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Introduction

About 50% of the population will suffer a urinary tract infection (UTI) at least once during their lifetime.¹ Complications such as urinary stones, indwelling catheters and surgeries considerably increase the risk of urosepsis, which has a high mortality (20%).¹ Furthermore, UTIs represent a considerable healthcare expenditure, costing about 2.8 billion USD in 2011 in the US alone.² The vast majority of UTIs originate from pathogens in the Enterobacteriaceae family, and in more than 80% of cases are caused by E. coli.3 Urine culture, which identifies the bacterium responsible for the infection as well as the pathogen load, is the gold standard method to diagnose symptomatic UTIs. Bacteriological culture can take several days to be completed, and therefore cannot be used to diagnose UTIs rapidly. Urine strips or dipsticks, which detect nitrite-producing bacteria as well as leukocytes (pyuria), are commonly used to aid in the rapid diagnosis of UTIs. However, the

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Mobile origami immunosensors for the rapid detection of urinary tract infections[†]

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Urinary tract infections (UTI) have a high prevalence and can yield poor patient outcomes if they progress to urosepsis. Here we introduce mobile origami biosensors that detect UTIs caused by *E. coli* at the bedside in less than 7 minutes. The origami biosensors are made of a single piece of paper that contains antibody-decorated nanoparticles. When the urine sample contains *E. coli*, the biosensors generate colored spots on the paper strip. These are then quantified with a mobile app that calculates the pixel intensity in real time. The tests are highly specific and do not cross-react with other common uropathogens. Furthermore, the biosensors only yielded one false negative result when queried with a panel containing 57 urine samples from patients, which demonstrates that they have excellent sensitivity and specificity. This, along with the rapid assay time and smartphone-based detection, makes them useful for aiding in the diagnosis of UTIs at the point of care.

results of these tests are not always reliable. Studies in symptomatic female patients have shown that the nitrite test has a low diagnostic sensitivity (53–66%) and variable specificity (75–95%) when performed at the point of care.³ Detecting the leukocyte esterase yields better sensitivity for diagnosing UTIs (88–96%), but very low specificity (16–37%).³ Furthermore, the urine strip cannot identify the pathogen responsible for the infection. This information, which is standardly obtained through longsome culture methods, could help clinicians personalize treatments according to local antimicrobial resistance patterns.^{4,5}

In this manuscript we introduce a new tool for diagnosing UTIs based on the rapid detection of bacteria in urine with mobile biosensors (i.e. biosensors interfaced with a mobile device). The biosensors consist of a lab-on-chip device made of a single piece of paper. This paper biosensor is folded following a simple origami pattern in order to enable all of the analytical steps required to detect the target⁶⁻⁹ Our device is one of the few origami immunosensors that is entirely made of cellulose, it does not require using other materials in order to store antibody-decorated nanoparticles.^{10,11} Instead, it uses polystyrene sulfonate (PSS) in order to fabricate nanoparticle reservoirs directly on filter paper.¹² This reservoir is separated from the detection area by a wax barrier, which is easily drawn manually without using wax printers. The immunosensors were designed in order to detect pathogens above the infectious threshold as soon as possible and with the easiest analytical procedure, since these are the key points that make a biosensor useful in the doctor's office, where there is limited time to diagnose and prescribe therapies. Accordingly, all

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analytical steps were designed to reduce the turnaround assay time below 10 min and to detect pathogens with a concentration of 10^5 cells per mL or higher (clinical infectious threshold¹) without diluting the sample, since this procedure would be cumbersome at the bedside.

The origami sequence that made this possible is shown in Fig. 1. First, a drop of urine is added and dried for 30 s (Fig. 1A). This method physically adsorbs the pathogens and other components of the urine to the cellulose matrix. After adding a blocking solution for less than 10 s, the biosensor is

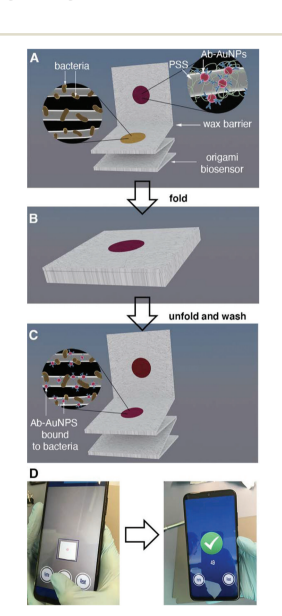


Fig. 1 Schematic representation of the mobile origami biosensors. (A) Urine samples are dried on a paper strip with a reservoir containing antibody-decorated gold nanoparticles (Ab-AuNPs), and polystyrene sulfonate (PSS) to avoid irreversible interactions with the paper; (B) after folding the paper the nanoparticles are transferred to the detection area where they specifically recognize *E. coli*; (C) after washing, a colored spot remains on the paper whose pixel intensity depends on the concentration of *E. coli* in the sample; (D) the pixel intensity is calculated with a smartphone app within seconds.

