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Two-step formulation of magnetic nanoprobe for microRNA capture

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MicroRNAs (miRs) belong to a family of short non-coding endogenous RNAs. Their over-expression correlates with various pathologies: for instance, miRNA-155 (miR-155) is over-expressed upon the development of breast cancers. However, the detection of miRs as disease biomarkers suffers from insufficient sensitivity. In the present study, we propose a protocol for a rapid and efficient generation of magnetic nanoprobe able to capture miR-155, with the aim of increasing its concentration. As a nanoprobe precursor, we first synthesized superparamagnetic iron oxide nanoparticles (SPIONs) coated with covalently attached polyethylene glycol carrying a free biotin terminus (PEG-bi). Using streptavidin–biotin interactions, the nanoprobe were formulated by functionalizing the surface of the nanoparticles with the miR sequence (CmiR) complementary to the target miR-155 (TmiR). The two-step formulation was optimized and validated using several analytical techniques, in particular with Size-Exclusion High Performance Liquid Chromatography (SE-HPLC). Finally, the proof of the nanoprobe affinity to TmiR was made by demonstrating the TmiR capture on model solutions, with the estimated ratio of 18 : 22 TmiR : CmiR per nanoprobe. The nanoprobe were confirmed to be stable after incubation in serum.

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1. Introduction

MicroRNAs (miR) are small non-coding RNAs with lengths ranging from 19 to 23 nucleotides. MiRs act as regulators in post-transcriptional gene expression by targeting specific messenger RNAs. They play a major part in various biological processes, which are influenced by dysregulation in miR expression levels.^{1,2}

As such, miRs are considered as novel biomarkers for diagnostics and therapeutics of various diseases (*e.g.*, numerous types of cancer, neurological and muscle degenerative diseases, cardiovascular disorders, *etc.*). A large part of the scientific research is focused on the atypical expression of miRs present in different types of cancer. Depending on their altered expression, miRs can be oncogenic (increased expression leading to inhibition of tumor suppressor genes and subsequent malignancy), or tumor-suppressing (obstructing the expression of oncogenes).³ The disruption of the balance between these two types impacts cellular development, leading to cancer development. As the expression levels of multiple miRs either increase or decrease compared to the healthy tissue, the focus is put on developing new methods with better sensitivity and detection limits.^{4,5}

One of the popular analytical methods widely used for purity determination and batch-to-batch comparison of compounds (miRs included) in pharmaceutical use is Size Exclusion High Performance Liquid Chromatography (SE-HPLC). It is built on separating the mixture in a solution into smaller components. A liquid mobile phase passes through the column filled with particles of a certain size, *i.e.*, stationary phase. As the solution is pumped through the column, the components of the solution interact differently with the surface of the particles of the stationary phase and are separated in the process, coming out of the column at different and specific retention times. Commercially available miRs are purified by HPLC, in order to eliminate shorter lengths species originating from the possible degradation. Precise determination of separation, isolation, and purification of nucleotides from biological samples still requires a lot of effort, and there are limits for qualitative analysis abilities by the means of quantification and identification.^{6–8}

Beyond generally used methods for miR detection (qRT-PCR,^{9,10} Northern blotting,¹¹ next-generation sequencing,^{12,13} microarrays,^{14,15} *etc.*), other approaches based on nanomaterials are becoming progressively more popular, either in combination with a well-known method or by developing altogether new ones.¹⁶

Methods and approaches of miR detection based on nanotechnologies offer an alternative path to a faster and less complicated detection process. Nanomaterials used can be in

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a form of nanoparticles, colloids, substrates, *etc.* They provide unique optical, electrical, and magnetic properties,¹⁷ and most importantly, high surface area, leaving room for a plethora of surface-active chemical groups, which can be utilized in the subsequent immobilization of target biomolecules. The sensing application and efficiency of nanomaterials have been demonstrated.^{18,19} Specific challenges include reproducibility, standardization, normalization, and data processing. Surprisingly, a significant part of the reported studies did not deal with natural bodily fluids (serum, blood, saliva, *etc.*) in the presence of possibly interfering biomolecules.²⁰

One possible way of using nanoprobe is to capture on their surface the target miR (TmiR) present in the biological fluid. The TmiR capture efficiency and specificity can be enhanced by using the nanoprobe surface decorated with a complementary miR sequence²¹ (hereafter called CmiR), double/single-stranded DNA,²² or sequence of oligonucleotides.²³ Thus, successful design, production, and use of the TmiR-specific nanoprobe depend on several important considerations such as (i) synthesis of the nanomaterial (core), (ii) surface modification (platform), and (iii) surface functionalization (specific activation).

Hybrid nanoprobe (inorganic core–organic shell) are of high interest since their utilization allows to take benefit from unique physical properties of the inorganic core while controlling the stability due to the organic shell.²⁴

The core–shell nanoprobe chosen for this study are based on the hybrid nanoparticles developed by our group and made of superparamagnetic iron oxide nanoparticles (SPIONs) coated with polyethylene glycol (PEG).²⁵

Our working hypothesis was that the TmiR present in biological samples can be efficiently captured on the surface of the nanoprobe functionalized with specific CmiR and then the TmiR can be reconcentrated by magnetic sorting (Fig. 1A), thus favoring their more sensitive detection. PEGylation of SPIONs is used for enhancement of their biocompatibility and colloidal stability in aqueous media while preserving relative surface neutrality. We have selected a PEG chain modified with *N*-succinimidyl ester (NHS) on one end, and with biotin on the other (hereafter called PEG-bi), for further conjugation *via* streptavidin–biotin affinity interactions.²⁶ Streptavidin- and biotin-modified materials are widely commercially available, for either the nanomaterial matrix or the biomolecule shell. Streptavidin-coated magnetic nanoparticles,²⁷ microparticles, or beads are very popular: for instance, they were exploited for detection of miR-144 (ref. 28) (*via* fluorescence signal amplification), or miR-21 (ref. 29) (*via* liquid chromatography-electrospray ionization tandem mass spectrometry). In our study, we formulated the nanoprobe starting from the PEGylated SPIONs and using streptavidin as a linker between the biotinylated nanoparticle surface and biotinylated CmiR (Fig. 1B). The idea is that the PEGylated SPIONs play a role of a versatile nanoplateform that can be used to easily formulate nanoprobe decorated with various CmiRs and therefore adaptable to various TmiRs.

