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# 1 Targeted nanomedicine with anti-EGFR scFv for siRNA 2 delivery into triple negative breast cancer cells

3 Phuoc Vinh Nguyen<sup>1</sup>; Katel Hervé-Aubert<sup>1</sup>; Stéphanie David<sup>1</sup>; Nolwenn Lautram<sup>2</sup>; Catherine  
4 Passirani<sup>2</sup>; Igor Hourpa<sup>1</sup>; Nicolas Aubrey<sup>3</sup>; Emilie Allard-Vannier<sup>1</sup>

5  
6 1 EA6295 Nanomédicaments et Nanosondes, Université de Tours, Tours, France  
7 2 INSERM U1066 / CNRS 6021, équipe MINT, Université d'Angers, Angers, France  
8 3 ISP UMR1282, INRA, équipe BioMAP, Université de Tours, Tours, France

## 10 Abstract

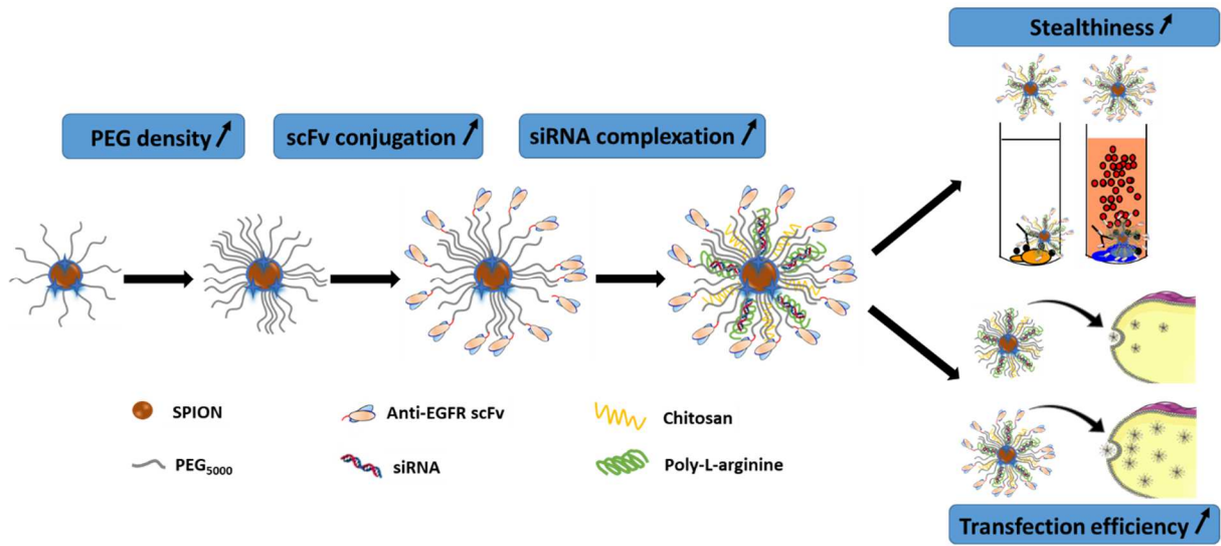
11 A targeted nanomedicine with humanized anti-EGFR scFv (NM-scFv) was developed for siRNA  
12 delivery into triple negative breast cancer (TNBC) cells. NM-scFv consisted of i) targeted nanovector  
13 (NV-scFv): nano-cargo with targeting properties; ii) siRNA: pharmacological agent and iii) cationic  
14 polymers (chitosan, poly-L-arginine): for siRNA complexation and endosomal escape. NV-scFv was  
15 based on superparamagnetic nanoparticle (SPION) labeled with Dylight<sup>TM</sup>680, a PEG layer and a  
16 humanized anti-EGFR scFv. The PEG density was optimized from 236±3 to 873±4 PEGs/NV-scFv  
17 and the number of targeting ligands per NV-scFv was increased from 9 to 13. This increase presented  
18 a double benefit: i) enhanced cellular internalization by a factor of 2.0 for a 24h incubation time and ii)  
19 few complement protein consumption reflecting a greater stealthiness (26.9 vs 45.3% of protein  
20 consumption at 150µg of iron/mL of NHS). A design of experiments was performed to optimize the  
21 charge ratios of chitosan/siRNA (CS) and PLR/siRNA (CR) that influenced significantly: i) siRNA  
22 protection and ii) gene silencing effect. With optimal ratios (CS=10 and CR=10), anti-GFP siRNA was  
23 completely complexed and the transfection efficiency of NM-scFv was 69.4% vs 25.3% for non-  
24 targeted NM. These results demonstrated the promising application of our NM-scFv for the targeted  
25 siRNA delivery into TNBC cells.

26 **Keywords:** TNBC, humanized scFv, gene delivery, targeted nanovector, cationic polymer, active  
27 targeting

## 28 Abbreviations:

29 TNBC: triple negative breast cancer; siRNA: small interfering RNA; PEG: polyethylene glycol; NV-  
30 scFv: targeted nanovector; NM-scFv: targeted nanomedicine; SPION: superparamagnetic iron oxide  
31 nanoparticle; PLR: poly-L-arginine; MS: mass ratio; CS: molar ratio of the positive charges of chitosan  
32 amine groups and the negative charges of siRNA's phosphate groups; CR: molar ratio of the positive  
33 charges of poly-L-arginine amine groups and the negative charges of siRNA's phosphate groups;  
34 EGFR: epidermal growth factor receptor; scFv: single chain variable fragment; NHS: normal human  
35 serum.

36 **Graphical abstract**



37

## 38 1 Introduction

39 Triple negative breast cancer (TNBC) is a complex subtype of breast cancer which is defined by the  
40 lack of expression of three receptors: estrogen, progesterone, and human epidermal growth factor 2  
41 (HER2) [1,2]. TNBC is a real challenge for oncologists and considered as the most aggressive subtype  
42 of breast cancer due to its highest rate of metastasis, risk of recurrence, and the poorest overall  
43 survival [1,3]. Currently, chemotherapy remains the only choice for TNBC systemic treatment but  
44 unfortunately, it is limited by poor bioavailability, toxicity, and the emergence of multidrug resistance  
45 [3,4]. Therefore, there is an urgent need for new treatment modalities to better manage TNBC.

46 With more knowledge on the molecular mechanisms of endogenous RNA interference (RNAi), nucleic  
47 acid medicines have emerged as innovative modalities for the treatment of incurable diseases such as  
48 cancers [5,6]. RNAi's mechanism is based on the interfering activity of double-stranded RNA onto the  
49 expression of a particular gene containing a homologous sequence [5]. Three strategies of RNAi  
50 including small hairpin RNA (shRNA), micro RNA (miRNA) and small interfering RNA (siRNA) have  
51 been largely exploited and are getting more and more interest of worldwide researchers due to its high  
52 specificity, significant effect, minor side effects and ease of synthesis [6,7]. Among these strategies,  
53 siRNA sequences of 21-23 nucleotides are the most used in the development of anticancer treatment.  
54 In fact, most cancers are caused by certain genes encoding for overexpressed proteins, which involve  
55 in cancer progress (so-called oncogenes). Rational siRNA sequences can be used to suppress cancer  
56 via specific oncogenes' suppression [6,7]. Although the therapeutic potency of siRNA for cancer  
57 treatment has been generally accepted, the clinical application of siRNAs remains limited due to extra-  
58 and intracellular barriers. Firstly, as the ideal administration route for siRNA is the systemic  
59 administration, siRNA is challenged by nuclease activity, kidney clearance, phagocyte uptake, and  
60 serum protein's aggregation. Secondly, the highly negative charge and the hydrophilicity of siRNA  
61 prevent them from crossing biological membranes. The off-target effect is another challenge for siRNA  
62 in *in vivo* applications. This phenomenon refers to the unexpected changes in gene expression  
63 sharing partial homology with the siRNA and causes potential toxicity. The last barrier is the  
64 recognition of siRNA by the innate immune system that results in the production of inflammatory  
65 cytokines. Taking these challenges into account and in order to better exploit the therapeutic potency  
66 of siRNA, safe and effective siRNA delivery systems are required [7].

67 One promising strategy for siRNA delivery is siRNA nanovectorization. This strategy consists in  
68 associating siRNA to suitable materials to obtain a nanomedicine (NM) that will be able to i) protect  
69 siRNA from nuclease degradation and ii) effectively deliver siRNA to its target. By loading siRNA in  
70 nanomedicine, several advantages can be achieved: i) provide serum stability by hiding the charge  
71 and the hydrophilicity of siRNA; ii) prolong the blood circulation time by covering the NM with a  
72 polymer layer; iii) increase the cellular internalization and intracytoplasmic siRNA release via active  
73 targeting and the use of pH-sensitive materials [6]. In this study, siRNA was complexed with cationic  
74 polymers surrounding an inorganic core. The presence of the inorganic core was shown to be  
75 advantageous for the stability and the size control of the final nanomedicine (NM) [6].

