

Kylt[®] MS1 DIVA

Real-Time PCR Detection



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| Revision No. | Amendments | | |
|--------------|--|--|--|
| 006 | supplemental information on detected strains | | |
| 005 | Amendments in Test Evaluation, Deletion of alternative storage at +2°C to +8°C | | |
| 004 | Brown lid on Reaction-Mix (from batch 20MS1:01 on) | | |
| 003 | Layout | | |

A. General

- Kylt[®] MS1 DIVA kits are intended for the specific differentiation of bacterial DNA of *Mycoplasma synoviae* (MS) and MS live vaccine strain MS1 (Nobilis(R) MS1live). The kits are suitable for the analysis of samples from birds such as swab samples, tissues and organs and pure or mixed colony material / isolates derived from cultural processes of the aforementioned samples.
- The qualitative testing with Kylt[®] MS1 DIVA kits is based on a triplex Real-Time PCR: In one reaction setting, the target genes for MS and MS1 as well as for the exogenous control (Internal Amplification Control (IAC)) are amplified in parallel by respective primer pairs in the Polymerase Chain Reaction (PCR). Amplified target gene fragments are detected via fluorescently labeled probes during the PCR reaction in real-time (Real-Time PCR). The probes specific for detection of amplified MS and MS1 and the exogenous control target genes are labeled with fluorescent dyes Cy5, FAM and HEX, respectively, and their emitted fluorescence is separately optically measured by the Real-Time PCR thermal cycler. By means of the three individual analyses in one reaction vessel per sample and the Negative Control and Positive Control per run the MS and MS1-specific status of a sample can be evaluated in the end. This way, results can be achieved within a few hours after sample receipt.
- These kits were developed for use by trained laboratory personnel following standardized procedures. This Direction For Use must be followed strictly.

B. Reagents and Materials

• The following Kylt® MS1 DIVA kits are available and comprise the following reagents:

| Reagent | Colour of Lid | 100 Reactions Article No 31412 | 25 Reactions Article No 31413 | Store at |
|------------------|---------------|--|--|----------|
| Reaction-Mix | brown | 4 x 450 µl | 1 x 450 µl | ≤ -18 °C |
| Positive Control | ered | 4 x lyophilizate (final 50 μl each) | 2 x lyophilizate (final 50 µl each) | ≤-18 °C |
| Negative Control | blue | 1 x 1 ml | 1 x 1 ml | ≤ -18 °C |

- After receipt, the components are immediately stored at ≤ -18 °C. Avoid repeated freezing and thawing of all the reagents and keep them thawed as short as possible. If occasional processing of few samples only is expected you may prepare appropriate aliquots of reagents before storage at ≤ -18 °C. Prepare aliquots in such a way that freeze-thaw-cycles are reduced to a maximum of three. The Negative Control can alternatively be stored at +2°C to +8°C.
- The components are to be used within the indicated shelf life (see box label). The components of different batches may not be mixed.
- Before its first use, rehydrate the <u>Positive Control</u>: add 50 μ l of Negative Control per vial, briefly incubate at room temperature and mix thoroughly by repeated vortexing. It is recommended to generate aliquots of suitable volumes and store them at \leq -18 °C.
- The <u>Reaction-Mix</u> needs to be stored protected from abundant light. Do not expose to direct (sun)light.

C. Equipment and Reagents not included

- This detection method can be used on all commercially available Real-Time PCR thermal cyclers that detect the emitted fluorescence of the fluorescent dyes FAM, HEX and Cy5 (emission 520, 550 and 670 nm, respectively). Note that default normalization option against ROX (e.g. using ABI cyclers) must be deactivated.
- Apart from the disposables, the following further devices are needed and are not included in the Kylt[®] MS1 DIVA kits:
 - DNA preparation kit / protocol (e.g. Kylt® DNA Extraction-Mix II or Kylt® RNA / DNA Purification products)
 - Table top microcentrifuge
 - Vortex
 - Micropipettes covering volumes of 1 µl to 1000 µl
 - Centrifuge for PCR tubes or plates
- Accessory Kylt[®] products: see chapter F "Related and Accessory Products".
- We recommend the exclusive use of certified Nuclease-free disposables as well as powder-free protective gloves. Please wear gloves during the entire experimental procedure. Gloves need to be changed frequently, especially after spillage or suspected contaminations.

D. Control Reactions

- The <u>Positive Control</u> allows for control of the specificity and efficiency of the reagents and the reaction itself, including the performance of the Real-Time PCR and of the Real-Time PCR thermal cycler.
- The <u>Negative Control</u> allows for exclusion of contaminations. The sample testing is only valid if both, Positive and Negative Controls, are used and verified for validity in every Real-Time PCR run.
- The Internal Amplification Control is included in the Reaction-Mix in a defined copy number; it is co-amplified (channel HEX) with every single reaction to detect possible inhibitory effects of the DNA preparation on the Real-Time PCR itself and thus to verify true-negative results.
- If appropriate sampling is unsure we recommend to analyze the samples in parallel with Kylt[®] Host Cells Real-Time RT-PCR Detection for presence of amplifiable nucleic acids derived from host cell material, see chapter F "Related and Accessory Products".

