gene 6000 Corbett (software v. 1.7 Build 87)

List of detectable mutations:

KRAS c	odon 12
•	G12R (34G>C)
	G12S (34G>A)
	G12C (34G>T)
	G12A (35G>C)
	G12D (35G>A)
	G12V (35G>T)
KRAS c	odon 13
•	G13D (38G>A)
KRAS c	odon 59
(not distir	nguishable between them)
•	A59T (175G>A)
•	A59E (176C>A)
•	A59G (176C>G)
	odon 61
(not distir	nguishable between them)
•	Q61K (181C>A)
•	Q61L (182A>T)
•	Q61R (182A>G)
•	Q61H (183A>C)
	Q61H (183A>T)
	odon 117
(not distir	nguishable between them) K117E (349A>G)
	K117E (349A>G) K117R (350A>G)
	K117R (350A>G) K117N (351A>T)
:	K117N (351A>T) K117N (351A>C)
	odon 146
	nguishable between them)
	A146T (436G>A)
	A146P (436G>C)
	A146V (437C>T)

#### PRINCIPLE OF THE ASSAY

The "Easy<sup>®</sup> KRAS" Kit is designed to selectively amplify mutant specific sequences in samples that contain a mixture of wild-type and mutated DNA. The detection is achieved using fluorescent probes labelled with FAM and HEX.

The "Easy<sup>®</sup> KRAS" Kit is composed of 11 assays for the detection of the KRAS mutations and a control assay for the assessment of DNA content in the sample.

Each assay contains primers and probes for the detection of the target (FAM) as well as an endogenous control gene (HEX). The amplification of the endogenous control gene enables to verify the amplification procedure and the possible presence of inhibitors, which may cause false negative results.

- 1. KRAS G12A: the assay detects the G12A (35G>C) mutation
- 2. **KRAS G12D**: the assay detects the G12D (35G>A) mutation
- 3. **KRAS G12V**: the assay detects the G12V (35G>T) mutation
- 4. **KRAS G12R**: the assay detects the G12R (34G>C) mutation
- 5. **KRAS G12S**: the assay detects the G12S (34G>A) mutation
- KRAS G12C: the assay detects the G12C (34G>T) mutation
   KRAS G13D: the assay detects the G13D (38G>A) mutation
- KRAS A59x: the assay detects the A59T (175G>A), A59E (176C>A), A59G (176C>G) mutations but does not distinguish between them
- KRAS Q61x: the assay detects the Q61K (181C>A), Q61L (182A>T), Q61R (182A>G), Q61H (183A>C), Q61H (183A>T) mutations but does not distinguish between them
- 10. KRAS K117x: the assay detects the K117E (349A>G), K117R (350A>G), K117N (351A>T), K117N (351A>C) mutations but does not distinguish between them
- 11. KRAS A146x: the assay detects the A146T (436G>A), A146P (436G>C), A146V (437C>T) mutations but does not distinguish between them
- 12. KRAS ctrl: the assay detects a KRAS region without any known polymorphism/mutation

The "Easy<sup>®</sup> KRAS" kit includes a DNA reference standard Horizon KRAS G12V 1% containing a defined ratio between wild-type and mutant DNA to check the analytical process and the assay performances.

#### KIT CONTENTS

COMP	QUAN	Color	
KRAS G12A mix (1)	3 x 10 µl	BLUE	Mixture of specific primers and probes targeting both
		00551	KRAS G12A mutation and the internal control.
KRAS G12D mix (2)	3 x 10 µl	GREEN	Mixture of specific primers and probes targeting both KRAS G12D mutation and the internal control.
KRAS G12V mix (3)	3 x 10 µl	PINK	Mixture of specific primers and probes targeting both
	• · · · • P.		KRAS G12V mutation and the internal control.
KRAS G12R mix (4)	3 x 10 µl	RED	Mixture of specific primers and probes targeting both
	0 40!		KRAS G12R mutation and the internal control.
KRAS G12S mix (5)	3 x 10 µl	WHITE	Mixture of specific primers and probes targeting both KRAS G12R mutation and the internal control.
KRAS G12C mix (6)	3 x 10 µl	ORANGE	Mixture of specific primers and probes targeting both
	с л то р.	0	KRAS G12C mutation and the internal control.
KRAS G13D mix (7)	3 x 10 µl	YELLOW	Mixture of specific primers and probes targeting both
			KRAS G13D mutation and the internal control.
KRAS A59x mix (8)	3 x 10 µl	BROWN	Mixture of specific primers and probes targeting both
KRAS Q61x mix (9)	3 x 10 µl	TRANSPARENT	KRAS A59T, A59E, A59G and the internal control. Mixture of specific primers and probes targeting both
	σχισμι		KRAS Q61K, Q61L, Q61R, Q61H (183A>C), Q61H
			(183A>T) and the internal control.
KRAS K117x mix (10)	3 x 10 µl	PURPLE	Mixture of specific primers and probes targeting both
			KRAS K117E, K117R, K117N (351A>T), K117N
KRAS A146x mix (11)	2 x 10 ul	BLACK	(351A>C) and the internal control. Mixture of specific primers and probes targeting both
	3 x 10 µl	DLACK	KRAS A146T, A146P, A146V and the internal
			control.
KRAS ctrl mix (12)	6 x 10 µl	GREY	Mixture of specific primers and probes targeting both
			region of the KRAS gene free from any known
			polymorphism/mutation and the internal control.
Easy KRAS pos ctrl	<b>CONTROL</b> + 3 x 200 µl		Positive control DNA containing a mixture of synthetic DNA sequences that correspond to each
			mutation detected by this kit in a background of wild-
			type genomic DNA.
Horizon KRAS G12V 1%	1 x 12 µl		Horizon DNA reference standard KRAS G12V 1% to
			check the analytical process.
WATER	CONTROL - 2 x 1.5 ml		DNase-, RNase-free water to be use exclusively for
			the preparation of the PCR mix and as negative control.
Water (diluent)	1 x 1.5 ml		DNase-, RNase-free water to be use exclusively as
			samples diluent.
Taq Premix 920	4 x 920 µl		Solution containing hot start Taq DNA polimerase,
	4 40 1		reaction buffer, Mg2+ and dNTP Mixture.
Dye R-I	4 x 40 µl		Inert fluorophore to be used for the amplification on the ABI 7300 instrument.
Dye R-II	4 x 40 µl		Inert fluorophore to be used for the amplification on
			the ABI 7500 instrument.
12 strip tubes & caps	1 x 5 strips		0.2 ml 12-tube strip DNase-, RNase-free to be used
			for the preparation of reaction mixture.

The kit contains sufficient reagents to carry out 24 tests for each assay for a maximum of 3 runs on ABI 7300, ABI 7500, CFX96 and Mx3000P/3005P (6 samples and two controls per run) and of 4 runs on Rotor-Gene (4 samples and two controls per run).

#### DOCUMENTS AVAILABLE ON-LINE

Easy® KRAS User Manual and Material Safety Data Sheet (MSDS) are available at <u>www.diatechpharmacogenetics.com/en/reserved-area</u>.

#### MATERIALS REQUIRED BUT NOT PROVIDED

#### Genomic DNA extraction

The "Easy<sup>®</sup> KRAS" kit does non contain reagents for DNA extraction.

#### Recommended kits:

- "QIAamp<sup>®</sup> DNA FFPE Tissue kit" (cod. 56404, Qiagen)
- "QIAamp<sup>®</sup> DNA Mini kit" (cod. 51304, Qiagen)
- "Genomic DNA FFPE One-Step Kit" (cod. MGF-03, RBC); to use with MagCore Automated Nucleic Acid Extractor (RBC Bioscience) automatic systems
- "Genomic DNA Tissue Kit" (cod. MGT-02, RBC); to use with MagCore Automated Nucleic Acid Extractor (RBC Bioscience) automatic systems
- In case you are using FFPE tissues, you will also need:
  - Xilene (e.g.: "Xylenes, histological grade" cod. 534056, Sigma Aldrich)
  - Absolute Ethanol (quality of analytical degree)

① In case you employ kits which are different from those recommended, it is the user's responsibility to use standardized samples (e.g.: VEQ – EQAS quality schemes, Horizon Diagnostics samples) to verify that this does not imply a reduction of the performance of the system under analysis.

#### Amplification

Real-Time PCR Instruments:

- CFX96 Bio-Rad (software v. 3.1)
- ABI 7300- Applied Biosystems (software v. 1.4.1)
- ABI 7500 Applied Biosystems (software v.2.0.5)
- Stratagene Mx3000P, Mx3005P Agilent Technologies (software v. 4.10 Build 389)
- Rotor-Gene Q Qiagen (software v. 1.7 Build 87)
- Rotor-Gene 6000 Corbett (software v. 1.7 Build 87)

Detection channels for FAM and HEX fluorescence. Range of environmental temperature: 15-30°C.