After synthesis and characterization of SPION–PEG-bi nanoplateforms, we formulated and characterized several

batches of the nanoprobe decorated with a model CmiR, specific to TmiR-155 upregulated in breast cancer cells and one of the most commonly reported miRs in body fluids.^{1,30}

Several physicochemical characteristics of the nanoprobe were established by means of a spectrum of analytical techniques: nanoparticle size (hydrodynamic diameter – D_H by DLS-dynamic light scattering) and chemical composition (iron concentration by AAS – atomic absorption spectrophotometry, organic shell identification by IR spectrophotometry, miR concentrations by SE-HPLC). The results confirmed the successful synthesis of the SPION–PEG-bi nanoplateforms and allowed us to optimize the nanoprobe formulation in terms of their surface saturation with streptavidin (formulation step 1) and of their functionalization with CmiR (formulation step 2). The function of synthesized nanoprobe was confirmed by trying out their ability to capture TmiR-155, and their stability was tested by an incubation in serum.

2 Experimental

2.1 Materials

Deionized water (18 M Ω) was used in all experiments. α -Biotinyl- ω -NHS PEG, M_w 5000 Da, (NHS–PEG₅₀₀₀–biotin, PEG-bi) was purchased from Rapp Polymere (Tübingen, Germany). Streptavidin in lyophilized form, CmiR-155 modified with biotin ([Bt_n]-5'-ACCCCUAUCACGAUAGCAUUA-3', 7674 g mol⁻¹) and TmiR-155 (5'-UUAUUGCUAUCGUGAUAGGGGU-3', 7391 g mol⁻¹) were purchased from Sigma-Aldrich (Saint Quentin-Fallavier, France). Standard miR solutions at required concentrations were prepared by adding RNase-free deionized water.

2.2 Synthesis of the nanoprobe precursors – SPIONs coated with PEG-bi (NP)

The synthesis of PEG-biotin-coated SPIONs was divided into 3 steps: (i) the coprecipitation of initial SPIONs, (ii) the silanization of SPION's surface, and (iii) the covering of silanized SPIONs with the PEG-bi layer.

The synthesis of initial SPIONs by improved Massart's method and the silanization of SPIONs *via* surface amination with 3-aminopropyl-trimethoxysilane (APS) were performed according to the protocol published by our group.²⁵ Afterwards, the surface of the particles was PEGylated by the following procedure.

A mixture of 1.6 mL of silanized SPIONs (0.14 mmol of iron) and 10 mL of DMSO was prepared, and the solution was dehydrated in a vacuum for 5 h with a rotary evaporator. Lyophilized NHS–PEG₅₀₀₀–biotin (220 mg, 0.04 mmol) was dissolved in 5 mL of dehydrated DMSO and added to the SPIONs mixture. The suspension was stirred for 24 h at the room temperature and afterward purified by dialysis (MWCO 1000 kDa) against water at the room temperature for 48 h.

2.3 Formulation of the nanoprobe

The formulation was made in two steps: (i) modification of the NP surface with streptavidin to result in NP–STR; and (ii)



