

# User Guide

## nPLEX Food Total

Catalog # TAGN01001



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**I. PRODUCT DESCRIPTION AND INTENDED USE**

The nPLEX Food Total kit is a multiplex real time PCR kit intended to avoid the delivery of contaminated food products into the market. This kit detects any of the following bacteria: *Listeria monocytogenes*, *Salmonella spp.*, *Escherichia coli* and *Staphylococcus aureus*; in a single reaction from enriched food sample.

**2. KIT CONTENT, STORAGE AND STABILITY**

The kit contains 3 bags (32 reactions each). The kit was designed for 96 reactions with a final reaction volume of 20

µL each. Up to 93 samples (single sample preparation) plus both positive and negative control reactions can be analyzed per run.

- Store the kit at -20 °C through the expiration date printed on the label.
- Avoid repeated freeze-thaw cycles.
- Avoid the direct exposure of the solutions to the light.
- Once the kit is opened, store the kit components as described in the table I:

**Table I.** nPLEX Food Total content

Vial	Volumen	Descripción
Master mix 5X	140 µl	Ready-to-use “hot start” reaction mix for PCR (without primers). It contains Taq DNA Polymerase, reaction buffer, dNTP mix and fluorescent dyes. • Store at -20°C. • Refreeze max. 2 more times. • Protect from light. • Avoid repeated freezing and thawing.
Primers and probe	420 µl	Ready-to-use primers and fluorescent dyes for specific DNA amplification of <i>L. monocytogenes</i> , <i>Salmonella spp.</i> , <i>S. aureus</i> , and <i>E. coli</i> and an artificial plasmid DNA used as PCR internal control • Store at -20°C. • Refreeze max. 2 more times. • Protect from light. • Avoid repeated freezing and thawing.
Negative control	110 µl	• Nuclease-free, PCR-grade water for dilution of reaction mixtures for use as a PCR run negative control. • Store at -20°C.
Internal control	352 µl	• Stabilized solution of plasmid DNA used as a PCR run positive control • Store at -20°C • Refreeze max. 2 more times. • Avoid repeated freezing and thawing
Positive control	110 µl	• Stabilized solution of plasmid DNA used as a PCR run positive control • Store at -20°C • Refreeze max. 2 more times. • Avoid repeated freezing and thawing

**3. ADDITIONAL EQUIPMENT AND REAGENTS REQUIRED**

- Real-time PCR thermal cycler with a FAM detection channel (We recommend using Open qPCR real-time PCR instrument from ChaiBio).
- Universal enrichment broth (MultiEbroth Food, provided by TAAG Genetics).
- DNA extraction kit from enriched food samples (ZyBa, provided by TAAG Genetics).
- Sterile reaction (Eppendorf) tubes for preparing master mixes and dilutions.
- Sterile PCR tubes with optically transparent plastic caps.
- Pipettes.
- Nuclease-free, aerosol-resistant pipette tips.
- Standard benchtop microcentrifuge containing a rotor for 2.0 ml reaction tubes.
- Thermoregulated bath or dry bath.

**4. BEFORE YOU BEGIN**

In order to achieve reliable results, the entire assay procedure must be performed under nuclease free conditions. Follow the instructions below to avoid nuclease-, carry-over-, or cross-contamination:

- Please don't use the kit if the package is damaged.
- Keep kit components separate from other reagents in the laboratory.
- Use nuclease-free labware (e.g., pipettes, pipette tips, reaction vials).
- Wear gloves when performing the assay.
- To avoid cross-contamination of samples and reagents, use fresh aerosol-preventive pipette tips.
- To avoid carry-over contamination, transfer the required solutions for one experiment into a fresh tube, rather than directly pipetting from stock solutions.
- Physically separate the workplaces for DNA preparation, PCR setup, and PCR to minimize the risk of carry-over contamination. Use a PCR hood for all pipetting steps.
- In order to avoid cross-contamination, pipetting of positive controls should be the last step.
- Keep the nPLEX Food Total kit away from light and moisture.

**4.1 Enrichment**

The food enrichment must be performed as follows. 25 gr. of food sample is incubated for 24 hrs at 35°C with 225 mL of autoclaved MultiEbroth Food.

Afterwards, 1 mL of the enriched sample is transferred to a 1,5 mL tube (avoid transferring food debris or oils from the enrichment broth into the tube. For samples that have a fatty supernatant, be sure to take the necessary volume below that layer). This sample will be used for DNA extraction and PCR using the nPLEX Food total kit (TAAG Genetics). It is recommended to save another tube with 1 mL of the same enriched sample to be used as counter sample.

**4.2 DNA extraction**

DNA extraction from enriched food samples must be performed using the ZyBa kit (TAAG Genetics). For more product information please contact TAAG Genetics ([www.taag-genetics.com](http://www.taag-genetics.com)).

**5. PROTOCOL**

**5.1 PCR Program Setup**

The following procedure was optimized for a real-time PCR instrument with a FAM detection channel. Program the PCR instrument before preparing the samples and PCR mixes. Use the following real-time PCR protocol for the nPLEX Food Total kit (for details about how to program the PCR protocol, see the Instrument Operator's Manual of your real-time PCR thermal cycler):

**Table 2.** PCR Program setup

PCR Setup				
Step	Temperature	Time	Cycles	Acquisition
Denaturation	95°C	10 min	1	
Denaturation	95°C	30 seg		
Anealing	63°C	30 seg	40	FAM
Extension	68°C	30 seg		

Melting curve step				
Step	Temperature	Time	Cycles	Acquisition
Denaturation	95°C	60 seg	1	
	72°C	30 seg		
Dissociation	98°C	30 eg	1	FAM

**5.2 PCR Mix Preparation**

Before the PCR Mix preparation, the sample DNA must be mixed with the Internal control solution. Transfer 10 µl of sample DNA and add 10 µl of the internal control solution. Use 4 µl of that solution as template DNA for the PCR reactions.

