Optomagnetic biosensor system for DNA and bacteria detection based on rolling circle amplification and immunomagnetic strategies

Tian, Bo; de la Torre, Teresa Zardán Gómez; Donolato, Marco; Hansen, Mikkel Foug; Svedlindh, Peter; Strömberg, Mattias

Publication date:
2016

Document Version
Peer reviewed version

Citation (APA):
Title: **Optomagnetic biosensor system for DNA and bacteria detection based on rolling circle amplification and immunomagnetic strategies**

Authors & affiliations:

Bo Tian,† Teresa Zardán Gómez de la Torre,† Marco Donolato,§ Mikkel Foug Hansen,‡ Peter Svedlindh,† Mattias Strömberg‡

† Department of Engineering Sciences, Division of Solid State Physics, Uppsala University, The Ångström Laboratory, Uppsala University, Box 534, SE-751 21 Uppsala, Sweden

‡ Department of Engineering Sciences, Division of Nanotechnology and Functional Materials, Uppsala University, The Ångström Laboratory, Box 534, SE-751 21 Uppsala, Sweden

§ Department of Micro- and Nanotechnology, Technical University of Denmark, DTU Nanotech, Building 345 East, DK-2800 Kongens Lyngby, Denmark

Abstract: (Your abstract must use **Normal style** and must fit in this box. Your abstract should be no longer than 300 words. The box will "expand" over 2 pages as you add text/diagrams into it.)

Preparation of Your Abstract

1. The title should be as brief as possible but long enough to indicate clearly the nature of the study. Capitalise the first letter of the first word ONLY (place names excluded). No full stop at the end.

2. Abstracts should state briefly and clearly the purpose, methods, results and conclusions of the work.

Introduction: Clearly state the purpose of the abstract.

Methods: Describe your selection of observations or experimental subjects clearly.

Results: Present your results in a logical sequence in text, tables and illustrations.

Discussion: Emphasize new and important aspects of the study and conclusions that are drawn from them.
Important notes:

Do NOT write outside the grey boxes. Any text or images outside the boxes will be deleted.

Do NOT alter the structure of this form. Simply enter your information into the boxes. The form will be automatically processed – if you alter its structure your submission will not be processed correctly.

Do not include keywords – you can add them when you submit the abstract online.

Benefiting from their rapid read-out, highly flexible devices and low-cost portable systems, optomagnetic biosensors have drawn increased attention in recent years as bioassay technologies for small molecules, DNA, and bacteria.

Herein, optomagnetic bioassay strategies are presented utilizing the binding of functionalized magnetic nanoparticles, with Brownian relaxation behaviour, to detect DNA molecules and bacteria. Presence of target changes the dynamic behaviour of the magnetic nanoparticles and thus the optomagnetic response of the sample, which is measured by a 405 nm laser-based optomagnetic setup. The output signal used for target quantification is the in-phase, \( V'_2 \), and the out-of-phase, \( V_2^* \), components of the complex second harmonic signal \( V_2 = V'_2 + iV_2^* \) of the transmitted light.

A turn-on competitive *Salmonella* immunoassay using two differently sized magnetic particles (micron-sized particles acting as capture particles and nano-sized particles acting as detection particles) is presented resulting in a limit of detection of \( 8 \times 10^4 \) CFU/mL (20 times lower than of volumetric magnetic stray field detection device based immunoassays) and a total assay time of 3 h. The improvement of sensitivity is enabled by the formation of immuno-magnetic aggregates providing steric hindrance protecting the interior binding sites from interaction with the magnetic nanoparticle labels. Additionally, a qualitative and homogeneous biplex immunoassay for *E. coli* and *S. typhimurium* is demonstrated.

Using 250 nm “ghost” magnetic nanoparticles, which widen the linear detection range of 100 nm magnetic nanoparticles, the performance of the optomagnetic sensor system for the quantitative analysis of target DNA sequences (through binding of DNA-tagged magnetic nanoparticles to rolling circle amplification products) and bacteria (direct immunoassay protocol) is improved by a factor of 15 compared with the classical approach excluding ghost nanoparticles.

The optomagnetic read-out platform is considered to be a potential candidate for low-cost and easy-to-operate biosensor devices for food safety and veterinary medicine applications.