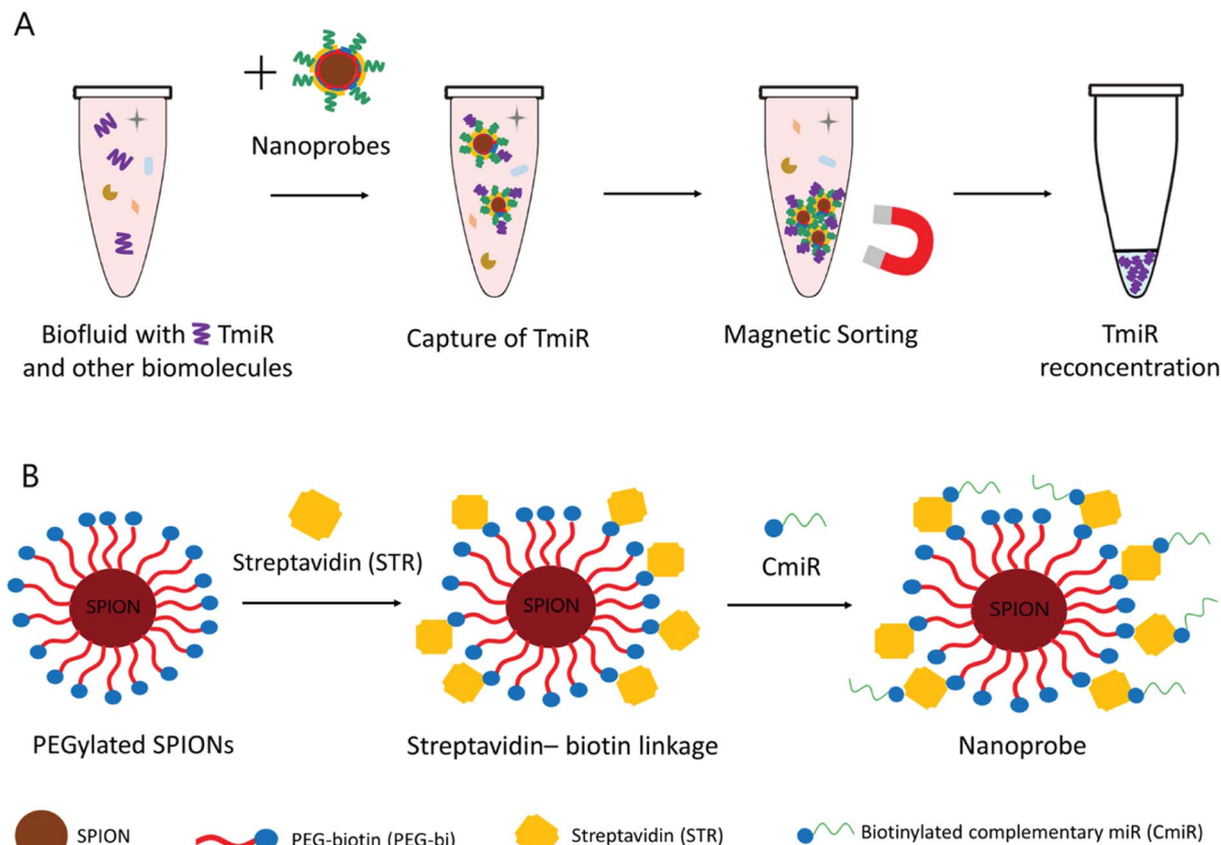


Fig. 1 (A) TmiR capture and reconcentration in a biofluid. (B) Schematic representation of the nanoprobe formulation from the PEGylated SPIONs.

functionalization of the NP-STR with CmiR to result in NP-STR-CmiR.

2.3.1 Streptavidin-modified nanoparticles (NP-STR). For modification of PEGylated SPIONs with STR, we used the interaction between STR and the biotin functional groups already present on the PEG-bi shell of the NP. To optimize the STR coating, we studied three batches of NP-STR with Fe/STR molar ratios of 280 (NP-STR1), 1400 (NP-STR2), and 2100 (NP-STR3).

NP-STR1 (Fe/STR = 280 molar ratio) was generated by mixing 400 μL of NP suspension at an iron concentration of 350 mg L^{-1} with 500 μL of 1 mg mL^{-1} STR and 100 μL of PBS 10X. The other two batches, NP-STR2 (Fe/STR = 1400) and NP-STR3 (Fe/STR = 2100) were generated accordingly.

All the samples were shaken overnight at room temperature. The resulting NP-STR were purified by magnetic separation (protocol described in Section 2.6) in order to eliminate an excess of streptavidin.

2.3.2 NP-STR functionalized with C-miR (NP-STR-CmiR). The batch NP-STR1 was selected as optimally modified with STR and was further functionalized by incubating it for 2 hours with 1, 5, or 10 μM aqueous solution of biotinylated complementary CmiR sequence. According to the CmiR concentration used, the resulting batches were named respectively: NP-STR-CmiR1 (1 μM CmiR); NP-STR-CmiR5 (5 μM CmiR); NP-STR-

CmiR10 (10 μM CmiR). The final molar ratio Fe/CmiR in those batches was respectively 1.9, 4.75, and 9.5. The resulting NP-STR-CmiR were purified by magnetic separation (protocol in Section 2.6) in order to eliminate an excess of CmiR.

The number of CmiR molecules per nanoparticle (N) was determined with the following formula:

$$N = C \times \frac{\pi D^3}{6} \times d \times 6.10^{23}$$

with C : the amount of CmiR in mol per g of iron; D : diameter of SPION core (m , as defined by TEM); d : mass density of magnetite = $5.2 \times 10^6 \text{ g m}^{-3}$; Avogadro's number = $6 \times 10^{23} \text{ mol}^{-1}$.

2.4. TmiR capture by nanoprobes

The batch of nanoprobes NP-STR-CmiR5 (5 μM CmiR) was combined with the solution of TmiR to reach a 5 μM final concentration of TmiR and incubated for 2 hours. The resulting sample NP-STR-CmiR-TmiR was purified by magnetic separation (protocol in Section 2.6) in order to eliminate any free TmiR.

2.5 Stability of nanoprobes

The batch of nanoprobes NP-STR-CmiR5 (5 μM CmiR) was combined with the solution of fetal bovine serum (FBS)



(Eurobio, Les Ulis, France) in the ratio 1 : 1, and incubated at room temperature for 4 and 12 hours. The resulting samples (S4)-NP-STR-CmiR5 and (S12)-NP-STR-CmiR5 were purified by magnetic separation (protocol in the Section 2.6).

2.6 Magnetic separation

Magnetic separation (MS) was performed with MS columns (MACS® Column Technology, Miltenyi Biotec GmbH, Germany). The volume of the SPIONs suspension (up to 500 μL , depending on the initial volume of the sample) was applied into a MS column held in a magnetic MACS separator, and the separated liquid was collected. The column was then washed three times with the buffer (PBS 1X). The column was removed from the magnetic separator and flushed with the buffer (75% of the initial volume) resulting in a sample, utilized in the following synthetic procedures. MS was used for purification of the batches after each step of the synthesis, *i.e.*, after saturation with STR, after functionalization with CmiR, and after TmiR capture.

2.7 Analytical methods

2.7.1 Size and zeta potential measurements. A Nanosizer instrument (Zetasizer®, Malvern Instrument, UK) was used for the determination of the hydrodynamic diameter (D_{H}), the polydispersity index (PDI), and the zeta potential (ZP) of suspension nanoparticles. All the measurements were performed in triplicate and the mean value \pm standard deviation is presented. The measurements were performed in H_2O and PBS 1 \times , for the batches adjusted at the iron concentration of 50 mg L^{-1} . For ZP measurements, the ionic strength was fixed by adding NaCl to a final concentration of 10 mM.

2.7.2. IR spectroscopy characterization. FT-IR measurements were carried out in ATR mode on a Bruker Vector 22 FT-IR spectrometer (Bruker, Germany) equipped with a Golden Gate single reflection diamond (Specac). The spectral window from 4000 to 500 cm^{-1} was typically recorded as an average of 32 scans.

