Silica nanoparticles with encapsulated DNA (SPED) – a novel surrogate tracer for microbial transmission in healthcare

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Abstract

Background

The increase in antimicrobial resistance is of worldwide concern. Surrogate tracers attempt to simulate microbial transmission by avoiding the infectious risks associated with live organisms. We evaluated silica nanoparticles with encapsulated DNA (SPED) as a new promising surrogate tracer in healthcare.

Methods

SPED and *Escherichia coli* were used to implement three experiments in simulation rooms and a microbiology laboratory in 2017-2018. Experiment 1 investigated the transmission behaviour of SPED in a predefined simulated patient-care scenario. SPED marked with 3 different DNA sequences (SPED1-SPED3) were introduced at 3 different points of the consecutive 13 touch sites of a patient-care scenario that was repeated 3 times, resulting in a total of 288 values. Experiment 2 evaluated SPED behaviour following hand cleaning with water and soap and alcohol-based handrub. Experiment 3 compared transfer dynamics of SPED versus *E. coli* in a laboratory using a gloved finger touching two consecutive sites on a laminate surface after a first purposefully contaminated site.

Results

Experiment 1: SPED adhesiveness on bare skin after a hand-to-surface exposure was high, leading to a dissemination of SPED1-3 on all consecutive surface materials with a trend of decreasing recovery rates, also reflecting touching patterns in concordance with contaminated fingers versus palms.

Experiment 2: Hand washing with soap and water resulted in a SPED reduction of 96%, whereas hand disinfection led to dispersal of SPED from the palm to the back of the hand. Experiment 3: SPED and *E. coli* concentration decreased in parallel with each transmission step – with SPED showing a trend for less reduction and variability.

Conclusions

SPED represent a convenient and safe instrument to simulate pathogen spread by contact transmission simultaneously from an infinite number of sites. They can be further developed as a central asset for successful infection prevention in healthcare settings.
Background
The increasing spread of antimicrobial resistance and the burden of healthcare-associated infections represent substantial threats to patient health and life. (1-3) Healthcare workers’ (HCW) hands play a key role in transmission of pathogens through bi-directional exchange of microbes between hands and surfaces, (4, 5) especially considering that hand hygiene performance is commonly substandard in almost all healthcare settings despite widely accepted concepts and guidelines. (6, 7) A recent head-camera-aided study uncovered that many more opportunities for microbe transmission might exist in acute healthcare than anticipated based on more traditional hand hygiene observation methods. (8-10) Tracing events leading to microbial transmission, however, is difficult, especially using living microorganisms. It might therefore be beneficial to use surrogate tracers to increase our understanding of transmission pathways. Prior studies have attempted to trace transmission events using microbial surrogates, (11-16) and/or DNA. (11, 17-21) Use of microbial surrogates is limited to benign organisms, like MS2 phage, and therefore limits their widespread use in transmission studies. DNA transmission behaviour is unlikely to mimic that of microorganisms. Cauliflower mosaic virus DNA, (11, 17-21) bacteriophages, and fluorescent lotion have been used to this aim in the past. (11-16, 19, 22) Yet, using surrogates with physiochemical properties more similar to microorganisms may allow simulating transmission events without the need for microbial contamination. Thus, we investigated the behaviour of silica nanoparticles with encapsulated DNA (SPED) as potential surrogate tracers for microbial transmission pathways in health care.