folded in order to transfer antibody-covered gold nanoparticles from a reservoir to the capture area for 5 min (Fig. 1B). After a quick washing procedure, a colored spot appears on the paper. The pixel intensity of the colored spot correlates to the biospecific recognition of the pathogen by the antibody (Fig. 1C). In this detection scheme the color is generated by the localized surface plasmon resonance (LSPR) of the gold nanoparticles, which is centered around 531 nm (nanoparticle size ca. 40 nm, Fig. S1[†]). Finally, the pixel intensity is quantified with a mobile app previously developed in our laboratory.¹³ The app automatically finds the region of interest and compensates for changes in illumination while at the same time guiding the user to the correct positioning of the smartphone with respect to the paper biosensor, all within a few seconds (Fig. 1D). With our app, there is no need to use external attachments or lighttight boxes for quantifying color consistently.¹⁴ The resulting origami mobile immunosensors were able to detect E. coli above the clinical infectious threshold in patient samples with a total assay time under 7 min. This rapid assay time, along with the minimal infrastructure requirements, make the biosensors ideal for clinical decision-making in a wide array of healthcare environments, from diagnosing UTIs in primary care to identifying the pathogen responsible for a urosepsis in the ICU. The biosensors were tested against 5 other uropathogens as well as with a panel of 57 different patient samples. Compared to other uropathogen immunosensors proposed in the literature, our biosensors yield results faster and only require an unmodified smartphone as a reader.^{15–21} Furthermore, they do not require a sample precondition step.¹⁴ They are also faster and easier to use than biosensors based on detecting nucleic acids.²²⁻²⁵ Moreover, the biosensors provide more information about the type of pathogen causing the UTI compared to the traditional urine strip. While this information is limited to a single pathogen at the moment, it has the potential to be expanded into a multisensor design including multiple reservoirs with antibodydecorated nanoparticles against different types of pathogens.

Experimental

Materials

Gold(m) chloride, sodium citrate tribasic dihydrate, poly(ethylene glycol) 2-mercaptoethyl ether acetic acid (thiol-PEG-acid) 2100, *N*-hydroxysulfosuccinimide sodium salt, *N*-(3-dimethylaminopropyl)-*N'*-ethylcarbodiimide hydrochloride (EDC), avidin from egg white, poly(sodium 4-styrenesulfonate) (PSS), *O*-(2-aminoethyl)polyethylene glycol 3000 (biotin-PEG), carboxyfluorescein diacetate succinimidyl ester (CFSE), and Tween-20 were purchased from Sigma Aldrich. Biotinylated polyclonal anti-*E. Coli* serotype O/K developed in goat was obtained from Thermo Fisher Scientific. Albumin from Bovine Serum (BSA, protease-free) was purchased from VWR Chemicals. PBS is phosphate buffered saline pH 7.4. PBST refers to PBS modified with 0.1% Tween. PBS-BSA is PBS containing 5 mg mL⁻¹ BSA.

Biosensor fabrication

The protocol for growing gold nanoparticles with a diameter of ca. 40 nm has been described elsewhere.¹³ The nanoparticles were stabilized against salt-induced aggregation by substituting citrate molecules on their surface for thiol-PEG-acid ligands, and the resulting carboxylate moieties were used to covalently bind avidin molecules through amidation (EDC/ sulfo-NHS coupling) with a previously described method.13 Antibody-decorated nanoparticles were obtained by adding 10 μ L of biotinylated antibodies (4–5 mg mL⁻¹) to 100 μ L of avidin-modified gold nanoparticles ([Au] = 125 mM) for 1 hour.¹² When required, biotin-PEG was added at final concentration of 0.1 mM in order to block free avidin binding sites. Then the nanoparticles were centrifuged (7000 rpm, 6 min), the supernatant was removed, and the pellet was resuspended in 100 µL of PBST twice. Finally, the nanoparticles were resuspended in 20 µL of PBST in order to yield antibodydecorated nanoparticles ([Au] = 625 mM).

