

INSTRUCTIONS FOR USE

Imegen® Equine ID Kit

Ref. IMG-263

Manufactured by

imegenagro



Guide overview

The information in this guide is subject to change without notice.

Health in Code, S.L. guarantees that its products are free from defects, both in used materials as 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 research use only. The user of the product is responsible for validating the usefulness of the protocol proposed by Health in Code, S.L. These protocols are considered a guide only. Health in Code, S.L. does not offer any other warranty, express or implied, which extends beyond the proper functioning of the components of this set. Health in Code S.L., 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 test, or in the manufacture of its products. Health in Code, S.L. 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, S.L. are subjected to rigorous quality control (App. A). The **Imegen® Equine ID 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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NOTE: ImegenAgro® is a trademark of Health in Code, S.L.

Instructions for Use (IFU) modifications

V.06	OCT 2024	Transcription errors: In section 1.3, modification of the positive control's description; modification of table 5 title. Content revision in 2.1 and addition in section 2.3
V.05	SEP 2023	Contents review. Modification of the storage temperature of the General Master Mix
V.04	AUG 2022	Change of the manufacturer's identification, going from Imegen to Health in Code, S.L.
V.03	OCT 2020	Contents review



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Product information

01.1 General description

The identification of meat species presence in food samples is an essential step in order to verify the origin of the used and traceability of the used raw materials, as well as to evaluate the quality control for the handling and cleaning process of production lines by which it passed.

The Health in Code species ID product works by PCR amplification of a specific tag located in the mitochondrial genome of the evaluated species. The particularity of this procedure is due to the fact that the mitochondrial genome is an ideal target since it can be analysed to ensure a specific detection of the desired species and at the same time excluding the detection of other related species. Since there are multiple copies of the mitochondrial genome in each cell, the sensitivity that this detection will have is up to 100 times greater than a test that only target a specific locus in a single copy within the nuclear DNA genome.

During the last decade there had been publicly known cases related to the deceptions that consumers suffer at the time of buying meat and receive other species that are not the ones that they are paying for. EU authorities revealed the presence of uncontrolled meat in food products, and because of this, the food industry authorities have developed food safety management systems to improve the resilience of supply chain to food fraud, mostly directed to prevent the fraud opportunity. Despite the fact, food fraud does not impose a health hazard, but in some ways, they are more dangerous because the raw materials and quality control actions are unknown and untraceable.

The possibility to have a fast and accurate method to determine the authenticity of the ingredient used for food preparation is now available but the precision of the results will be something important to take in consideration at the time of evaluating the food processing. The importance of this phenomenon also lies in economic and commercial problems for both the consumer and the production company Customers want to be sure about the origin of the product they are consuming, also the concern of the contained risk for health. DNA analysis allows a valuable and conscious identification of plants and animal derivatives, by efficiently detecting contaminations or fraud related to inaccurate declaration on the label of the species constituting the food.



01.2 Intended use

Imegen® Equine ID Kit allows determining the presence of DNA of Equine (Equus Caballus) in any food samples.

Equine DNA detection is done by real time PCR using hydrolysis probes. One of them, labelled with FAM $^{\text{\tiny{M}}}$ dye, specifically detects one mitochondrial DNA sequence of Equine. The second probe is labelled with VIC $^{\text{\tiny{M}}}$ and detects an Internal Positive Control, which is used to rule out inhibitors in the sample and check the correct functioning of the assay.

01.3 Content and storage conditions of the kit

Imegen® EquineID Kit contents the necessary reagents to perform 48 reactions:

Equine Master Mix	Master Mix with specific oligonucleotides, fluorophore- labelled hydrolysis probes (FAM™ probe for Equine detection and VIC™ probe for the Internal Positive Control detection, IPC), synthetic plasmid including the specific IPC sequence and nuclease-free water.
General Master Mix	Master Mix of PCR with nucleotides, MgCl ₂ , DNA polymerase and buffer needed to carry out RT- PCR.
Positive Control	Represents 0.1% of <i>Equus caballus</i> specific-DNA.

Table 1. IMG-263 Imegen® Equine ID Kit components and description.