76 Targeted nanovector (NV) which was based on superparamagnetic iron oxide nanoparticle (SPION)  
77 coated with polyethylene glycol (PEG) has been developed in our group as a promising inorganic core  
78 for efficient siRNA delivery [2,8]. In addition to its safety in *in vivo* application, siRNA loaded with  
79 SPION can combine therapeutic functions and MRI-mediated diagnosis that we call a theranostic  
80 approach [8]. Moreover, Dylight™680, a near-infrared fluorescent dye was covalently attached to our  
81 SPION core that helped to monitor our nanomedicine in cells and tissues by following its fluorescence  
82 signal. Nowadays, the *in vitro* and *in vivo* NM tracking remains a big challenge for NM's  
83 commercialization and few studies have focused on this aspect. Thus, this labeling will allow us to  
84 overcome this problem. For a successful siRNA delivery system, the stability and stealthiness must be  
85 optimized [6]. Therefore, our SPION was covered by PEG<sub>5000</sub> shell and one of the main objectives of  
86 this study was to optimize the PEG layer's density to obtain the optimal stealthiness. In addition to the  
87 passive targeting via the enhanced permeability and retention (EPR) effect, active targeting utilizing  
88 biological ligands could provide further advantages such as increased cellular uptake, reduced side  
89 effects, and better therapeutic efficacy both *in vitro* and *in vivo* [9].

90 EGFR overexpression has been observed in almost 70% cases of TNBC and is strictly correlated to  
91 tumor proliferation, invasion, and metastasis [16]. Moreover, several studies have been focusing on  
92 EGFR targeted nanomedicines for TNBC treatment and have presented preliminary success  
93 [10,12,17].

94 In this study, a specifically designed humanized single chain variable fragment (scFv) directed against  
95 epidermal growth factor receptor (EGFR) was chosen as active targeting ligand and conjugated  
96 directly on the PEG layer. According to the literature, several ligands have been exploited to target  
97 EGFR such as: entire antibodies [10], peptides [11] and aptamers [12].

98 Compared to the whole monoclonal antibodies (mAbs), this humanized scFv which retains at least one  
99 antigen-binding region presents the following advantages: (i) lower molecular weight (25-27kDa vs  
100 150kDa) that keeps our NM small, stable and stealthy; (ii) decreased immunogenicity due to the  
101 absence of the Fc constant domain; (iii) possibility to be engineered for a specific conjugation and (iv)  
102 ease in synthesis [9,13]. In addition, scFv may overcome the challenge of the peptides' weak binding  
103 affinity and the aptamers' degradation by nuclease [14,15].

104 The targeted nanovector (NV-scFv) was characterized in terms of physicochemical properties and was  
105 subsequently associated with siRNA and other cationic polymers to obtain a targeted nanomedicine  
106 (NM-scFv) for triple negative breast cancer theragnosis. This inorganic core was necessary for the  
107 stability, the size control of the final formulation and the tumor targeting properties. Among the cationic  
108 polymers, chitosan and poly-L-arginine (PLR) were chosen for our NM-scFv. These polymers carrying  
109 amino groups ( $\text{NH}_3^+$ ) were complexed with the phosphate groups ( $\text{PO}_4^{3-}$ ) of siRNA via electrostatic  
110 interactions. This kind of formulation has several advantages such as the ease and rapidity of NM  
111 formulation and the prevention from siRNA chemical modification [6]. On the other hand, chitosan is  
112 commonly used in siRNA delivery due to its safety, biocompatibility, biodegradability, and its important  
113 role in siRNA release into the cytoplasm [2,8]. The second polymer-PLR was added to compensate

114 the low stability of siRNA-chitosan complex and to increase transfection efficiency at physiological pH  
115 [8].

116 Two siRNA including one control siRNA and one anti-GFP siRNA were used in this study. For further  
117 *in vivo* application, siRNA sequences targeting TNBC oncogenes such as anti BCL-xl would be used.

118 In summary, the targeted nanovector (NV-scFv) with humanized anti-EGFR scFv antibody fragment  
119 was firstly developed, physicochemically characterized, and optimized in terms of the PEG layer's  
120 density. Secondly, the targeted nanomedicine (NM-scFv) formulation was performed between NV-  
121 scFv, siRNA, chitosan, and PLR. The ratios between components were optimized to obtain the  
122 maximal siRNA protection and transfection efficiency. Finally, the improvement of our optimized NM-  
123 scFv was evaluated concerning the *in vitro* stealthiness and anti-GFP siRNA transfection efficiency.

## 124 **2 Materials and methods**

### 125 **2.1 Preparation of targeted nanomedicine with anti-EGFR scFv for siRNA** 126 **delivery**

#### 127 **2.1.1 Synthesis of anti-EGFR scFv**

128 The humanized scFv fragment resulted from the association between the heavy (VH) and light (VL)  
129 variable domains of an antibody via the (Gly<sub>4</sub>Ser)<sub>3</sub> peptide link and the inclusion of peptide flag  
130 composed of a hexahistidine tag, a GS spacer and a terminal cysteine at the C-terminus [13]. The  
131 design of this humanized scFv fragment was based on the sequences of the variable domains of the  
132 chimeric antibody Cetuximab. However, the variable domains have been humanized by removing the  
133 N-glycosylation site which is present in the VH and conferring recognition to the protein L for the kappa  
134 light chain [18]. So, a gene encoding for humanized scFv was synthesized with an optimized codon for  
135 cricetulus griseus and cloned in pCDNA3.4 (GeneArt/ Thermo Fisher Scientific). ScFv was produced  
136 in ExpiCHO-S™ Cells in a defined, serum-free medium (Thermo Fisher Scientific). ScFv was purified  
137 by loading the supernatant onto a HiScreen™ Capto™ L column (GE Healthcare Bio-Science, 17-  
138 5478-14). Fractions containing the recombinant proteins were selected at 280nm, pooled, dialyzed  
139 against PBS (pH 7.4) overnight, and centrifuged (10,000g, 4°C, 10min). ScFv molecular mass, pI, and  
140 molar extinction coefficient data were all generated by the Protparam tool from  
141 <http://web.expasy.org/protparam/>. The fragment was purified in a pure and homogeneous approach  
142 according to an analysis on SDS-PAGE (sodium dodecyl sulfate-polyacrylamide gel electrophoresis).

#### 143 **2.1.2 Synthesis of the targeted nanovector (NV-scFv)**

144 The NV-scFv's synthesis was based on the protocol developed by our group [9] and included three  
145 steps: (i) the coupling of fluorescent dye onto silanized SPION's surface, (ii) the covering of  
146 fluorescent silanized SPION with a PEG layer and (iii) the functionalization of PEGylated SPION with  
147 the humanized anti-EGFR scFv. In the fluorescence labelling, 0.5mg of Dylight™680 Amine-Reactive  
148 Dye (Thermo Scientific, Rockford, U.S.A) was dissolved in 1mL of anhydrous DMSO and  
149 subsequently added into a suspension of 10mL (16mg or 0.287mmol of iron) of silanized SPION

150 dispersed in DMSO. The suspension was kept under magnetic stirring in dark at room temperature for  
151 24h. NHS-PEG<sub>5000</sub>-Maleimide (Rapp Polymer, Tuebingen, Germany) was chosen for the polymer  
152 layer. NHS-PEG<sub>5000</sub>-Maleimide was dissolved in anhydrous DMSO and added directly into the  
153 fluorescent silanized SPION. The mixture was remained under stirring in dark for 24h. A purification  
154 was made by dialysis (MWCO 1000KDa) in the dark against distilled water at 4°C for 48h to eliminate  
155 free polymer. Afterward, the conjugation of anti-EGFR scFv (2.1.1) was performed in PBS at pH=7.0  
156 for 24 hours. In order to eliminate all non-conjugated scFv and fluorescent dye, the resulted  
157 nanoparticles were purified with size-exclusion chromatography (SEC) using AKTA purifier FPLC  
158 system equipped with a prepacked Superdex 200pg column (600x16mm<sup>2</sup>) (GE Healthcare Bio-  
159 Science AB, Uppsala, Sweden) and a PBS 1X solution as the mobile phase (flow rate of 1.6mL/min).  
160 The injected volume was 5mL at 0.4g of iron/L and the suspension was detected using a UV/Vis  
161 detector at 280nm. At the end of the purification, the NV-scFv was collected and re-concentrated if  
162 necessary, with Vivaspin® (cut-off 30kDa, Fisher Scientific, Illkirch, France).