#### Materials:

- 1,5 ml polypropylene twist-lock tubes (DNase-, RNase-, DNA-, PCR inhibitor-free)
- Micropipettes (volumes from 1 to 1.000 µl)
- Sterile filter tips DNase-, RNase-free (volumes from 1 to 1.000 µl)
- 96 well plates and foil or caps compatibles with the thermal-cycler used (check the compatibility in the user manual of the instrument) for instance:
  - 1. Bio-Rad CFX96: Hard Shell PCR plates 96-well WHT/CLR cod. HSP 9601; MICROSEAL B SEALS cod. MSB 1001
  - ABI 7300/ ABI 7500: MICROAMP OPTICAL 96 WELL RNX PLATE cod. N8010560; OPTICAL ADHESIVE COVERS cod. 4360954
  - 3. Stratagene Mx3000P/Mx3005P: Optical caps (8x strip) cod. 401425; QPCR 96-Well Plates, Non-Skirted cod. 401333
  - DNase-and RNase-free, thin-wall, PCR tubes with flat cap or 0.1 ml tubes in strip, suitable for use on Rotor-Gene, for instance:
  - 1. Qiagen, cat. no. 981103
- 2. LTF-Labortechnik GmbH & Co, no. 102.0170
- Powder-free disposable gloves

#### STABILITY AND STORAGE

Store all the reagents according to the instructions on the packages, in particular:

- Store all the reagents at -35/-20°C in the original package immediately upon receipt.
- After thawing, store **Taq PreMix 920** at +2/+8°C and use it within 6 months or within the expiration date.
- Avoid thawing and re-freezing the reagents more than twice, as this could lead to poor performance.
- Protect all mixes containing probes from light to avoid degradation of the fluorescent dyes.
- If properly stored, the reagents remain stable until the expiration date displayed on the individual label.

#### SYMBOLS

REF	Catalog number (product code)	CONTROL -	Negative control
LOT	Lot number	Í	Consult the instruction
Σ	Content sufficient for <n> tests</n>	НВ	User manual (handbook)
IVD	For in vitro diagnostic use	$\leq$	Expiration date
CONT	Content	J.	Storage conditions
COMP	Components		Manufactured by
NUM	Number of aliquots	í	Important Note
QUAN	Quantity per aliquot	GTIN	Global Trade Item Number
CONTROL +	Positive control		

#### PRODUCT USE LIMITATIONS

- The "Easy<sup>®</sup> KRAS kit can only be used by specialized personnel, properly instructed and trained.
- It is necessary to operate in compliance with the general guidelines of Good Laboratory Practice (GLP) and the instructions contained in this manual.
- Do not use expired or incorrectly stored reagents.
- The "Easy<sup>®</sup> KRAS" kit is designed to be used with the instruments "Rotor-Gene Q" (Qiagen), "Rotor-Gene ™ 6000" (Corbett Research), CFX96 (Bio-Rad), ABI 7300/ABI 7500 (Applied Biosystems) and Stratagene Mx3000P/Mx3005P (Agilent Technologies).
- The reliability of the results also depends on the procedures carried out in the pre-amplification stages, including the selection of starting biological speciments, the preservation of the samples and DNA extraction.
- Any diagnostic results generated by this procedure must be interpreted with reference to other clinical or laboratory findings.

#### QUALITY CONTROL

- The "Easy<sup>®</sup> KRAS" Kit was designed, developed and validated in accordance with D.Lgs n. 332 of 08/09/2000 ("Implementation of Directive 98/79 / EC on in vitro diagnostic medical devices") and subsequent legislative changes.
- In accordance with the company system of full quality assurance (certified according to European standards EN ISO 9001 and ISO 13485), to ensure consistent product quality, each batch of "Easy<sup>®</sup> KRAS" is subjected to functional quality control according to technical specifications and is released only if it is compliant with the quality control criteria.

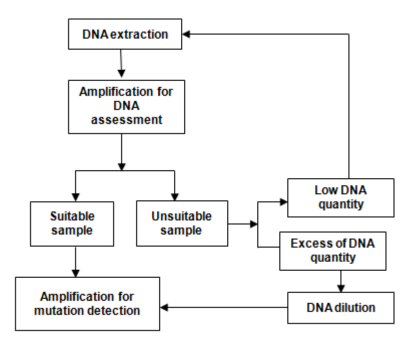
#### WARNINGS AND PRECAUTIONS

- The kit may only be used by specialist personnel, properly instructed and trained to perform *in vitro* laboratory techniques.
  - Handle all samples as potentially infectious material inside a laminar flow hood (class II biological safety cabinet or higher).
- Follow the laboratory safety procedures described in "Biosafety in Microbiological and Biomedical Laboratories" (Richmond, JY and McKinney, RW (eds) 5th edition (2009) and in the NCCLS (National Committee for Clinical Laboratory Standards) Document M29-T. Protection of Laboratory Workers from Infectious Disease Transmitted by Blood, Body Fluids and Tissue. Tentative guidelines. Villanova, PA:NCCLS, 1989).
- Do not eat, drink or smoke in the laboratory. When handling biological samples, disposable gloves, gowns and goggles or face
  masks should be worn to protect against biological agents.
- ① Constantly check that the gloves are free from contamination by the biological material being treated. If not, replace them immediately to avoid the possibility of cross-contamination between samples and contamination of the workplace. Wash hands thoroughly after handling samples and reagents.
- The Material Safety Data Sheet (MSDS) is available in the reserved area of the web-site Diatech Pharmacogenetics <u>www.diatechpharmacogenetics.com</u>.
- Perform the procedure in accordance with Good Laboratory Practice (GLP) general guidelines.
- It is recommended to ensure that the laboratory work flow proceeds in a unidirectional manner, preparing, if possible, two separate work areas for:
  - extraction of nucleic acids;

o amplification reaction;

- Organize the laboratory so that dedicated pipettes, tips and materials are used for each activity.
- Use sterile filter tips. Avoid aerosols.
- Use tubes with twist-lock caps during the extraction of nucleic acids in order to avoid the leakage of the samples and potential contamination.
- During the procedures for nucleic acid extraction and amplification, avoid contamination of reagents with airborne microbes by only opening the reagents within the hood.
- Change the pipette tip before each extraction of reagents and every time you move from one sample to another in any stage of the procedure.
- The precision pipettes used should have an accuracy of within 3% of the set volume.
- Periodically check the calibration status of the dispensing instruments.
- Do not use reagents after the expiration date shown on each container.
- All reagents supplied in the "Easy<sup>®</sup> KRAS" kit are intended to be used solely with the other reagents in the same "Easy<sup>®</sup> KRAS" kit. Do not substitute or mix reagents in the kit from different batches, in order to maintain optimal performance.
- Only use the Taq PreMix 920 that is provided in the kit. Do not substitute with Taq PreMix 920 from other kits or with similar reagents from other suppliers.
- Discard unused reagents and the expired kit and waste in accordance with current national laws and local regulations.
- <u>Extraction area</u>: at the end of the procedure, decontaminate the pipettes and the laboratory surfaces on which work has been carried out, by cleaning with appropriate products (e.g. FD 322, Dürr Dental, Germany) and UV irradiate the work surface of the biological cabinet where the pipettes should be carefully placed after decontamination.
- <u>Amplification area</u>: at the end of the procedure, decontaminate the pipettes and the laboratory surfaces on which work has been carried out, by cleaning with appropriate products to eliminate nucleic acids and amplicons (e.g. "DNA Cleaner" - code DC001, Diatech Pharmacogenetics) and subsequent UV irradiation, if available.
- Avoid contamination of samples and reagents.
- Store reagents and samples separately.
- In order to avoid possible contamination from carry-over, do not open the reaction tubes after amplification.
- Before use all reagents need to be thawed at room temperature, mixed by inverting 10 times and centrifuged briefly.
- All reagents contained in the kit are ready-to-use and don't need to be diluted. The reagent dilution may result in a loss of performance.
- Include in each run at least 1 negative control (WATER) and 1 positive control (Easy KRAS pos ctrl).
- In order to avoid any mixing up of samples pay particular attention to samples dispensation, placement of tubes into the instrument, editing the sample name in the software.
- Carefully read this User Manual.
- Check that the version of this User Manual corresponds to the one described on the "Easy<sup>®</sup> KRAS" kit box label.
- The right to contest the kit before the expiration date becomes void if the product is used in violation of GLP guidelines and the manufacturer's recommendations.
- The registered names and trademarks indicated in this document are to be considered protected by law, even when not
  explicitly stated.

#### ANALYTICAL PROCEDURE



#### DNA EXTRACTION

① Perform this step in the area dedicated to DNA isolation and dilution, using dedicated materials and instruments.

