



Kylt[®]



For Feed and Food Testing Applications only.

Kylt[®] *Salmonella* spp.

DNA Extraction and Real-Time PCR
Detection Kit for *Salmonella* spp.

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Kylt® *Salmonella* spp.

Kylt® *Salmonella* spp. DNA Extraction and Real-Time PCR Detection Kit

Validated according to international standards / DIN EN ISO 16140

100 reactions, for feed and food testing applications only

A. General

- This Real-Time PCR test is based on methods described in ASU L00.00-98 according to German feed and food regulations (§64 LFGB). After bacterial pre-enrichment and DNA extraction, *Salmonella* spp. is qualitatively detected via specific and real-time detected amplification of a *Salmonella* spp. target gene. This way, results can be achieved within a few hours after bacterial enrichment.
- Kylt® *Salmonella* spp. is validated according to international standards / DIN EN ISO 16140. The validated testing spectrum includes the product categories feed and food as well as pure and mixed bacterial culture material originating from the product categories feed and food.
- This kit was developed for use by trained laboratory personnel following standardized procedures.
- It is recommended to follow the general requirements of EN ISO 22174:2005 (Microbiology of food and feed – Polymerase-chain-reaction (PCR) for detection of pathogenic microorganisms in food – General requirements and terms).

B. Reagents and Materials

- Kylt® *Salmonella* spp. DNA Extraction and Real-Time PCR Detection Kit contains the following reagents:

Reagent	Color code of lid	Quantity in kit with 100 reactions	Storage conditions
DNA Extraction-Mix II	○ white	1 x 20 ml	+2 °C to +8 °C
Reaction-Mix	● yellow	4 x 500 µl	+2 °C to +8 °C
Positive Control	● red	4 x lyophilisate (final 20 µl each)	+2 °C to +8 °C lyophilized +2 °C to +8 °C rehydrated (max. 2 days) -18 °C to -20 °C rehydrated (max. 6 months)
Negative Control (Nuclease-free water)	● blue	1 x 1 ml	+2 °C to +8 °C

- All reagents are stored at +2 °C to +8 °C. Reaction-Mix has to be stored dark, do not expose to direct (sun)light! The reagents are to be used within the indicated shelf life.
- Before the first use the lyophilized Positive Control is rehydrated: 20 µl of the Negative Control (Nuclease-free water) are added per vial of Positive Control, briefly incubated at room temperature and mixed thoroughly by repeated vortexing. Storage of aliquots with 5 – 10 µl volume (depending on the expected number of Positive Control reactions per kit) at -18 °C to -20 °C is recommended.
- This kit can be used on all commercially available Real-Time PCR thermal cyclers that are able to detect the emitted fluorescence of fluorescent dyes FAM and HEX (except glas-capillary based thermal cyclers). The following Real-Time PCR thermal cyclers have e.g. been verified for routine diagnostics with this kit: Mastercycler® RealPlex2 (Eppendorf), Rotor-Gene® 3000, 6000 & Q (Corbett / QIAGEN), Chromo4™ & CFX96 Touch™ (BIO-RAD), Applied Biosystems® 7500 & 7500 Fast (Life Technologies), Mx3005P (Stratagene / Agilent Technologies), qTOWER rapidPCR & qTOWER 2.0 (Analytik Jena), LightCycler® 96 (Roche), Eco™ (Illumina®).
- We recommend to exclusively use certified Nuclease-free materials and powder free protective gloves. Pipette tips have to be changed between samples to avoid cross-contamination. Gloves have to be changed frequently, especially after spillage or other suspected contamination.
- Apart from the disposables, the following devices are needed (not included in this kit):
 - Stomacher for homogenization of samples
 - Incubators for sample pre-enrichment (+37 ± 1 °C & +41,5 ± 1 °C)
 - Table top microcentrifuge
 - Dry heating block (+100 °C ± 3 °C)
 - Vortex
 - Magnet stirrer
 - Micropipettes volume range 1 – 1000 µl
 - Centrifuge for PCR tubes or plates
 - Real-Time PCR thermal cycler

C. Control Reactions

- The Positive Control included in this kit allows for control of the specificity and efficiency of the reagents and the reaction itself. The Negative Control included in the kit allows exclusion of contaminations. The test is only valid if both, Positive and Negative Controls, are used and verified for validity in every Real-Time PCR run.
- An Internal Amplification Control is included in the Reaction-Mix in defined copy number; it is co-amplified (channel HEX) in every single reaction to detect possible inhibitory effects of the DNA-Extract and to verify true negative results.