### 163 **2.1.3 Formulation of the targeted nanomedicine (NM-scFv)**

164 The formulation protocol of NM-scFv was developed and optimized in our group [8,19]. This  
165 formulation refers to the siRNA loading onto NV-scFv with the help of two cationic polymers, chitosan  
166 (MW=110-150kDa; the degree of acetylation: ≤ 40mol.%, Sigma-Aldrich Chimie GmbH, St. Quentin  
167 Fallavier, France) and poly-L-arginine (PLR, MW 15-70kDa, Sigma-Aldrich Chimie GmbH, St. Quentin  
168 Fallavier, France). Briefly, control siRNA (Ambion®, New-York, U.S.A) or anti-GFP siRNA (Ambion®,  
169 New-York, U.S.A) was precomplexed with PLR while chitosan was mixed with NV-scFv. The  
170 complexed siRNA/PLR was then added to the mixture of NV-scFv/chitosan and homogenized using  
171 micropipette mixing and vortexing. The final siRNA concentration was fixed at 50nM for transfection  
172 experiments, 2000nM for physicochemical characterization, and 190nM for internalization assay. Mass  
173 ratio (MS) was used to determine the NV-scFv/siRNA ratio and was fixed at 10. The cationic polymers'  
174 content was defined as the charge ratio or the molar ratio of the positive charges of polymers and the  
175 negative charges of siRNA. For the optimization of NM-scFv, the charge ratio of chitosan/siRNA (CS)  
176 was varied from 10 to 50 and that of PLR/siRNA (CR) from 2 to 10.

## 177 **2.2 Physicochemical characteristics**

### 178 **2.2.1 Size and zeta potential analysis**

179 The hydrodynamic diameter ( $D_H$ ), the polydispersity index (PDI), and the zeta potential ( $\zeta$ ) were  
180 determined using a Nanosizer apparatus (Zetasizer®, Malvern Instrument, UK). For NV-scFv, the  
181 measurement of  $D_H$  and  $\zeta$  was made in PBS 1X or NaCl solution (0.01M) respectively at the  
182 concentration of 50mg of iron/L. For the NM-scFv, the measurement was performed after dilution at  
183 1:25 (v/v) of NM-scFv in NaNO<sub>3</sub> 0.01M to fix the ionic strength. The  $D_H$  was based on intensity. All the  
184 measurements were achieved at 25°C in triplicate and presented in mean values ± SD.

## 185 **2.2.2 Determination of NV-scFv and NM-scFv concentration**

186 The concentration of NV-scFv and NM-scFv was expressed in iron concentration (mg of iron/L) or in  
187 siRNA concentration (nM). The total iron concentration was determined by atomic absorption  
188 spectrophotometry (iCE 3000 spectrometer, Thermo Instruments, France). NV-scFv and NM-scFv  
189 were digested by adding concentrated hydrochloric acid (6M) for at least 2h and then diluted with  
190 hydrochloric acid (0.12M). Measurements were performed at 248.3nm, and the concentration of the  
191 samples was determined using a calibration curve (iron concentrations of 0.25; 0.5; 1.0; 2.0 and  
192 5.0mg/L).

## 193 **2.2.3 Determination of the polymer layer's density**

194 The grafted polymer density was interpreted via the number of polymer chains per NV. Firstly, the  
195 concentration of PEG (mg/L) was quantified using the modified Dragendorff method [20] based on the  
196 formation of PEG-Bil<sub>4</sub><sup>-</sup> complex. The reaction was made between 5mL NV suspension at 10mg of  
197 iron/L and Dragendorff's reagent in excess - a solution of potassium bismuth iodide in acetic acid at  
198 3.669M (pH=2.1) for a final volume of 10mL during 15min at room temperature. The absorbance at  
199 520nm of the resulted solution was measured. The NHS-PEG<sub>5000</sub>-Maleimide concentration of the  
200 samples (C<sub>PEG</sub>) was determined using a calibration curve (NHS-PEG<sub>5000</sub>-Maleimide concentrations of  
201 0; 12.5; 25.0; 37.5; 50.0; 62.5 and 75mg/L) and the silanized SPION was used as the negative control.  
202 The number of polymer chains per NV was then calculated based on the following formula:

203 Concentration of PEG/NV (mole of PEG/g of iron):  $C_{PEG/NV} = \frac{2.C_{PEG}}{10.M_{PEG}}$

204 Number of polymer chains per NV =  $C_{PEG/NV} \cdot \frac{\pi D_H^3}{6} \cdot d \cdot 6 \cdot 10^{23}$

205 with M<sub>PEG</sub>= 5311g/mole; d: masse density of SPIONs and determined at 5.2x10<sup>6</sup>g/m<sup>3</sup>; D<sub>H</sub>:  
206 hydrodynamic diameter of NV (m), Avogadro's number= 6. 10<sup>23</sup>mole<sup>-1</sup>.

## 207 **2.2.4 Quantification of grafted antibody fragments**

208 The anti-EGFR scFv concentration conjugated onto NV-scFv was determined by a modified Bradford  
209 assay [21] using the Coomassie Plus Assay Kit (Thermo Scientific, Rockford, U.S.A) according to  
210 manufacturer's instructions. The absorbance of the samples was measured at 630nm and the  
211 concentration of scFv (µg/mL) was determined using a calibration curve (scFv concentrations of 0; 2.5;  
212 5; 10; 15; 20; 25µg/mL) and NV without scFv was used as the negative control. This concentration  
213 was transferred into mole of scFv per g of iron and the number of scFv/NV was finally calculated using  
214 the same formula for the calculation of the number of polymer chains.

## 215 **2.2.5 Functionality test of grafted antibody fragments**

216 The functionality of the grafted anti-EGFR scFv was evaluated by indirect enzyme-linked  
217 immunosorbent assay (ELISA). In this experiment, EGFR recombinant protein (Sino Biologicals,  
218 Beijing, P.R. China) was coated in a 96-well plate at 2µg/mL in PBS and incubated overnight at 4°C.  
219 The wells were then saturated with 3% of BSA-PBS for 1h at 37°C. Afterward, PBS for the negative

220 control, NV-scFv, or NV (from 0.003 to 100mg of iron/L) was incubated in wells for 1h at 37°C. The  
221 wells were subsequently washed with PBS-Tween 20 (0.05%; m/v) and incubated with 100µL of  
222 protein L-peroxidase (Pierce®, Thermo Fisher Scientific) diluted at 1.25µg/mL for 1h at 37°C. An  
223 enzymatic reaction was made by the addition of 100µL of 3,3',5,5'-Tétraméthylbenzidine substrate  
224 (TMB; Sigma, St Louis, USA) and stopped with 50µL of H<sub>2</sub>SO<sub>4</sub> 1M. Finally, the absorbance was  
225 measured at 450nm using an absorbance microplate reader (Bio-Tek® instruments, Inc., USA). The  
226 presence of scFv was revealed by the well coloration and its content is proportional to the absorbance  
227 at 450nm.

## 228 **2.2.6 siRNA protection with agarose gel electrophoresis**

229 To verify the siRNA protection capacity, the electrophoresis technique on agarose gel was used. An  
230 agarose gel at 1% (m/v) was prepared containing 0.01% (v/v) ethidium bromide (EtBr) to visualize free  
231 siRNA. NM and NM-scFv were formulated in double for each type of NM or NM-scFv. The first sample  
232 was diluted with water and the second was diluted with heparin 10g/L (Sigma-Aldrich Chemie GmbH,  
233 Steinheim, Germany) at a dilution factor of 2:1 (v/v) in order to destabilize the formulation and release  
234 the formulated siRNA. A loading buffer (Agarose gel loading dye 6X, Fisher, Bioreagents®, Illkirch,  
235 France) was added and a final content corresponding to 16pmol of siRNA per well was deposited. The  
236 migration of samples on the gel was conducted in a Tris-acetate-EDTA (TAE) 1X buffer (Acros  
237 Organics, Geel, Belgium) for 15min at 150V. The visualization of free siRNA was made with UV-  
238 imaging using the EvolutionCapt software on a Fusion-Solo.65.WL imager (Vilbert Lourmat, Marne-la-  
239 Vallée, France).

