A Simple and Fast Assay Based on Carboxyfluorescein-Loaded Liposome for Quantitative DNA Detection

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Supporting Information

ABSTRACT: The development of an innovative and easy way to run assays for the quantitative detection of DNA present in biological fluids (i.e., blood, urine, and saliva) is of great interest for early diagnosis (e.g., tumors) and personalized medicine. Herein, a new quantitative assay based on the use of highly sensitive carboxyfluorescein-loaded liposomes as signal amplification systems is reported. The method has been tested for the detection of low amounts of DNA sequences. The reported proof of concept exploits a target DNA molecule as a linker between two complementary oligonucleotides. One oligonucleotide is biotinylated at its 3′ end and binds to streptavidin-coupled magnetic beads, whereas the other one is conjugated to a cholesterol molecule incorporated in the phospholipidic bilayer of the fluorescent liposomes. In the presence of the target fragment, the correct formation of a construct takes place as witnessed by a strong fluorescence signal, amplified by dissolving lipidic nanoparticles with Triton X-100. The system is able to detect specific nucleotide sequences with a very low detection threshold of target DNA (tens of picomolar). The assay allows the detection of both single- and double-stranded DNA. Studies performed in human blood serum show the correct assembling of the probe but with a reduction of limit of detection (up to ∼1 nM). This liposome signal amplification strategy could be used not only for the detection of DNA but also for other nucleic acids (mRNA; microRNA) that are difficult to be quantified by currently available protocols.

INTRODUCTION

The access to new, fast, and low-cost molecular diagnostic platforms is crucial to enable more effective monitoring and personalized treatments in cancer patients. In fact, the rapid and accurate detection of genomic alterations is at the basis of both the early diagnosis and the selection of the correct targeted therapy. Moreover, in addition to neoplastic diseases, nucleic acids may represent potential markers for a wide array of infectious diseases, including metabolic, inflammatory, and infectious pathologies. Despite the fact that standard protocols are still based on tissue biopsies to identify possible genomic alterations, these methodologies are limited by the difficulties related to the low accessibility of tumor tissue and by the risks associated with the repetition of many tissue sampling. Recently, the sampling of biological liquids (i.e., blood, serum, urine, and saliva) has been recognized as an alternative source of nucleic acids for diagnosis of diseases. In a liquid biopsy technique, great attention is devoted to the detection of the so-called circulating tumor DNAs (ctDNAs), a class of tumor-derived DNAs that are not associated with cells and are fragmented in the bloodstream.

In general, only very low amounts of ctDNA are present in healthy patients. Conversely, significant levels of ctDNA can be found in cancer. The methods routinely used for DNA/RNA detection involve two steps, namely, (i) the in vitro amplification of the sequence of interest (by polymerase chain reaction, PCR, or reverse transcription PCR, RT-PCR) and (ii) the detection of target DNA/RNA by electrophoresis, southern/northern blots, sandwich hybridization assays, or next-generation sequencing. The most innovative and best performing methodologies are represented by digital droplet PCR (ddPCR) and beads, emulsification, amplification, and magnets (BEAMing). Both of them show several advantages and disadvantages. The ddPCR methodology utilizes a droplet generator that allows the partition of single pieces of DNA into droplets using an oil/water emulsion. The routine detection of ctDNA is still hampered in the clinical practice not only for an intrinsic lack of sensitivity/speciﬁcity for ctDNA but also because the currently available procedures require a high level of expertise and a long execution time. Moreover, the facilities necessary for the analysis, as flow cytometry and sequencing machineries, are present in specialized and certiﬁed laboratories for this kind of testing but are not always available in ordinary diagnostic laboratories.

Alternatively, there are many examples of sandwich hybridization protocols based on the use of highly sensitive radioactive probes or gold or fluorescent nanoparticles. The latter ones are preferred to the radioactive probes for...
obvious safety reasons, although they are less sensitive.\textsuperscript{18} Hence, the development of highly sensitive fluorescence-based methods for ctDNA detection may be attractive but challenging. In fact, this could help overcome the drawbacks of classical molecular biology assays (i.e., the need of a facility and expert staff)\textsuperscript{19} without the need to use hazardous radioactive probes.

