

## Instructions for use

Imegen® BCR-ABL1 Screening

Ref. IMG-108



Manufactured by:

HEALTH IN CODE, S.L.

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healthincode



Health in Code guarantees that its products are free from defects, both in the used materials and in its manufacturing process. This warranty is extended to the expiration date, as long as the storage conditions specified in this manual are met.

Our products are designed for in vitro diagnostics. The user of the product is responsible for validating the usefulness of the protocol proposed by Health in Code. Health in Code does not offer any other warranty, express or implied, which extend beyond the proper functioning of the components of this kit. Health in Code sole obligation in respect of the preceding guarantees, will be to replace the product or return the purchase price thereof, as desired by the customer, as long as the existence of a defect in the materials or in the manufacture of its products is identified. Health in Code will not be responsible for any damage, direct or indirect, resulting in economic losses or damages resulting from the use of this product by the purchaser or user.

All products sold by Health in Code are subjected to rigorous quality control. The Imegen® BCR-ABL1 Screening kit has passed all internal validation tests, ensuring the reliability and reproducibility of each assay.

For any questions about the applications of this product or its protocols, please contact our Technical Department:



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Imegen® is a trademark registered in Spain, which belongs to the Health in Code Group.

		Modifications to the Instructions for Use (IFU)
Version 06	NOV 2022	Change of manufacturer's address: HEALTH IN CODE, S.L., Calle de la Travesía s/n, 15E Base 5, Valencia 46024, España.
Version 05	SEP 2022	Change of manufacturer's identification from Imegen to HEALTH IN CODE, S.L.
Version 04	MAR 2021	Adaptation to the Regulation (EU) 2017/746 of the European Parliament and of the Council of 5 April 2017 on in vitro diagnostic medical devices

PAGE 2 OF 18 IMG-108 V.06 HIC-PT-KIT 03-F-03 V.01



## index

01	General information	4
02	Intended use	5
03	Technical characteristics	6
04	Safety warnings and precautions	7
05	Content and storage conditions of the kit	8
06	Necessary equipment, reagents and materials not included in the kit	9
07	Assay protocol	10
	07.1   Preparation of the PCR reagents	10
	07.2   Preparation of the PCR assay	10
	07.3   Setup of the real-time PCR program	11
80	Analysis of results	13
09	Troubleshooting	16
10	Limitations	17
	10.1   Equipment	17
	10.2   Reagents	17
	10.3   Product stability	17



## 01 General information

Translocation between BCR-ABL1 or the Philadelphia chromosome translocation is a genetic disorder that is usually present in most patients with chronic myeloid leukemia (CML) and in some patients with acute lymphoblastic leukemia (ALL).

Fusion between BCR gene, located on chromosome 22 and ABL1, located on chromosome 9, produces an oncogene (BCR-ABL1), and consequently, an abnormal protein. This abnormal protein results in an increased tyrosine kinase activity that produces the abnormal and uncontrolled growth of lymphocytes, generating leukemia.

Depending on where the breakpoint is generated within the BCR gene, different rearrangements might occur. There are three prominent forms of the chimeric oncogene BCR / ABL1:

- M-BCR-ABL1 (major region): when b3/a2 or a3 and b2/a2 or a3 are fused a chimeric tyrosine kinase of 210 kDa (p210) is generated.
- m-BCR-ABL1 (minor region): when e1/a2 or a3 are fused a chimeric tyrosine kinase of 190 kDa (p190) is generated.
- p230: when e19/a2 or a3 are fused a chimeric tyrosine kinase of 230 kDa is generated.

In addition, two other much less frequent rearrangements, which are not studied in this assay, have been described in the literature:

- e8a2 o e8a3

#### References

- > https://crm.jrc.ec.europa.eu/p/q/bcr-abl/ERM-AD623-BCR-ABL-pDNA-CALIBRANT/ERM-AD623
- > Leukemia. 2003; Volume 17: 2318-2357. doi:10.1038/sj.leu.2403135



## 02 Intended use

Imegen® BCR-ABL1 Screening employs a combination of oligonucleotides and fluorescent hydrolysis probes in a diagnostic assay by real-time PCR directed to amplify and detect the most frequent variants of the BCR-ABL1 rearrangement, including M-BCR-ABL1 and m-BCR-ABL1 and the reference gene ABL1. This genetic analysis will enable the user to detect the presence or absence of such translocations in a multiplexed reaction, but it does not distinguish between the mayor and the minor translocations as it is a qualitative analysis.