- The "Easy<sup>®</sup> KRAS" kit does not include the reagents for DNA extraction.
- The quantity of biological material required for the DNA extraction depends on protocols.
- Refer also to the extraction kit manual for selection and treatment of FFPE slides.
- Commercial kits working with silica filters or magnetic beads are recommended. Avoid methods that use phenol or boiling in basic solution without purification.
- Perform the DNA extraction following the instructions of the extraction kit in use.
- If the extraction protocol involves the use of wash buffers containing ethanol, it is advisable to perform a further centrifugation before final elution to remove any possible traces of ethanol. This will prevent illubilition of the reaction by the ethanol.
- After the extraction, proceed immediately with the quali-quantitative evaluation of the DNA and the amplification reaction, or store the extracted DNA at ≤-20°C, divided into aliquots in order to maintain the experimental conditions constant in case of repetition.
- Just as an indication, for non-paraffin embedded samples like fresh/frozen tissue, plasma, blood, the recommended DNA amount in each test tube is 5-10 ng; for paraffin embedded samples, the recommended DNA amount in each reaction tube is 15-20 ng.
- ① As absorbance reading cannot distinguish between fragmented and not fragmented DNA and therefore it can overestimate the concentration of template, DNA assessment should be based on the KRAS ctrl mix. The evaluation of quality and quantity of the DNA in the samples with the KRAS ctrl mix before the mutation analysis is highly recommended. DNA assessment based on the KRAS ctrl mix may differ from spectrophotometric quantification.
- ③ All assays in the "Easy<sup>®</sup> KRAS" kit amplify short DNA sequences. However heavily fragmented DNA can generate no amplification product.

#### AMPLIFICATION

General recommendations for amplification (valid for all instruments)

- ① Perform this step in the area dedicated to amplification mixes preparation, using dedicated materials and instruments. Before starting decontaminate pipettes, benches and wood in order to degrade any trace of DNA and possibly radiate with UV light for at least 30 minutes.
- Switch on the instrument and the software at least 20-30 minutes before starting the reaction to allow the heating of the lamps where necessary.
- Thaw all necessary reagents before use.
- Thoroughly mix the reagents in a vortex, or inverting each tube ten times, and spin them briefly before use.
- Prepare and mark an appropriate number of tubes or wells of the plate to use.
- Each run must include at least one amplification negative control (WATER) and one amplification positive control (Easy KRAS pos ctrl) for each mix.

#### INSTRUMENT SETUP

#### Rotor-Gene Q, Rotor-Gene 6000

- Follow the instructions reported in the user manual of the instrument to set up the following fluorescence acquisition channels and thermal profile:
- "Green": source 470 nm detector 510 nm Gain 8
- "Yellow": source 530 nm detector 555 nm Gain 10
- "Green 2": source 470 nm detector 510 nm –- "Gain Optimisation" 58°C Before 1st acquisition, "Tube Position":51 (KRAS A146x mix), "Target Sample Range" 20-30 FI
- "Yellow 2": source 530 nm detector 555 nm "Gain Optimisation" 58°C Before 1st acquisition, "Tube Position":51 (KRAS A146x mix), "Target Sample Range" 20-30 FI

#### **Thermal Profile**

Hold 95°C for 2 minutes

40 cycles 95°C for 10 seconds / 58°C for 60 seconds (acquire fluorescense in channels "Green", "Green 2", "Yellow", Yellow 2")

Reaction volume: 20 µl.

#### Stratagene Mx3000P, Mx3005P

- Select "New" "Quantitative PCR (Multiple standards)", then "OK".
- Check for the presence of the message "Lamp Warm-up".
- Follow the instructions reported in the user manual of the instrument to set up the following fluorescence acquisition channels and thermal profile:
  - "FAM": source 492 nm detector 516 nm Filter gain factor x8
    - "HEX": source 535 nm detector 555 nm Filter gain factor x8

## Thermal Profile Hold 95°C for 2 minutes 40 cycles 95°C for 10 seconds / 58°C for 60 seconds (acquire fluorescence in channels "FAM" and "HEX" setting up END 1)

Reaction volume: 20 µl.

#### CFX96

• Follow the instructions reported in the user manual of the instrument to set up the following fluorescence acquisition channels and thermal profile:

Therm	nal Profile
Step	
1	95°C for 2 minutes
2	95°C for 10 seconds
3	58°C for 60 seconds (Plate read – All Channels)
4	GO TO 2 39 more times

Reaction volume: 20 µl.

① Select the options "All Channels" to acquire the signal in both FAM and HEX.

#### <u>ABI 7300</u>

- Select "Create new document" On Assay: "Standard Curve (Absolute Quantitation)", then "Next".
- Follow the instructions reported in the user manual of the instrument to set up the following fluorescence acquisition channels and thermal profile:

#### Thermal Profile

Hold	95°C for 2 minutes
40 cycles	95°C for 10 seconds / 58°C for 60 seconds (acquire fluorescence in channels FAM, JOE; Quencer – None; Passive
	Reference Dye - ROX)

Reaction volume: 20 µl.

#### ABI 7500

- Select "New Experiment" then "7500 (96 Wells)", "Quantitation Standard Curve", "TaqMan<sup>®</sup> Reagents", "Standard (-2 hours to complete a run)".
- Follow the instructions reported in the user manual of the instrument to set up the following fluorescence acquisition channels and thermal profile:

# Thermal Profile Hold 95°C for 2 minutes 40 cicli 95°C for 10 seconds / 58°C for 60 seconds (acquire fluorescence in channels FAM, JOE; Quencer – None; Passive Reference Dye - ROX)

Reaction volume: 20 µl.

#### DNA ASSESSMENT

• For each sample and control prepare an amplification mixture (**Amp-Mix**), according to the following table, where N is the total number of samples and controls to be tested:

Rotor-Gene, Stratagene, CFX96											
Amp-Mix	Reagent volume for 1 reaction (µI)	Reagent volume for N reactions +1 (µl)									
Taq Premix 920	10										
WATER CONTROL -	4										
KRAS ctrl mix (12)	1										
Total Volume	15										

ABI 7300		
Amp-Mix	Reagent volume for 1 reaction (µl)	Reagent volume for N reactions +1 (µl)
Taq Premix 920	10	
Dye R-I	0.4	
WATER CONTROL-	3.6	
KRAS ctrl mix (12)	1	
Total Volume	15	

ABI 7500		
Amp-Mix	Reagent volume for 1 reaction (µI)	Reagent volume for N reactions +1 (µl)
Taq Premix 920	10	
Dye R-II	0.2	
WATER CONTROL-	3.8	
KRAS ctrl mix (12)	1	
Total Volume	15	

• Mix the **Amp-Mix** thoroughly by repeated pipetting or rapid vortexing, then centrifuge briefly.

Pipette 15 µl of the Amp-Mix in all the marked reaction test tubes or wells.

Add to the respective test tubes or wells:

negative control	5 μl <b>WATER</b>	CONTROL -
samples	5 µl DNA	
positive control	5 µl Easy KRAS pos ctrl	CONTROL +

- Final volume: 20 µl.
- Briefly centrifuge the plate.
- Check that the thermal profile is setted up correctly and start the run.
- ③ Before starting the run, please pay attention to the plate orientation (well A1 on the upper left position) or to the Rotor-Gene 0.1ml strips of tubes orientation (mark the first tube of each strip).
- ① Proceed with the analysis following the instructions of the section "Data analysis".

#### **MUTATION DETECTION**

- ① Only samples that after DNA assessment are suitable can be analyzed for mutations detection.
- (i) Each sample must be amplified with 12 different mixes: KRAS G12A (1), KRAS G12D (2), KRAS G12V (3), KRAS G12R (4), KRAS G12S (5), KRAS G12C (6), KRAS G13D (7), KRAS A59x (8), KRAS Q61x (9), KRAS K117x (10), KRAS A146x (11), KRAS ctrl (12).
- ① The DNA reference standard Horizon KRAS G12V 1% must be amplified only with the mixes KRAS G12V (3) and KRAS ctrl (12).
- i The kit content is optimized to analyze six clinical samples and two controls Easy KRAS pos ctrl and WATER in each run performed on CFX96, ABI7300/7500, Stratagene and four clinical samples and two controls Easy KRAS pos ctrl and WATER in each run performed on Rotor-Gene:

#### Mx3000P/3005P, ABI 7300/7500, CFX96 (RT001 sample grid A)

	1	2	3	4	5	6	7	8	9	10	11	12
А	DNA1											
В	DNA2											
С	DNA3											
D	DNA4											
E	DNA5											
F	DNA6											
G	POS CTRL											
н	WATER											



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#### Rotor-Gene (RT001 sample grid B)

DN	IA1	DN	IA2	DN	IA3	DN	IA4	POS	CTRL	WA	TER	DN	IA1	DN	IA3	POS	CTRL
1	•	9	•	17	•	25	•	33	•	41	•	49	/./	57	/./	65	
2	•	10	•	18	•	26	•	34	•	42	•	50	•	58	•	66	•
3	•	11	•	19	•	27	•	35	•	43	•	51	•	59	•	67	•
4	•	12	•	20	•	28	•	36	•	44	•	52	•	60	•	68	•
DN	IA1	DN	IA2	DN	IA3	DN	IA4	POS	CTRL WATER DNA2		A2	DN	IA4	WA	TER		
5	•	13	•	21	•	29	•	37	•	45	•	53		61	·	69	
6	•	14	•	22	•	30	•	38	•	46	•	54	•	62	•	70	•
7	•	15	•	23	•	31	•	39	•	47	•	55	•	63	•	71	•
8	•	16	•	24	•	32	•	40	•	48	•	56	•	64	•	72	•

Prepare, for each sample and control, 12 different amplification mixtures (Amp-Mix), one for each assay, as indicated in the following scheme, where N is the numer of samples and controls to be tested.