2.7.3 Iron concentration determination. Iron concentrations of suspensions were measured by Atomic Absorption Spectrophotometry (AAS) (iCE 3000 spectrometer, Thermo Instruments, France). The synthesized samples were mineralized for a minimum of 2 hours with concentrated hydrochloric acid (6 M), then diluted with hydrochloric acid (1%). The concentration of the samples was determined by using a calibration curve (iron absorption at 248.3 nm, standard iron solutions at 0.25; 0.5; 1.0; 2.0 and 5.0 mg L^{-1}).

2.7.4 Analysis by size exclusion HPLC (SE-HPLC). SE-HPLC measurements were performed with the Shimadzu system (Kyoto, Japan), using LC Column AdvanceBio SEC 300A, 2.7 μm , 7.8 \times 300 mm, 1.0 mL min^{-1} flow rate. Software LC Solution was used for data acquisition and analysis.

All SE-HPLC measurements were performed in phosphate buffer mobile phase (0.15 mol L^{-1} ; the combination of KH_2PO_4 at 0.075 mol L^{-1} and $\text{Na}_2\text{HPO}_4 \cdot 2\text{H}_2\text{O}$ at 0.075 mol L^{-1}), with the flow rate 1.0 mL min^{-1} , injection volume 10 μL , temperature 30 $^\circ\text{C}$, duration 15 min and UV detection at 260 nm and 280 nm.

All samples for SE-HPLC measurements containing PEGylated SPIONs were magnetically separated, and the liquid from the sample was measured, in order to not inject a large concentration of magnetic nanoparticles into the column.

3 Results and discussion

3.1 Synthesis and characterization of the nanoprobe precursors

Coating nanosystems with neutral hydrophilic molecules such as PEG is important for biomedical applications, since it provides colloidal stability in aqueous media, independently from the pH (due to a sterical hindrance). In our study, the PEG chains were covalently attached to the silanized SPION surface, and the process involved three synthetic stages (Fig. 2), as described in the Experimental section. After the synthesis of initial SPIONs by coprecipitation of iron salts, the nanoparticles were silanized, *i.e.* coated with 3-aminopropyltrimethoxysilane (APS) molecules. The latter is bound to the hydroxyl groups abundant on the surface of SPIONs. The primary amines of APS remain available to form amide bonds with the activated esters of α -biotinyl- ω -NHS PEG. The resulting amide bond is well-known for its strength and stability.²⁵ The PEGylated SPIONs were then extensively purified from the residual free PEG molecules by dialysis against water for 48 hours and stored in aqueous suspensions.

The nanoparticles were characterized by a spectrum of analytical techniques: IR spectrometry to confirm the molecular composition and DLS to measure the NP D_{H} and ZP. The iron concentration was determined by means of AAS and used for calculating the molar ratio in the next steps of the nanoprobe formulation.

Successful PEGylation of SPIONs was firstly confirmed by IR bands (Fig. 3A) characteristic of the PEG molecules: the strong bands at 3444 cm^{-1} (O–H stretching), 2921 cm^{-1} (C–H stretching of in CH_2 and methylene groups), and 2853 cm^{-1} (C–H stretching). The amide bonds formed between PEG and silane were seen due to the C=O stretching band at 1651 cm^{-1} . More importantly, strong bands at 1099 cm^{-1} (O–H and C–O–H stretching, siloxane Si–O–Si stretching) characteristic for PEG molecules were observed and assigned to polycondensation of silanes.

Likewise, the successful formation of the polymeric shell was confirmed by the DLS results with the D_{H} increase, from 36.9 \pm 1.4 nm for initial SPIONs to 43.5 \pm 3.4 nm after their silanization and 63.0 \pm 4.0 nm after their PEGylation.

Similarly, ZP also changed after PEGylation: being positively charged for silanized SPIONs (18 \pm 2.5 mV, pH 7) it became nearly neutral (–3.47 \pm 0.97 mV, pH 7). The curves of ZP as a function of pH changed after the nanoparticle modification (Fig. 3B). The ZP of the initial SPIONs at lower pH is positive due to the presence of OH_2^+ on the surface of nanoparticles. The ZP decreases to nearly zero at pH 7 (isoelectric point), due to deprotonation of OH_2^+ to OH. At higher pH, the surface of SPIONs is covered with O^- and the ZP is negative. For silanized SPIONs, the isoelectric point shifted closer to pH 9. As expected, the ZP of the PEGylated SPIONs was less sensitive to pH change



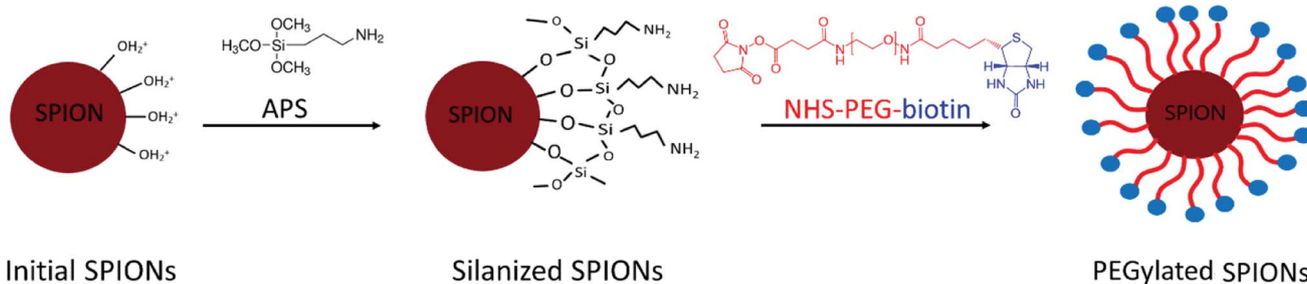


Fig. 2 Schematic representation of the PEGylated SPIONs synthesis in 3 steps: (i) synthesis of the initial SPIONs; (ii) silanization of the SPIONs surface; (iii) PEGylation of the silanized SPIONs.