E. Protocol (see also "Protocol At A Glance" at the end of this Direction For Use)

- The overall protocol of the analysis consists of the following main workflow:
 - 1. Sample Preparation
 - 2. DNA Preparation
 - 3. Reaction Setup and Amplification (Real-Time PCR)
 - 4. Data Analysis Validity and Qualitative Result
- We recommend proceeding through the protocol without interruption to avoid potential degradation of the processed samples and reagents. If necessary, you may store the final DNA preparation at ≤ -18 °C until further processing. Avoid repeated freezing and thawing of the DNA preparations.

1. Sample Preparation

- We recommend <u>pooling</u> of at most five samples or samples from five individuals, respectively, per DNA preparation.
- Pool <u>swabs</u> in a sufficient volume of sterile buffer (e.g. 1 ml of Normal Saline or 0.1 x TE), let the swabs soak for an adequate period of time and finally wash out the swabs by thorough pulse-vortexing.
- The supernatant is used for DNA preparation.
- For Kylt[®] DNA Extraction the supernatant is (fully) transferred to a conical screw cap tube (please refer to 2 "DNA Preparation").
- Small swabs may directly be immersed in Kylt® DNA Extraction-Mix II or lysis buffer, if applicable.
- Tissue and organ samples are homogenized thoroughly in sterile buffer (see above) and a suitable volume is used for the DNA preparation.
- Material derived from cultural processes, i.e. <u>colony material</u>, is directly transferred into respective tubes for Kylt[®] DNA Extraction (please refer to 2 "DNA Preparation"), such as conical screw cap tube; therefore a little amount of a single colony is picked with a sterile loop wire or sterile pipette tip and transferred to the tube.

2. DNA Preparation

a) Kylt® DNA Extraction (requires Kylt® DNA Extraction-Mix II)

For detailed information, please refer to the Direction For Use of Kylt® DNA Extraction-Mix II.

b) Kylt® RNA/DNA Purification products

- All kinds of sample matrices, including pure isolates, swabs, tissues and organs may be processed with Kylt[®] RNA/DNA Purification products (please refer to chapter F "Related Products").
- For detailed information on the DNA preparation process, please refer to the respective Direction For Use.

c) Alternative methods

- All kinds of sample matrices, including pure isolates, swabs, tissues and organs may be processed with appropriate DNA preparation kits or appropriate in-house methods.
- For detailed information on the DNA preparation process, please refer to the Direction For Use or Standard Operating Procedure of the specific kit or in-house method, respectively.

3. Reaction Setup and Amplification (Real-Time PCR)

- Before each use, briefly vortex and spin down the Reaction-Mix and Negative Control.
- To determine the total number of reactions needed, count the number of samples and add two more for the Negative Control and the Positive Control.
- The <u>Reaction-Mix</u> is ready-to-use, add 16 µl to each of the PCR tubes or plate wells ("cavities").
- Keep exposure of the Reaction-Mix to (sun)light as short as possible and return it back to appropriate storage temperature right after application. Avoid the formation of bubbles when pipetting samples and controls.
- Add 4 µl of the <u>Negative Control</u> to the corresponding cavity and seal it individually, if possible.
- Add 4 µl of each <u>DNA preparation</u> to the corresponding cavities and seal them individually, if possible.
- To minimize risk of potential cross-contaminations, 4 µl of the <u>Positive Control</u> are added to the corresponding cavity after all previous samples and control reactions are set up. Before each use, briefly vortex and spin down the rehydrated Positive Control (see also chapter B "Reagents and Materials").
- If not already done, finally seal the cavities. It is recommended to briefly spin them down before the start of the Real-Time PCR run.
- Place the cavities in the Real-Time PCR thermal cycler and run the test with Kylt[®] Profile II as given below.

| Kylt® Profile II | | | | |
|------------------|--------------------------|---------------------------|----------|-----------|
| Step No | Description | Temperature | Duration | |
| 1 | Activation of Polymerase | 95 °C | 10 min | |
| 2 | Denaturation | 95 °C | 15 sec |) |
| 3 | Annealing & Extension | 60 °C | 1 min | 42 cycles |
| 4 | Fluorescence Detection | channels FAM, Cy5 and HEX | | J |

• Kylt® Profile II allows for combined run of this and most other Kylt® qPCR detection methods.