If you are working with a 96 well plate, it is possible to preprare the Amp-Mix in the strip provided with the kit, so you can í dispense the all Amp-Mix with a multichannel pipette.

Rotor-Gene, Stratagene, CFX96					
Amp-Mix	Reagent volume for 1 reaction (µl)	Reagent volume for N reactions +1 (µI)			
Taq Premix 920	10				
WATER CONTROL-	4				
KRAS G12A mix (1) or	1				
KRAS G12D mix (2) or					
KRAS G12V mix (3) or					
KRAS G12R mix (4) or					
KRAS G12S mix (5) or					
KRAS G12C mix (6) or					
KRAS G13D mix (7) or					
KRAS A59x mix (8) or					
KRAS Q61x mix (9) or					
KRAS K117x mix (10) or					
KRAS A146x mix (11) or					
KRAS ctrl mix (12)					
Total Volume	15				

ABI 7300					
Amp-Mix	Reagent volume for 1 reaction (µI)	Reagent volume for N reactions +1 (µl)			
Taq Premix 920	10				
Dye R-I	0.4				
WATER CONTROL -	3.6				
KRAS G12A mix (1) or	1				
KRAS G12D mix (2) or					
KRAS G12V mix (3) or					
KRAS G12R mix (4) or					
KRAS G12S mix (5) or					
KRAS G12C mix (6) or					
KRAS G13D mix (7) or					
KRAS A59x mix (8) or					
KRAS Q61x mix (9) or					
KRAS K117x mix (10) or					
KRAS A146x mix (11) or					
KRAS ctrl mix (12)					
Total Volume	15				

ABI 7500		
Amp-Mix	Reagent volume for 1 reaction (µl)	Reagent volume for N reactions +1 (µl)
Taq Premix 920	10	
Dye R-II	0.2	
WATER CONTROL-	3.8	
KRAS G12A mix (1) or	1	
KRAS G12D mix (2) or		
KRAS G12V mix (3) or		
KRAS G12R mix (4) or		
KRAS G12S mix (5) or		
KRAS G12C mix (6) or		
KRAS G13D mix (7) or		
KRAS A59x mix (8) or		
KRAS Q61x mix (9) or		
KRAS K117x mix (10) or		
KRAS A146x mix (11) or		
KRAS ctrl mix (12)		
Total Volume	15	

- Mix all **Amp-Mix** thoroughly by repeated pipetting or rapid vortexing, then centrifuge briefly.
- Pipette 15 µl of each Amp-Mix in all the tubes/wells previously marked.
- Add to the respective tubes/wells for each of the twelve assays:

negative control	5 µl <b>WATER</b>	CONTROL -
<u>sample</u>	5 µl DNA	
positive control	5 µl Easy KRAS pos ctrl	CONTROL +

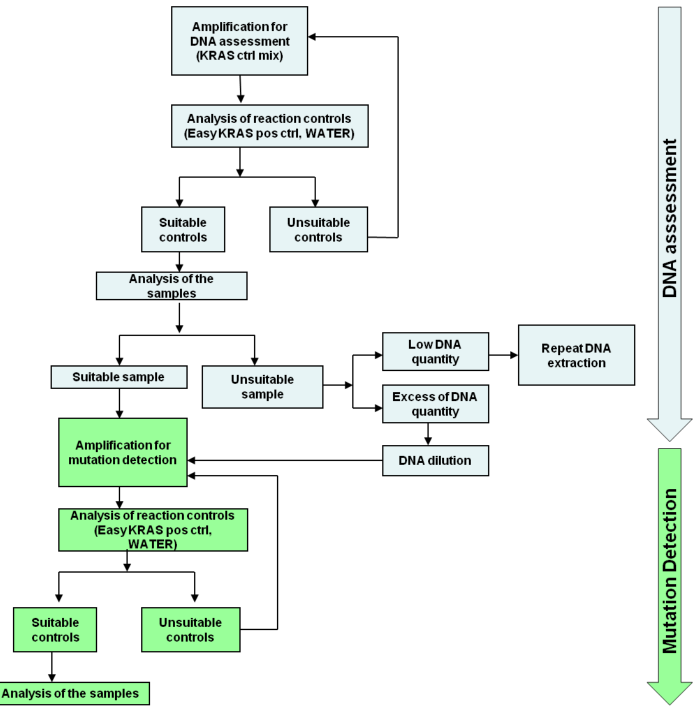
- Reaction volume: 20 µl.
- Briefly centrifuge the plate.
- Check that the thermal profile is setted up correctly and start the run.
- ③ Before starting the run, please pay attention to the plate orientation (well A1 on the upper left position) or to the Rotor-Gene 0.1ml strips of tubes orientation (mark the first tube of each strip).
- ① Proceed with the analysis following the instructions of the section "Data analysis".

#### DATA ANALYSIS

General reccomandations pertain to all the instruments

- ① Analyze first the negative control WATER and the positive control Easy KRAS pos ctrl. If they are in the range of expected values, proceed with the analysis of the samples, otherwise the session should be considered invalid and the results of the samples should be rejected.
- It is necessary to verify that the Ct values obtained are generated from a real amplification reaction (sigmoidal fluorescense curve) and not from an artifact (linear fluorescence curve), checking the normalized fluorescence graphs.
- ① Only the samples that after the DNA assessment reaction (KRAS ctrl mix) are suitable can be analyzed for the presence of mutations.

Proceed with the analysis as indicated by the following scheme:



#### Rotor-Gene

- At the end of the run, click <u>Analysis</u> and select <u>Quantitation</u>. Highlight <u>Dynamic tube</u> and <u>Slope correct</u> for all the channels.
- Set Threshold 0.04 for the Green and Yellow channels. .
- If it is not possible to perform the analysis in the Green and/or Yellow channel because of the fluorescence level is too high, see i Troubleshooting section.
  - 1. Analysis of the reaction controls

	KRAS ctrl mix	G12A, G12D, G12V, G12R, G12S, G12C, G13D, A59x, Q61x, A117x, A146x mix	All mixes	
	Green	Green	Yellow	Results
	Ct > 35	Ct > 35	Ct > 32	Proceed with analysis of the samples.
WATER	Ct ≤ 35	Ct ≤ 35	Ct ≤ 32	Possible contamination: it is not possible to analyze the samples (see Troubleshooting).
	23.5 ≤ Ct ≤ 26.5	15≤ Ct ≤ 21	19 ≤ Ct ≤ 24	Proceed with analysis of the samples.
Easy KRAS pos ctrl	Ct<23.5 or Ct >26.5	Ct<15 or Ct >21	Ct<19 or Ct >24	Possible error in the set up of the reaction/run: it is not possible to analyze the samples (see Troubleshooting).