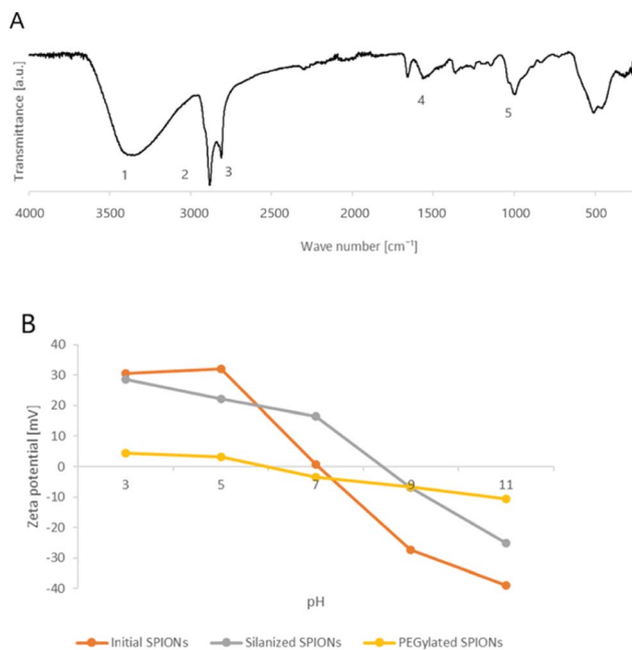


Fig. 3 Physicochemical properties of SPIONs. (A) IR spectra of PEGylated SPIONs, with the major bands labelled as follows: (1) 3444 cm⁻¹; (2) 2921 cm⁻¹; (3) 2853 cm⁻¹; (4) 1651 cm⁻¹; (5) 1099 cm⁻¹. (B) Evolution of zeta potential (ZP) of initial SPIONs (red line), silanized SPIONs (grey line), and PEGylated SPIONs (yellow line) as a function of pH.

and remained nearly zero. It can be concluded that the PEGylated SPIONs remained colloidally stable at neutral pH due to steric hindrance.

3.2 Nanoprobe formulation and characterization

In our study, we have chosen a method of conjugation of miR to the PEGylated SPIONs *via* biotin (Bi)–streptavidin (STR) interaction, since the protocol is simple and cost-effective, requiring no additional reagents. Biotin-modified polymers and miRs are widely commercially available, and biotin insertion into the biomolecules is not intended to affect their properties or activity. Interaction between streptavidin and biotin is irreversible under physiologic conditions since the resulting bond is approaching in strength the covalent one and is unaffected by

variation of temperature, pH, organic solvents, and denaturing agents. Furthermore, STR has four available binding sites, one in each of the four subunits of this protein, that allows each STR molecule to complex several molecules of biotin and thus of biotinylated molecules like PEG or miRs. Void of carbohydrate residues, STR has a low capability of forming non-specific bonds with other molecules. Once bound to the PEG-bi shell of the nanoparticles, STR should still have sites available for binding of biotinylated CmiR and thus enable the formation of nanoprobe described in Fig. 1.^{22,27}

3.2.1 SE-HPLC method for analysis of the formulation ingredients. We used SE-HPLC which appears appropriate for confirmation of the successful formulation of the nanoprobe. SE-HPLC is known as a routine analytical method compared to XPS or element analysis. SE-HPLC enables the confirmation that each separate component present in the synthesis is able to bind with its intended match.

First of all, the SE chromatograms were obtained for the ingredients taken separately (nanoparticles, STR, CmiR) then we analyzed their binary complexes (STR–CmiR), and finally the samples from nanoprobe synthesis steps. Fig. 4 presents chromatograms of PEGylated SPIONs, STR, CmiR, and of complex STR–CmiR. While all the chromatograms were also recorded with UV detection at 280 nm, CmiR has stronger absorbance at 260 nm, ergo herein mentioned results are obtained at this wavelength.

PEGylated SPIONs had a relatively strong chromatographic peak at *ca.* 4.4 min (Fig. 4A). The peak of 1 g L⁻¹ solution of streptavidin was observed at *ca.* 8.2 min (Fig. 4B). The strong peak of CmiR at concentration 5 μM was detected at *ca.* 8.8 min (Fig. 4C). The fact that the NP are eluted earlier than molecules is concomitant with their much bigger size. On the other hand, the close retention time about 8.2 and 8.8 min observed respectively for STR (*M_w* of *ca.* 55 kDa) and biotinylated CmiR (*M_w* of *ca.* 7.6 kDa) is much less expectable.

The formation of the complex STR–CmiR at molar ratio 3.78 : 1 was visible as the new peak rose at *ca.* 6.8 min (Fig. 4D) since the complex was larger than separate molecules. The uniform peak of the complex is in favor that all possible binding places on STR are probably occupied by the biotinylated CmiRs. In the mixed sample, the peak of free CmiR (*ca.* 8.8 min) decreased in intensity almost by a half (37.1 vs. 76.5 AU) compared to the free CmiR taken at the same concentration.