Useful routes to pursue the signal amplification may be provided by the use of nanoparticles.\textsuperscript{20} For example, fluorescein-encapsulating liposomes were reported to provide a 1000-fold decrease of limit of detection (LOD), with respect to fluorophore-labeled antibodies, in a sandwich hybridization-based flow injection analysis system.\textsuperscript{21} Baemunner et al. proposed an assay based on dye-loaded liposomes, bearing oligonucleotides (complementary to the target DNA) on their outer surface and polyethersulfone membranes with streptavidin immobilized in the detection area.\textsuperscript{22} The target DNA of interest acts as a linker between the two shorter oligonucleotide capture probes. Upon loading high amounts of sulforhodamine B (SR-B) inside liposomal nanoparticles, the detection threshold was as low as 1 nM analyte.

Starting from these results, herein, we propose a reproducible, versatil, and fast assay for DNA detection and quantification based on the use of highly stable liposomes loaded with fluorescent dyes that can be stored at 4 °C for 4 months without degradation. The proposed method does not require neither a high expertise level nor the use of radioactive molecules. It is based on the use of carboxyfluorescein (CF)-loaded liposomes functionalized in the lipidic bilayer with a cholesterol-tagged DNA sequence, which is complementary to a portion of the target DNA.

The proof of concept of the proposed assay deals with the capture of both single- and double-stranded DNA (ssDNA and dsDNA, respectively) molecules using complementary oligonucleotides, exploiting the target ssDNA or dsDNA as a linker between two artificial strands. These strands are bound to dye-encapsulating liposomes through a cholesterol molecule and to streptavidin-coated magnetic beads via a biotin molecule. To observe the fluorescent signal from the highly concentrated liposome payload, the lipodic nanoparticles were disrupted with Triton X-100 obtaining a high increase of signal intensity.

\section*{RESULTS AND DISCUSSION}

\subsection*{Assay Description}

The assay consists of the following four steps:

1) Commercially available magnetic beads functionalized with streptavidin molecules (Strep Mag) are incubated with a selected 18 base pair (bp) long ssDNA. This ssDNA, referred to as ANCHOR hereafter, was previously functionalized with a biotin-TEG group at its 5′ end to bind streptavidin. The ANCHOR fragment was specifically designed to be complementary with the first 18 nucleotides present on the 3′ region of the fragment to be detected (referred to as TARGET hereinafter) (Figure 1).

2) Magnetic beads, covered with ANCHOR fragments, are incubated for 30 min with the TARGET containing solution. In the herein reported proof of concept, this fragment was designed to be 36 bp long. After three washing steps with Hepes/NaCl buffer, the unbound TARGET molecules and every other component possibly present in the matrix were washed out.

3) The 5′ portion of the TARGET fragments bound to the ANCHOR can recognize a third fragment hereinafter called PROBE. This third fragment, functionalized with a cholesterol moiety at its 3′ end, was previously intercalated in the phospholipidic bilayers of a liposome loaded with a highly concentrated solution of a fluorescent dye. The high concentration of the dye inside the liposome aqueous cavity causes an extensive “quenching” of the detectable fluorescence.\textsuperscript{23}

4) After washing, to amplify the fluorescence signal, liposomes are destroyed with Triton X-100. The signal detected is directly proportional to the amount of the TARGET fragment bound to magnetic beads through ANCHOR.

\subsection*{Liposome Characterization}

The used liposomes are prepared with 100% DPPC, a saturated phospholipid that ensures good membrane stability, avoiding leakage of the fluorescent molecule from the liposome core. The average diameter measured by dynamic light scattering (DLS) was of 31 nm with a PDI value of 0.106 ± 0.007. Among the commercially available dyes, CF was chosen due to its high solubility in water, its high quantum yield (\(\Phi = 0.92\)),\textsuperscript{24} its low

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Assay proof of concept: step 1: Streptavidin Mag Sepharose beads interact with the biotinylated ANCHOR fragment. Step 2: 18 nucleotides of the ANCHOR fragment recognize specific complementary sequence on the TARGET fragment. Step 3: cholesteryl-tagged PROBE fragment, conjugated with 50 mM CF liposome, interacts with the free 18 bp long portion of the TARGET fragment.}
\end{figure}
commercial cost, and the compatibility with a wide array of detectors commonly present in analytical laboratories. The maximum loading CF concentration was set at 50 mM. Higher CF concentrations were tested (e.g., 75 and 100 mM), but the resulting liposomal formulations, which display a PDI value higher than 0.5, were not used because of the occurrence of high instability. When a liposome membrane was destabilized by Triton X-100, the CF core content is released, and the fluorescent signal markedly increases (Figure 2A).}