#### 2. Analysis of KRAS ctrl mix for the DNA assessment

KRAS ctrl mix	Green	Yellow	Results
	21≤ Ct ≤ 30	18≤ Ct ≤30	Suitable sample. Proceed with the mutation analysis <sup>1</sup> .
	Ct<21	Ct <18	Excess of DNA. Samples must be diluted with <b>Water</b> (diluent) so that Ct fall in the ranges indicated above. Consider that the dilution 1:2 of the DNA increases the Ct of 1 unit.
DNA samples	Ct >30	Ct >30	<ul> <li>Suboptimal amount of starting DNA or PCR inhibition; YOU CAN NOT CONTINUE WITH THE ANALYSIS OF MUTATION:</li> <li>If it is possible, amplify a DNA volume &gt; 5μl (max 9μl) or proceed with a new DNA extraction to obtain an higher concentration of template and/or a DNA of higher quality.</li> <li>If the presence of inhibitors is suspected, dilute the sample with Water (diluent). Consider that the dilution reduces the presence of inhibitor, but also decreases the concentration of the target DNA.</li> </ul>

#### 3. Analysis of the mutation assays

- ① Only samples that following the assessment of DNA are suitable can be analyzed to search for mutations.
- Compare ΔCt values of the samples with those reported in the following table. The specified values are in the range and include extremes. The ΔCt values should be calculated with the following formula, taking care that the Ct value in Green for the mutation and the equivalent for the control assay belong to the same sample:

Assay	Amplification of Internal Control (Yellow)	ΔCt	Results	
G12A		≤ 10.6	G12A positive	
G12D		≤ 7.0	G12D positive	
G12V		≤ 9.5	G12V positive	
G12R		≤ 9.2	G12R positive	
G12S	OK	≤ 6.8	G12S positive	
G12C	(similar Ct value for all the mixes)	≤ 9.4	G12C positive	
G13D		≤ 7.0	G13D positive	
A59x		≤ 4.3	A59x positive	
Q61x		≤ 5.0	Q61x positive	
K117x		≤ 10	K117x positive	
A146x		≤ 5.0	A146x positive	
G12A		> 10.6		
G12D		> 7.0		
G12V		> 9.5		
G12R		> 9.2		
G12S	OK	> 6.8		
G12C	(similar Ct value for all the mixes)	> 9.4	Wild-type sample or beneath the LOD <sup>1</sup>	
G13D		> 7.0		
A59x		> 4.3		
Q61x		> 5.0		
K117x		> 10		
A146x		> 5.0		
G12A		١		
G12D		١		
G12V		١		
G12R		١.		
G12S		١	Foiled: not sufficient template/DCD inhibition /	
G12C	NO	1	Failed: not sufficient template/PCR inhibition / mistake during samples dispensation	
G13D		1	mistare during samples dispensation	
A59x		1		
Q61x		1		
K117x		\		
A146x		1		
<ol> <li>LOD = Limit C</li> <li>Any value</li> </ol>	Of Detection.			

If multiple assays show a  $\Delta$ Ct equal to or below the cut-off value, the signal giving the higher  $\Delta$ Ct is probably due to crossreactivity. Although double mutants have been observed, these are rare. In this case the sample should be considered positive only for the mutation with the lowest  $\Delta$ Ct.

#### Stratagene Mx3000P, Mx3005P

- At the end of the run click <u>Set-up</u>, <u>Plate Setup</u>, and enter the sample names. .
- In the section <u>Analysis, Analysis Selection/Setup</u>, deselect the function <u>Amplification-based threshold</u> and select only the reactions controls **WATER** and **Easy KRAS pos ctrl.** .
- In the section Analysis, Results, click on the lock icon in the box Threshold Fluorescence (in this way the automatically selected values cannot be modified).
- Go back to the section <u>Analysis</u>, <u>Analysis Selection/Setup</u>, and select all the samples. In the section <u>Analysis</u>, <u>Results</u>, select Text report to see the results in both channels with their respective threshold and Ct values.
- Click save.
  - 1. Analysis of reaction control

	KRAS ctrl mix	G12A, G12D, G12V, G12R, G12S, G12C, G13D, A59x, Q61x, A117x, A146x mix	All mixes	
	FAM	FAM	HEX	Results
	Ct > 35	Ct > 35	Ct > 32	Proceed with analysis of the samples.
WATER	Ct ≤ 35	Ct ≤ 35	Ct ≤ 32	Possible contamination: it is not possible to analyze the samples (see Troubleshooting).
Easy KRAS pos ctrl	23 ≤ Ct ≤ 28	15≤ Ct ≤ 22	19 ≤ Ct ≤ 25	Proceed with analysis of the samples.
	Ct<23 or Ct >28	Ct<15 or Ct >22	Ct<19 or Ct >25	Possible error in the set up of the reaction/run: it is not possible to analyze the samples (see Troubleshooting).

#### 2. Analysis of KRAS ctrl mix for the DNA assessment

KRAS ctrl mix	FAM <sup>1</sup>	HEX <sup>1</sup>	Results
	$(x-2.5) \le Ct \le (x+5.5)$	(x-2.5) ≤ Ct ≤ (x+5.5)	Suitable sample. Proceed with the mutation analysis <sup>2</sup> .
	Ct<(x-2.5)	Ct<(x-2.5)	Excess of DNA. Samples must be diluted with <b>Water</b> (diluent) so that Ct fall in the ranges indicated above. Consider that the dilution 1:2 of the DNA increases the Ct of 1 unit.
DNA samples	Ct > (x+5.5)	Ct > (x+5.5)	<ul> <li>Suboptimal amount of starting DNA or PCR inhibition; YC CAN NOT CONTINUE WITH THE ANALYSIS OF MUTATION:</li> <li>If it is possible, amplify a DNA volume &gt; 5µl (max 9µ or proceed with a new DNA extraction to obtain an higher concentration of template and/or a DNA of higher quality.</li> <li>If the presence of inhibitors is suspected, dilute the sample with Water (diluent). Consider that the diluti reduces the presence of inhibitor, but also decrease the concentration of the target DNA</li> </ul>

2. If the results of the HEX channel are not in the expected range, consider only the FAM channel results to value the sample suitability.

#### 3. Analysis of the mutation assays

- ① Only samples that following the assessment of DNA are suitable can be analyzed to search for mutations.
- Compare ΔCt values of the samples with those reported in the following table. The specified values are in the range and include extremes. The ΔCt values should be calculated with the following formula, taking care that the Ct value in FAM for the mutation and the equivalent for the control assay belong to the same sample:

Assay	Amplification of Internal Control (HEX)	ΔCt	Results
G12A		≤ 9.0	G12A positive
G12D		≤ 7.5	G12D positive
G12V		≤ 9.0	G12V positive
G12R		≤ 9.0	G12R positive
G12S	ОК	≤ 6.0	G12S positive
G12C	(similar Ct value for all the mixes)	≤ 8.4	G12C positive
G13D		≤ 6.0	G13D positive
A59x		≤ 3.0	A59x positive
Q61x		≤ 3.5	Q61x positive
K117x		≤ 6.0	K117x positive
A146x		≤ 4.5	A146x positive
G12A		> 9.0	
G12D		> 7.5	
G12V		> 9.0	
G12R		> 9.0	
G12S	OK	> 6.0	
G12C	(similar Ct value for all the mixes)	> 8.4	Wild type sample or beneath the LOD <sup>1</sup>
G13D		> 6.0	
A59x		> 3.0	
Q61x		> 3.5	
K117x		> 6.0	
A146x		> 4.5	
G12A		١	
G12D		١	
G12V		١	
G12R		١	
G12S	NO	1	Failed: not sufficient template/PCR inhibition /
G12C		1	mistake during samples dispensation
G13D		1	
A59x		1	
Q61x		1	
K117x		1	
A146x		1	
1. LOD = Limit C \ Any value	Of Detection.		

 $\Delta Ct = Ct FAM Mutation - Ct FAM KRAS ctrl mix$ 

0 If multiple assays show a  $\Delta$ Ct equal to or below the cut-off value, the signal giving the higher  $\Delta$ Ct is probably due to cross-reactivity. Although double mutants have been observed, these are rare. In this case the sample should be considered positive only for the mutation with the lowest  $\Delta$ Ct.

#### CFX96

- At the end of the run, click <u>Plate Setup View/Edit Plate</u>, select the wells used, pick the FAM and HEX fluorophores and enter samples name.
  - In the section <u>Settings</u> set up the following analysis criteria as default:
    - Cq determination mode: Single Threshold
    - Baseline settings: Baseline Subtracted Curve fit
    - Analysis Mode: Fluorophore
    - Baseline Threshold: Auto Calculated for every channel
  - In the page Quantification data, the results are displayed for both channels with the corresponding values of Cq.
- Click <u>Save</u>.
  - 1. Analysis of reaction controls

	KRAS ctrl mix	G12A, G12D, G12V, G12R, G12S, G12C, G13D, A59x, Q61x, A117x, A146x mix	All mixes	
	FAM	FAM	HEX	Results
	Cq > 35	Cq > 35	Cq > 32	Proceed with analysis of the samples.
WATER	Cq ≤ 35	Cq ≤ 35	Cq ≤ 32	Possible contamination: it is not possible to analyze the samples (see Troubleshooting).
	23 ≤ Cq ≤ 28	15≤ Cq ≤ 21	19 ≤ Cq ≤ 25	Proceed with analysis of the samples.
Easy KRAS pos ctrl	Cq <23 or Cq >28	Cq <15 or Cq >21	Cq <19 or Cq >25	Possible error in the set up of the reaction/run: it is not possible to analyze the samples (see Troubleshooting).