**IMPORTANT:** Before using any reagent of the nPLEX Food Total kit, it is necessary to homogenize each tube on the vortex for at least 10 seg, or failing that, homogenize with micropipette, taking approximately 50% of the tube volume, and mixing carefully 10 times for pipetting. This step is a milestone to ensure confident results. The homogenization of each reagent must be performed every time the kit will be used.

Away from light, prepare a final PCR mix containing Master Mix 5x, and Primers and probes, according to table 3.

Master mix = 4\*n\*1.1

Primers and probe= 12\*n\*1.1

n: reactions number

**Table 3.** Volume for PCR mix

Reactions	Primers and probe (uL)	Master mix (uL)
1	13.2	4.4
2	26.4	8.8
3	39.6	13.2
4	52.8	17.6
5	66	22
6	79.2	26.4
7	92.4	30.8
8	105.6	35.2
9	118.8	39.6
10	132	44
11	145.2	48.4
12	158.4	52.8
13	171.6	57.2
14	184.4	61.6
15	198	66
16	211.2	70.4

It is important to homogenize the final mix in vortex for 10 seg or in their replace, homogenize with micropipette taking approximately 50% of the tube volume, and mix carefully 10 times by pipetting. After complete the preparation of PCR Mix, use this solution immediately.

Add 16 µL of the PCR Mix solution to each PCR tube, and then add 4 µL of each template DNA (DNA sample with Internal control solution).

For control samples, transfer 10 µL of Positive control and 10 µL of the Negative control, to two different PCR tubes. Mix each of them with 10 µL of Internal control. Use 4 µL of the solutions as PCR controls.

Finally, close the tubes, strips or PCR plates with optically transparent plastic caps (do not write on top of the tube caps, since it may interfere with the fluorescence reading). Bubble formation inside the PCR tube must be prevented. To eliminate possible bubbles, it is recommendable to concentrate tube contents at the bottom of the tube by performing a spin at the tubes, strips or PCR plate before placing them in the real time PCR equipment.

Transfer the PCR tubes to the real-time PCR equipment and start PCR run as described in section 5.1.

**6. DATA INTERPRETATION**

When the PCR run is over, export the PCR data to a CSV file. Users of the real-time PCR instrument Chai Open qPCR must proceed as described below:

To export the PCR data, click the menu button at the top left of the page. Select “Export” to download the PCR data. A .zip file will be generated and saved to a default directory. After extracting the ZIP file, you will find the following .csv files:

- amplification
- cq
- temperature\_log
- melt\_curve\_analysis
- melt\_curve\_data
- temperature\_log

**7. TROUBLESHOOTING**

<b>Observation</b>	<b>Possible reason</b>	<b>Recommendation</b>
<b>Negative results in positive controls or Internal control</b>	Pipetting errors or omitted reagents	Check for correct pipetting scheme and reaction setup. Repeat the PCR run.
	Insufficient vortex or centrifugation of the reagents	Always homogenize all reagent before using them
	Air bubble is trapped in the PCR tube	Always centrifuge PCR tubes before loading the PCR machine
	Outer surface of the PCR tube cap is dirty (e.g., by direct skin contact)	<ul style="list-style-type: none"> <li>• Always wear gloves when handling the PCR tubes</li> <li>• Do not write on the PCR tube cap</li> </ul>
<b>No signal in Internal control (PCR inhibition)</b>	Inappropriate storage of kit components	Keep the kit components at -20 °C, away from light. Avoid repeated freezing and thawing. Replace positive control tubes for new tubes
	Internal control solution was not mixed with sample DNA	Check for correct protocol scheme and be sure of mixing Internal control solution with sample DNA. Repeat the PCR run.
	Inhibitory effects of the sample material	<ul style="list-style-type: none"> <li>• Perform a DNA extraction starting from the counter sample, using a bacterial DNA purification kit with silica columns, resins or magnetic particles.</li> <li>• Perform a 1:10 dilution from the extracted DNA to decrease the inhibitor concentration in the reaction. The drawback of diluting the sample is that the actual target may be removed if it is present in low copy number.</li> </ul>
<b>Negative control samples are positive</b>	Inappropriate storage of kit components	Keep the kit components at -20 °C, away from light. Avoid repeated freezing and thawing. Replace positive control tubes for new tubes.
	Carry-over contamination	<ul style="list-style-type: none"> <li>• Exchange all critical solutions.</li> <li>• Repeat the complete experiment with fresh aliquots of all reagents.</li> <li>• Always handle samples, kit components and consumables in accordance with commonly accepted practices to prevent carry-over contamination.</li> <li>• Add positive controls after sample DNA and negative control tubes have been sealed.</li> <li>• Carry out decontamination procedures in the work zone, either through cleaning with sodium hypochlorite and/or with some commercial reagent to eliminate DNA fragments.</li> </ul>