Paper biosensors were made of Whatman #41 paper sheets cut into 2×8 cm strips. The strips were subdivided into four 2 × 2 cm squares that could be folded like an accordion as shown in Fig. 1A.²⁶ Next a wax barrier was added between the top square and the rest of the strip by placing the paper substrate on a hot plate and drawing a line with a piece of paraffin. The heat from the plate melts the paraffin and helps it penetrate the paper therefore creating a hydrophobic wax barrier after cooling (Fig. S3[†]). Next, 30 µL of 30% PSS was added in the center of the top square and left to dry at room temperature for at least 30 min. Finally, reservoirs containing anti E. coli-decorated nanoparticles were obtained by adding $1 \ \mu L$ of the colloidal suspension in the middle of the dried PSS spot and letting it dry at room temperature for 10 min.¹² The resulting biosensors could be stored in vacuum sealed bags at 4 °C for at least 14 days without noticeable loss in performance (Fig. S4 in ESI[†]).

Patient samples and bacteriological culture

Anonymized patient samples were obtained from the Microbiology Department at Son Espases University Hospital after obtaining approval from the local ethics committee (protocol IB 4005/19 PI). Informed consent was waived by the ethics committee as these samples were leftovers from clinical diagnosis that would otherwise be discarded. The reference strains *Escherichia coli* ATCC25922 and Enterococcus faecalis ATCC 29212, along with *Klebsiella pneumoniae, Enterobacter cloacae, Proteus mirabilis, and Streptococcus viridans* group from a collection of clinical isolates were used in this study. First, each strain was grown in blood agar to check purity. Isolates were then inoculated in 50 mL tubes containing 10 mL of Luria Bertani (LB) broth and incubated overnight at 37 °C with shaking at 180 rpm in aerobic conditions.

Detection of bacteria

The following experiments were performed to demonstrate that the paper substrate captures *E. coli* cells in Fig. 2. First, a

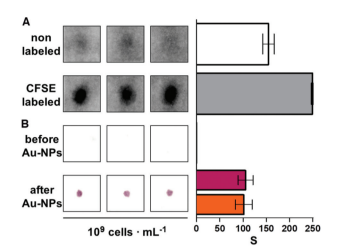


Fig. 2 Adsorption of bacteria to paper substrates; scanned images (triplicate) and densitometric signal (S) for *E. coli* cells trapped in the paper before and after labelling them with (A) CFSE or (B) antibody-decorated gold nanoparticles. In (B), pink bars were quantified by scanning the paper biosensor with a desktop scanner and quantifying the pixel intensity with Image J whereas the orange bar was obtained by means of real-time detection with our smartphone app. Error bars are the standard deviation (n = 3).

 10^7 cells per mL bacterial suspension in Luria–Bertani broth (LB) was stained with 50 μ M CFSE for 30 min at 37 °C. Then bacteria were pelleted by centrifugation at 4000 rpm for 10 min. After washing 3 times with PBS by centrifugation at 7000 rpm for 4 min, bacteria were finally resuspended in PBS to obtain CFSE-labeled *E. coli* with a concentration of 10^9 cells per mL. Subsequently, 10 μ L of non-labeled or CFSE-labeled bacteria at 10^9 cells per mL were spotted on the receiving paper substrate and left to dry at room temperature for 10 min. Finally, paper substrates were rehydrated with distilled water and the CFSE fluorescence was measured immediately with a Typhoon FLA 9500 laser scanner (General Electric) using the blue LD laser (473 nm).

Detection of bacteria with the proposed origami biosensors proceeded as follows. First, 10 μ L of the sample was added to the second square of the strip and dried with a hairdryer for 30 seconds (Fig. 1A and Fig. S2†). Afterwards, the last three squares of the paper strip were folded and 1 mL of PBS-BSA was added. Subsequently, antibody-decorated nanoparticles were transferred by folding the top square (Fig. 1B). This brings together the nanoparticle reservoir and the detection area where the bacteria are bound. After pressing with a clamp for 5 min, the reservoir is peeled off of the detection platform and the receiving paper is washed 3 times with 1 mL of PBST. The colorimetric signal is measured immediately afterwards with the mobile app.¹³

Experiments to determine whether bacteria interact specifically with avidin or streptavidin were conducted as follows. Solutions containing *E. coli* and *E. faecalis* (100 μ L) at different concentrations in PBS were dried onto 96-well ELISA plates (Nunc MaxiSorp, Thermo Scientific) by means of overnight incubation at 37 °C on a heating plate. Next, plates were

washed 3 times with PBST, blocked during two hours at room temperature (RT) with PBS-BSA and washed again 3 times. Then, 100 μ L of streptavidin-HRP diluted 1:3000 in PBST was added for 30 minutes at RT. Subsequently, the plates were washed 5 times with PBST and 100 μ L of 100 mg mL⁻¹ of TMB supplemented with 1.2 mM of hydrogen peroxide in 50 mM acetate buffer (pH 5.0) was added for 10 minutes at RT. Finally, the colorimetric reaction was stopped with 100 μ L of 2N H₂SO₄ and optical density (OD) was measured at 450 nm.