The type of sample required for this analysis is complementary DNA (cDNA). Prior to the synthesis of cDNA, total RNA must be extracted from the peripheral blood cells or a bone marrow sample, from which the retrotranscription to cDNA will take place.

Health in Code recommends to consult the local guidelines for the measurement of BCR-ABL1 transcripts for an optimal preparation of the cDNA sample required for the study and the setup of the assay.

The results obtained by this assay will orientate the clinician in the diagnostic of the type of leukaemia suffered by the patient.

Imegen® BCR-ABL1 Screening has been designed for *in vitro* diagnostics and it is directed to professionals from the molecular biology sector.

IMG-108 V.06 REVISION DATE 16.11.2022 HIC-PT-KIT 03-F-03 V.01 PAGE 5 OF 18

## 03 Technical characteristics

Imegen® BCR-ABL1 Screening has been validated using cDNA samples synthesised from the retrotranscription of total RNA extracted from peripheral blood samples of healthy patients and patients diagnosed with chronic myeloid leukemia, and it reliably detects the fusions products and the reference gene ABL1 in Section 2 of this manual (Intended Use).

The validation of Imegen® BCR-ABL1 Screening has been carried out using the following reagents not included in this kit:

☐ M-MLV RT (Moloney murine leukemia virus reverse transcriptase). Retrotranscription performed using 1 µg of total RNA.

☐ TagMan Environmental Master Mix 2.0 (ThermoFisher Scientific)

The limit of detection (LOD) for this assay has been established following the protocol specified on Section 7 (Assay protocol). Hence, the LOD of this assay has been established in 5 total copies for ABL1, M–BCR–ABL1 and m–BCR–ABL1 using a synthetic DNA vector calibrated with the *IRMM Certified Reference Material (ERM®–AD623)*.

Health in Code is certified, in accordance with the standard UNE-EN ISO 13485:2018

Medical devices – Quality management systems – Requirements for regulatory
purposes, by THE AGENCIA ESPAÑOLA DE MEDICAMENTOS Y PRODUCTOS SANITARIOS
for the Design, development and manufacturing of "in vitro" diagnostic medical devices:

- + Kits for genetic testing
- Software for bioinformatic analysis of genetic data

 IMG-108 V.06
 REVISION DATE 16.11.2022
 HIC-PT-KIT 03-F-03 V.01
 PAGE 6 OF 18



# O4 Safety warnings and precautions

- Strictly follow the instructions of this manual, especially regarding the handling and storage conditions.
- O Do not pipette by mouth.
- On not smoke, drink, or eat in areas where specimens or kit reagents are being handled.
- You must properly protect any skin condition, as well as cuts, abrasions and other skin lesions.
- Avoid discharge of reagents waste to the sink drinking water. Use waste containers established by the legislation and manage their treatment through an authorized waste manager.
- In case of an accidental release of any of the reagents, avoid contact with skin, eyes and mucous membranes and clean with abundant water
- The materials safety data sheets of all hazardous components contained in this kit are available on request to Health in Code.
- This product requires the handling of samples and materials of human origin. You should consider all human source materials as potentially infectious and handled in accordance with OSHA Biosafety Level 2 of bloodborne pathogens or must use other relevant biosafety practices for materials containing or suspect that they may contain infectious agents.
- The reagents included in this kit are not toxic, explosive, infectious, radioactive, magnetic, corrosive not biological or environmental pollutants.
- This kit has been validated using specific equipment and conditions which might vary from the conditions in other laboratories. Thus, it is recommended that each laboratory performs an internal validation prior to the utilization of the kit.
- The manufacturer is not responsible for the malfunction of the assay when one or more reagents included in the kit are replaced by other reagents not supplied by Health in Code
- The manufacturer does not guarantee the reproducibility of the assay when the user employs reagents not validated by Health in Code, considering them equivalent to those provided in the Kit.

IMG-108 V.06 REVISION DATE 16.11.2022 HIC-PT-KIT 03-F-03 V.01 PAGE 7 OF 18



## O5 Content and storage conditions of the kit

The kit contains the following reagents required to perform 48 real-time PCR reactions for each of the two targets analyzed in this assay:

- **BCR Screening Master Mix**: specific oligonucleotides and hydrolysis probe (FAM<sup>™</sup>) that simultaneously detect the M-BCR-ABL1 y m-BCR-ABL1 rearrangements.
- ► ABL1 Master Mix: Specific oligonucleotides and hydrolysis probe (FAM<sup>TM</sup>) that detect the reference gene ABL1.
- Positive Control: Positive control of a BCR-ABL1 transcript and ABL1