## 240 **2.3 Cell culture experiments**

### 241 **2.3.1 Cell culture**

242 Triple negative breast cancer cells MDA-MB-231 (ECACC, Salisbury, U.K.) and MDA-MB-231  
243 expressing GFP (MDA-MB-231/GFP) (Euromedex, Souffelweyersheim, France) were cultured at 37°C  
244 in an atmosphere containing 5% CO<sub>2</sub>. The culture medium was made of DMEM supplemented with  
245 10% fetal bovine serum, 1% non-essential amino acid (Hyclone Laboratories, Logan, Utah) and 1%  
246 penicillin/streptomycin (Gibco®, Life Technologies, Paisley UK). The cell harvesting was made with  
247 trypsin/EDTA (0.05%) (Gibco®, Life Technologies, Paisley UK) at 80% of confluence.

### 248 **2.3.2 Transfection assay**

249 MDA-MB-231/GFP cells were seeded at 3.10<sup>4</sup> cells/well in a 12-well plate for 24h before the  
250 transfection. The day of transfection, NM-scFv, Oligofectamine™, and Lipofectamine™ (Invitrogen,  
251 Thermo Fisher Scientific, Paisley, UK) were prepared with anti-GFP siRNA at a final siRNA  
252 concentration of 50nM in respect of the formulation protocol and the manufacture recommendation.  
253 Cells were treated with NM-scFv in serum-free Opti-MEM (Gibco®, Life Technologies, Paisley UK) or  
254 in complete medium (10% of serum) and maintained for 4h. Afterward, the NM-scFv was removed or  
255 maintained in normal growth conditions (the supplementation in serum was made for cells treated with  
256 Opti-MEM) for 68h until the analysis. Non-treated cells were used as the negative control. Cells were

257 removed using trypsin and then analyzed with a flow cytometer (Gallios flow cytometer, Beckman  
258 Coulter). Data were analyzed using Flowing Software 2.5.1. The siRNA transfection efficiency (gene  
259 silencing effect) was calculated by the percentage of the cells with reduced GFP fluorescence intensity  
260 over the analyzed cells using the GFP fluorescence histogram.

### 261 **2.3.3 Cytotoxicity test**

262 Cell viability was evaluated by MTT assay. MDA-MB-231 cells were seeded at  $3 \cdot 10^4$  cells/well in a 12-  
263 well plate for 24h. Formulations of siRNA, NV-scFv, and PLR at different charge ratios of PLR/siRNA  
264 were prepared and incubated with cells for 72h in normal growth conditions. Non-treated cells or cells  
265 treated with H<sub>2</sub>O<sub>2</sub> at 20mM were used as the negative and positive control respectively. The culture  
266 medium was then replaced by the mixture of 190µL of fresh medium and 10µL of an aqueous solution  
267 of MTT (5g/L) and incubated for 4h at 37°C. The medium/MTT mixture was removed and 200µL of  
268 DMSO was added in each well. The plate was agitated until the homogeneity and the absorbance was  
269 measured at 540nm using KC-junior V1.40 software on a BIOTEK EL800 microplate reader. The  
270 number of viable cells was directly proportional to the absorbance value and the percentage of viable  
271 cells was calculated according to the followed equation:

$$272 \quad \frac{As - Apc}{Anc - Apc} \times 100$$

273 As, Apc and Anc are the absorbance of the sample, the positive control and the negative control.

### 274 **2.3.4 Internalization assay**

275 The internalization of NV and NM into MDA-MB-231 cells was evaluated by following the Dylight™680  
276 fluorescence with flow cytometry. MDA-MB-231 cells were seeded onto a 12-well plate at  $1.5 \times 10^5$   
277 cells/well. After 24h, NV-scFv or NM-scFv prepared with control siRNA in Opti-MEM was added to  
278 cells at a final concentration of 25mg of iron/L (or 190nM in siRNA for NM) for 4h. After 4h, a cell  
279 culture medium of 20% of serum was added at a dilution factor of 1:1 (v/v) for 20h. Non-treated cells  
280 were used as the negative control. After incubation time, cells were washed, removed using trypsin,  
281 and analyzed by flow cytometry.

## 282 **2.4 Complement activation test (CH50 test)**

283 Complement consumption was assessed in normal human serum (NHS) (Établissement Français du  
284 Sang, Pays de la Loire, Nantes, France) by measuring the residual hemolytic capacity of the  
285 complement system after contact with NM-scFv. The final dilution of NHS in the mixture was 1:4 (v/v)  
286 in 400µL of reactive media. The technique consisted in determining the amount of serum that was able  
287 to lyse 50% of a fixed number of sensitized sheep erythrocytes with rabbit anti-sheep erythrocyte  
288 antibodies (CH50) [22]. Complement activation was expressed as a function of iron concentration to  
289 estimate the impact of the polymer coating for a similar quantity of SPIONs. The *in vivo* iron  
290 concentration of interest was in the range of 125-188µg/mL of NHS. This range was calculated based  
291 on the blood-serum volume per mouse (1,4-1,7mL blood or 0.55-0.83mL serum for a mouse of 20-

292 25g) and the ferrofluid quantity that would be injected per mouse (125 $\mu$ L of NM-scFv at 0.83 $\mu$ g/ $\mu$ L or  
293 103.75 $\mu$ g of iron per mouse).

## 294 **2.5 Experimental design**

295 A full factorial design  $2^2$  was created by Minitab<sup>®</sup>16 (Minitab, Inc.) to determine the effect of charge  
296 ratios (CR and CS) on the transfection efficiency of anti-GFP siRNA loaded in NM-scFv. This  
297 experimental design composed of two independent parameters and one variable response (dependent  
298 variable). The independent parameters including CR and CS were studied at two levels. Besides,  
299 three central points were included to get an indication of curvature and an estimation of pure error.  
300 The transfection efficiency was evaluated as the dependent variable. Each experiment was performed  
301 in triplicate and an overview of the experimental design including the results is shown in Table SI 2. To  
302 determine the optimal parameters, the optimization function of Minitab was used.

## 303 **2.6 Statistics**

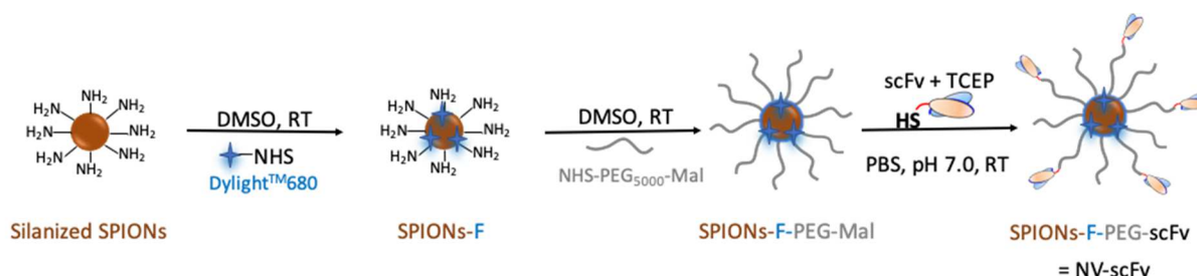
304 Values were expressed as mean  $\pm$  standard deviation (SD). Fisher's ANOVA test was used to  
305 evaluate the effect of charge ratios on the transfection efficiency. The effect of charge ratio on  
306 transfection efficiency was considered significant when p-value<0.05 (\*). For the internalization assay,  
307 Student's t-test was used to compare the results and the difference was considered significant when  
308 p-value<0.05.

## 309 **3 Results and discussion**

### 310 **3.1 Development and optimization of the targeted nanovector (NV-scFv)**

311 The synthesis of our targeted nanovector as shown in Fig. 1, was based on the protocol developed in  
312 our group for HER2-positive breast cancer with some adaptations [9,13]. Briefly, the three-step  
313 synthesis allowed us to modify the surface of initial silanized SPION according to our purposes. It  
314 started with the direct introduction of a near-infrared (NIR) fluorescent dye onto the silanized SPION  
315 core via the reaction between NHS groups of the fluorophore and the primary amines on the SPION  
316 core. In this study, the fluorophore was Dylight<sup>™</sup>680 with fluorescence emission in the very near  
317 infrared. This property was particularly well adapted for *in vivo* optical analysis due to the high  
318 transparency of tissues in this spectral region. For a systemic administration, the colloidal stability at  
319 physiological conditions and the stealthiness against the immune system are required [23]. One of the  
320 most common strategies is to cover our NV-scFv with a PEG layer that increases: i) the stability  
321 thanks to steric hindrance [24] and ii) the circulation time by avoiding the recognition by the  
322 mononuclear phagocyte system (MPS) via opsonization process [22]. For this purpose, the bi-  
323 functional NHS-PEG<sub>5000</sub>-Maleimide was chosen. The NHS groups reacted with the primary amines of  
324 silanized SPION core and the maleimide groups allowed the subsequent conjugation with active  
325 targeting ligands. Furthermore, the PEG layer would protect the fluorophore from the quenching by  
326 external interactions. Finally, to achieve the active targeting for TNBC, a humanized recombinant anti-

327 EGFR scFv of the Cetuximab antibody was conjugated to our NV. This conjugation implied the  
 328 carbodiimide chemistry by creating disulfide bridges between the C-terminal cysteines introduced into  
 329 the scFv sequence and the maleimide groups of the polymer layer. Our anti-EGFR scFv had been  
 330 also rationally designed to facilitate its orientation and was coupled selectively onto the polymeric  
 331 surface of the NV via a site-specific covalent conjugation.