Reagents	Color indicator	Quantity	Conservation
Equine Master Mix*	Red pad	360 μl	-20 °C
General Master Mix*	White pad	600 μl	-20 °C upon receipt. 2 - 8 °C after initial use. Store protected from light
Equine Positive control*	Orange cap	60 μl	-20 °C

^(*) See the expiration date on the box and tubes.



01.4 Equipment, reagents and materials required but not supplied

Equipment

- Real-Time PCR Thermal Cycler with channels for detection of FAM™ (520 nm) and VIC™ (550 nm)
- Micropipettes (10 μl, 20 μl and 200 μl)
- Tabletop centrifuge with adaptors for 96 well PCR plates and/or 0.2 ml tubes
- Vortex

Materials

- Optical 96-well reaction plates or 0.2 ml optical tubes
- Optical adhesive film for 96 well plates or optical caps for 0.2 ml tubes
- \bigcirc Disposable micropipette filter tips (10 μL, 20 μL and 200 μL)
- 1.5 ml sterile tubes
- Powder-free latex gloves

Reagents

Nuclease-free water





Methods

02.1 Preparation of the amplification reactions

<u>Imegen® Equine ID Kit</u> is designed to determine, in a single PCR reaction, the presence or absence of Equine DNA and the internal positive control.

The PCR Master Mix contains:

- Equine Master Mix
- General Master Mix (2X)

The recommended protocol for preparation of amplification reactions is showed below:

- 01. Thaw the Equine Master Mix, the Positive Control vial and samples DNA.
- 02. Vortex each reagent and keep cold.
- O3. Add into a 1.5 mL tube (one for each PCR master mix preparation), the following reagents (Table 2). To estimate the amount of necessary reagents, we recommend to make calculations taking into account the number of samples to be simultaneously analyzed, and then considering one more reaction.

Table 2. Reagents and volumes needed to perform PCR reactions.

Reagents	Amount per reaction
Equine Master Mix	7.5 μL
General Master Mix	12.5 μL

- 04. Vortex and spin the 1.5 mL tube and dispense 20 μ l per well or tube of 0.2 ml.
- 05. Add 5 μ l of each DNA sample at 10 ng/ μ l, 5 μ l of Positive Control and 5 μ l of the Negative Controls* into the appropriate wells.
- **06**. Cover the well plate with optical film or the tubes with optical cover and spin in the centrifuge.



(*) We strongly recommend using an **extraction negative control** for each run of extractions carried out. This control consists in one tube to which no sample is added, and which is summited to the same extraction process as the other samples. Likewise, we recommended using a **PCR negative control** for each PCR run; this tube contains no DNA but all PCR reagents.

02.2 Settings for the Real-Time PCR program

This kit is compatible with the Real-Time PCR platforms 7500 FAST, StepOne Real-Time PCR System (Thermo Fisher Scientific) and QuantStudio™ 5 Real-Time PCR system.

Table 3. IMG-263 Imegen® Equine ID Kit probes and specifications.

Target	Receptor	Quencher
Equine DNA	FAM™	MGB
IPC	VIC™	MGB

The following instructions should be taken into account in order to setup the amplification program:

- Reaction volume: 25 μL.
- ◆ Targets: FAMTM and VICTM.
- In case the quencher has to be defined, select MGB for all probes. If the real time PCR system does not take into account the quenchers, select only the receptors (FAM™ and VIC™).
- If the Real-Time PCR system is a 7500 Fast, a StepOne Real-Time PCR system (Thermo Fisher Scientific) or a QuantStudio™ 5 Real-Time PCR system, select Quantitation - Standard curve as a type of experiment and include ROX™ as a reference.
- Ramp rate: standard
- Optimal program:

Table 4. Optimal PCR program.

Fields Step 1 Enzyme activation		Step 2 PCR	
		36 (cycles
No. of cycles	1 initial cycle	Denaturation	Primers binding/extension
Temperature	ure 95°C		60°C
Time	10 minutes	15 seconds	1 minute *

^(*) Fluorescence detection.



02.3 Analysis of results

To analyze the results, it is recommended to establish the Threshold at 0.1 and to keep the default Baseline value to minimize the residual signal in the detection channels.

Ct settings	Threshold	0.1
	Baseline	AUTO

Before analyzing the samples results, it should be checked if obtained results in controls are as expected:

- Positive control: the result must always be positive in all amplification reactions, both in the FAM™ channel as VIC™.
- Negative controls: amplification should only be detected in the VIC™ channel. In this channel an internal positive control (IPC) is detected, which determines the absence of inhibition in the sample.