D. Protocol (> see also „Protocol At A Glance“ at the end of this Direction For Use)

- The overall protocol of this *Salmonella* spp. analysis consists of the following steps: cultural pre-enrichment, DNA extraction, Real-Time PCR setup and amplification and final analysis.

1. Cultural Pre-Enrichment

- The sample preparation and handling should follow the rules of Good Laboratory Practice with sterile instruments to avoid external contamination. All samples are homogenized with nine times the sample volume of Buffered Peptone Water (BPW) in a Stomacher bag (as a rule e.g. 25 g or 25 ml sample material in 225 ml BPW). To enable sufficient propagation of potentially present *Salmonellae* during incubation time, the BPW has to be pre-warmed to at least room temperature prior to the start of incubation time. The pre-enrichment is incubated at $+37 \pm 1$ °C for 18 ± 2 hours without agitation with the following exceptions.
- For samples that are expected to show a very rapid onset of bacterial growth during incubation – for example directly sampled native pork[#] and poultry meat* collected during slaughter process, or neck skin samples from poultry* – the incubation time can be reduced to 15 ± 2 hours ([#]) and 13 ± 2 hours (*), respectively.
- At least 3 ml of supernatant of the pre-enrichment are transferred to a sterile test tube with sterile transfer pipettes and the Stomacher bag is discarded.
- **Attention:** Avoid shaking / agitation of the pre-enrichment; avoid to transfer solid or fatty ingredients. An aliquot should be taken directly below the surface, avoiding floating particles.
- A 1 ml aliquot is transferred from the test tube into a sterile, conical screw cap microcentrifuge tube. The remaining pre-enrichment in the test tube or Stomacher bag can be stored in the refrigerator for potential subsequent cultural examination.
- **Suggestion:** Storage in test tubes is just a recommendation, the 1 ml sample for DNA extraction may also be transferred directly from the Stomacher bag into the conical screw cap tube and the Stomacher bag may be retained in the refrigerator.
- For samples that are too difficult to separate from the debris after incubation, Stomacher bags with filters can be used for pre-enrichment in BPW.
- In case of inhibition of the Real-Time PCR, a second enrichment step is performed and the entire process of DNA extraction and Real-Time PCR is repeated (see also chapter D.5. »Analysis«). Alternatively, for samples known to have an inhibitory effect on the Real-Time PCR, the second enrichment step may directly be performed subsequently to the pre-enrichment step without Real-Time PCR-testing in between.

- For the second enrichment step the selective medium Rappaport-Vassiliadis-Soy Broth (RVS) is used. The readily incubated pre-enrichment (sample in BPW) is diluted at a ratio of 1:100 in RVS (e.g. 100 µl in 10 ml). It is incubated for 5 ± 1 hours at $+41,5 \pm 1$ °C without agitation. Alternatively or additionally, you may also use Brain-Heart-Infusion bouillon (BHI) as a non-selective media: In this case the readily incubated pre-enrichment (sample in BPW) is diluted at a ratio of 1:100 in BHI (e.g. 100 µl in 10 ml) and incubated at $+37 \pm 1$ °C for 5 ± 1 hours without agitation.

2. Colony Material from Solid Media

- In case that pure or mixed cultures on solid media are available, a small amount of colony material can be transferred to a sterile, conical screw cap tube.