332

333 Fig. 1. Schematic representation of targeted nanovector (NV-scFv) synthesis.

### 334 3.1.1 Improvement in the polymer density and the number of grafted scFv onto the targeted 335 nanovector

336 The former results for HER2+ breast cancer showed a perspective of using NV-scFv for tumor active  
 337 targeting [9]. However, its stealthiness needed to be optimized to achieve a long circulation time  
 338 allowing a higher *in vivo* tumor accumulation. The developed HER2-targeted NV-scFv was quickly  
 339 captured and eliminated by the liver and spleen [9]. Facing this problem, we hypothesized that the  
 340 polymer layer used for HER2 targeted NV-scFv (molar ratio between PEG/iron 0.3) was not sufficient  
 341 to obtain a relevant stealthiness. The impact of PEGylation on the NV's stealthiness is highly  
 342 dependent on PEG molecular weight (MW), polymer chain architecture, and surface density of the  
 343 PEG coating [25]. As the PEG<sub>5000</sub> is appropriate in terms of molecular weight [25,26], the  
 344 enhancement in the density of the PEG layer may be a potential solution [27]. Thus, this research  
 345 aimed to increase the PEG density by optimizing the ratio of PEG/iron in the synthesis. Three  
 346 nanovectors were synthesized with different molar ratios of PEG/iron at 0.3 (NV1); 0.6 (NV2); and 0.9  
 347 (NV3) respectively using the same initial SPION and the number of polymer chains per NV was  
 348 determined using the modified Dragendorff method. For the same type of NV (shape) with a similar  
 349 size, the number of polymer chains per NV was used to evaluate the polymer density [27]. The results  
 350 showed that by increasing the molar ratio of PEG/iron, an increase in the number of polymer chains  
 351 per NV was observed between NV2 and NV1:  $873 \pm 4$  PEGs vs  $236 \pm 3$  PEGs/NV respectively. The  
 352 higher polymer density of NV2 compared to NV1 could consequently improve its stealthiness.  
 353 Nevertheless, there was no enhancement in the number of polymer chains for NV2 compared to NV3  
 354 ( $873 \pm 4$  vs  $850 \pm 10$  PEGs/NV). This result suggested a possible saturation in the number of PEGs that  
 355 can be grafted onto the NV's surface. For this reason, the molar ratio of PEG/iron superior to 0.9 was  
 356 not further considered.

357 The increase in the number of polymer chains per NV did not change the physicochemical properties  
 358 of the nanovector. The size of all evaluated nanovectors was around 80nm with a highly

359 monodispersed population (PDI inferior to 0.3) and the nanovectors were almost neutral in charge that  
 360 is required for an injectable form (Table I).

361 Table I. Number of grafted PEGs, scFv molecules and physicochemical properties of different  
 362 nanovectors.

	NV1	NV2	NV3	NV1-scFv	NV2-scFv
PEG/iron molar ratio	0.3	0.6	0.9	0.3	0.6
Number of PEG per NV	236±3	873±4	850±10	236±3	873±4
D <sub>H</sub> (nm)	76.6±5.3	78.8±0.2	82.8±3.4	78.0± 1.7	74.4 ± 4.0
PDI	0.15±0.01	0.22±0.01	0.10±0.01	0.15±0.02	0.19±0.02
ζ (mV, pH 7.4)	-2.8±2.4	-5.2±1.3	-4.5±1.2	-1.1±1.5	-3.5±1.0
Number of scFv per NV	0	0	0	9	13

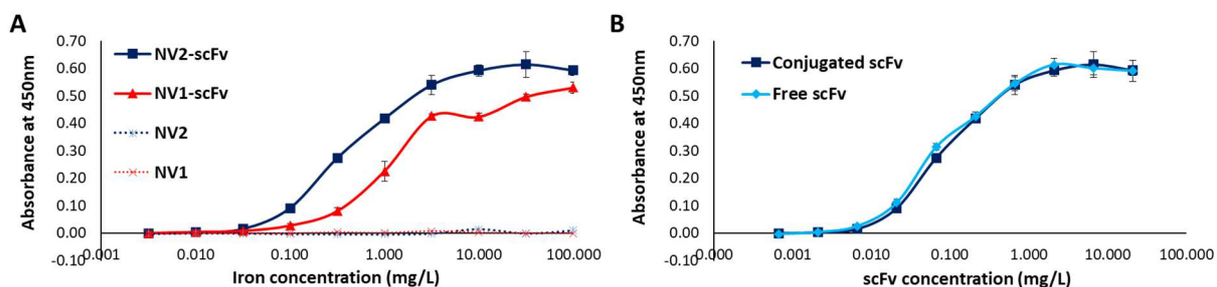
363

364 Also, the possibility of increasing the polymer density layer led us to a hypothesis: the higher the  
 365 density of polymer layer is, the more functional groups the nanovector can have on its surface  
 366 resulting in a higher scFv grafting rate onto the nanovector. To verify this hypothesis, two targeted  
 367 nanovectors were synthesized from NV1 and NV2 at two corresponding molar ratios of scFv/iron at  
 368 1/2000 and 1/1000 respectively. The number of scFv grafted onto NV-scFv was then quantified with a  
 369 modified Bradford assay. This method exploits the strong red shift of Coomassie G-250 dye (from 465  
 370 to 595nm) when it binds to protein and is suitable for SPION-grafted scFv quantification [13].  
 371 Consistent with our previous studies, no change in physicochemical characteristics was observed after  
 372 the conjugation of scFv (Table I). As predicted, more antibody fragments were successfully conjugated  
 373 onto NV's surface when the polymer density was higher: 13 scFv vs 9 scFv that might improve the  
 374 targeting properties of our NV.

375 To verify the functionality of the conjugated scFv, an ELISA assay on increasing concentrations of NV  
 376 and NV-scFv was carried out with the target protein EGFR and detected by Protein L (PpL). This  
 377 protein presents a great advantage to detect antibody fragments such as scFv thanks to its ability of  
 378 binding to some kappa light chain variable domains without interfering with the antigen-binding site  
 379 [9,18].

380 As shown in Fig. 2A, while the absorbance of PpL's substrate for the whole range of NV's  
 381 concentration remained negligible for non-targeted NV, both NV-scFv presented a gradually increasing  
 382 absorbance. At the same NV's concentration, the absorbance of PpL's substrate for NV2-scFv was  
 383 higher than NV1-scFv (~0.6 vs ~0.4 at 10mg of iron/L for example). To compare the binding affinity  
 384 between grafted scFv and free scFv, this experiment was performed also for the same quantity of free  
 385 scFv as the scFv grafted on NV2-scFv. For the whole studied range of concentrations, there was no  
 386 difference in the absorbance of PpL's substrate between free scFv and conjugated scFv on NV2-scFv  
 387 (Fig. 2B).

388 All the above results indicated i) the possibility to increase the PEG density on NV's surface, ii) the  
 389 enhancement in PEG density resulted in the increased number of scFv conjugated onto NV-scFv and  
 390 iii) the preserved functionality of conjugated scFv.