![Figure 2. (A) Fluorescence intensity measured at increasing concentrations of SR-loaded (\(\lambda_{ex} = 565\) nm, \(\lambda_{em} = 585\) nm) and CF-loaded liposomes (\(\lambda_{ex} = 495\) nm, \(\lambda_{em} = 517\) nm) before and after the Triton X-100 addition. (B) Bar plot comparing fluorescence intensity at the lowest liposome concentrations. Error bars indicate standard deviation (\(N = 4\)).](image)

CF-loaded liposomes where compared with liposomes of the same membrane composition but carrying up to 150 mM sulforhodamine B (SR-B) inside their aqueous core. Liposomes loaded with 150 mM SR-B have been previously reported as suitable probes for sandwich hybridization assays for nucleic acid markers.\(^{20}\) Figure 2 shows that a 3-fold increase in fluorescence intensity was observed when CF liposomes were treated with Triton X-100. The concentration of the liposomal nanoparticles was determined by using the following equation

\[
[\text{liposomes}] (\text{nM}) = \frac{[\text{CF}]_{\text{tot}} \times 3.97 \times 10^8}{[\text{CF}]_{\text{hydr}} \times (r - S)^3}
\]

Further information about this equation is reported in the Supporting Information.

Although sulforhodamine B-loaded liposomes showed a higher fluorescence increase when treated with Triton X-100, their minimum detectable liposome concentration was 2 orders of magnitude higher than the detection threshold determined for CF-loaded liposomes (10\(^{-15}\) vs 10\(^{-17}\) M for SR-B liposomes and 10\(^{-17}\) M for CF liposomes, as determined through subsequent dilutions in Hepes/NaCl buffer; Figure 2B). Even though SR-B concentration inside liposomes is three times higher than CF, the improved performance shown by CF-loaded liposomes might rely on the different quantum yields of the internalized fluorescent compounds.\(^{18,25}\) These findings state that CF-loaded nanoparticles are more sensitive at lower concentrations, giving the possibility to investigate a very low amount of the target ssDNA and possibly detecting nucleic acids without the need of an amplification procedure. For these reasons, CF was used in the further experiments to set up the proposed assay.

The occurrence of a stable intercalation of cholesteryl-tagged ssDNA (PROBE) into the phospholipid bilayer was assessed using high-resolution nuclear magnetic resonance (NMR).

The specific peaks, characteristic of nucleotide sequences, appeared only in the \(^1\)H-NMR spectrum of liposomes incubated with the PROBE segment, while any nonspecific signals were detected in the control ones (Figure S1).

Moreover, the number of dialysis cycles needed to eliminate the nonencapsulated CF from liposome preparation was evaluated. It was reported that six dialysis cycles (4 h at 4 °C for each cycle) are necessary (Figure S2). In concentrated suspension, the stability of liposomes has been assessed by measuring the fluorescence signal intensity of intact liposomes and Triton X-100-treated liposomes. When CF is released from liposomes, a fluorescence enhancement occurs (because of the “dequenching” of leaked CF). As reported in Figure S3, liposomes are stable up to 4 months.

**Procedure Setup and Assay Detection Limit.** The experimental setup takes less than 2 h (Figure S4). During the first 30 min, magnetic beads, ANCHOR, and TARGET fragments were mixed to carry out the first annealing step. During the following 30 min, incubation in the presence of the PROBE fragment allowed for the fluorescent labeling of the TARGET. Figure 3 shows the linear range of correlation between fluorescence intensity and TARGET concentration (Pearson correlation coefficient = 0.966), with an LOD of 7 × 10\(^{-11}\) M.\(^{26}\) The entire curve of fluorescence intensity versus

![Figure 3. CF-loaded liposome fluorescence intensity measured after the addition of increasing TARGET concentrations to the Strep Mag and ANCHOR mixture. The blue line indicates the background fluorescence intensity measured without the TARGET (purple lines show standard error of the mean relative to the noise signal). The red line was obtained by linear fitting of the data. Error bars indicate standard deviation (\(N = 10\)).](image)
[DNA Target] is reported in Figure S5, and the parameters of the calibration curve are reported in Table S1.

