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For research use only

Calcium Metabolism REAL-TIME PCR Genotyping Kit

REF

R1-H913-N3/4EU

Package: N (bulk solution)

General information

Intended use:

Calcium Metabolism REAL-TIME PCR Genotyping Kit is intended for detection and allelic discrimination of genetic polymorphisms associated with inherited risk of calcium exchange impairment.

Calcium Metabolism REAL-TIME PCR Genotyping Kit can be used in scientific research practice.

Method:

Real-time PCR followed by melting curve analysis, qualitative analysis.

Samples:

Peripheral blood.

DNA extraction:

The DNA-Technology's PREP-GS Genetics or PREP-RAPID Genetics extraction kits are recommended.

Features:

Two alleles are detected simultaneously in single tube.

PCR-mix contains an internal control (IC). IC is intended for PCR quality and sufficiency of DNA assurance. We also recommend including in assay the negative control (C-) which is not supplied but very helpful for contamination control purposes. Use deionized water or sterile buffered saline instead of sample, starting from extraction step.

Devices:

The automatic analysis for **Calcium Metabolism REAL-TIME PCR Genotyping Kit** is available on "DNA-Technology" made DTlite¹, DTprime² REAL-TIME Thermal Cyclers; the latest version of the software is available for download at https://www.dna-technology.com/software.

Time of analysis (excluding sample preparation procedure):

from 2 hours.

The number of tests:

48 (including one negative control in each run).

Kit contents:

Reagent	Organoleptic parameters	Quantity	
PCR-mix:	Transparent colorless liquid		
 VDR: 283 A>G (Bsml) PCR-buffer 	Transparent colorless liquid	960 μL 480 μL	1 tube 1 tube
Taq-AT-polymerase	Transparent colorless viscous liquid	24 μL	1 tube
Mineral oil	Transparent colorless viscous oily liquid	960 μL	1 tube

Dye label detection channels

PCR-mix	Fam	Hex	Rox	Cy5	Cy5.5
VDR: 283 A>G (Bsml)	G	Α	1	IC	-

¹ - supported by 4S1, 4S2, 5S1, 5S2, 6S1, 6S2 instruments

² - supported by 4M1, 4M3, 4M6, 5M1, 5M3, 5M6, 6M1, 6M3, 6M6 instruments

Procedure

1 PCR amplification



- 1. The quantity of DNA to be analyzed must be greater than or equal to 1.0 ng per reaction (the Cp parameter for IC must not be more than 32.0). The violation of this requirement will affect the validity of analysis and void the manufacturer guarantee.
- 2. The reagents and tubes should be kept away from direct sunlight!
- **1.1** Mark the required number of 0.2 mL PCR-tubes for polymorphism to be tested (one tube for each sample and one extra for negative control "C-").
- 1.2 Vortex the tubes containing PCR-mix for 3-5 s, then spin for 1-3 s to collect the drops.
- 1.3 Add 20 µL of PCR-mix into the marked tubes.
- 1.4 Vortex the tubes with PCR-buffer and Taq-AT-polymerase for 3-5 s, then spin for 1-3 s to collect the drops.



Taq-AT-polymerase must be stored at temperatures from minus 18 $^{\circ}$ C to minus 22 $^{\circ}$ C. Room temperature exposure is permitted only for a short time. Remove from freezer just prior to use and place on ice.

- Prepare the mixture of PCR-buffer and Taq-AT-polymerase. Add into one tube:
- 10×(N+1) µL of PCR-buffer;
 - 0.5×(N+1) μL of Taq-AT-polymerase;
 - N number of the marked tubes including "C-".

Example: For simultaneous testing of 5 samples and 1 $^{\circ}$ C- $^{\circ}$ (resulting number of marked tubes is 6), prepare mixture of PCR-buffer and Taq-AT-polymerase for 7 tubes, i.e. mix 70 μ L of PCR-buffer with 3.5 μ L of Taq-AT-polymerase.



Vortex the tube for 3-5 s, then spin for 1-3 s to collect the drops.



The mixture of PCR-buffer and Tag-AT-polymerase must be prepared just prior to use.

