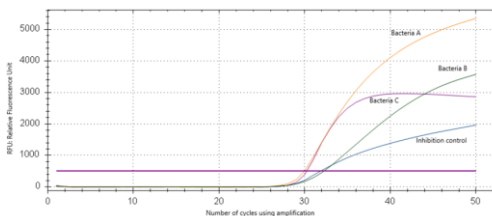


Mixed AF sample containing 3 bacteria (A, B, C) and inhibition control confirming the functionality of one of the 3 rt-PCR panels for MIAC detection



Technology owner

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Inventor (s)

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IPR status

Know-how licensing

Stage of Development

Proof of concept

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Bacterial Detection in Amniotic Fluid

Background

Preterm Premature Rupture of Membranes (PPROM) is a **pregnancy complication**. In this condition, the sac (amniotic membrane) surrounding the fetus breaks (ruptures) before week 37 of pregnancy. Once the sac breaks, pregnant woman has increased risk for infection. PPRM complicates 3 – 4% of all pregnancies and is up to **1/3 complicated by microbial invasion of the amniotic cavity (MIAC)** leading to infection in amniotic fluid (AF) and development of intra-amniotic inflammation. Although this complication is usually asymptomatic, it is a major cause of preterm birth and neonatal morbidity and mortality worldwide. Neonates from these pregnancies are at **increased risk of developing neonatal sepsis and impaired psychomotor development** and other sometimes lifelong health consequences. Currently, determining the MIAC in patients with PPRM is very time-consuming and technically demanding. Main disadvantage is that results are available in days, which is already clinically irrelevant for the initiation of targeted antibiotic treatment.

Description of the Invention

We have developed **multiplex Real Time - PCR (Real - Time Polymerase Chain Reaction) assay for simultaneous detection of the most common targeted pathogens in AF** collected by amniocentesis. The method will **reduce the time required to diagnose specific pathogen and help clinicians make quick and accurate treatment decision**. The assay consists of 3 panels for DNA detection of 8 most common microorganisms. Each panel contains specific sets of primers and probes. Assay results are determined by PCR fluorescence signal. Panels were validated retrospectively on 20 samples of AF of PPRM patients and have reached 90 % sensitivity so far. Further prospective validation study is currently underway.

Advantages

High sensitivity of the method - panel of selected microorganisms in multiplex assay was defined on the basis of research in nearly 700 patients with PPRM and it covers 88 % of microorganisms responsible for MIAC in PPRM patients. The multiplex approach **determines the selected microbes in several hours after sampling** in semi-quantitative way compare to the current approach (combination of specific multiplex Real Time - PCR for 3microorganisms in combination with PCR assay targeting 16S rRNA regions followed by Sanger sequencing for the rest of targeted microbes and cultivation techniques). **Personalized approach** to the clinical management and therapeutic intervention of the patients with PPRM based on the assay results. The technology **does not require specialized laboratory equipment** or specialized personnel and thus is suitable for molecular laboratory of perinatology centres equipped with standard Real-Time PCR cyclers.

Potential Applications

IVD test to be used by medical facilities that take care of pregnant women esp. regional hospitals or perinatology centers.