After the washing steps, liposomes bound to Strep Mag beads through the DNA linkage showed a fluorescence intensity that is not distinguishable from the fluorescence signal of buffer (Figure 3, green dots). When treated with 0.5% Triton X-100, fluorescence intensity of the samples showed an increment directly correlated with the TARGET concentration (Figure 3, black diamonds). Few liposomes were able to bind to Strep Mag beads in a nonspecific manner when the TARGET fragment is not present, producing a low interference (Figure 3, blue line).

Next, the possibility to further improve the detection limit using microcuvettes was explored. The advantage of using microcuvettes relies on an ~7times reduction of the final volume, thus allowing the increase of the dye concentration, which can allow the discrimination of lower concentrations of TARGET (e.g., $2 \times 10^{-11}$ M; Figure 4) not detectable by using the standard cuvettes (as shown by P values obtained by using Dunnett’s one-way ANOVA).

![Figure 4](image)

**Figure 4.** Microcuvettes assay. Error bars indicate standard deviation ($N = 4$).

The same experiment was repeated using SB-R-loaded liposomes to assess if the use of this dye makes it possible to reduce the background fluorescence signal. The results, as reported in Figure S6, confirmed the lower performance of this dye. In fact, the LOD was 2 orders of magnitude higher than that of CF-loaded liposomes ($2 \times 10^{-10}$ M).

Figure 5 shows that SCRAMBLE fragments are not able to link the liposome and the magnetic components. In fact, the observed fluorescence signal was not distinguishable from the background in the absence of the TARGET. Furthermore, by maintaining the scramble-36 and scramble-100 in a 1:1 ratio and varying the TARGET molarity, no significant interference in the binding of TARGET to the ANCHOR fragment was noted, even when the SCRAMBLE:TARGET ratio was 1000:1. In fact, the linearity of the detection of the assay calibration curve did not vary.

**Assay Behavior in the Presence of Double-Stranded TARGET.** The assay response was also evaluated with double helix conformation of the TARGET fragment (d$\text{S}$TARGET). To obtain d$\text{S}$TARGET, the single-stranded TARGET was incubated with its complementary reverse fragment, heated to 85 °C, and then let to cool at room temperature to favor the correct annealing of the two fragments. The formation of the dsDNA has been assessed through agarose gel electrophoresis (Figure S7).

![Figure 5](image)

**Figure 5.** (A) Fluorescence intensity of the solution containing increasing amount of TARGET and a fixed $2 \times 10^{-11}$ M scramble mix, added to the Strep Mag ANCHOR mixture. (B) TARGET concentration is normalized, and signal intensity of the solution obtained after incubation of TARGET alone is set to 100%. Signal intensities in the presence of scramble mix are proportionally calculated and compared to the corresponding TARGET fluorescence signal. Error bars indicate standard deviation ($N = 3$).

To allow the assembly of the ANCHOR and PROBE components to the transient ssTARGET fragments, the samples were heated at 82 °C for 5 min, and then, the temperature was rapidly decreased in liquid nitrogen for 10 s. To avoid the annealing of the two complementary fragments and favor the binding of the assay components, the d$\text{S}$TARGET was incubated together with the ANCHOR and PROBE fragments in a 1:100 molar ratio. Finally, the liposome was added as the latest component due to its instability at high temperatures. No significant differences in signal intensity among the samples incubated at room temperature with ssTARGET and the ones incubated with d$\text{S}$TARGET were observed nor among the samples heated to 82 °C (Figure 6). Adding the Strep Mag beads after the rise in temperature did not alter the obtained signal intensity, thus suggesting that the overall decrease might be related to different causes and not the degradation of the magnetic beads (data not shown).