Reagents	Color	Vials	Storage
BCR Screening Master Mix	Red pad	2 x 24 reactions	4°C
ABL1 Master Mix	Yellow pad	2 x 24 reactions	4°C
Positive Control	Red cap	1 vial	4°C

Tabla 1. Imegen® BCR-ABL1 Screening kit contents

\* The reagents contained in this kit are lyophilized. Once the reagents are rehydrated, the vials should be stored at -20°C

 IMG-108 V.06
 REVISION DATE 16.11.2022
 HIC-PT-KIT 03-F-03 V.01
 PAGE 8 OF 18

# Equipment, reagents and materials not included in the kit

#### Equipment:

- Real-time PCR thermocycler
- Micropipette (10 μL, 20 μL and 200 μL)
- Vortex

#### Reagents:

- 2x PCR Master Mix (HotStart DNA Polymerase)
- Nuclease free water

 $\underline{\text{NOTE}}$ : In addition, this kit does not include the reagents required to perform the retrotranscription from RNA to cDNA.

#### Materials:

- Optical PCR tubes 0.2 mL
- Optical lids for the PCR tubes
- Filter tips (10 µL, 20 µL and 200 µL)
- Sterile tubes 1.5 mL
- Dust-free gloves

#### Related kits

If the results obtained for the BCR-ABL1 rearrangement is positive in the screening for any of the samples analysed, the quantification assays for the M-BCR-ABL1 transcripts, or the m-BCR-ABL1 transcripts are available as

- IMG-121 Imegen® M-BCR-ABL1
- → IMG-122 Imegen® m-BCR-ABL1

In addition, Imegen® M–BCR–ABL1 consists of a precise and sensitive assay calibrated using the IRMM Reference Material (ERM–AD623a–f) for the international standardisation of the Molecular Response (MR).

## 07 Assay protocol

#### 07.1 | Preparation of the PCR reagents

All reagents included in the kit are lyophilized. The first step before using any of our kits consists of rehydrating the reagents by adding the amount of nuclease–free water indicated in the following table. To facilitate resuspension of each component, we recommend vortexing well, spinning the tubes and allow them to rest at them at 4°C for one hour before use.

Reagents	Rehydration volume
BCR Screening Master Mix	130 µL water/vial *
ABL1 Master Mix	130 µL water/vial *
Positive Control	100 µL water/vial *

Table 2. Volume of nuclease-free water needed to rehydrate the reagents

(\*) If the reagents are not going to be used immediately after rehydration, it is recommended to store them at -20°C.

#### 07.2 | Preparation of the PCR assay

The protocol for preparation of amplification reactions is showed below:

- O1 Thaw all the reagents needed for the analysis including:
  - Master Mix BCR Screening / Master-Mix ABL1
  - Control BRC-ABL1
  - Undiluted cDNA samples
  - Nuclease-free water for the negative controls (no template controls, NTC)
  - 2x HotStart DNA Polymerase (not provided)
- O2 Vortex and spin each reagent to mix thoroughly and keep on ice.
- O3 PCR master mixes to perform the analysis of the reference gene ABL1 and the oncogenes BCR-ABL1 will be performed separately. For this, independent master mixes have to be previously prepared accordingly:
  - BCR Screening Master Mix (oncogene)

Reagents	Volume per reaction
BCR Screening Master Mix	5 μL
2x Hot Start DNA Polymerase	10 µL



#### ABL1 Master Mix (reference gene)

Reagents	Volume per reaction
ABL1 Master Mix	5 μL
2x Hot Start DNA Polymerase	10 µL

The volumes required of each mix have to be scaled up based on the number of samples that will be analyzed. In addition, extra reagents must be prepared for each mix to enable the inclusion of a negative control (NTC) and a positive control.

<u>NOTE</u>: To estimate the number of necessary reagents, we recommend making the calculations taking into account the volume of reagents needed to analyze all the samples to be included in the same PCR analysis and add 10% extra of each reagent.