3. DNA Extraction

- Pre-heat heating block to set temperature of +100 °C, the block must have an actual temperature of $+100 \pm 3$ °C at use.
- The aliquoted pre-enrichment in the conical screw cap tube is pelleted via centrifugation at 10.000 g to 12.000 g for five minutes.
- Remove the supernatant using a 1000 µl pipette tip (not by decantation) and discard it.
- The DNA Extraction-Mix II is stirred on a slow moving magnet stirrer, so that aliquots of a homogenous suspension can be taken. The pellet is resuspended by repeated up-and-down pipetting with 200 µl of DNA Extraction-Mix II. Use a 1000 µl pipette with filtered tips. Avoid formation of bubbles and aerosols.
- Screw cap tight, vortex thoroughly and incubate for 10 min to 15 min at $+100 \pm 3$ °C.
- **Optional:** Subsequently, place screw cap tube for two minutes on a massive, precooled block at -18 °C to -20 °C to induce quicker cooling; do not freeze sample! This optional step can be done with very greasy / fatty samples.
- Vortex sample thoroughly and centrifuge at 10.000 g to 12.000 g for five minutes; the supernatant is the DNA-Extract and can be used for Real-Time PCR immediately. Short-term storage (few hours) of the DNA-Extract at +2 °C to +8 °C is possible. For long-term storage of the DNA-Extract at -18 °C to -20 °C, transfer supernatant to a new (screw cap) tube. Before next use in Real-Time PCR, incubate DNA-Extracts that were stored at -18 °C to -20 °C for few minutes at $+100 \pm 3$ °C, vortex and spin down.

4. Real-Time PCR Setup and Amplification

- Reaction-Mix and Negative Control should be vortexed and spun down before each use.
- Determine the number of PCR-reactions needed: number of samples + Positive and Negative Control.
- Pipette 18 µl of Reaction-Mix to each PCR-reaction tube / each PCR-plate well (»cavity«). Keep exposure of Reaction-Mix to light as short as possible!
- Add 2 µl of Negative Control to corresponding cavity and seal.
- Add 2 µl of sample DNA-Extract to corresponding cavity and seal. Solely use clear supernatant, avoid to transfer debris.

- **Suggestion:** In case of a greasy / fatty layer on top of the DNA-Extract, punch pipette tip through the layer in the center of the surface and use only clear DNA fluid for PCR. Should debris adhere to the outside of the pipette tip, gently wipe it off the pipette on the inner surface of the screw cap tube.
- Once all sample cavities are sealed, 2 µl of the Positive Control are added to the corresponding cavity and sealed. The Positive Control is vortexed and spun down before each use.
- Avoid formation of bubbles when pipetting Reaction-Mix, samples and controls. It is recommended to always centrifuge cavities before the PCR run.
- Place cavities in Real-Time PCR thermal cycler and amplify using the following parameters:

Step	Temperature	Duration
Activation Polymerase	95 °C	10 min
Denaturation	95 °C	15 sec
Annealing & Extension	60 °C	1 min
Fluorescence Detection	channels HEX and FAM	

} 42 cycles

- Please follow the specific instructions of your Real-Time PCR thermal cycler as recommended by the manufacturer.
- These amplification parameters allow the combination of this Kylt® *Salmonella* spp. Real-Time PCR with other Kylt® Real-Time PCRs for detection of bacteria. When combining Kylt® detection tests for different pathogens, make sure all necessary channels are used!

5. Analysis

General

- The data of the amplification reactions can be automatically processed using specific software for the Real-Time PCR thermal cycler. Alternatively a threshold can be set manually, with following considerations: The threshold should be at least a ten-fold standard deviation above the average fluorescence value of cycles 3 to at least 15. The threshold should cross the HEX-curve of the Negative Control and the FAM-curve of the Positive Control, respectively, in the linear area of its slope. By setting the thresholds, the crossing points with the HEX- and FAM-curves determine the respective cycle threshold (Ct), which is negatively correlated with the initial concentration of copies of the target genes in the Real-Time PCR reaction.
- Only curves with the typical exponential amplification, meaning the curve of the raw data shows a flat baseline at the beginning, followed by a clear (exponential) slope in fluorescence and possibly reaching a plateau-phase (y-axis in logarithmic scaling), should be regarded as positive.
- For the test analysis the validity of the run is determined with the Negative and Positive Controls. After that the validity of each single sample is verified by Internal Amplification Control reaction and finally their *Salmonella* spp.-specific status is checked.

Test Evaluation

- The **test** is **only valid** if the FAM-curve of the Negative Control is negative, if the HEX-curve of the Negative Control is positive and if the FAM-curve of the Positive Control is positive. For a valid test the FAM-Ct-value of the Positive Control has to be > 15 and ≤ 35 and the HEX-Ct-value of the Negative Control has to be > 10 and ≤ 40 .