**Assay Behavior in Serum.** The effect of different biological matrices, among which is human serum, was tested with the aim of investigating the efficiency of the proposed DNA assay. The ssTARGET was dissolved in human serum before performing the already described assay. In serum, the assembly of the probe appears to be greatly hampered (Figure 7 and Figure S8), and the assay detection limit was 2 orders of magnitude higher, setting the LOD to $0.8 \times 10^{-9}$ M. These findings led us to investigate the possibility of unwanted interactions among serum and assay components, in particular,
The herein described assay allows the detection of both single- and double-stranded DNA, measuring the signal arising from a fluorescent dye (carboxylfluorescein) encapsulated into liposomes at high concentration and released upon treatment with detergents (i.e., Triton X-100). The optimized liposomes are stable for a long period of time (~4 months) when stored at high concentration at 4 °C. The LOD of target DNA is $7 \times 10^{-11}$ M, making it possible to investigate specific nucleotide sequences at very low concentrations. Moreover, in comparison to multistep coupling procedures, the incorporation of a cholesteryl-modified ssDNA probe was allowed to obtain targeted liposomes without using expensive coupling reagents and time-consuming coupling procedures.21,28 Most of the commonly used techniques (among which are ddPCR and BEAMing) and sensors for DNA/RNA detection in liquid biopsies30,31 have received an in-depth validation for clinical and preclinical applications.30,31 Compared to these techniques, the herein proposed method shows the following advantages: (i) fast assay (execution time, approximately 2 h); (ii) simplicity of execution, while both ddPCR and BEAMing need a medium—high level of expertise to be performed and require the presence of loader and reader machineries, which have an elevated price, whose purchase may not be justified for laboratories not fully dedicated to routinely screen for DNA markers; (iii) the decrease of biases derived from multiple amplifications of the low amount of starting nucleotidic material. Moreover, it is worth to mention the improvements with respect to the method reported by Baumner et al.,22 namely, (i) a lower LOD (2 orders of magnitude); (ii) the use of a more efficient fluorescent dye (carboxylfluorescein); (iii) the use of Triton X-100 to obtain a fluorescence dequenching; and (iv) the use of magnetic beads instead of immobilized surfaces that have been proven to be very efficient for this kind of quantitative assays.3 As previously reported, plasma concentration values of cell-free DNA in melanoma and lung cancer patients are ∼36 and 31 ng/mL plasma, respectively.33 The fragment length range is 134–144 bp.3 Thus, using 650 g/mol as the average molecular weight for a base pair, we can theorize that the plasma ctDNA concentration range for these patients is $1 \times 10^{-12}$ to $6 \times 10^{-10}$ M. Considering that the fraction of mutant DNA ranges from 0.005 to 0.1% of the total DNA concentration in plasma,34 the expected amount of mutated DNA in serum is in the $10^{-13}$ to $10^{-15}$ M range. In the case of colon carcinoma, the amount of mutant ctDNA increases dramatically, reaching a value of ∼8%.31,35 This value is of the order of $10^{-11}$ M, that is, in the same order of magnitude of the LOD of the herein proposed assay, when the latter is performed in Hepes buffer. In principle, by exploiting a purification method to extract ctDNA from their biological matrices, it would be possible to detect and quantify nucleic acid markers without further amplification. In principle, the LOD of the herein reported method can be further reduced (i.e., enhanced assay performance) by using larger liposomes (e.g., medium unilamellar vesicles or giant liposomes), that is, containing a higher amount of fluorescent dye encapsulated in the inner core and/or by using a fluorescent dye with a higher quantum yield. Furthermore, other items have to be considered, especially for measurements in serum-containing specimens. First of all, the role of DNases in degradation of the target DNA has to be evaluated. For instance, serum DNA extraction protocols (based on phenol–chloroform extraction

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**CONCLUSIONS**

Figure 6. Comparison of assay behavior in the presence of double- and single-stranded TARGET DNA (means are not significant among temperature-treated and not treated samples, $P$ value > 0.05). Error bars indicate standard deviation ($N = 3$).

![Image](image-url)

**Figure 7.** Assay behavior in Hepes, Hepes + HSA, and human serum: fluorescence intensity (a.u.) was measured after the addition of two concentrations of DNA TARGET ($3.3 \times 10^{-10}$ and $2.0 \times 10^{-9}$ M) in three different media ($P$ value > 0.05). Error bars indicate standard deviation ($N = 3$).

with albumin, which is the most abundant serum protein. However, fluorescence measurements demonstrated that the presence of albumin did not affect the assembly of the components and the fluorescence signal intensity (Figure 7).