#### 2. Analysis of KRAS ctrl mix for the DNA assessment

DNA samples       Cq > (x+5.5)       Cq > (x+5.5)       Cq > (x+5.5)         Cq > (x+5.5)       Cq > (x+5.5)       Cq > (x+5.5)	KRAS ctrl mix	FAM <sup>1</sup>	HEX <sup>1</sup>	Results
DNA samples       Cq < (x-2.5)		(x-2.5) ≤ Cq ≤ (x+5.5)	(x-2.5) ≤ Cq ≤ (x+5.5)	Suitable sample. Proceed with the mutation analysis <sup>2</sup> .
DNA samples       CAN NOT CONTINUE WITH THE ANALYSIS OF MUTATION:         If it is possible, amplify a DNA volume > 5µl (max 9µl or proceed with a new DNA extraction to obtain an higher concentration of template and/or a DNA of higher quality.         Cq > (x+5.5)       Cq > (x+5.5)         If the presence of inhibitors is suspected, dilute the sample with Water (diluent). Consider that the dilution reduces the presence of inhibitor, but also decreases		Cq <(x-2.5)	Cq <(x-2.5)	(diluent) so that Ct fall in the ranges indicated above. Consider that the dilution 1:2 of the DNA increases the Cq
	DNA samples	Cq > (x+5.5)	Cq > (x+5.5)	<ul> <li>MUTATION:</li> <li>If it is possible, amplify a DNA volume &gt; 5µl (max 9µl) or proceed with a new DNA extraction to obtain an higher concentration of template and/or a DNA of higher quality.</li> <li>If the presence of inhibitors is suspected, dilute the sample with Water (diluent). Consider that the dilution reduces the presence of inhibitor, but also decreases</li> </ul>

2. If the results of the HEX channel are not in the expected range, consider only the FAM channel results to value the sample suitability.

#### 3. Analysis of the mutation assays

- ① Only samples that following the assessment of DNA are suitable can be analyzed to search for mutations.
- Compare ΔCq values of the samples with those reported in the following table. The specified values are in the range and include extremes. The ΔCq values should be calculated with the following formula, taking care that the Cq value in FAM for the mutation and the equivalent for the control assay belong to the same sample:

Assay	Amplification of Internal Control (HEX)	ΔCq	Results		
G12A		≤ 10.6	G12A positive		
G12D		≤ 6.5	G12D positive		
G12V		≤ 9.0	G12V positive		
G12R		≤ 9.0	G12R positive		
G12S	OK (similar Ct value for all the mixes)	≤ 5.5	G12S positive		
G12C		≤ 8.0	G12C positive		
G13D		≤ 7.5	G13D positive		
A59x		≤ 3.5	A59x positive		
Q61x		≤ 3.5	Q61x positive		
K117x		≤ 8.5	K117x positive		
A146x		≤ 3.5	A146x positive		
G12A		> 10.6			
G12D	ОК	> 6.5			
G12V		> 9.0			
G12R		> 9.0			
G12S		> 5.5	]		
G12C	(similar Ct value for all the mixes)	> 8.0	Wild type sample or beneath the LOD <sup>1</sup>		
G13D		> 7.5			
A59X		> 3.5			
Q61x		> 3.5			
K117x		> 8.5	]		
A146x		> 3.5	]		
G12A		1			
G12D		1	]		
G12V		1			
G12R		1	]		
G12S	NO	1	Failed: not sufficient template/PCR		
G12C	NO	1	inhibition / mistake during samples		
G13D		1	dispensation		
A59x		1			
Q61x		1			
K117x		1			
A146x		\			
1. LOD = Limit C \ Any value	Df Detection				

 $\Delta Cq = Cq FAM$  Mutations – Cq FAM KRAS ctrl mix

(1) If multiple assays show a  $\Delta Cq$  equal to or below the cut-off value, the signal giving the higher  $\Delta Cq$  is probably due to cross-reactivity. Although double mutants have been observed, these are rare. In this case the sample should be considered positive only for the mutation with the lowest  $\Delta Cq$ .

#### ABI 7300

- At the end of the run, click Results Plate and enter samples name. .
- In the page <u>Results Amplifcation plot</u> select all the samples and controls. .
- Select Detector All. .
- In <u>Analysis settings</u> select <u>Auto Ct</u> then <u>Analyze</u>. Select <u>Manual Ct</u>, <u>Manual baseline</u> Start cycle 3, End cycle 15 then <u>Analyze</u>.
- In the page Report Ct values are visualized for both channels. .
- Click Save. .
  - 1. Analysis of reaction controls

	KRAS ctrl mix	G12A, G12D, G12V, G12R, G12S, G12C, G13D, A59x, Q61x, A117x, A146x mix	All mixes	
	FAM	FAM	JOE	Results
	Ct > 35	Ct > 35	Ct > 32	Proceed with analysis of the samples.
WATER	Ct ≤ 35	Ct ≤ 35	Ct ≤ 32	Possible contamination: it is not possible to analyze the samples (see Troubleshooting).
	23 ≤ Ct ≤ 28	15≤ Ct ≤ 21	21 ≤ Ct ≤ 27	Proceed with analysis of the samples.
Easy KRAS pos ctrl	Ct<23 or Ct >28	Ct<15 or Ct >21	Ct<21 or Ct >27	Possible error in the set up of the reaction/run: it is not possible to analyze the samples (see Troubleshooting).

#### Analysis of KRAS ctrl mix for the DNA assessment 2.

KRAS ctrl mix	FAM <sup>1</sup>	JOE <sup>1</sup>	Results
	(x-2.5) ≤ Ct ≤ (x+5.5)	$(x-2.5) \le Ct \le (x+5.5)$	Suitable sample. Proceed with the mutation analysis <sup>2</sup> .
	Ct<(x-2.5)	Ct<(x-2.5)	Excess of DNA. Samples must be diluted with <b>Water</b> (diluent) so that Ct fall in the ranges indicated above. Consider that the dilution 1:2 of the DNA increases the Ct of 1 unit.
DNA samples	Ct > (x+5.5)	Ct > (x+5.5)	<ul> <li>Suboptimal amount of starting DNA or PCR inhibition; YOU CAN NOT CONTINUE WITH THE ANALYSIS OF MUTATION:</li> <li>If it is possible, amplify a DNA volume &gt; 5µI (max 9µI) or proceed with a new DNA extraction to obtain an higher concentration of template and/or a DNA of higher quality.</li> <li>If the presence of inhibitors is suspected, dilute the sample with Water (diluent). Consider that the dilution reduces the presence of inhibitor, but also decreases the concentration of the target DNA.</li> </ul>

If the results of the JOE channel are not in the expected range, consider only the FAM channel results to value the sample suitability. 2.

#### 3. Analysis of the mutations assays

- ① Only samples that following the assessment of DNA are suitable can be analyzed to search for mutations.
- Compare ΔCt values of the samples with those reported in the following table. The specified values are in the range and include extremes. The ΔCt values should be calculated with the following formula, taking care that the Ct value in FAM for the mutation and the equivalent for the control assay belong to the same sample:

Assay	Amplification of Internal Control (JOE)	ΔCt	Results
G12A		≤ 10.6	G12A positive
G12D		≤ 6.0	G12D positive
G12V		≤ 9.5	G12V positive
G12R		≤ 9.0	G12R positive
G12S	OK (similar Ct value for all the mixes)	≤ 6.0	G12S positive
G12C		≤ 8.0	G12C positive
G13D		≤ 6.5	G13D positive
A59x		≤ 3.5	A59x positive
Q61x		≤ 4.0	Q61x positive
K117x		≤ 9.0	K117x positive
A146x		≤ 4.0	A146x positive
G12A		> 10.6	
G12D		> 6.0	
G12V	ОК	> 9.5	
G12R		> 9.0	
G12S		> 6.0	
G12C	(similar Ct value for all the mixes)	> 8.0	Wild type sample or beneath the LOD <sup>1</sup>
G13D		> 6.5	
A59x		> 3.5	
Q61x		> 4.0	
K117x		> 9.0	
A146x		> 4.0	
G12A		١.	
G12D		\	
G12V		1	
G12R		1	
G12S	NO	1	Failed: not sufficient template/PCR
G12C	NO	١	inhibition / mistake during samples
G13D		1	dispensation
A59x		١	
Q61x		1	
K117x		1	
A146x		1	
<ol> <li>LOD = Limit C</li> <li>Any value</li> </ol>	Df Detection		

$\Delta$ Ct = Ct FAM Mutation – Ct FAM KRAS ctrl mix	

(1) If multiple assays show a  $\Delta$ Ct equal to or below the cut-off value, the signal giving the higher  $\Delta$ Ct is probably due to cross-reactivity. Although double mutants have been observed, these are rare. In this case the sample should be considered positive only for the mutation with the lowest  $\Delta$ Ct.

