Combining an Acoustic Trap with Attenuated Total Reflection Infrared Spectroscopy for Detection of Bacteria in Drinking Water

Stephan Freitag, Bettina Baumgartner, Andreas Schweighofer, Bernhard Lendl

Institute of Chemical Technologies and Analytics, Technische Universität Wien, Getreidemarkt 9/164, A-1060 Vienna, Austria
stephan.freitag@tuwien.ac.at

Introduction

• In this work, we present our efforts towards the combination of an acoustic trap [1,2] and attenuated total reflection (ATR) Fourier-transform infrared (FTIR) spectroscopy [3] to perform an ultrasound (US) enhanced assay for the rapid detection of bacteria in drinking water.
• By mounting an acoustic trap on top of a custom-built ATR setup, we were able to trap bacteria by relying on so-called ultrasonic radiation forces without the need of mechanical retention elements.
• To showcase the potential of the presented setup for sensing microbial pollution in water, we monitored *Escherichia coli* suspensions at different concentrations.
• Experiments were performed by pre-incubating bacteria with antibodies on a tube rotator. The bacteria-antibody suspension was then injected to the acoustic trap, where the bacteria were retained using ultrasound radiation forces. In the end, enzyme substrate was pumped into the cell and the conversion via the antibody was monitored by ATR-FTIR spectroscopy.
• Throughout the whole liquid-handling sequence, bacteria were stably retained in the cell. In contrast to common direct enzyme-linked immunosorbent assays (ELISA), no immobilization of *E. coli* on a surface is needed.

Ultrasound Enhanced Assay

**A** the top view of the assembled acoustic trap, (B) the bottom side and (C) trapped beads, encircled in red.
• The acoustic trap is made of aluminum. The liquid compartment has a volume of approximately 20 µL and a height of 500 µm. Above the sample compartment, a 8 mm piezo disc (US-source) is directly glued to the aluminum body.
• The assembled acoustic trap was mounted on top of a custom built ATR fixture, holding a multibounce zinc sulfide ATR element (17 x 10 x 1 mm³, 45°).
• The acoustic trap was operated using a sonic amp (USEPAT, Wien, Austria) US driver, set to a US frequency of 2.20 MHz and a gain of 65%. Prior to liquid handling, the US was turned on for 30 min to allow the system to thermally stabilize.

**D** Schematic of the acoustic trap on top of the multibounce ATR element, **E** Principle of the assay performed with acoustically trapped bacteria.
• TRIS buffered saline pH 7.6 containing 0.05% Tween 20 was used as carrier. Antibody was diluted in the same buffer containing 1% non-fat dry milk.
• 50 mM p-nitrophenylphosphate in 2 M diethanolamine buffer pH 9.8 was used as substrate.
• Antibody labelled bacteria were trapped and retained in the cell due to acoustic radiation forces.

Conclusions & Outlook

• We introduced the combination of an acoustic trap with ATR-FTIR spectroscopy, for monitoring bacterial load in water.
• The overall assay time could be reduced to 60 min and different bacteria concentrations could be distinguished.
• The presented results pave the way for ultrasound enhanced assays.
• Future efforts will focus on optimizing the overall assay procedure to measure lower concentrations of bacteria.

Experimental Setup

**FTIR spectrometer**
- ATR element
- ATR beam
- IR beam
- **camera**
- **waste**
- **syringe pump**
- **holding coil**
- **selection valve**

• ATR measurements were performed by guiding the IR beam of a FTIR spectrometer (Vertex 70, Bruker Optics) through a custom-made multibounce ATR setup with the acoustic trap on top.
• Automated liquid handling is performed via a sequential injection analysis manifold consisting of a selection valve and a syringe pump.

Results

**Schematic of the assay sequence performed with the sequential injection manifold (F).**
• Bacteria and antibody were pre-incubated for 1 h on a tube-rotator.
• Antibody-labelled bacteria suspension was aspirated and returned to the reservoir four times via the syringe pump (blue) to prevent inhomogeneity in the suspension.
• 60 µL of the bacteria suspension was injected into the acoustic trap (blue).
• In the next step, 300 µL carrier solution was injected into the cell (yellow) to remove excess antibody.
• In the end, enzyme substrate was pumped into the acoustic trap and the enzymatic conversion was monitored by recording 30 consecutive mid-IR spectra.
• After successfully monitoring the reaction, the US was switched off and the acoustic trap was rinsed thoroughly with carrier solution.
• Combining the acoustic trap with a dedicated fully-automated liquid handling procedure lead to an overall assay time of approximately 3 hours.

**G** Mid-IR band of the antibody induced enzymatic conversion by different bacteria suspensions originating from the NO₂ moiety that shifts upon cleavage of the enzyme. (H) band height used as analytical signal vs bacteria concentration.