- O4 Vortex the tubes containing the PCR master mixes and dispense 15 µL in each well.
- **O5** Once the master mixes have been dispensed, add the following into the corresponding wells:
  - ♦ 5 µL of the cDNA sample
  - ♦ 5 µL of the positive control
  - ♦ 5 µL of nuclease–free water (NTC)

BCR Screening Master Mix			
cDNA sample 1	Positive control		
cDNA sample 2	NTC		
cDNA sample 3			
cDNA sample 4	•		

ABL1 Master Mix			
cDNA sample 1	Positive control		
cDNA sample 2	NTC		
cDNA sample 3			
cDNA sample 4			

Figure 1. Example of the plate layout for the real-time PCR system

#### 07.3 | Setup of the real-time PCR program

The following instructions must be followed to setup the amplification program:

#### 7500 Fast or StepOne Real-Time PCR system (Thermo Scientific)

- Experiment: Quantitation Standard curve
- Ramp rate: Standard
- Reaction volume: 20 µL
- Reference ROX<sup>TM</sup>: Include
- TagMan® probes fluorophores:

Probe	Fluorophore	Quencher
BCR-ABL1	$FAM^TM$	TAMRA*
ABL1	$FAM^TM$	TAMRA*

Table 3. Hydrolysis probe information

(\*) In StepOne PCR System (ThermoFisher Scientific) this field should be filled as "None"



#### Optimal PCR program:

Fields		Step 1 atic activation	S	Step 2 PCR
No of cycles	1 initial cycle	1 initial cycle	50	O ciclos
No or cycles	Timilal Cycle		Denaturation	Annealing / Extension
Temperature	50°C	95°C	95°C	60°C
Time	2 minutes	10 minutes	15 seconds	1 minute*

Table 4. Optimal PCR program for 7500 FAST or StepOne PCR Systems

(\*) Fluorescence detection



## 08 Analysis of results

For the correct interpretation of the results, the following recommendations are given:

#### □ NEGATIVE CONTROLS

Confirm that there is not amplification in the **negative controls (NTC)**. The presence of an amplification curve is indicative of accidental contamination, and the assay should be repeated.

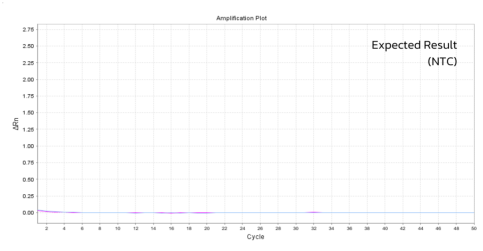


Figure 2. Expected result for the negative control (NTC)

#### → POSITIVE CONTROL

Confirm that there is amplification for ABL1 and for BCR-ABL1 in the **positive** control. If no amplification is detected in the positive control, see section 9 (Troubleshooting). The positive control should be detected between Ct16 and Ct20.

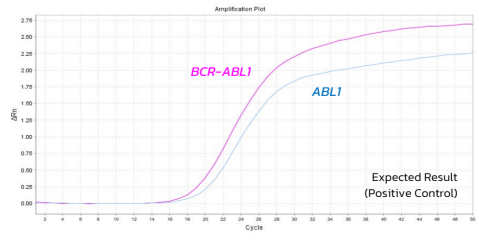


Figure 3. Expected result for the positive control. BCR-ABL1 and ABL1 transcripts are detected. The BCR Screening and the ABL1 systems are set up on different PCR reactions

IMG-108 V.06 REVISION DATE 16.11.2022 HIC-PT-KIT 03-F-03 V.01 PAGE 13 OF 18



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#### **ABL1 Master Mix**

Confirm that the reference gen (reactions prepared with ABL1 Master Mix) is detected in all the cDNA samples. ABL1 is a reference gene constitutively expressed, thus this reaction informs the user of the good quality and integrity of the cDNA sample.

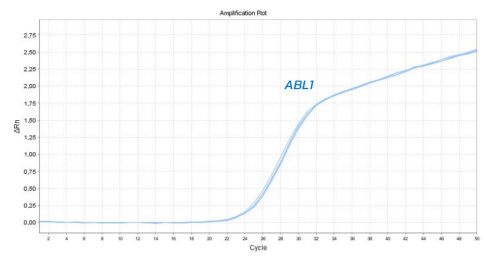


Figure 4. Expected result when a good quality cDNA sample is amplified with the ABL1 Master Mix.

#### BCR Screening Master Mix & ABL1 Master Mix

After verifying all the controls included in the analysis are correct, the cDNA samples are analysed. The sample analysed presents a BCR-ABL1 translocation if amplification is detected with the Master Mix BCR-ABL1 as indicated below.