#### ABI 7500

- At the end of the run, click Setup Plate setup Assign Targets and Samples and enter samples name. .
- In the section Analysis Amplification plot in the page View Plate layout select all the samples and controls and omit all empty wells.
- Select Reanalyse.
- In Analysis settings, for each channel, deselect Use Default Settings, Automatic Threshold and Automatic Baseline.
- .
- Click on <u>Apply Analysis Settings</u> and then <u>Reanalyse</u>. In the page <u>View Well Table</u> Ct values are visualized for both channels. .
- Click Save.
  - 1. Analysis of reaction controls

	KRAS ctrl mix	G12A, G12D, G12V, G12R, G12S, G12C, G13D, A59x, Q61x, A117x, A146x mix	All mixes	
	FAM	FAM	JOE	Results
	Ct > 35	Ct > 35	Ct > 32	Proceed with analysis of the samples.
WATER	Ct ≤ 35	Ct ≤ 35	Ct ≤ 32	Possible contamination: it is not possible to analyze the samples (see Troubleshooting).
	23 ≤ Ct ≤ 29	14≤ Ct ≤ 20	20 ≤ Ct ≤ 26	Proceed with analysis of the samples.
Easy KRAS pos ctrl	Ct<23 o Ct >29	Ct<14 o Ct >20	Ct<20 o Ct >26	Possible error in the set up of the reaction/run: it is not possible to analyze the samples (see Troubleshooting).

#### 2. Analysis of KRAS ctrl mix for the DNA assessment

KRAS ctrl mix	FAM <sup>1</sup>	JOE <sup>1</sup>	Results
	(x-2.5) ≤ Ct ≤ (x+5.5)	$(x-2.5) \le Ct \le (x+5.5)$	Suitable sample. Proceed with the mutation analysis <sup>2</sup> .
	Ct<(x-2.5)	Ct<(x-2.5)	Excess of DNA. Samples must be diluted with <b>Water</b> (diluent) so that Ct fall in the ranges indicated above. Consider that the dilution 1:2 of the DNA increases the Ct of 1 unit.
DNA samples	Ct > (x+5.5)	Ct > (x+5.5)	<ul> <li>Suboptimal amount of starting DNA or PCR inhibition; YOU CAN NOT CONTINUE WITH THE ANALYSIS OF MUTATION:</li> <li>If it is possible, amplify a DNA volume &gt; 5µl (max 9µl) or proceed with a new DNA extraction to obtain an higher concentration of template and/or a DNA of higher quality.</li> <li>If the presence of inhibitors is suspected, dilute the sample with Water (diluent). Consider that the dilutio reduces the presence of inhibitor, but also decreases the concentration of the target DNA.</li> </ul>

If the results of the JOE channel are not in the expected range, consider only the FAM channel results to value the sample suitability. 2.

#### 3. Analysis of the mutations assays

- ① Only samples that following the assessment of DNA are suitable can be analyzed to search for mutations.
- Compare ΔCt values of the samples with those reported in the following table. The specified values are in the range and include extremes. The ΔCt values should be calculated with the following formula, taking care that the Ct value in FAM for the mutation and the equivalent for the control assay belong to the same sample:

Assay	Amplification of Internal Control (JOE)	ΔCt	Results
G12A		≤ 11.0	G12A positive
G12D		≤ 8.0	G12D positive
G12V		≤ 12.0	G12V positive
G12R		≤ 10.5	G12R positive
G12S	OK	≤ 8.5	G12S positive
G12C	(similar Ct value for all the mixes)	≤ 9.5	G12C positive
G13D		≤ 8.5	G13D positive
A59x		≤ 4.0	A59x positive
Q61x		≤ 5.8	Q61x positive
K117x		≤ 12.5	K117x positive
A146x		≤ 6.5	A146x positive
G12A		> 11.0	
G12D	OK (similar Ct value for all the mixes)	> 8.0	
G12V		> 12.0	
G12R		> 10.5	
G12S		> 8.5	
G12C		> 9.5	Wild type sample or beneath the LOD <sup>1</sup>
G13D		> 8.5	
A59x		> 4.0	
Q61x		> 5.8	
K117x		> 12.5	
A146x		> 6.5	
G12A		\	
G12D		١.	
G12V		1	
G12R		1	
G12S	NO	1	Failed: not sufficient template/PCR
G12C	NO	1	inhibition / mistake during samples
G13D		١.	dispensation
A59x		1	
Q61x		1	
K117x		١	
A146x		1	
2. LOD = Limit C	Df Detection		
Any value			

$\Delta Ct = Ct FAM I$	Mutation –	Ct FAM K	RAS	ctrl mix
	nutation –			

If multiple assays show a  $\Delta$ Ct equal to or below the cut-off value, the signal giving the higher  $\Delta$ Ct is probably due to cross-reactivity. Although double mutants have been observed, these are rare. In this case the sample should be considered positive only for the mutation with the lowest  $\Delta$ Ct.

#### TROUBLESHOOTING

Problem	Possible reason	Recommendation
Low or absent amplification signal in the channel "Green/FAM" and/or in	Incorrect selection of the fluorescence acquisition	<ul> <li>Check the fluorescence acquisition channels and repeat amplification with the settings described in this manual.</li> </ul>
the channel "Yellow/HEX" for both	channels.	amplification with the settings described in this manual.
Easy KRAS pos ctrl and samples.	Incorrect selection of the Gain on the Rotor-Gene.	<ul> <li>Repeat amplification of Rotor-Gene setting the gain properly, as described in this manual.</li> </ul>
	Incorrect setting of the thermal-profile.	<ul> <li>Check the temperature profile and repeat amplification with the settings described in this manual.</li> </ul>
	Incorrect dispensation or	Mix reagents by vortexing or inverting the tubes ten times and
	manipulation of the reagents.	<ul> <li>briefly spin before use.</li> <li>Keep reagents on ice or refrigerated blocks during the preparation of the Amp-Mix.</li> </ul>
	Reagents improperly stored or expired.	<ul> <li>Protect all mixes from light.</li> <li>Store all reagents at -35/-20°C and avoid thawing and refreezing more than twice.</li> </ul>
		<ul> <li>Do not store the mixes containing primers and probes at +2/+8°C for more than 5 hours.</li> </ul>
		<ul> <li>Do not store the Taq Premix 920 at +2/+ 8°C for more than 6 months.</li> </ul>
		Do not use expyred reagents.
No amplification signal in both "Green / FAM" and "Yellow / HEX"	Insufficient amount of starting DNA and / or presence of PCR inhibitors.	<ul> <li>Check the quantity and quality of the extracted DNA and, if appropriate, repeate the extraction faithfully following the instructions of the extraction kit.</li> </ul>
channels for all assays. <b>Easy</b> <b>KRAS pos ctrl</b> is within the expected values.	PCR Infiliations.	<ul> <li>If the extraction protocol involves the use of washing buffers containing ethanol, it is advisable to carry out a further</li> </ul>
		centrifugation prior to final elution to remove any possible trace of alcohol.
		<ul> <li>If it is assumed that the amount of starting DNA is insufficient, repeat the reaction amplifying &gt; 5ul DNA (max volume 9 ul),</li> </ul>
		reducing the corresponding volume of <b>WATER</b> in the <b>Amp-Mix</b> . Otherwise repeat the DNA extraction by reducing the volume of elution.
		<ul> <li>If you suspect the presence of inhibitors, repeat the amplification diluting sample 1:5 or 1:10 with Water (diluent).</li> </ul>
	Incorrect or no dispensation	<ul> <li>Repeat the amplification dispensing the correct volume of DNA</li> </ul>
	of the samples.	and including positive and negative controls.
Amplification signal weak or absent in "Green / FAM" and "Yellow /	Degradation of the <b>Easy</b> <b>KRAS pos ctrl</b> .	<ul> <li>Repeat the amplification testing a new aliquot of Easy KRAS pos ctrl.</li> </ul>
HEX" only for the positive control <b>Easy KRAS pos ctrl</b> .	Incorrect or failure in the dispensation of the <b>Easy KRAS pos ctrl</b> .	<ul> <li>Repeat the amplification by pipetting the appropriate volume of Easy KRAS pos ctrl.</li> </ul>
You can not perform the analysis on Rotor-Gene in Cycling A. Green and/or Cycling A. Yellow because	Rotor-Gene gain too high.	<ul> <li>Perform analysis on Rotor-Gene selecting the channel Green 2 and/or Yellow 2 and the threshold value at 0.04.The ΔCt values should be calculated with the following formula: ΔCt = Ct Green 2</li> </ul>
of a fluorescence intensity too high or out of scale for one or more than		Mutation – Ct Green 2 KRAS ctrl mix.
one assay. The positive control <b>Easy KRAS</b>	Wrong tubes identification.	<ul> <li>Repeat the amplification after marking unambiguously the reaction tubes for complex and controls</li> </ul>
<b>pos ctrl</b> shows no amplification signal in the channel "Green / FAM" for account of mutations or the signal	Incorrect dispensing of the	<ul> <li>tubes for samples and controls.</li> <li>Repeat the amplification paying attention to the dispensation of the DNA and the Facu KPAS nos attel in reaction tubes (wells).</li> </ul>
for assays of mutations or the signal is detectable only for some mixes;	samples. Incorrect samples names set-	<ul> <li>DNA and the Easy KRAS pos ctrl in reaction tubes/wells.</li> <li>Check samples names set-up.</li> </ul>
while one or more samples show an amplification signal in all assays of mutations.	up in the software.	
One sample shows HEX/Yellow Ct values different from each other for the assays.	DNA dispensation error	<ul> <li>Repeat the amplification paying attention to the dispensation of the DNA in reaction tubes/wells.</li> </ul>
The negative control <b>WATER</b> , shows an amplification signal in both	Contamination.	<ul> <li>The results should be rejected and samples must be reamplified using new reagents.</li> </ul>
FAM/Green and HEX/Yellow channels.		<ul> <li>Prepare Amp Mix in a dedicated area. Carefully decontaminate benches, pipettes and instruments.</li> </ul>
Fluorescence intensity variable.	Cutaneous fat on the reaction tube.	<ul> <li>Wear gloves.</li> </ul>
<ul> <li>If the problems persist desp contact technical support of Diate</li> </ul>	bite the implementation of the re	ecommendations given and for any other questions or problems, please
e-mail <u>support@</u>	diatechpharmacogenetics.com	
<ul> <li>telephone +39 0731 2</li> <li>fax +39 0731 2</li> </ul>		
<u> </u>	215253	