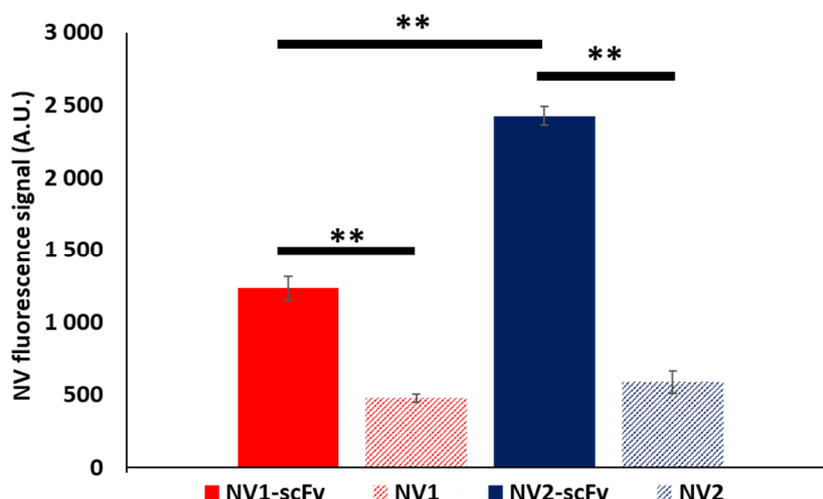
391  
 392 Fig. 2. Functionality interpreted by the absorbance at 450 nm of PpL's substrate obtained from ELISA  
 393 experiment of conjugated and free anti-EGFR scFv in suspension regarding EGFR protein. A)  
 394 Conjugated scFv in different NVs and B) Conjugated scFv in NV2-scFv vs free scFv.

### 395 3.1.2 Internalization assay into cancer cells of the optimized nanovector

396 To clarify the benefit of the scFv conjugation onto NV-scFv, cellular internalization of non-targeted or  
 397 targeted nanovector was performed on MDA-MB-231 TNBC cells. In this experiment, cancer cells  
 398 were incubated with four types of nanovector including NV1; NV1-scFv; NV2; and NV2-scFv for 24h.  
 399 The signal of the fluorescent dye of NVs in suspension was checked beforehand and the signal of  
 400 Dylight™680 was followed using flow cytometry.

401 As shown in Fig. 3, both functionalized nanovectors with anti-EGFR scFv internalized better into cells  
 402 than non-targeted NVs by a factor of 2.6 and 4.1 for NV1-scFv vs NV1 and NV2-scFv vs NV2  
 403 respectively ( $p < 0.01$ ). This result demonstrated the benefit of active targeting with the humanized anti-  
 404 EGFR scFv. In addition, the increased number of grafted scFv on the NV2-scFv was shown to be  
 405 favorable for the cellular internalization. The NV2-scFv internalized better than NV1-scFv by a factor of  
 406 2.0 ( $p < 0.01$ ).

407 These results led us to choose the NV2-scFv for further experiments.



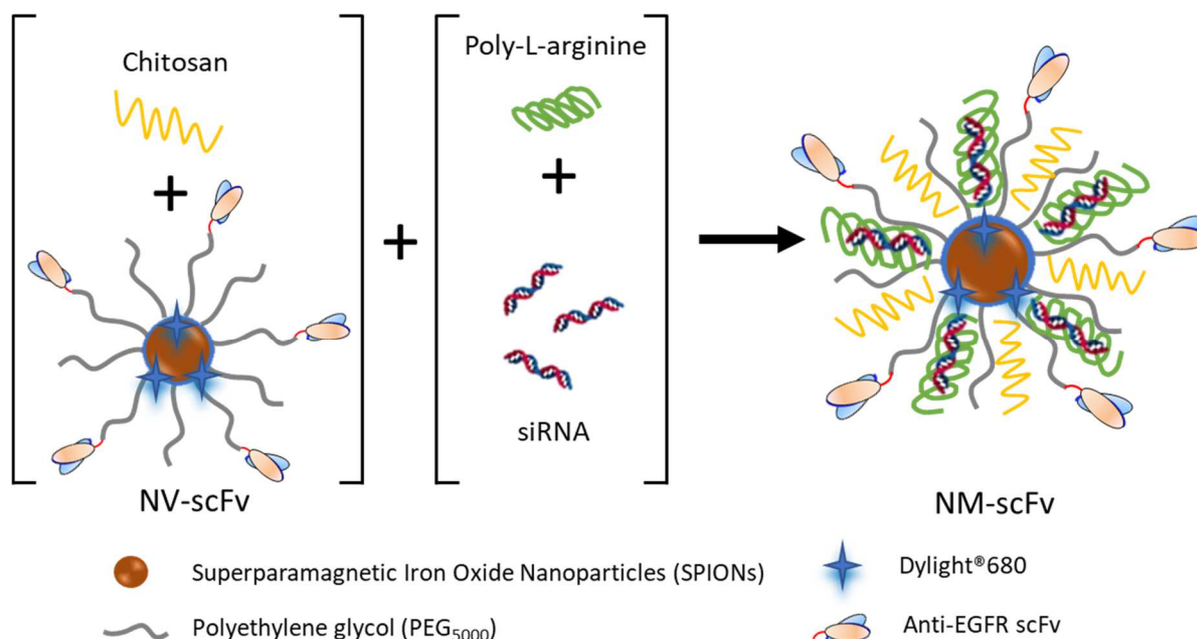
409 Fig. 3. NV fluorescence signal by following Dylight™680 fluorescence intensity of different NVs  
410 internalized into MDA-MB-231 cells. Double asterisk (\*\*) indicates statistically significant factors with  
411  $p < 0.01$ .

### 412 **3.2 Optimization of the targeted nanomedicine (NM-scFv) for siRNA delivery**

413 After optimizing the targeted nanovector, the next step was to associate this NV-scFv with siRNA to  
414 form the targeted nanomedicine (NM-scFv). Among the available methods, the electrostatic  
415 complexation was chosen due to its ease and speed of the preparation [6].

416 The NM-scFv comprised of three ingredients: i) siRNA to inhibit the expression of targeted oncogenes  
417 (in this study, a control siRNA and an anti-GFP siRNA were used as models); ii) NV-scFv which  
418 played a key role in the stability, control of the size and targeting properties of the final NM-scFv [6];  
419 and iii) two cationic polymers including chitosan and poly-L-arginine (PLR) which were essential for not  
420 only siRNA protection at physiological pH but also an endosomal escape for siRNA transfection  
421 efficiency. Concerning the protocol (Fig. 4), the siRNA and NV-scFv were precomplexed with PLR and  
422 chitosan respectively. By mixing negatively charged NV-scFv with chitosan, an electrostatic interaction  
423 may occur and the chitosan was trapped into the PEG coating. PLR, which produced a more compact  
424 complex with siRNA could enable more efficient shielding of the siRNA-PLR charges within the PEG  
425 coating of NV. Once siRNA-PLR was added into Chitosan-NV, the siRNA-PLR could trap low-charged  
426 chitosan on the NV resulting in a lower size and polydispersity. In cells, with the protonation of  
427 chitosan amine groups by the acidification of the endosome, the electrostatic repulsions between  
428 cationic charges triggered polymer expansion (umbrella effect) and led to the liberation of siRNA into  
429 cytoplasm.

430 To successfully complex siRNA, the ratios of components needed to be optimized. In this research,  
431 two types of ratios were taken into consideration including mass ratio and charge ratio. Mass ratio  
432 (MS) describes the ratio between the mass of the inorganic core and that of siRNA. MS was optimized  
433 in the previous study in our group and fixed at  $MS = 10$  [28]. The charge ratio represented the molar  
434 ratio between the number of positive charges of polymers (those of amino groups) and that of negative  
435 charges of siRNA (phosphate groups). Herein, there were two charge ratios to optimize including  
436 PLR/siRNA (CR) and chitosan/siRNA (CS). The following parts aimed to optimize these ratios by  
437 evaluating their impacts on the siRNA protection capacity and the siRNA transfection efficiency.



438

439 Fig. 4. Schematic representation of the targeted nanomedicine (NM-scFv) complexation between  
 440 targeted nanovector (NV-scFv), siRNA and cationic polymers (Poly-L-arginine and Chitosan).

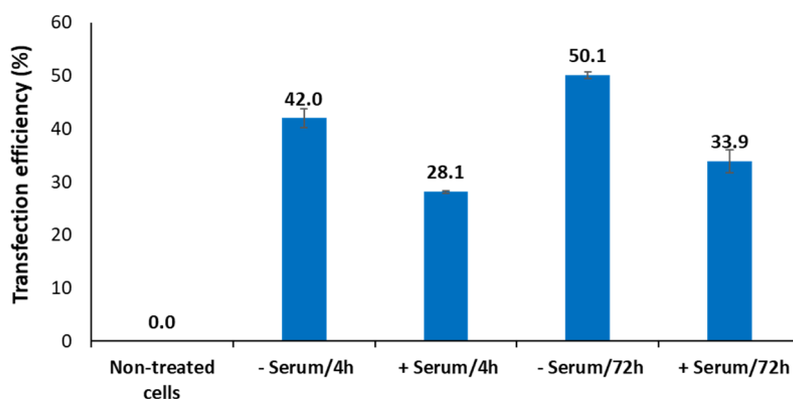
### 441 3.2.1 Optimization of transfection conditions

442 Before optimizing the charge ratios, relevant siRNA transfection conditions were required to clarify the  
 443 difference between the tested formulations. According to the previous studies, the period between the  
 444 treatment and the analysis was fixed at 72h to have enough time for the inhibition of the GFP protein  
 445 (analysis time) [8,29]. Here, two parameters were studied to obtain an efficient transfection: i) the  
 446 medium in which the transfection was performed (serum-free vs complete medium with 10% serum)  
 447 and ii) the contact time between our NM-scFv with cells (4h vs 72h). The tested formulation (NM<sub>0</sub>-  
 448 scFv) was formulated using the following component ratios: MS=10; CR=4; and CS=30.