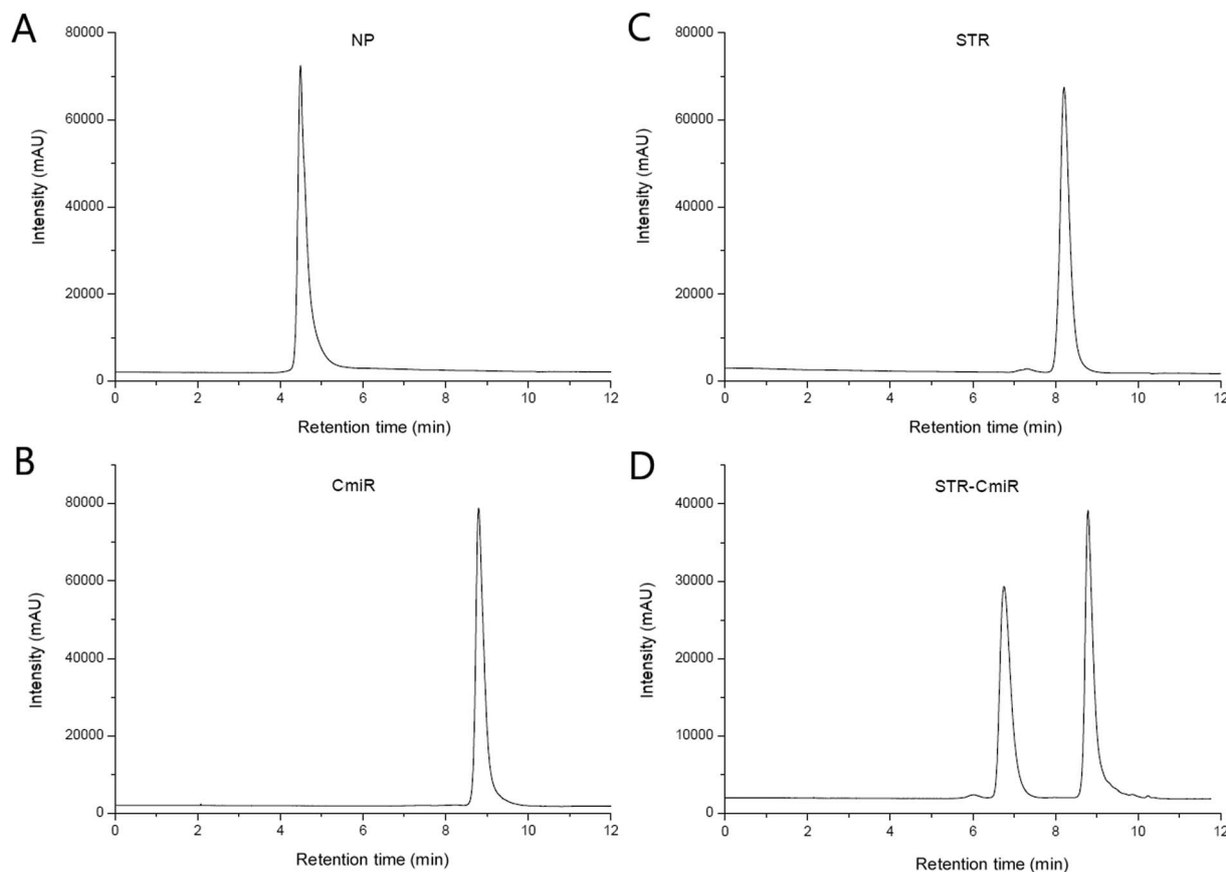


Fig. 4 SE-HPLC chromatograms. (A) PEGylated SPIONs (NP), t_R : 4.4 min, (B) STR, t_R : 8.2 min, (C) biotinylated CmiR, t_R : 8.8 min, (D) STR-CmiR complex, t_R : 6.8 and 8.8 min.

The absence of the free STR peak (around 8.2 min) allows to suppose that, at the conditions used, all the STR molecules were bound to CmiR.

3.2.2 Validation of the nanoprobe formulation steps. As explained above, the nanoprobe synthesis included 2 steps: (1) modification of the PEGylated SPIONs with STR to obtain the NP-STR; and (2) functionalization of the NP-STR with CmiR to produce the NP-STR-CmiR batches.

Formulation step 1 – modification of PEGylated SPIONs with STR. Aiming for a maximal coverage with STR at the biotin sites on the surface of the NP to achieve the best efficiency, we made the attempts with three Fe/STR molar ratios: 2100, 1400, and 280. At the lower Fe/STR molar ratio of 280 (sample NP-STR1), a weak peak of free STR was still present in the chromatograms (t_R 8.2 min), while at higher Fe/STR molar ratio of 1400 and 2100 (the samples NP-STR2 and NP-STR3), no free STR peak was detected (Fig. 5). Thus, one could suppose that at the highest concentration (sample NP-STR1), the STR was able to saturate the available biotin sites.

The weak peaks at 6.3, 6.5 and 6.9 min observed in those samples were possibly due to the presence of PEG-bi chains which were detached from the NP into solution after their complexation with the high molecular weight STR. Indeed, one can reasonably suppose that despite the purification *via* dialysis which was used to remove the excess of the free PEG-bi, it was

still possible to desorb from the nanoparticles some minor fraction of the PEG-bi bound to STR. In agreement with this hypothesis, the peak of the molecular complex STR-PEG (Fig. 5B) had a close retention time of *ca.* 6.7 min. The 3 peaks at 6.3, 6.5 and 6.9 min might correspond to complex fractions with different STR binding sites. This result demonstrates the importance of the purification between the two formulation steps.

The presence of the free STR and the STR-PEG complexes in the NP-STR samples was undesirable since they might compete with STR-modified NP for the binding to the CmiR upon the next step of the nanoprobe formulation. Thus, we applied magnetic separation to purify the NP-STR and successfully eliminate the free STR and the PEG-bi-STR complexes.

Ideally, all the biotin sites on the PEGylated SPIONs surface would be occupied with STR. However, the steric hindrance of the large STR molecules can possibly prevent achieving the saturation of all the biotin sites since some of them are not accessible.

Since the sample NP-STR1 was the best covered with STR, the corresponding batches were purified from unbound species and used for the second formulation step.