Methods
Silica Particles with Encapsulated DNA
SPED of known nucleotide sequences have been established by Paunescu et al. in order to create inert surrogate tracers that serve to track the presence and distribution of biological and chemical entities (e.g. wastewater, milk, pesticides). (23-26) Forming part of daily meals of many people through seafood, grains, vegetables and broadly utilized in the food industry, silica has been acknowledged as “safe” by the U.S. Food and Drug Administration 50 years ago. (27) In medicine (e.g. imaging) silica nanoparticles are widely used and investigations of their biorelated degradability and
clearance has been conducted. (27) The synthesis and characterization of SPED was conducted according to Paunescu et al. (23) (Figure 1) First, per batch, 4x4ml of silica nanoparticles (50mg/ml in isopropanol) were surface functionalised in 4 separate by adding 40µg of N-trimethoxysilylpropyl-N,N,N- trimethylammonium chloride (TMAPS; 50% wt in methanol) and stirred at 900 rounds per minute (rpm). for 12 hours at room temperature. The trimethylammonium on the surface gave the silica particles a positive surface charge. For DNA adsorption on the surface a 2 ml batch of corresponding annealed DNA molecules was added to 200ml ultrapure MilliQ (mQ) water (150ng dsDNA/µl, Nanodrop). Of the previously TMAPS functionalised particles, 0.4g were added to the solution and the batch was shaken for 10 seconds. Subsequently, 4µl TMAPS were added, the batch was shaken and then sonicated for 20 seconds. Next, 62.5µl of tetraethyl orthosilicate (TEOS) (≥99.0%, Aldrich) were added and the batch was shaken for 5 hours. In a next step, 10ml isopropanol and 5.9ml TEOS were mixed with 484.1ml mQ water before adding it to the previous mixture. The batch was then stirred at 600rpm for 4 days. To demonstrate the potential of multi-tracing, three different batches were produced, each containing a unique DNA sequence. The three particle batches named SPED1, SPED2 and SPED3 had a hydrodynamic size of 218±80nm, 144.6±46nm and 173.4±82nm and a DNA loading of 21µg, 23µg, and 26µg dsDNA/mg of particles, respectively. The suspension used for our experiments was prepared by diluting the particles to 0.1mg/ml or 1mg/ml in mQ water + 10% glycerine. We used glycerine to increase the probability of transfers in our experiments according to preliminary data showing higher transfer rates in the presence of glycerine (data not shown). Glycerine is viscous and increases therefore the stickiness of the test suspension, simulating roughly the appendage of bacteria, called pili.

**Escherichia coli bacterial culture**

Standard procedures were used to cultivate bacterial cultures. *Escherichia coli* were cultured overnight in tryptic soy broth (TSB; Sigma-Aldrich, St. Louis, Missouri, USA). An aliquot of 2ml of the bacterial cell culture was triple washed by spinning down the *E. coli* culture in a 2ml microcentrifugation tube (Eppendorf AG, Hamburg, Germany) for 5 minutes at 3500rpm. After each cycle, the supernatant TSB was removed, and the bacterial culture resuspended in 1ml phosphate-
buffered saline (PBS). The obtained bacterial concentration was $10^8$ CFU/ml. To prevent further bacterial growth during transfer experiments, microcentrifugation tubes were stored temporarily on ice.

**Study procedures**

We conducted three experiments to investigate the behaviour of SPED in a simulated care scenario (Experiment 1), while performing hand hygiene (Experiment 2), and comparing SPED transfer dynamics with those of *E. coli* (Experiment 3).

**Experiment 1 - SPED transmission characteristics in a simulated care scenario**

We defined a simulated patient care scenario featuring an examination stretcher with bed rails, a privacy screen, and an infusion stand with two infusion bags. A patient actor (henceforth patient) and a HCW trained as a physician (MS; henceforth HCW) conducted the following actions in chronological order: closing a door using the door handle (stainless steel), moving the privacy screen (polyvinylchloride), elevating the bed rail (polypropylene), auscultating the patient’s chest with a stethoscope (polyvinylchloride, PVC), taking the patient’s radial pulse (bare skin), checking pupil reactivity while pulling up the patient’s upper eyelid (bare skin), and changing the infusion bag (polyolefin) (Figure 2). Two video cameras positioned in a 90° angle on adjacent walls (GoPro® Hero 4 Black edition, GoPro Inc., San Mateo, CA) recorded the scene simultaneously to monitor the sequence.