Further investigations were carried out to determine the role of phosphates on the decrease of assay performance (i.e., increased LOD) in serum. Thus, the assay components were assembled in PBS buffer prepared with 1 and 10 mM phosphate concentrations. As seen from albumin, no significant decrease in the detected fluorescence intensity of the assay was observed, suggesting that phosphates do not play a role in the assembly of the deoxyribonucleic nor protein components (Figure S9). Finally, serum was inactivated through heating for 25 min at 55 °C and the assay performed as previously stated. No significant differences among not treated and treated sera have been observed, suggesting that the complement is not responsible for the decrease of the assay performance in this complex biological matrix (Figure S10). This can be due to the activity of DNase I, one of the components of the circulating free DNA (cfDNA) elimination system.27

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**ACS Omega**

DOI: 10.1021/acsomega.9b01457
ACS Omega 2020, 5, 1764−1772

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1768
or pull-down techniques) can be employed to check whether this may contribute to the loss of sensitivity.

The herein reported method appears suitable, upon proper optimization, for the screening of circulating cancer DNA. The developed assay is versatile. In fact, by maintaining the proposed scheme (liposomes formulation, magnetic beads, etc.), it is possible to properly design the DNA sequences of ANCHOR and PROBE to specifically recognize TARGET sequences. Moreover, considering previously reported works, it appears possible to expand the herein reported method to the detection of longer (>36 bp) target DNA molecules.36,37

This is, in principle, applicable to the detection of DNA circulating in biological fluids (not only circulating tumor DNA). As an example, one may think of a number of pathogenic microorganisms that release their DNA in the bloodstream, thus opening the possibility to exploit their nucleic acids as markers for characterizing infectious diseases. Setting the nucleotidic sequence of the ANCHOR and PROBE fragments on a specific target sequence, the possibilities in the different markers detectable are limited only by the assay LOD. Finally, future investigations could also lead to tune and validate the assay for the detection of RNAs, another class of diagnostic and prognostic markers rising in interest nowadays.58 This class of molecules is more difficult than DNAs to be analyzed since RNAs are more prone to enzymatic degradation and because of the complexity and biases present in RNA amplification protocols.

**EXPERIMENTAL SESSION**

**Materials.** DPPC (1,2-dipalmitoyl-sn-glycero-3-phosphatidylcholine) was purchased from Avanti Polar Lipids (Alabaster, Alabama, USA). Cholesterol, sodium chloride, sodium monobasic phosphate, sodium dibasic phosphate, potassium chloride, Hepes, chloroform, and all other chemicals were purchased from Sigma-Aldrich Inc. (Merck KGaA, Darmstadt, Germany). DNA oligonucleotides were purchased from Eurofin Genomics. Lyophilized human serum (Seronorm) was purchased from SERO AS, Norway.

**Liposome Preparation.** Large unilamellar vesicles (LUV) liposomes were prepared according to the classical thin lipid film hydration method as previously reported.39-41 Briefly, 45 mg of DPPC (1,2-dipalmitoyl-sn-glycero-3-phosphatidylcholine) were dissolved in a chloroform solution in a 250 mL glass balloon. The chloroform has been subsequently evaporated by using a rotavapor device in vacuum conditions at 60 rpm at room temperature (RT). The lipid film formed after the solvent evaporation was left stirring for 1.5 h at RT to remove the presence of any water contamination. Subsequently, 1.5 mL of 50 mM carboxylfluorescein solution, in Hepes buffer (0.15 M NaCl, 3.8 mM Hepes, pH 7.4, ~320 mM/s, L), was added to the lipid film. Then, it was heated through immersion in a water bath at a temperature of 56 °C, which corresponds to the transition phase temperature of DPPC phospholipids. The bottom of the balloon was vortexed at max speed to ensure the detachment of the film from the glass wall of the recipient. These heating and vibrating steps were performed multiple times until the complete lipid film detachment. The formed solution was then introduced in an extruder device, heated at 56 °C, and under argon pressure, the vesicles were forced to pass through a 400 nm filter membrane. After eight runs, the filter membrane was changed with a 200 nm one, and other eight runs were performed. To eliminate the unloaded CF molecules, the LUVs’ suspension (1.5 mL) underwent extensive dialysis cycles against isotonic Hepes/NaCl buffer solution (2 L) at 4 °C by using a 14 kDa dialysis cellulose membrane. The external Hepes solution was replaced every 4 h for six cycles. After purification, the liposome size was determined by using dynamic light scattering (DLS) technology with a Zetasizer Malvern instrument. Liposomes were then stored at 4 °C in the absence of light to avoid photobleaching of the fluorescent molecules.42 Under these storage conditions, liposomes remained stable for at least 4 months.