Formulation step 2 – functionalization of the NP-STR with CmiR. Here again, our aim was to functionalize the NP-STR with as many CmiR as possible since it is supposed to favor the



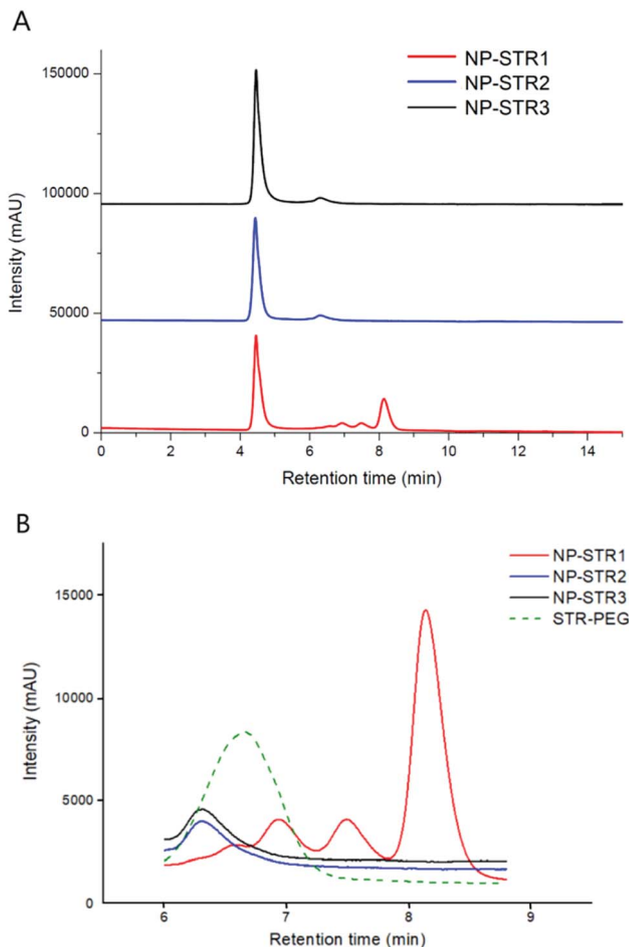


Fig. 5 SE-HPLC analysis of the NP-STR (A): the full chromatograms of the samples: NP-STR1 (molar ratio Fe/STR = 280, red line, t_R : 4.5, 6.9, 7.5 and 8.2 min), NP-STR2 (molar ratio Fe/STR = 1400, blue line, t_R : 4.4, 6.3 min), NP-STR3 (molar ratio Fe/STR = 2100, black line, t_R : 4.5 min and 6.3 min). (B) Zoom on the interval from 6 to 8.8 min with the overlaid peaks of the samples NP-STR1, NP-STR2, NP-STR3, and that of the STR-PEG-bi molecular complex (green dashed line).

efficiency of the final nanoprobe. We started with the CmiR concentration of 5 μM and generated the samples NP-STR-CmiR5. As the formation of STR-CmiR complex was proved to be relatively fast, the 2 hours of incubation of the NP-STR with CmiR (under stirring, at the room temperature) was supposed long enough. The conjugation of both STR and CmiR did not markedly change the physicochemical properties of the respective batches (Table 1).

The batches of PEGylated SPIONs showed a comparable D_H in both deionized water and in PBS 1X (Table 1): 61.6 ± 2.0 nm vs. 63.0 ± 4.0 nm. After the modification with STR, the D_H in PBS 1X slightly increased to 67.0 ± 0.5 nm (Table 1), presumably due to the presence of STR molecules on the surface of the nanoparticles. Functionalization with CmiR led only to a minor increase of D_H till 71.2 ± 0.4 nm. Zeta potential remained largely unchanged (Table 1), with values close to neutral. Interestingly, the PDI values diminished in going to the formulated nanoprobe compared to the PEGylated SPIONs.

Table 1 Physicochemical properties of the batches NP (PEGylated SPIONs), NP-STR1 and NP-STR-CmiR5

	D_H [nm]	PDI	Zeta potential [mV]
NP (PEGylated SPIONs)	61.6 ± 2.0	0.20 ± 0.00	2.7 ± 0.1
NP-STR1	67.0 ± 0.5	0.13 ± 0.01	3.1 ± 0.2
NP-STR-CmiR5	71.2 ± 0.4	0.11 ± 0.01	2.5 ± 0.4

The PDI value of 0.11 is typically considered as characteristic of a monodisperse nanoparticle population. To summarize, the formulation steps did not decrease the colloidal stability of the nanoprobe.

The amount of CmiR bound to the nanoprobe was determined by means of the SE-HPLC analysis. The NP-STR-CmiR5 sample (Fig. 6) revealed 2 peaks: that of nanoparticles at ca. 4.4 min and that of the free CmiR at ca. 8.7 min. The latter decreased in the intensity down to 73.9% compared to the chromatogram of the pure CmiR. Thus, after incubation with 5 μM CmiR solution, about 26% of CmiR were bound to the nanoprobe (the concentration of the bound CmiR was 1.3 μM).

Subsequently, we also prepared samples by incubating NP-STR with the lower and higher concentrations of CmiR: 1 μM (NP-STR-CmiR1) and 10 μM (NP-STR-CmiR10). According to the SE-HPLC data, at 1 μM and 10 μM of incubated CmiR, the nanoprobe-bound CmiR fraction was respectively 66% and 12% and corresponded to concentrations of 0.66 and 1.2 μM (Table 2). To summarize, using a 5 μM concentration of CmiR would be more advisable than 1 μM . In contrast, larger excess of 10 μM does not allow to increase the bound CmiR concentration and leads to use-less loss of the nucleic acid.