HEX-curve positive	yes	yes	no	no
FAM-curve positive	no	yes	yes	no
The sample is <i>Salmonella</i> spp.	negative	positive	positive	inhibited

- The **sample** is **negative** if its HEX-curve is positive ($10 < Ct \leq 40$), but its FAM-curve is negative.
- The **sample** is **positive** if its FAM-curve is positive ($10 \leq Ct \leq 42$), independent of the HEX-curve.
- The **sample** is **inhibited** if neither the HEX-curve nor the FAM-curve are positive.
- **Recommendation:** In case of an inhibited sample the pre-enrichment is incubated for additional 5 ± 1 hours in RVS and / or BHI (for details see chapter D.1. »Cultural Pre-Enrichment«). By that potentially present ingredients in the pre-enrichment that have inhibitory effects on the effectiveness of the Real-Time PCR are diluted. Also potentially present *Salmonellae* are selectively propagated. After the second enrichment the entire process of DNA extraction and Real-Time PCR is repeated (see chapters D.3. »DNA Extraction« and D.4. »Real-Time PCR Setup and Amplification«).
- Convenient and reliable sample data entry, Real-Time PCR start, final qualitative analysis and documentation can be conducted with the Kylt® PCR Software package, please inquire.

Production:

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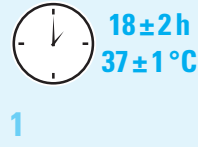
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PROTOCOL AT A GLANCE

Pre-Enrichment, DNA Extraction and Real-Time PCR

1. Pre-Enrichment of *Salmonella* spp.

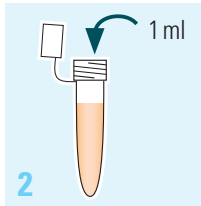
Buffered Pepton Water 18 ± 2 h* at 37 ± 1 °C



* for exceptions please refer to the Direction For Use

2. Bacterial Harvest

transfer 1 ml pre-enrichment



3. DNA Extraction

3.1 10.000-12.000g 5 min

3.2 Discard supernatant

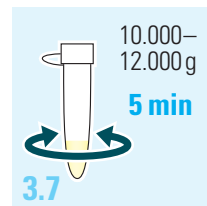
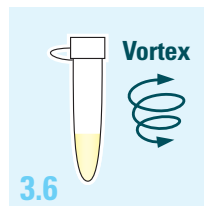
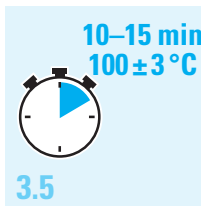
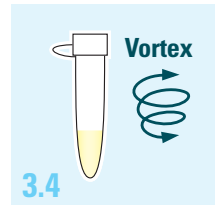
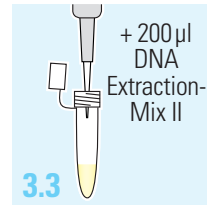
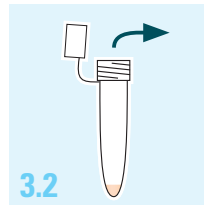
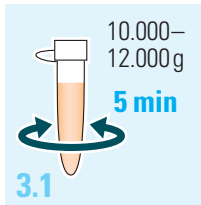
3.3 Add 200 µl DNA Extraction-Mix II

3.4 Mix by vortexing

3.5 Incubation 10–15 min 100 ± 3 °C

3.6 Mix by vortexing

3.7 10.000–12.000g 5 min

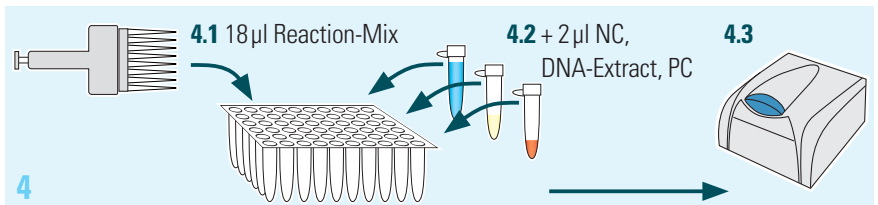


4. Real-Time PCR Setup

4.1 Mix Reaction-Mix and dispense

4.2 Add 2 µl NC, DNA-Extract, PC

4.3 Seal cavities and amplify



5. Analysis

Set threshold and analyse samples