#### PERFORMANCE VALIDATION

The performance validation has been performed using all the reagents supplied with the Kit.

The experiments have been performed according to the instructions reported in this user manual on the following real-time instruments:

- CFX96 Bio-Rad (software v. 3.1)
- ABI 7300- Applied Biosystems (software v. 1.4.1)
- ABI 7500 Applied Biosystems (con software v. 2.0.5)
- Stratagene Mx3000P, Mx3005P Agilent Technologies (software v. 4.10 Build 389)
- Rotor-Gene Q Qiagen (software v. 1.7 Build 87)
- Rotor-Gene 6000 Corbett (software v. 1.7 Build 87)

#### **Clinical specificity**

In order to evaluate the Kit specificity DNA samples isolated from FFPE tumor tissue have been tested. Samples were suitable in terms of starting DNA amount and for the presence of mutations detected by the Kit, and have been already genotyped through pyrosequencing technology (kit "Anti-EGFR MoAb response<sup>®</sup> (KRAS status)", cod. UP032 Diatech Pharmacogenetics), or with Mass Spectrometry using MassArray<sup>®</sup> platform ("Myriapod<sup>®</sup> Cancer Status" cod. SQ020; "Myriapod<sup>®</sup> Colon Status" cod. SQ010, "Myriapod<sup>®</sup> Lung Status" cod. SQ011, Diatech Pharmacogenetics), or with direct sequencing. If no FFPE samples were available, Horizon Diagnostics standards, cell lines or plasmids have been tested.

	Rotor-Gene		Stratagene Mx3000P, Mx30005P		CFX96		ABI 7300		ABI 7500	
	N° samples tested	N° samples correctly genotyped	N° samples tested	N° samples correctly genotyped	N° samples tested	N° samples correctly genotyped	N° samples tested	N° samples correctly genotyped	N° samples tested	N° samples correctly genotyped
KRAS G12A	8	8/8	4	4/4	*	*	*	*	1	1/1
KRAS G12D	11	11/11	8	8/8	4	4/4	4	4/4	1	1/1
KRAS G12V	12	12/12	5	5/5	3	3/3	3	3/3	4	4/4
KRAS G12R	2	2/2	2	2/2	*	*	*	*	*	*
KRAS G12S	4	4/4	2	2/2	*	*	*	*	*	*
KRAS G12C	12	12/12	6	6/6	2	2/2	2	2/2	*	*
KRAS G13D	8	8/8	2	2/2	2	2/2	2	2/2	1	1/1
KRAS A59X	*	*	*	*	1	1/1	*	*	*	*
KRAS Q61X	9	9/9	9	9/9	6	6/6	8	8/8	1	1/1
KRAS K117X	2	2/2	2	2/2	2	2/2	2	2/2	*	*
KRAS A146X	4	4/4	4	4/4	4	4/4	4	4/4	*	*
KRAS wild-type	41	41/41	23	23/23	10	9/10 <sup>(1)</sup>	10	9/10 <sup>(1)</sup>	13	13/13
Totale	113	113/113	67	67/67	34	33/34	35	34/35	21	21/21

\* FFPE sample not available: Horizon Diagnostics standard, cell lines or plasmids have been tested.

1. Sample genotyped as wild-type with a different method, is positive for KRAS G12D on different real-time instruments. This can be due to an higher sensitivity of the real-time assay.

#### Limit of detection (LOD)

The LOD of the Kit is defined as the lowest amount of mutant DNA in a background of wild-type DNA at which a mutant sample will provide mutation-positive results in 95% of tests.

To determine the LOD, samples with different percentage of mutation and a medium input DNA concentration have been tested. For each real-time instrument (Rotor-Gene, Mx3000P/Mx3005P, ABI 7300, ABI 7500, CFX96) three independent experiments have been performed. In each experiment mutated samples have been tested in duplicates.

If available Horizon Diagnostics standards have been tested to determinate the LOD, otherwise cell lines or plasmids have been used.

The LOD of the "**Easy<sup>®</sup> KRAS**" Kit, considering all the instruments tested is:

Saggio	LOD C <sub>95</sub> a concentrazione media di DNA input
G12A	0.5-1%
G12D	1-5%
G12V	0.5%
G12R	0.5%
G12S	2-5%
G12C	0.5-1%
G13D	1-5%
A59x	5-7.5%
Q61x	5-7.5%
K117x	2-5%
A146x	5%

#### **Reproducibility**

System reproducibility (*inter-assay* variability) has been evaluated analyzing the data deriving from three independent runs with standard DNA samples. Results were reproducible in terms of genotyping for all assays and samples analyzed.

#### **Repeatability**

System repeatability (*intra-assay* variability) has been evaluated analyzing the data deriving from three independent runs with standard DNA samples. Results were reproducible in terms of genotyping for all assays and samples analyzed.

#### **Robustness**

#### Lot to lot consistency

Different batches of Taq PreMix have been tested with the same DNA samples. Results from the different lots are comparable. Two different batches of primers and probes have been tested with the same DNA samples. Results from the different lots are comparable.

## diatech pharmacogenetics

## RT001 sample grid A – Easy<sup>®</sup> KRAS

DAT	E				RUN NA	ME						
	1	2	3	4	5	6	7	8	9	10	11	12
Α												
В												
С												
D												
E												
F												
G												
н												
	G12A mix (1)	G12D mix (2	2) G12V mix (3)	G12R mix (4	4) G12S mix (5)	G12C mix (6)	G13D mix (7)	A59x mix (8)	Q61x mix (9)	K117x mix (10)	A146x mix (11)	ctrl mix (12)
INSTRUMENT n. USER MANUAL version												
	CODE LOT EXPIRY DATE											
NOT	TES											
Easy <sup>®</sup>	Easy <sup>®</sup> KRAS 30/34											

Code: RT001

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## RT001 sample grid B – Easy<sup>®</sup> KRAS

OPERATOR			SIGN	
DATE		RUN NAME		

1	•	9	•	17	•	25	•	33	•	41	•	49		57	///•///	65	(//•//)
2	•	10	•	18	•	26	•	34	•	42	•	50	•	58	•	66	•
3	•	11	•	19	•	27	•	35	•	43	•	51	•	59	•	67	•
4	•	12	•	20	•	28	•	36	•	44	•	52	•	60	•	68	•
5	•	13	•	21	•	29	•	37	•	45	•	53		61		69	
6	•	14	•	22	•	30	•	38	•	46	•	54	•	62	•	70	•
7	•	15	•	23	•	31	•	39	•	47	•	55	•	63	•	71	•
8	•	16	•	24	•	32	•	40	•	48	•	56	•	64	•	72	•

G12A mix (1	G12D mix (2)	G12V mix (3)	G12R mix (4)	G12S mix (5)	G12C mix (6)	G13D mix (7)	A59x mix (8)	Q61x mix (9)	K117x mix (10)	A146x mix (11)	ctrl mix (12)	
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INSTRUMENT n.		USER MANUAL version	
CODE	LOT	EXPIRY DATE	
NOTES			
OPERATOR		SIGN	


Diatech Pharmacogenetics srl a Socio Unico via Ignazio Silone, 1 b - 60035 Jesi AN Italy Tel +39-0731-213243 Fax +39-0731-213239 <u>info@diatechpharmacogenetics.com</u> <u>www.diatechpharmacogenetics.com</u>