Add 10 µL of PCR-buffer and Tag-AT-polymerase mixture into each PCR-tube with amplification mix.



Follow the steps listed in pp 1.8 - 1.14 within two hours after addition of PCR-buffer and Taq-AT-polymerase mixture to amplification mix.

- 1.8 Add one drop (~20 µL) of mineral oil in each PCR-tube.
- 1.9 Vortex the tubes with samples and C^{-} for 3-5 s and spin down the drops on vortex mixer for 1-3 s.



- 1. In case of using **PREP-GS Genetics DNA Extraction Kit**. After vortexing centrifuge the tubes with the DNA preparation at RCF(g)16000 for one minute at room temperature (from 18 °C to 25 °C) to precipitate the sorbent. If, after isolation, the supernatant containing the isolated DNA was transferred to new tubes, centrifugation is carried out for 1-3 s in a vortex mixer.
- 2. Open the cap of the tube, add DNA sample, then close the tube before proceeding to the next tube to prevent contamination. Use filter tips. Close tubes tightly.
- **1.10** Add 5.0 μ L of the DNA sample corresponding PCR-tubes. Do not add DNA into the "C-" tubes.
- 1.11 Add 5.0 µL of negative control (C-), which passed whole DNA extraction procedure into corresponding tube.
- **1.12** Spin the tubes for 1–3 s to collect the drops.
- 1.13 Set the tubes to real-time PCR thermal cycler.
- 1.14 Launch the operating software for DT instrument¹. Add corresponding test², specify the number and ID's of the samples and negative control samples. Specify the position of the tubes in the thermal unit (see 1.13) and run PCR.



The type of the negative control tubes must be specified as "Sample".

Data collection and data analysis.

Registration and interpretation of the PCR results are operated by software and held in automatic mode.

For samples containing a sufficient quantity of DNA for correct analysis, the software defines the genotype. The samples containing an insufficient quantity of DNA (less than 1.0 ng per reaction or Cp>32.0) will be analyzed as invalid).

¹ Please, apply to Operation Manual for DTprime and DTlite Real-Time PCR instruments PART II.

² Instructions for uploading "files with test parameters" can be found on "DNA-Technology's" website https://www.dna-technology.com/assaylibrary.

Storage, shipping and handling requirements

All components of the **Calcium Metabolism REAL-TIME PCR Genotyping Kit**, except the Taq-AT-polymerase, must be stored at temperatures from 2 °C to 8 °C during the storage period. PCR-mix must be stored at temperatures from 2 °C to 8 °C and out of light during the storage period. The Taq-AT-polymerase must be stored at temperatures from minus 18 °C to minus 22 °C during the storage period.

Excessive temperature and light can be detrimental to product performance.

The kit has to be transported in thermoboxes with ice packs by all types of roofed transport at temperatures corresponding to storage conditions.

Transportation of the kit, except the Taq-AT-polymerase, is allowed in termobox with ice packs by all types of roofed transport at temperatures from 2 $^{\circ}$ C to 25 $^{\circ}$ C but no more than 5 days and should be stored at temperatures from 2 $^{\circ}$ C to 8 $^{\circ}$ C immediately on receipt.

It is allowed to transport the Taq-AT-polymerase in termobox with ice packs by all types of roofed transport at temperatures up to $25\,^{\circ}\text{C}$ but no more than $5\,\text{days}$ and should be stored at temperatures from minus $18\,^{\circ}\text{C}$ to minus $22\,^{\circ}\text{C}$ immediately on receipt.

Shelf-life - 12 months if all the conditions of transportation, storage and operation are met.

Contact our customer service department regarding quality issues with the kit:

8 800 200-75-15 (toll-free call for Russia)

+7 (495) 640-16-93 (chargeable call for CIS and foreign countries)

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Key to symbols

X	Temperature limit	i	Consult instructions for use	REF	Catalogue number
\subseteq	Use-by date	3	Manufacturer	LOT	Batch code
	Date of manufacture	Σ	Contains sufficient for <n> tests</n>	** ***	Keep away from sunlight
\triangle	Caution	NON	Non-sterile	**	

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