The scenario featured 13 experimentally pre-defined hand-to-surface exposures, of which 10 were fomites and 3 were patient skin.(8) We marked swabbing areas of 1x3cm on each of the 13 sites. The sites and both hands and fingers of the HCW were swabbed prior to every simulation to demonstrate they were free of SPED. The swabbing technique included with 20% glycerine solution premoistened sterile cotton swabs (Naturaline Wattstäbchen, Steinfels Swiss, Winterthur, Switzerland) that were rolled three times over the area and then stored in a 2ml microcentrifugation tube at room temperature. This swabbing procedure applied to all following swabs in our experiments. We inoculated three of the 13 sites with 0.2ml of the SPED test suspensions (1mg/ml) and left to air dry for 5 minutes, namely the door handle with SPED1, the bed rail with SPED2, and the left forearm
of the patient with SPED3 (Figure 2). Throughout the entire simulated care scenario, the patient was laying on his back on the examination stretcher with his bare arms placed beside him without contact to his body or the examination stretcher. Subsequently, the HCW actor performed the care scenario while exclusively touching the 13 marked sites in a natural way. Once the simulation scenario was concluded, we swabbed the 13 marked test areas as well as both of the HCW’s palms and the fingers of her dominant right hand, resulting in 48 specimens overall. After the experiment, the microcentrifugation tubes were immediately transported to the laboratory for processing.

The experiment was repeated three times on different days and different simulation rooms, each, referred to as ‘runs’.

**Experiment 2 - Effect of hand hygiene action on SPED concentration**

We tested the impact of hand disinfection with alcohol-based handrub and hand washing using soap and water on SPED skin concentration. Three groups of five participants, each, were formed: a control group (CTRL), a hand disinfection group (DISINF), and a hand washing group (WASH). First, swabs were taken from the right thenar of each participant to demonstrate no SPED were present. Subsequently, a previously marked area of 2x2cm on the palmar side of the right thenar of each participant was inoculated with 0.2ml of the SPED1 test solution (0.1mg/ml). The fluid was dispensed over the palms of both hands by rubbing both hands against each other for 10 seconds. Once dry, swabs were taken from the previously marked area of the right thenar of each person, and additionally from the dorsal hypothenar of DISINF participants. CTRL participants held both their hands in the air without any contact for 1 minute. DISINF participants performed hand hygiene according to the norm EN1500 using 3ml of a commercially available alcohol-based handrub (80% Ethanol with 1% Glycerine, B. Braun Medical AG, Sempach, Luzern, Switzerland). WASH participants washed their hands for 30 seconds with common soap (Liquid Soap®, Sterisol AB, Vadstena, Sweden) and water using the norm EN1499 procedure. The right thenar of all participants – and additionally the dorsal hypothenar in DISINF participants – were swabbed again after the non-contact waiting time in the CTRL group and hand hygiene actions in the DISINF and WASH group. Thereafter the test tubes were immediately transported to the laboratory for
analysis.

**Experiment 3 - Transmission dynamics of SPED compared with *Escherichia coli***

We analysed transmission tendencies of *E. coli* and SPED. Prior to the experiment, a piece of laminate (melamine resin) was rinsed with 70% ethanol, wiped with a sterile cloth and allowed to air dry. Forty series of four squares of 2x2cm were defined. The four squares were marked as origin surface (ORIGS), 1st transfer surface (1TS), 2nd transfer surface (2TS) and control surface (CTRLS), respectively. Next, samples of the surface and gloves (GLOVES) were taken by either rinsing the surface with 100µl of PBS or by cutting off the glove’s index fingertip and adding it to a 1.5ml micro-centrifugation tube filled with 100µl of PBS to prove sterility. Then, 20µl of the *E. coli* solution was spread onto ORIGS, and with intervals of 2 minutes, onto CTRLS. The 2-minute interval allowed for continuous sampling of surfaces with identical times between surface inoculation and subsequent sampling. A two-step transmission was performed from ORIGS → 1TS → 2TS by pressing the gloved index finger onto the corresponding areas in that order. Samples were taken as described above. The CTRLS served as a reference for the number of *E. coli* or nanoparticles recovered from an inoculated surface in the absence of a transfer event, and so not part of the transfer. This procedure was repeated until there were 20 replicates. The same procedure was reproduced for 20 replicates with 20µl of the SPED solution (0.1mg/ml).