#### ○ Negative sample:

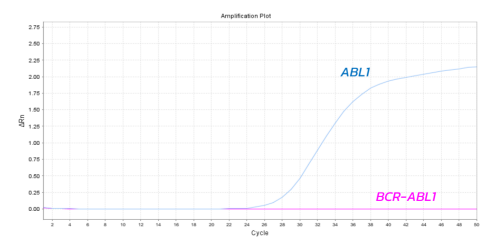


Figure 5. Expected result when a non-pathogenic cDNA sample is analysed. The ABL1 master mix detects the presence of the ABL1 wild type, but the BCR Screening master mix does not amplify the BCR-ABL1 oncogenes

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#### O Positive sample:

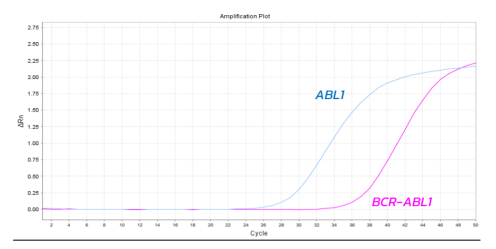


Figure 6. Expected result when a pathogenic cDNA sample is analysed. The ABL1 master mix detects the presence of the ABL1 wild type, and the BCR Screening master mix does amplify the BCR-ABL1 oncogenes



## 09 Troubleshooting

The table below represents the results that could be obtained using the positive and negative controls and the cDNA samples. In case an unexpected result is obtained, the interpretation of the result and the cause most likely reason for such result is given in the table below.

Control	BCR-ABL1	ABL1	Result / Interpretation	
Positive control	+	+	Expected result	
Positive control	_	ı	Fail in the PCR setup <sup>1</sup>	
	_	+	Expected result	
cDNA sample	+	+		
	-	-	Fail to amplify the cDNA sample <sup>2</sup>	
	-	-	Expected result	
Negative control (NTC)	+	+	Contamination with human DNA or with the positive control <sup>3</sup>	

Table 5. Interpretation of the possible results obtained using Imegen® BCR-ABL1 Screening

- (1) Fail in the PCR setup: an error in the amplification might be due to a technical issue during the configuration of the PCR setup. Check the amplification program and the setup of the fluorescence detection.
- (2) Fail to amplify the cDNA sample: an error to amplify the reference gene in the cDNA sample might suggest the quantity or the quality of the cDNA sample is compromised. In this situation, a second analysis, extraction of RNA or synthesis of a fresh cDNA samples would be recommended before an interpretation of the results is made.
- (3) Contamination with human DNA or with the positive control: PCR contamination might be caused by an inappropriate handling of the sample, the use of contaminated reagents or caused by an environmental contamination. To solve this issue, a thorough cleanse of the laboratory where the PCRs are prepared, including the equipment and material used is recommended. If necessary, use fresh aliquots of the PCR reagents and prepare last, the PCR reactions containing the positive controls to avoid any cross contamination.

IMG-108 V.06 REVISION DATE 16.11.2022 HIC-PT-KIT 03-F-03 V.01 PAGE 16 OF 18



### 10 Limitations

#### 10.1 | Equipment

Imegen® BCR-ABL1 Screening has been validated using the following real-time PCR systems:

- 7500 FAST Real-Time PCR System (ThermoFisher Scientific)
- StepOnePlus™ Real-Time PCR System (ThermoFisher Scientific)

Technically, this kit is compatible with any real-time PCR systems that enable the detection of the fluorescence emitted by  $FAM^{TM}$  fluorophore.

If a real-time PCR cycler different from the systems described in this section is going to be used for the screening of BCR-ABL1 with this kit, it is possible that the PCR program might need to be readjusted. In this case, please contact our Technical Support Team for more details.

#### 10.2 | Reagents

Imegen® BCR-ABL1 Screening has been validated using the reagents included in the kit and the DNA polymerase recommended by the supplier of the real-time PCR systems used in the validation as follows:

- M-MLV RT (Moloney murine leukemia virus reverse transcriptase). Retrotranscription performed using 1 µg of total RNA.
- TagMan Environmental Master Mix 2.0 (ThermoFisher Scientific)

If a PCR master mix (DNA polymerase) different from the DNA polymerase used in the validation is going to be used to perform the analysis, a validation with the new reagents is recommended beforehand. Please, contact our Technical Support Team if you request any further information.

In addition, this kit does not include the reagents for the RNA extraction or the retrotranscription of RNA to cDNA. It is recommended that a protocol that uses 1  $\mu$ g of RNA be used for reverse transcription.

#### 10.3 | Product stability

The optimal analytical functioning of this product is confirmed if the recommended storage conditions are applied as specified on Section 5 (Contents and Storage Conditions) from the reception of the kit until the expiry date assigned to each production batch.

For any questions about the applications of this product or its protocols, please contact our Technical Department:



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