The estimated number of CmiR molecules grafted per nanoprobe was 11, 22 and 20 for the samples NP-STR-CmiR1, NP-STR-CmiR5, and NP-STR-CmiR10 respectively. To the best of our knowledge, very few publications reported the quantity of grafted CmiR^{31,32} or other oligonucleotides²⁷ per nanoprobe or

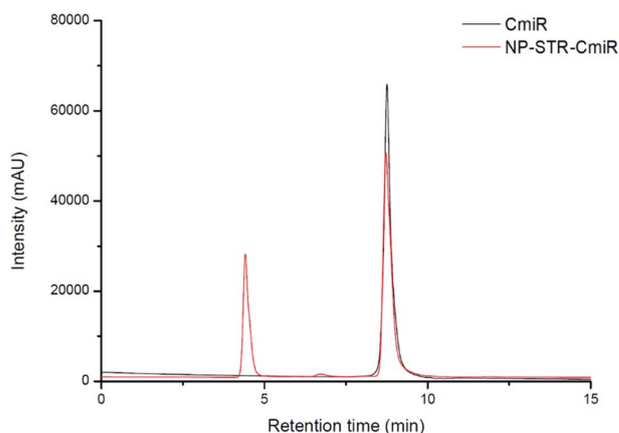


Fig. 6 SE-HPLC results for the second (final) step of the nanoprobe formulation. Typical chromatograms of NP-STR-CmiR5 (red curve, peaks at 4.4 and 8.7 min) compared to those of the pure CmiR at the same concentration of 5 μM (black curve, peak at ca. 8.8 min).



Table 2 Comparative results of functionalization of NP-STR-CmiR batches

Sample	Fraction of bound CmiR	Fraction of bound CmiR (concentration)	Number of bound CmiRs per NP
NP-STR-CmiR1	66%	0.66 μM	11
NP-STR-CmiR5	26%	1.3 μM	22
NP-STR-CmiR10	12%	1.2 μM	20

even the concentration of grafted CmiR.^{33,34} Although less explored in the literature, these values are important since they allow to evaluate both the conjugation method used and the expected efficiency of TmiR capture.³⁵ It has been reported that only one CmiR per nanoprobe may be sufficient for TmiR detection.³²

3.3 Proof of the nanoprobe's affinity to TmiR-155

To determine the ability of nanoprobe to capture the target miR, the purified NP-STR-CmiR5 samples were incubated with the 5 μM solution of the TmiR-155, then were magnetically sorted. The chromatography data indicate that after the sorting, 21% ($\approx 1 \mu\text{M}$) of TmiR were bound to the nanoprobe, which corresponds to approximately 18 TmiRs per nanoprobe. This result is promising since it confirms that the nanoprobe was able to adsorb TmiR and they allowed to remove a significant fraction of it. The 18 : 22 molar ratio for TmiR : CmiR is also promising, while has not been considered as definitive, since it depends on the application protocols.

3.4 Proof of the stability of the nanoprobe in serum

To confirm the stability of the nanoprobe NP-STR-CmiR5 (5 μM CmiR), a new batch was prepared and combined with the solution of fetal bovine serum to be incubated for 4 and 12 hours, resulting in the samples (S4)-NP-STR-CmiR5 and (S12)-NP-STR-CmiR5 respectively. The sample (S4)-NP-STR-CmiR5 was visually monitored for signs of aggregation after 1, 2 and 3 hours, with no apparent change to the solution visible by the eye. We measured the physicochemical properties of the nanoprobe prior to the mixing with serum, and after the incubation and purification *via* magnetic separation (Table 3). The batch NP-STR-CmiR5 showed similar D_H to the one after the incubation in the serum for both 4 h and 12 h, the size increased only slightly, $72.8 \pm 3.3 \text{ nm}$ vs. $78.2 \pm 6.1 \text{ nm}$ vs. $78.2 \pm 1.7 \text{ nm}$, with a bigger standard deviation in the case of the sample (S4)-NP-STR-CmiR5. The polydispersity remained stable for all three batches. The zeta potential of the new batch NP-STR-CmiR5 shifted to the negative value of $-7.1 \pm 2.0 \text{ mV}$,

and after the incubation showed no significant variation. The small changes in physicochemical properties could be explained by a small-scale absorption of proteins present in the serum to the surface of the nanoprobe. The nanoprobe however remain colloidal stable, with their properties not influenced even by a long incubation in the serum.

4. Conclusion

In this study, CmiR-functionalized magnetic nanoprobe were designed for TmiR-155 capture. As precursors of the nanoprobe, we synthesized SPIONs coated with covalently bound PEG biotinylated on its external terminus. Then, the precursors were used to formulate nanoprobe in two steps: (i) modification of the biotinylated external shell with streptavidin and (ii) functionalization of the STR-modified shell with biotinylated CmiR. Saturation of the surface of NP with streptavidin was achieved with molar ratio $\text{Fe}/\text{STR} = 280$. The corresponding batches of the NP-STR were then incubated with biotinylated CmiR (5 μM), and 26% of the latter were bound. The nanoprobe coated with 22 CmiR per nanoprobe were able to capture 18 TmiR per nanoprobe from the 5 μM TmiR solution. The nanoprobe were proved to be stable after an incubation in the serum. This result is promising, while in real samples the nanoprobe concentration has to be certainly used in excess compared to that of TmiR. The capture protocols optimization is a subject of an independent study we are starting.

In close future, the developed nanoprobe will be also studied concerning their ability to capture the target miR (miR-155) in serum from patients and thus to monitor their concentration before and after an anti-cancer treatment.

Conflicts of interest

There are no conflicts to declare.

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Table 3 Physicochemical properties of the batches NP-STR-CmiR5, (S4)-NP-STR-CmiR5 and (S12)-NP-STR-CmiR5

	D_H [nm]	PDI	Zeta potential [mV]
NP-STR-CmiR5	72.8 ± 3.3	0.18 ± 0.01	-7.1 ± 2.0
(S4)-NP-STR-CmiR5	78.2 ± 6.1	0.19 ± 0.02	-6.6 ± 1.0
(S12)-NP-STR-CmiR5	78.2 ± 1.7	0.18 ± 0.01	-4.7 ± 1.3



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