**Quantification of SPED**

To quantify the amount of DNA particles from each swab after having performed the experiments, the cotton swab was placed in a 2ml microcentrifugation tube (Eppendorf AG, Hamburg, Germany) with 200µl of mQ water (**Figure 1**). Each sample was ultrasonicated for 1 minute and vortexed for 10 seconds. To release the DNA from the silica coating, 1% vol of a highly diluted solution of buffered oxide etch (BOE) with 0.03wt% ammonium hydrogen difluorid (NH4FHF, pure, Merck) and 0.02wt% ammonium fluoride (NH4F, puriss., Sigma-Aldrich, St. Louis, Missouri, USA) were added to the sample. The resulting suspension was then analysed by quantitative PCR (LightCycler® 96) in a multiplex setup with three sequence-specific fluorescent probes (SPED1: Hex, SPED2: Texas Red, SPED3: FAM). All three fluorescent probes could be detected simultaneously and allowed to quantify each particle.
separately. The qPCR total reaction volume was 12.5µl consisting of 2.5µl sample solution, 0.2µl of each primer (3x forward and reverse), 0.125µl of each probe (Microsynth AG) and Mastermix 2x (GoTaq® Probe, Promega). The qPCR program consisted of a preincubation step for 600 seconds at 95°C, followed by a 2-step cycling for 15 seconds at 95°C and for 60 seconds at 56°C. For quantification, a dilution series with known concentration of particles was performed.

**Sampling of *Escherichia coli***

Once the transfer procedures were completed, the samples were processed within 2 hours. For sampling, 100µl of PBS were utilised to rinse the with *E. coli* contaminated areas, added to a 1.5ml micro-centrifugation tube containing 100µl of PBS, and vortexed. As for gloves (Nitrile latex free, LLG Labware, Meckenheim, Germany), the index fingertip was cut off after performing the transmissions, placed in a 1.5ml micro-centrifugation tube containing 200µl of PBS and subsequently vortexed. Aliquots of 100µl of undiluted specimen solution and 100µl from 1:10–1:100’000 dilutions were plated onto TBX Agar (Sigma-Aldrich, St. Louis, Missouri, USA) and incubated at 36±1°C for 24 hours.

**Statistical analysis**

Descriptive analysis of our data in Experiment 1 and 2 was performed with Microsoft® Excel® 2016. To estimate the detection limit of the nanoparticles in our experiments, we used the Method Detection Limit (MDL). The MDL is defined as “the minimum measured concentration of a substance that can be reported with 99% confidence that the measured concentration is distinguishable from method blank results.” (28) Values that are below MDL are considered to be negative, values higher than the limit as positive. The MDL was calculated as the mean concentration of the blank samples plus three times the standard deviation.

For Experiment 3, test statistics including t-test were conducted with R Statistical Software (version 3.6.0, R foundation, Vienna, Austria). We defined transfer percent as the fraction of *E. coli* or SPED recovered from 1TS, 2TS, and glove as a percentage of the seeded inoculum recovered from the surface by swabbing, defined by CTRLS.

**Results**

Experiment 1 - SPED transmission characteristics in a simulated care scenario
According to the video analysis, the sequence of touched surfaces was maintained as scheduled for all runs. Overall, we collected 96 swabs, each analysed for the three different DNA sequences, resulting in 288 values. All 39 control swabs on the 13 test sites for the three scenario runs before SPED were negative. Equally, none of the swabs tested positive for a given SPED type ahead of its deposition site in the care sequence (Fig. 3). For runs 1, 2 and 3, the MDL determined from the recovered sample for SPED1 was $1.49^{-7}$ mg/ml; $3.70^{-7}$ mg/ml, $9.95^{-8}$ mg/ml, for SPED2, $1.80^{-8}$ mg/ml; $5.20^{-9}$ mg/ml; $3.75^{-9}$ mg/ml and for SPED3 $4.24^{-8}$ mg/ml, $6.74^{-8}$ mg/ml; $1.25^{-8}$ mg/ml, respectively. Accordingly, overall, the number of positive results among all sites except the deposition site and hands were 26 of 36 (72.2%) for SPED1, 28 of 30 (93.3%) for SPED2, and 8 of 21 (38.1%) for SPED3. As for hands, right hand fingers revealed positive SPED results in 9 of 9 (100%), right hand palm in 6 of 9 (66.6%) and left-hand palm in 2 of 9 (22.2%). Detection rates for the tracers varied among the touch-sites. Consistently positive samples were found on the left and right forearm of the patient, on the infusion bag valve, and on HCW’s fingers. No differences were found in detectability regarding the material on which the test suspension was deposited.