IPC	It must be checked that the IPC (VIC™) is positive in all samples, with a Ct similar to the Positive Control. A negative result in the IPC indicates the presence of inhibitors in the sample. It should be noted that IPC result may be negative ir samples where a lot of Equine DNA (FAM™) is detected, because the PCR reagents are exhausted before amplification of the IPC begins.
Equine	Amplification in the FAM™ channel indicates presence of Equine DNA in the sample

It is necessary to check if sample Ct is less than the $Ct_{cut\text{-off}}$ in order to determine if one reaction of amplification is positive. Any reaction of amplification with Ct upper than $Ct_{cut\text{-off}}$ may be considered as negative. The $Ct_{cut\text{-off}}$ is equal than the positive control Ct (0.1%) plus 3.32.

Establish the positive cut-off value for the test samples and assign results:

$$Ct_{(cut-off)} = Ct_{(Positive\ Control)} + 3.32$$

Table 5. Cut-off values. (1) For fresh or minimally processed meats samples, the cut-off value corresponds to approximately 0.01% of Equine DNA, when DNA concentration is 10ng/uL.

Sample Ct value	Sample result
Ct > Ct _(cut-off)	Negative
$Ct \le Ct_{(cut-off)}$	Positive



NOTE: Any sample with a Ct equal than Ct_{cut-off} contains approximately 0.01% of *Equus Caballus* DNA.

In samples where no amplification in the FAM™ channel is seen, we can conclude that no Equine DNA is detected or that their amount in the sample is below than the detection limit.

The following table shows graphically the results that may be obtained from one sample analysis, as well as the interpretation that should be done from the obtained result:

Table 6. Results interpretation.

Equine Ma	aster Mix		
Equine (FAM™ channel)	IPC (VIC™ channel)	Interpretation	
-	+	No Equine DNA is detected	
+	+	Equine DNA is detected	
-	-	PCR inhibitors presence in the sample*	
+	-	Sample with big amount of Equine DNA	

^(*) If presence of inhibitors in the sample is detected, we recommend checking whether there has been an excess of DNA in the reaction (the recommended maximum is 250 ng). If the amount of DNA is right, we recommend repeating DNA extraction. If the problem persists, please contact our technical department.





Troubleshooting

The following table shows graphically the results that may be obtained from the analysis of different assay controls, as well as the interpretation that should be done from the obtained result:

Table 7. Possible results and their interpretation.

Control	Equine Master Mix		lutatatia
Controls	Equine	IPC	- Interpretation
Desitive control	+	+	Expected result
Positive control	-	-	¹ PCR Amplification Failure
Extraction	-	+	Expected result
Negative Control	+	+	² Contamination in the DNA extraction procedure
PCR Negative	-	+	Expected result
Control	+	+	³ PCR contamination with Equine DNA

- (1) PCR Amplification Failure: check amplification program and configuration of fluorescence capture. Amplification failure may be due to a setup technical problem.
- (2) Contamination in the DNA extraction procedure: contamination may be due to some error made in the process of sample handling, reagents contamination, or environmental contamination. Check DNA extraction protocol, wipe the laboratory where DNA extraction process was performed and take care to avoid any contamination during sample homogenization. If necessary, use new aliquots of the reagents used in DNA extraction.
- (3) PCR contaminations with Equine DNA: contamination of PCR reactions may be due to an error made in the process of sample handling, contamination of the reagents or environmental contamination. Thoroughly clean the laboratory where the PCR process was performed, as well as equipment. If necessary, use new aliquots of the reagents used in the PCR. Prepare the PCR reaction containing the Positive Control last to avoid cross contamination.





Limitations

04.1 Equipment

Imegen® Equine ID Kit has been validated using the following Real-Time PCR systems:

- 7500 FAST Real-Time PCR System (Thermo Fisher Scientific)
- StepOne™ Plus Real-Time PCR System (Thermo Fisher Scientific)
- QuantStudio5™ Real-Time PCR System (Thermo Fisher Scientific)

Technically, this kit is compatible with any Real-Time PCR systems that enable the detection of the fluorescence emitted by FAM^{TM} and VIC^{TM} fluorophores.