**Cholesteryl—DNA-Tagged Liposome Preparation.** DPPC liposomes (1 × 10^-8 M) containing 50 mM CF were diluted 1:10 in sterile Hepes buffer (150 μL of liposome solution in 1.35 mL of Hepes buffer). PROBE DNA was then added to this solution in a 300:1 PROBE/liposome ratio, and the resultant solution was incubated for 1.5 h on a shaking ThermoMixer at 4 °C in slow speed in the absence of light.

Subsequently, the solution was dialyzed overnight against 2 L of Hepes buffer in a 16 kDa MWCO dialysis membrane at 4 °C in dark conditions to ensure the elimination of unbound PROBE molecules. After the dialysis, the functionalized liposomes were stored at 4 °C in dark conditions.

**NMR Characterization.** Liposome specimens (20 μL) have been dissolved in CHCl3 into a 125 mL glass balloon. This procedure allows destroying liposomes and dissolving all the apolar molecules. Then, CHCl3 was evaporated using a rotavapor, and the specimen dissolved into deuterated methanol.

3H NMR spectra of specimens have been acquired at 4 °C using a Bruker 600 MHz spectrometer (AVANCE 600, 14 T).43

**ssDNA Molecule Design.** ssDNA molecules were purchased from Eurofin Genomics. A quality certificate describing yield and purification steps of the ordered oligonucleotide was delivered by the company together with the product. ssDNAs were artificially designed to be representatives of DNA sequences occurring in biological fluids. They have been designed through MeltCalculator, a Biophysics Integrated DNA Technologies online tool, through which the presence of self-dimers, hairpins structures, and annealing temperatures have been calculated.

ssDNAs used in our experiments have the following sequences:

TARGET: 5′-CTG GAC TGT TGA TTG GAT TGA TAG GGC GAC TGC 3′

ANCHOR: 5′-biotin-TGG-GCA GTC GCC CTA TCA ATC-3′.

PROBE: 5′-CCA CAA TCA ACA GTC CAG-cholesterol-3′.

RT (reverse target): 5′-GCA GTC GCC CTA CTA ATC CCA CAA TCA ACA GTC CAG-3′.

S36 (scramble-36): 5′-AGC CGA TCT TAA TGG ATG GCC GCA GGT GTG ATG GAA-3′.

S100 (scramble-100): 5′-CAA CGG CAT GGG CAA CTG GTG AAG TGC CTA CTA TCC TAA AAC GCA TAT CTC GCA CAG TAA TCC CCC AAT ATG TGA GCA TCT GAT GTT GCC CGG GCC GAG T-3′.

Lyophilized ssDNAs were rehydrated with sterile Tris-EDTA solution to obtain a final concentration of 100 μM. Purity of the shipped ssDNA samples was guaranteed by the
manufacturer company and subsequently double-checked in our lab through Nanodrop analysis.

Double-stranded TARGET DNA has been obtained through mixing the two complementary fragments TARGET and REVERSE TARGET in equimolar amounts, heating to 82 °C, and letting it cure at room temperature to assure the correct annealing of these two ssDNAs.

**Assay Protocol.** Sterile Hepes buffer (3 mL) was added to a 15 mL Falcon tube, and 3 μL of streptavidin-coated magnetic beads were added (Streptavidin Mag Sepharose, GE Healthcare S.P.A.). A 3 μL ANCHOR ssDNA 100 μM solution was added to this solution together with the desired amount of TARGET ssDNA. The Falcon tubes were placed on a shaker, and the beads and DNA mix were incubated for 30 min at room temperature to permit the binding of all the probe components. The tubes were then centrifuged for 2 min at 3000 rpm, and the magnetic beads were retained through a magnet. The solution was then eluted together with all the unbound probe components, and three washing steps were performed using Hepes buffer. Cholesterol–PROBE-tagged liposome (100 μL) was added to the 3 mL Hepes buffer solution containing the beads and the components bound from the previous step. This solution was placed for 30 min on a shaker at medium speed and 4 °C in dark conditions. After this incubation, the tubes were again centrifuged for 2 min at 3000 rpm. Washing steps were performed to ensure that the elimination of all the elements was not bound to the magnetic beads until basal fluorescence intensity was comparable with the one present in Hepes buffer.