Experiment 2 - Effect of hand hygiene procedure on SPED concentration

All samples taken before applying SPED were negative. The mean SPED1 DNA-concentration before and after the experiment was $8.25^{-5}$ mg/ml (standard deviation, ± $4.84^{-5}$ mg/ml) and $6.88^{-5}±7.60^{-5}$ mg/ml for DISINF of the right thenar, $9.34^{-8}±5.35^{-8}$ mg/ml and $2.41^{-5}±2.35^{-5}$ mg/ml for DISINF of the hand back, $1.1^{-4}±7.61^{-5}$ mg/ml and $3.96^{-6}±2.52^{-6}$ mg/ml for WASH, and $6.69^{-5}±6.53^{-5}$ mg/ml and $8.32^{-5}±5.61^{-5}$ mg/ml for CTRL, respectively. This corresponds to a 24% increase of SPED concentration for CTRL, a reduction of 17% on the thenar and 25.670% increase on the back of the hand for DISINF, and a reduction of 96% for WASH (Fig. 4).

Experiment 3 – Transmission dynamics of SPED compared with *Escherichia coli*

In total, 40 two-step transfers were conducted, 20 for *E. coli* and 20 for SPED. Samples taken from gloves and surface prior to the experiment were negative. The results are displayed in Fig. 5. The median (interquartile range) recovery fraction of ORIGS$_E. coli$ was $-0.34$ (-0.44; 0.02), -1.10(-1.52;
-0.96) for 1TS\textsubscript{\textit{E. coli}}, -1.68 (-2.19; -1.48) for 2TS\textsubscript{\textit{E. coli}} and – 2.66 (-3.11; -2.07) for GLOVES\textsubscript{\textit{E. coli}} (n = 20).

For the SPED, the median resulted in -0.16 (-0.24; -0.04) for ORIGS\textsubscript{SPED}, in -0.71 (-0.80; -0.65) for 1TS\textsubscript{SPED}, in -1.19 (-1.35; -1.10) for 2TS\textsubscript{SPED} and in -2.47 (-2.53; -2.23) for GLOVES\textsubscript{SPED} (n = 5). This correlates to a median (range) recovered fraction of \textit{E. coli} from ORIGS against CONTLS of 57% (0.1%-540%), from 1TS against CONTLS of 8% (0.1%-150%), from 2TS against CONTLS of 2.1% (0.01%-13%) and 0.4% (0.005 -11%) from GLOVES. The median recovered SPED fraction from ORIGS was 70% (28%-287%), 16% (0.3%-40%) from 1TS, 5.3% (0.1%-16%) from 2TS and 0.7% (0.1%-55%) from GLOVES.

**Discussion**

We explored SPED as a promising agent to trace microbial transmission in healthcare. First, we were able to establish the suitability of SPED to trace the transmission pathways of microbes in a simulated care scenario featuring a typical sequence of hand-to-surface exposures. They showed decreasing trends with consecutive touched surfaces and discriminative patterns on hands and surfaces reflecting the way objects were manipulated. Second, we found that presence of SPED following handwashing with soap and water was similar to that of microorganisms but are inert to alcohol-based handrub, as expected. Third, and most importantly, SPED showed a nearly parallel decrease in concentration on surfaces compared to \textit{E. coli} in a head-to-head two-step transmission experiment.

Experiment 1 demonstrated successfully the transmission of three different SPED markers in a simulation room. The markers SPED1 and SPED2 were transmitted from their deposit site to the palm of the HCW’s right hand and fingers, and subsequently spread onto almost all surfaces except eyelid and one of the two infusion bags. In contrast, the deposition site of SPED3 – the left forearm of the patient – was touched by the HCW’s right hand fingers, which resulted in a specific transmission to consecutive contact-sites that were touched by fingers only (Fig. 3). This finding suggests that SPED could be used for a detailed understanding of transmission pathways in healthcare settings, especially benefitting from the possibility to use identifiable SPED in parallel and avoid the problem of contamination in case of repeated investigations in the same environment.