Results and discussion

One of the key factors for detecting bacteria rapidly with the proposed origami biosensors is to dry a drop of sample on the substrate, i.e. filter paper (Fig. 1A). The sample is quickly dried by positioning a hair dryer 5 cm away and blowing warm air for 30 s, which generates a temperature gradient that renders the pathogen physically adsorbed within the cellulose matrix, where they are specifically detected by antibody-coated nanoparticles in posterior steps (Fig. 1B and C). To demonstrate this point, E. coli cells were dyed with carboxyfluorescein diacetate succinimidyl ester (CFSE) and spotted on the filter paper. In Fig. 2A, images of three independent experiments obtained with a fluorescent scanner show that the CFSE-labeled cells are trapped within the cellulose matrix. When the same experiment was performed with unlabeled bacteria, the signal was significantly lower, which indicates that the fluorescent signal is originated by the CFSE-labeled cells and not by the spotting procedure. When the cells were immunostained with antibodydecorated nanoparticles, a red-colored spot appeared in the area where the bacteria were trapped, which was not present before the addition of the colloids (Fig. 2B). Densitometric analysis of the color in the spots with a flatbed scanner and with our mobile app yielded nearly identical results, further validating the smartphone-based approach for signal quantification at the point of care (Fig. 2B).¹³ In summary, the results shown in Fig. 2 demonstrate that the filter paper captures bacteria efficiently, and that these cells can subsequently be detected with antibody-decorated nanoparticles, which generate colored spots that are then able to be automatically quantified with an app. Below we apply these concepts for detecting bacteria at different concentrations with the origami biosensors shown in Fig. 1.

Next we calibrated the mobile biosensors with solutions containing different known concentrations of *E. coli*. Gram positive *S. viridans* was used as a control to test the specificity of the antibody-antigen interaction. In Fig. 3A, samples containing *E. coli* yield dose-dependent signals that followed the typical sigmoidal shape seen in immunodetection methods. The limit of detection, expressed as the sample that yields a signal higher than 3 times the standard deviation of the blank (99% confidence), is 10^5 cells per mL. Control experiments with *S. viridans* yielded signals that are below the limit of detection in all the concentrations assayed, which demonstrates that the dose-dependent signals observed during the

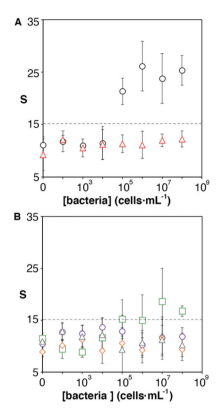


Fig. 3 Calibration plots representing the colorimetric signal S with respect to the concentration of pathogens. (A) *E. coli* (black dots) and *S. viridans* (red triangles); (B) *E. faecalis* (green squares); *K. pneumoniae* (purple dots); *P. mirabilis* (orange diamonds), *E. cloacae* (grey triangles) (semi-logarithmic scale). Dotted lines show the signal above 3 times the standard deviation of the *E. coli* blank. Error bars are the standard deviation (n = 3).

analysis of *E. coli* samples are originated by the specific recognition of the pathogen with antibody-decorated nanoparticles. We then tested the selectivity of the biosensors with samples containing some the most prevalent pathogens in UTIs. In Fig. 3B, solutions containing *P. mirabilis, E. cloacae* or *K. pneumoniae* always yield signals that are lower than the limit of detection for *E. coli*. However, *E. faecalis* is detectable at a concentration of 10^5 cells per mL or higher with the same biosensors, which indicates that the antibody-decorated nanoparticles are establishing non-specific interactions with these bacteria. This came as a surprise because *E. faecalis* is Gram positive, and therefore, no cross-reactivity was expected with the antibodies.

It has been shown that some microorganisms generate biotinylated proteins, and because of this they interact with streptavidin-modified probes.²⁷ In our biosensor design antibodies are bound to the nanoparticles by means of avidin-biotin interactions. Since avidin has 4 biotin binding sites, these are likely not fully occupied by biotinylated antibodies. With this in mind, we hypothesized that avidin could be responsible for the selectivity issues shown in Fig. 3B. To test this hypothesis, we physically adsorbed bacteria at different concentrations on an ELISA plate and added streptavidin-peroxidase. After