449 The NM<sub>0</sub>-scFv was incubated with the cells either in Opti-MEM™ – a popular serum-free medium (-  
 450 Serum) or in complete medium (+Serum). Two contact times, 4h or 72h were tested. For the short  
 451 contact time, the NM<sub>0</sub>-scFv was removed after 4h. For the contact time of 72h, the NM<sub>0</sub>-scFv remained  
 452 in contact with cells until the analysis. The cells in Opti-MEM™ were supplemented with serum after  
 453 4h to obtain a final concentration of 10%. This supplementation in serum helped to preserve cells from  
 454 cell-death in serum-free medium.

455 The results (Fig. 5) showed that the transfection made in Opti-MEM™ was more efficient than in  
 456 normal culture medium: 42.0±1.8% vs 28.1±0.9% for 4h and 50.1±0.7% vs 33.9±2.1% for 72h of  
 457 incubation time. On the other hand, the contact time between NM<sub>0</sub>-scFv and cells played also a  
 458 relevant role. In the same condition of transfection (-Serum or +Serum), the transfection efficiency was  
 459 better for the contact time of 72h compared to the short contact time of 4h (50.1±0.7% vs 42.0±1.8%  
 460 for -Serum and 33.9±2.1% vs 28.1±0.9% for +Serum).

461 Taking into account these results, all the subsequent transfections in this study were performed in  
462 serum-free medium (Opti-MEM™) for 4h and the NM-scFv was kept in contact with the cells in normal  
463 medium for additional 68h until the analysis.



464  
465 Fig. 5. Transfection efficiency of the NM<sub>0</sub>-scFv at different conditions of transfection: with (+) or without  
466 (-) serum and at different contact times: 4h or 72h with MDA-MB-231 cells.

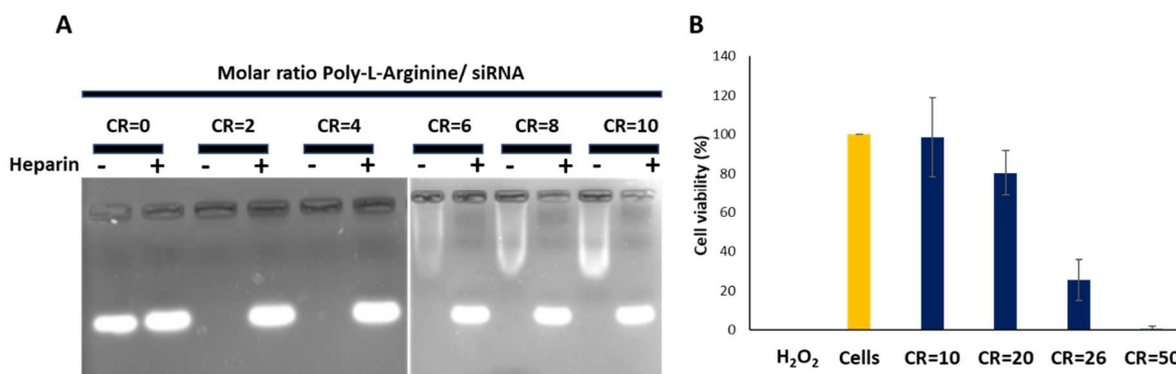
### 467 3.2.2 Optimization of the formulation parameters for an efficient siRNA transfection

468 The next step was to optimize the charge ratios including CR and CS. Both CR and CS were shown to  
469 be important for an efficient siRNA transfection [8,19,28]. A design of experiments was used to  
470 optimize these ratios. To construct this design of experiments, it was necessary to determine the  
471 levels of each ratio in our design.

472 In the case of CR, as PLR played a major role in siRNA complexation, the levels of CR had to be  
473 optimized to obtain a maximal siRNA protection capacity [8]. In this experiment, siRNA was complexed  
474 with PLR at different CR values from 0 to 10 and subsequently added to NV2-scFv. siRNA retention  
475 was evaluated using agarose gel electrophoresis. After the electrophoresis, free siRNA migrated and  
476 appeared as fluorescent bands, whereas protected siRNA remained in the loading wells. Without  
477 heparin addition, no fluorescent band was observed for formulations prepared with CR $\geq$ 2, and with  
478 heparin addition, the fluorescent band of released siRNA from the formulations was detected (Fig. 6A).  
479 This result indicated that CR $\geq$ 2 was required to obtain a complete siRNA complexation.

480 Nevertheless, as a strong cationic polymer, poly-L-arginine is susceptible to cause a cytotoxicity. To  
481 ensure the safety of our NM-scFv for further applications, a MTT cytotoxicity test was performed on  
482 MDA-MB-231 cells. In this experiment, the formulations of siRNA, PLR, and NV2-scFv at different CR  
483 were incubated with cancer cells for 72h. Non-treated cells and cells treated with H<sub>2</sub>O<sub>2</sub> were used as  
484 controls. Cytotoxicity was observed for formulations with CR $\geq$ 20 compared to non-treated cells and no  
485 cytotoxicity was recorded for formulations with CR $\leq$ 10 (Fig. 6B). The levels of CR in our design of  
486 experiments were thus chosen between 2 and 10.

487 Concerning CS, the research of Bruniaux et al. showed that GFP down-regulation efficiency of non-  
488 targeted nanomedicine increased following CS up to its maximum at CS=30 and then decreased. The  
489 levels of CS in this study were varied from 10 to 50 to provide a maximal design space [8].



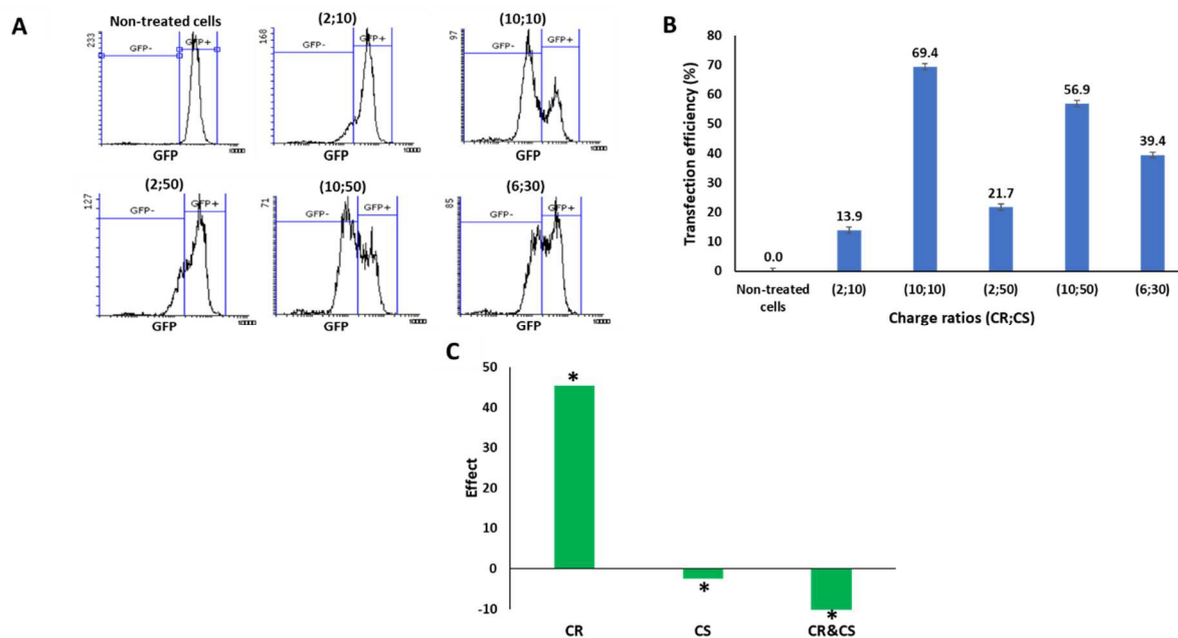
490

491 Fig. 6. Optimization of the PLR/siRNA charge ratio (CR). A: Gel retardation assay to detect free siRNA  
 492 in formulations of siRNA, poly-L-arginine and NV2-scFv at different CR with (+) or without (-) heparin.  
 493 B: Cellular viability of MDA-MB-231 cells treated with different formulations of siRNA, poly-L-arginine  
 494 and NV2-scFv at different CR.