In Experiment 2 hand washing with soap and water resulted in a nearly complete removal of SPED
from hands and thus ideally simulated the mobilisation and dilution of microorganisms. In contrast, hand cleaning with alcohol-based handrub led to the dispersion of SPED to the back of the hands – simulating the behaviour of partially inert microorganisms such as spores (e.g. *Clostridium difficile*) and non-enveloped virus (e.g. Norovirus).(29, 30)

By comparing transmission behaviour of SPED to those of *E. coli* in Experiment 3 we found similarities in transfer with SPED exhibiting a trend to a lower decrease from one consecutive surface to the other. This suggests that, when used as surrogate markers in real healthcare settings, transmission will be rather over- than underestimated.

Previous studies have evaluated the use of harmless surrogate tracers to simulate contact transmission pathways of potential pathogens in healthcare settings.(11–22) These studies employed three surrogate markers, namely DNA sequences of the cauliflower mosaic virus DNA,(11, 17–21) bacteriophage MS2, a non-pathogenic non-enveloped RNA virus,(11–16) and fluorescent agents.(13, 14, 16, 19, 22) Some of these studies compared fluorescent agents against MS2 bacteriophages (13, 14, 16) or against cauliflower virus DNA (19); or used the two viral DNA markers in parallel.(11) Only one of these studies compared the behaviour of cauliflower virus DNA and MS2 bacteriophage against *Clostridioides difficile* spores and found a similar environmental transmission behaviour,(11) as did we in our comparison between SPED and *E. coli*. Two studies used visible fluorescent and flashlight reflective markers, respectively, as part of an educational intervention on personal protective equipment (PPE) handling and MS2 bacteriophages as outcome measure.(15, 16) These studies all show an impressively widespread transmission of surrogate markers across the healthcare setting in a short interval of time, even if one considers that their persistence in the environment might be longer than viral or bacterial pathogens.

We used video to verify the correct execution of the experimentally predefined hand-to-surface exposure sequence in a simulated care scenario. One study assessed PPE handling quality by video and related it to transmission of MS2 bacteriophage and fluorescent liquid.(13) Two studies outside of healthcare settings estimated the risk of bacterial cross-contamination in farming work with human excreta by combining videography with environmental microbial contamination data into a stochastic-
mechanistic simulation of bacterial transmission.(31, 32)

In contrast to other methods to investigate pathogen transmission in healthcare settings, using SPED has several advantages. First, silica particles are easy to handle, and their management requires only little specialised expertise. The equipment for the analysis, i.e. qPCR analysers, are common equipment in hospital microbiology laboratories. Second, SPED are widely available, safe, and inexpensive. As an example, the raw materials for three repetitions of the Experiment 1 amounted to ~200 USD, including DNA, primers and other reagents. The costs of SPED are mainly determined by the price of the synthetic DNA amplicon, which currently equals 500 USD for 100 mg of newly synthesized DNA. Third, the physical characteristics of SPED, such as electric charge, hydrophobic versus hydrophilic properties, size and weight can be modified to match those of pathogens.(33–35) Forth, and most importantly for a clinical application, the fact that SPED solutions with multiple DNA-codes can be applied and analysed by PCR simultaneously provides the opportunity to study an almost infinite number of scenarios and allows for repeated experiments in the same environment with minimal risks of cross-contamination.

This proof of concept study has limitations. The conducted experiments were limited in replications and simulated patient care was a simplified predefined scenario. However, we do not expect a categorically different behaviour of SPED in real-life care settings. The concentration of SPED on consecutive sites was high enough to rise confidence that even much wider dispersal can still be traced. Differences in hand-to-surface exposures in addition to inhomogeneous distribution of the SPED in the test solution and slight variations in the swabbing technique are may explain the non-linear decline of recovery rates from consecutive touch sites. Future work on SPED tracking should attempt to decrease the technical variation in recovery. Considering that concentrations of pathogens (e.g. Norovirus in faeces) appear to be in the same range (36), our results represent an extreme but not unrealistic scenario. Future dilution series could help to establish concentration that are more congruent to daily healthcare conditions. Finally, given the variation in SPED recovery in our study, future work could be invested in standardising the sampling method.