Triton X-100 (0.5%; Sigma-Aldrich) was added to the liposome solution. After 15 min, the total fluorescence intensity was analyzed with a spectrofluorimeter (FluoroMax-4, Horiba Jobin Yvon). Spectrofluorimetric excitation and emission parameters were set, respectively, at wavelengths of 492 and 517 nm, wavelengths for carboxyfluorescein molecules, and at 565 and 586 nm for sulforhodamine B.

In the experiments performed in serum, Strep Mag beads, ANCHOR, and TARGET ssDNA were directly added to the serum (human serum, Seronorm SERO AS) and positioned on a shaker in slow speed at RT for 1 h. Then, the samples were treated as described above with a centrifugation step, three washing steps in Hepes buffer, and the addition of DNA–cholesterol-tagged liposomes.

**Data Analysis and Statistic.** Data have been obtained by using at least three independent experiments. The exact number of replicates has been inserted in each graph.

Data have been reported as mean ± SD.

Pearson correlation coefficient of data in Figure 3 has been evaluated.

Unpaired two-tailed Student t test has been carried out in Figures 5 and 6.

Dunnett’s (one-way ANOVA) statistical test has been applied for the data in Figures 2B and 4. One-way ANOVA test with Bonferroni correction has been applied for the data in Figure 7.

The following legend has been added to the figures: *P value < 0.05; **P value < 0.01; and ***P value < 0.001.

LOD has been calculated by using the following equation: LOD = 3σ/m, where σ is the SD of blank and m is the slope of linear fitting of data in the linear range.

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**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.9b01457.

Calculation of liposomes’ concentration; NMR spectra of liposomes (Figure S1); assessment of the optimal number of dialysis cycles for purification of CF liposomes (Figure S2); assessment of liposomes stability over months (Figure S3); assessment of fluorescence intensity by using variable incubation time (Figure S4); CF-loaded liposome fluorescence intensity measured after the addition of increasing TARGET concentrations to the Strep Mag and ANCHOR mixture, entire dataset (Figure S5); DNA assay carried out by using 150 mM sulforhodamine containing liposomes (Figure S6); 3% agarose electrophoresis gel of TARGET DNA (Figure S7); calibration curve performed in human serum (Figure S8); comparison among DNA assay carried out in Hepes buffer, 1 mM PBS buffer, and 10 mM PBS buffer (Figure S9); comparison among DNA assay carried out in Hepes buffer, in normal serum, and in inactivated serum (Figure S10); calibration curve parameters (Table S1); comparison of LOD by using buffer, serum or buffer, and microcuvettes (Table S2); and supplementary references (PDF).

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**Author Contributions**

The manuscript was written through contributions of all authors. All authors have given approval to the final version.

**Notes**

The authors declare no competing financial interest.

**ACKNOWLEDGMENTS**

This project has received funding from the European H2020-INFRADEV-2014-2015 project CORBEL (grant no. 654248).

**ABBREVIATIONS**

BEAMing, beads, emulsification, amplification, and magnets; bp, base pair; CF, carboxyfluorescein; cDNA, circulating free DNA; ctDNA, circulating tumor DNA; ddPCR, digital droplet PCR; DLS, dynamic light scattering; DPPC, 1,2-dipalmitoyl-sn-glycero-3-phosphatidylcoline; dsTARGET, double-stranded TARGET; LUV, large unilamellar vesicles; NMR, nuclear magnetic resonance; PCR, polymerase chain reaction; rpm, rotation per minute; RT, room temperature; RT-PCR, reverse transcription PCR; SR-B, sulforhodamine B; ssDNA, single-stranded DNA; ssTARGET, single-stranded TARGET

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