If a PCR system different from the systems described in this section is going to be used, it is possible that the PCR program might need to be readjusted. In this case, please contact our Technical Support Team for more details.

04.2 Reagents

<u>Imegen® Equine ID Kit</u> 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:

TagMan Environmental Master Mix 2.0 (Thermo Fisher 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.



04.3 Product Stability

The optimal analytical functioning of this product is confirmed as long as the recommended storage conditions are applied as specified on Section 01.3 (Contents and Storage Conditions) from the reception of the kit until the expiry date assigned to each production batch.





Supplemental information

AA.1 Sensitivity and specificity

The specificity of the kit was tested through comparison with the NCBI sequence database and was also experimentally tested with success on a collection of reference DNAs. See the results in the table below:

Table 8. Specificity of IMG-263 Imegen® Equine ID Kit.

Meat species	Result
Sheep (Ovis aries)	Not detected
Goat (Capra aegagrus hircus)	Not detected
Horse (Equus caballus)	Detected
Mule (Equus asinus x Equus caballus)	Detected
Donkey (Equus asinus)	Detected*
Beef (Bos taurus)	Not detected
Water buffalo (Bubalus bubalis)	Not detected
Fallow deer <i>(Dama dama)</i>	Not detected
Chicken (Gallus gallus)	Not detected
Turkey (Meleagris gallopavo)	Not detected
Duck (Genus <i>Anas</i>)	Not detected
Ostrich (Struthio camelus)	Not detected
Goose (Anser anser)	Not detected
Human (Homo sapiens)	Not detected

^(*) Donkey is detected but with less sensitivity than horse or mule.



AA.2 Detection limit

The detection limit was calculated with standard samples consisting of mixtures of raw Equine meat and other species. The Imagen® Equine ID Kit can detect blends with as little as 0.01% (w/w) of Equine meat. The limit of detection in processed samples varies depending on the composition and food processing.

To ensure the representativeness of the results, we recommend the use of a DNA extraction method that allows you to process a large amount of sample (10-20 g). If you do not have a procedure with these features, we recommend the use of Imagen® Food Extraction Kit (Ref No.: IMG-262).

AA.3 Quality certifications

- Health in Code, S.L. is certified against the standard UNE-EN ISO 9001:2015 "Quality management systems" for the design, development, manufacture, and commercialization of kits for genetic analysis.
- Health in Code, S.L. is certified against the standard UNE-EN ISO 14001:2015 "Environmental Management Systems" for the design, development, manufacture, and commercialization of kits for genetic analysis.



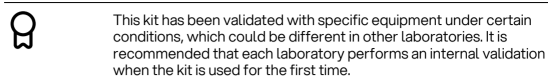


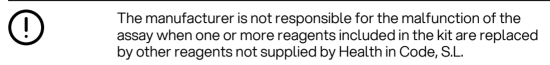
Safety warnings and precautions

<u>(1)</u>	Strictly follow the instructions of this manual, especially regarding the handling and storage conditions.
Ø	Do not pipette by mouth.
\bigcirc	Do not smoke, drink, or eat in areas where specimens or kit reagents are being handled.
ري کئ	You must properly protect any skin condition, as well as cuts, abrasions and other skin lesions.
ري	Send down the drain only those materials found on the safe list. Compounds not listed are not suitable for drain disposal. Use waste containers according to the local legislation and manage their treatment through an authorised waste manager.
ô	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, S.L.
&	This product could require the handling of samples and materials of human and animal origin. You should consider all human and animal 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.
$\overline{\Theta}$	Reagents included in this kit are non-toxic, neither explosive, infectious, radioactive, magnetic, corrosive nor environmental

polluters.







The manufacturer does not guarantee the reproducibility of the assay when the user employs reagents not validated by Health in Code, S.L., considering them equivalent to those provided in the kit.





Documentation and support

AC.1 Food safety support

Please, visit our website for the latest services, orders and support information:



imegenagro.es

Health in Code certificates of analysis and other product documentation:



portal.imegen.es/en/certificate-of-analysis/

AC.2 Customer and technical support

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



+34 963 212 340



tech.support@healthincode.com

NOTE: For SDSs for reagents and chemicals from other manufacturers, please contact the appropriate manufacturer.