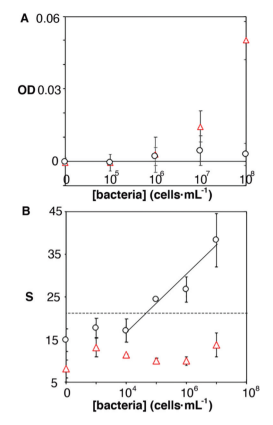


Fig. 4 Origin of non-specific interactions between *E. faecalis* and antibody-decorated nanoparticles. Optical density (OD) or colorimetric signal (S) with respect to the concentration of *E. faecalis* (red triangles) or *E. coli* (black dots) after incubating cells with streptavidin-HRP (A); or after repeating the calibration experiment in Fig. 3 with nanoparticles capped with biotin-PEG (B) (semi-logarithmic scale). Dotted lines show the signal above 3 times the standard deviation of the *E. coli* blank.

washing away excess reagents, we quantified the presence of bacteria-streptavidin interactions by adding a chromogen and measuring the generation of color by horseradish peroxidase (HRP). In Fig. 4A, E. faecalis yields signals that are higher than those obtained with E. coli, which indicates that the streptavidin-HRP complex interacts specifically with E. faecalis. We then sought to eliminate this interference in our biosensors by repeating the calibration curve shown in Fig. 3 with nanoparticles modified with anti-E. coli and capped with biotinylated polyethylene glycol (biotin-PEG). In Fig. 4B, E. coli shows a dose-dependent behavior whereas E. faecalis always yields very low signals independently of the concentration of bacteria in the sample. These experiments demonstrate that the selectivity issues in Fig. 3B are caused by avidin-biotin interactions, and that these issues can be alleviated by capping free biotinbinding sites with biotin-PEG. Furthermore, E. coli signals fit well to a linear regression model in the concentration range between 10^4 and 10^7 cells per mL ($y = 6.6x - 9.5, r^2 = 0.93$).

After studying the selectivity of our biosensors towards the recognition of different uropathogens, we sought to determine whether they could be used to detect UTIs in real samples. To this end, a panel of 57 patient samples that included UTIs by

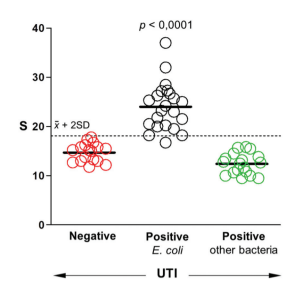


Fig. 5 Biosensor response when queried with a panel of patient samples that were *E. coli* positive ($\geq 10^5$ cells per mL, black), *E. coli* negative, or positive due to an infection by another pathogen (green). Horizontal bars represent the mean. *P*-value was obtained with a Kruskall-Wallis test.

E. coli, UTIs by other pathogens, and negative samples was queried with our biosensors. All samples were obtained from the Microbiology Unit at Son Espases University Hospital. They were analyzed using bacteriological culture following clinical guidelines. According to these guidelines, bacteriuria is defined by a positive urine culture with a bacteria concentration equal or higher than 10⁵ cells per mL.^{28,29} Quantification above this threshold is not associated with disease severity or a different diagnosis, and therefore it is not performed in the standard clinical routine. Fig. 5 shows the comparison between our biosensors and the gold standard bacteriological culture following this criterion. Samples that yield signals above 2 times the standard deviation of the mean value of negative samples are considered E. coli positive. Following this criterium, only one sample from E. coli UTI vielded a false negative. Samples containing other bacteria always yielded signals below the threshold value, which corroborates the excellent selectivity of our biosensor. The test diagnostic sensitivity (true positive rate) is 95.5%, and the specificity (true negative rate) is 100%. These results demonstrate that our mobile biosensors are useful for detecting pathogens in real samples rapidly and with high accuracy, which paves the way for their implementation in clinical practice.

Conclusions

In conclusion, we introduced origami biosensors with mobile readouts for detecting UTIs at the point of care. The whole assay takes less than 7 min. Selectivity issues due to unforeseen interactions between avidin and *E. faecalis* were overcome by capping biotin-binding sites with biotin-PEG. After this

treatment, the biosensor shows an excellent selectivity and specificity towards *E. coli*. When queried with a panel of patient samples the sensors yielded only one false negative. Biosensors for detecting other pathogens could readily be fabricated by changing the antibodies around the nanoparticles. Moreover, multi-sensors containing reservoirs with nanoparticles designed to detect different bacteria could be used to identify the pathogen causing a UTI. These features, along with the short assay time and mobile detection scheme, make our biosensors promising for guiding antibiotic prescriptions at the point of care.

Conflicts of interest

R. R., and C. A. have filed a patent application (PCT/EP2020/ 075013) in order to protect the intellectual property of the method for fabricating nanoparticle reservoirs.

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