Alternatively, the Kylt[®] Profile I given below can be applied. Kylt[®] Profile I allows for combined run of this and most other Kylt[®] qPCR detection methods as well as Kylt[®] RT-qPCR detection products that need Reverse Transcription, such as those for detection of viral RNA.

| Kylt [®] Profile I | | | | |
|-----------------------------|--------------------------|---------------------------|----------|-----------|
| Step No | Description | Temperature | Duration | |
| 1 | Reverse Transcription | 50 °C | 10 min | |
| 2 | Activation of Polymerase | 95 °C | 1 min | |
| 3 | Denaturation | 95 °C | 10 sec |) |
| 4 | Annealing & Extension | 60 °C | 1 min | 42 cycles |
| 5 | Fluorescence Detection | channels FAM, Cy5 and HEX | | |

In the event of a combined Real-Time (RT-)PCR run, make sure all necessary channels are detected.

Please follow the specified instructions of your Real-Time PCR thermal cycler as recommended by the manufacturer.

<u>4. Data Analysis – Validity and Qualitative Result</u>

General

- The amplification data can be processed automatically using the specific software tool of your Real-Time PCR thermal cycler. Alternatively, the threshold can be set manually considering the following directions: The threshold should cross the FAM-curve, the Cy5-curve and the HEX-curve in the linear increase of their slope (linear scaling of the y-axis). By setting the threshold, the crossing points with the HEX-, Cy5- 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 log scaling), should be regarded as positive.
- The actual test analysis starts with the validity check of the entire Real-Time PCR run. Afterwards, by means of the Internal Control the validity of each sample reaction and its true test result can be verified according to the Ct-value of the Internal Control channel (HEX). Finally, the MS and MS1-specific status of each sample is analyzed (Cy5 and FAM).

Test Evaluation

■ The Real-Time PCR test run is only valid if the FAM- and Cy5-curve of the Negative Control are negative, the HEX-curve of the Negative Control is positive and the FAM-, Cy5- and HEX-curve of the Positive Control are positive. For a valid test the FAM- and Cy5-Ct-value of the Positive Control has to be > 15 and ≤ 35 and the HEX-Ct-value of the Negative Control has to be ≤ 40.

| Target | Channel | Signal | | | |
|-------------------|---------|----------|------------------------|------------------------|-----------|
| Internal Control | HEX | positive | positive / negative | positive / negative | negative |
| MS | Cy5 | negative | positive | positive | negative |
| MS1 | FAM | negative | negative | positive | negative |
| The sample is MS | | negative | positive | positive | inhibited |
| The sample is MS1 | | negative | negative | positive | minoiteu |

■ A sample is negative for MS and MS1 if its HEX-curve is positive (Ct ≤ 40), but its Cy5- and FAM-curves are negative.

- A sample is positive for MS if its Cy5-curve is positive (Ct ≤ 42), independent of the HEX-curve.
- A sample is positive for MS and MS1 if its Cy5- and FAM-curves are positive (Ct ≤ 42), independent of the HEX-curve. Note that strain MS ATCC 25204 (also known as MS WVU 1835 or NCTC 10124), from which the MS1 strain was derived from, and potentially derivatives of it may be detected as MS1.
- A sample is inhibited if neither the Cy5- and FAM-curves nor the HEX-curve are positive.
- Recommendation: In the case of an inhibited sample the test may be repeated with a dilution of the DNA preparation at e.g. 1:10 (9 volumes Negative Control + 1 volume DNA Extract or eluted DNA). The Negative Control is used as the diluting agent. Preferably, the entire DNA preparation process is repeated: in case of inhibited DNA Extracts derived from Kylt[®] DNA Extraction-Mix II, the original sample or the DNA Extract can be utilized for DNA preparation using appropriate alternative systems, such as Kylt[®] RNA/DNA Purification.
- Convenient and reliable sample data entry, Real-Time PCR start, final qualitative analysis and documentation can be conducted with the Kylt[®] Software, please inquire.

F. Related and Accessory Products

| Product | Article No | Reactions | Description |
|----------------------------------|---------------|-----------|---|
| Kylt® DNA Extraction-Mix II | 31398 | 100 | Simplified and economic DNA extraction |
| Kylt® RNA / DNA Purification | 31314/31315 | 250 / 50 | Combined RNA and DNA purification from veterinary samples |
| Kylt® RNA / DNA Purification HTP | 31826 | 4x96 | Combined, magnetic beads-based purification of RNA and DNA from veterinary samples, suitable for automated high throughput processing |
| Kylt® Host Cells | 31106 / 31107 | 100 / 25 | Kit to detect animal host cells; to verify sample taking process |
| Kylt® Purifier | 31436 | | Purification system for magnetic beads. Up to 96 samples in under 30 minutes. |

G. Ordering information

For a fast and efficient service please send your order to orders@kylt.eu and please provide the following information:

- Delivery address
- Invoice address
- Purchaser contact telephone number
- End user name and telephone number (if different)
- Purchase order number
- Product name and cataloge number
- Quantity and size of products
- Indicate if your account is VAT exempt

Production:

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Development, manufacturing and distribution of Kylt[®] *In-Vitro* Diagnostica is certified according to ISO 9001:2015.



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PROTOCOL AT A GLANCE Real-Time PCR Setup