Conclusions
This proof-of-concept study is, to our knowledge, the first investigation using SPED as a surrogate tracer to study pathogen transmission in healthcare. Further studies in various real-life care settings – and modifying SPED characteristics to align even better with pathogen features – will show their potential to investigate outbreaks and endemic infectious risks at a system level.

In conclusion, SPED spread quickly over all the touch-sites through hand contact and allow to determine their origin and mimic the transmission of live bacteria. They represent a convenient and safe tool to explore pathogen spread from an infinite number of sites simultaneously. They can be developed further to become a central asset for a more effective infection prevention in healthcare and beyond.

Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| SPED         | Silica nanoparticles with encapsulated DNA |
| E. coli      | Escherichia coli |
| HCW          | Healthcare worker |
| TMAPS        | N-trimethoxysilylpropyl-N,N,N- trimethylammonium chloride; |
| TEOS         | Tetraethyl orthosilicate |
| TSB          | Tryptic soy broth |
| PBS          | Phosphate-buffered saline |
| CTRL         | Control group in Experiment 2 |
| DISINF       | Disinfection group in Experiment 2 |
| WASH         | Hand washing group in Experiment 2 |
| ORIGS        | Origin in Experiment 3 |
| ITS          | First transfer in Experiment 3 |
| 2TS          | Second transfer in Experiment 3 |
| CTRLS        | Control in Experiment 3 |
| BOE          | buffered oxide etch |
| MDL          | Method Detection Limit |

Declarations

Ethics approval and consent to participate

The three experiments were formally declared as not being subject to the Swiss law on research on humans by the cantonal ethics board (Rec-2017-00889). All participants received comprehensive information about the experiment and gave oral consent to their voluntary participation. Withdrawal was possible at any moment without having to communicate a reason.

Consent for publication

Not applicable.

Availability of data and materials

Not applicable. Please contact author for data requests.

Competing interests

The authors declare that they have no competing interests.
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None.

Authors’ contributions
All authors contributed at least to one of the following fields: conception and study design (MS, JK, LC, AW, TJ, NRG, HS), data acquisition and analysis (MS, JK, TJ, APK), interpretation of data (MS, JK, TJ, NRG, HS) and writing of the manuscript (MS, JK, TJ, HS). All authors read and approved the final manuscript.

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Figures

Figure 1

Scheme of synthesis of SPED and analysis with qPCR quantification. Legend: TMAPS, N-trimethoxysilylpropyl-N,N,N-trimethylammonium chloride; TEOS, tetraethyl orthosilicate.
Legend: 1-13, hand-to-surface exposures (1, door handle; 2, privacy screen; 3, bed rail; 4&5, stethoscope; 6, left forearm; 7, eyelid; 8, infusion bag valve; 9&10, infusion bags; 11, bed rail; 12, privacy screen; 13, right forearm)

👶, SPED1 introduction (door handle); 👦, SPED2 introduction (bed rail); 👦, SPED3 introduction (patient forearm); →→→, transmission pathway with numbers representing the consecutive hand-to-surface exposures

Figure 2

SPED deposition and screening points in the patient care simulation in Experiment 1
Figure 3

Detection of SPED1-3 on environmental and patient body sites and healthcare worker’s hands in Experiment 1
Figure 4

SPED1 skin concentration before and after hand cleaning procedures in Experiment 2.

Legend: Disinfection, hand cleansing with alcohol-based handrub; Washing, washing hands with soap and water; Control, no hand hygiene action; Error bars = Standard deviation
Figure 5
Log10 percent transfers for the original, 1TS, 2TS, and GLOVES in Experiment 3. Legend:
ORIGS, origin surface; 1TS, first surface touched after the origin surface; 2TS, surface touched after 1TS; GLOVES, nitrile disposable glove index finger tested after all three surfaces have been touched

Supplementary Files
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