495 Based on these results, a full factorial design of experiments was created with the levels varied from 2  
 496 to 10 for CR and from 10 to 50 for CS. Three center points were included in the model and one  
 497 response, the transfection efficiency, was analyzed (Table SI 2). The tested nanomedicine was made  
 498 of NV<sub>2</sub>-scFv to deliver a model siRNA (anti-GFP siRNA) into MDA-MB-231/GFP+ cells. All tested NM-  
 499 scFv were controlled in terms of size (Table SI 2). Cells were analyzed using flow cytometry to  
 500 measure the fluorescence intensity of GFP. Successful transfection of anti-GFP siRNA resulted in  
 501 reduced GFP fluorescence compared to non-treated cells that was presented by two populations of  
 502 cells following the GFP fluorescence intensity (Fig. 7A). The transfection efficiency was calculated  
 503 based on the percentage of the cells with reduced GFP fluorescence intensity over the analyzed cells  
 504 (Fig. 7B).

505 The regression analysis of the model indicated a R<sup>2</sup> of 99.76% with a predicted R<sup>2</sup> of 99.45% and an  
 506 adjusted R<sup>2</sup> of 99.66%. These values, as well as a non-significant lack of fit (p=0.226), indicated a  
 507 good fitting of the model, permitting further interpretation of the data. Together with the normalized  
 508 effects chart (Fig. 7C), CR was remarked to have the most significant impact on the transfection  
 509 efficiency (p<0.001). The transfection efficiency increased from 13.9±1.1% to 69.4±0.5% while CR  
 510 varied from 2 to 10. On the other hand, a significant impact but less than that of CR (Table SI 3) on  
 511 transfection efficiency was recorded for CS (p=0.01). The transfection efficiency increased when CS  
 512 decreased (effect<0). The interaction between CS and CR displayed also a significant impact on  
 513 transfection efficiency (p<0.001). This impact was less important than that of CR alone but more  
 514 important than CS alone (Table SI 3). When CR was low (CR=2), the transfection efficiency increased  
 515 with the increase of CS (13.9±1.1% for CS=10 vs 21.7±1.2% for CS=50). On the contrary, when CR  
 516 was high (CS=10), the transfection efficiency decreased following the increase of CS (69.4±0.5% for  
 517 CS=10 vs 56.9±1.9% for CS=50). These results indicated that there was an additive effect between  
 518 chitosan and PLR and an equilibrium of both polymers was necessary for a good transfection  
 519 efficiency. In summary, a high PLR amount and a low amount of chitosan were more favorable to high  
 520 transfection efficiency.

521 Finally, the optimal ratios were determined using the optimization function of Minitab. To obtain a  
 522 maximal transfection efficiency, the optimal CR and CS should be both fixed at 10. With these  
 523 parameters, the predicted transfection efficiency was  $69.4 \pm 0.5\%$ . This value corresponded to the value  
 524 obtained in the performed experiments. Comparison of this result with the size distribution of the  
 525 different formulations (between  $100.0 \pm 0.5\text{nm}$  and  $193.4 \pm 5.4\text{nm}$ ) suggested a correlation between the  
 526 maximal transfection efficiency and the lowest size (Table SI 2). In the literature, a similar  
 527 phenomenon has been observed in the study of Huang et al. for the internalization of gold  
 528 nanoparticles [30]. Concerning the charge ratios, our CR was higher than that in the study of Ben  
 529 Djemaa et al. (CR=2) but similar to the study of Zhao et al. for siRNA delivery [2,31]. Nevertheless, the  
 530 transfection efficiency was similar in three mentioned studies (around 70%). These experiments  
 531 demonstrated not only the perspective of our NM-scFv as a siRNA delivery system but also the major  
 532 effect of charge ratios and the size on the final gene silencing efficiency.



533  
 534 Fig. 7. Anti-GFP siRNA transfection efficiency. A: MDA-MB231 population following the GFP signal  
 535 after transfection. B: Mean transfection efficiency of NM-scFv made of NV2-scFv with different molar  
 536 ratios of Poly-L-Arginine/ siRNA (CR) and Chitosan/ siRNA (CS). C: Normalized effects chart for the  
 537 main factors (CS or CR) and two-factor interaction (CR&CS) on transfection efficiency. Asterisk (\*)  
 538 indicates statistically significant factors with  $p < 0.05$ .

### 539 3.3 Efficiency of the optimized targeted nanomedicine

#### 540 3.3.1 *In vitro* stealthiness

541 As shown in 3.1.1, the enhancement in PEG density improved the anti-EGFR scFv conjugation onto  
 542 NV and resulted in a better functionality for our NV-scFv. However, the initial purpose was to improve  
 543 the NM-scFv stealthiness that remained in question. Activation of the complement experiment (CH50

544 test) was performed to evaluate the impact of i) anti-EGFR scFv conjugation and ii) improvement in  
 545 PEG density on the NM stealthiness.

546 As the stealthiness of NM is also dependent on the physicochemical properties [26], all the  
 547 nanomedicines' physicochemical properties were controlled (Table II). There was an increase in the  
 548 size of nanomedicines compared to nanovectors ( $97.4 \pm 0.7$  nm for NM1 vs  $76.6 \pm 5.3$  nm for NV1) and  
 549 the charge was lightly positive due to the presence of exceeded cationic polymers ( $+4.5 \pm 3.5$  mV for  
 550 NM1 vs  $-2.8 \pm 2.4$  mV for NV1). Nevertheless, the size around 100 nm with a narrow size distribution  
 551 (PDI < 0.3) remained interesting for siRNA nanomedicines associated by electrostatic interactions.

552 Furthermore, the increase in PEG density might result in less condensed packaging of siRNA with  
 553 cationic polymers. It was correlated to an increase in the size of NM2 and NM2-scFv compared to  
 554 NM1 and NM1-scFv ( $109.6 \pm 4.5$  and  $100.0 \pm 2.3$  nm vs  $97.4 \pm 0.7$  and  $92.2 \pm 1.3$  nm) [32,33].

555 Table II. Physicochemical properties of the evaluated nanomedicines.

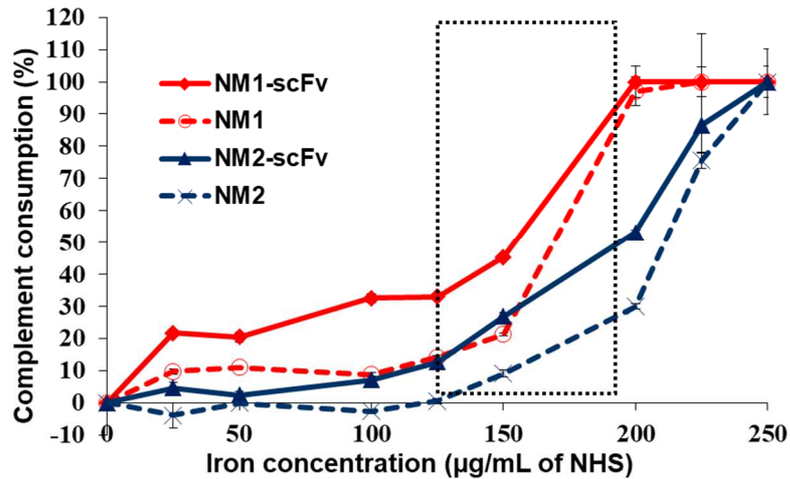
	NM1	NM1-scFv	NM2	NM2-scFv
D <sub>H</sub> (nm) (in intensity)	$97.4 \pm 0.7$	$92.2 \pm 1.3$	$109.6 \pm 4.5$	$100.0 \pm 2.3$
PDI	$0.27 \pm 0.02$	$0.23 \pm 0.01$	$0.27 \pm 0.01$	$0.25 \pm 0.02$
ζ (mV, pH 7.4)	$+4.5 \pm 3.5$	$+6.7 \pm 1.2$	$+5.4 \pm 1.3$	$+9.2 \pm 0.7$

556

557 Complement consumption was evaluated as the lytic capacity of the serum toward 50% of antibody-  
 558 sensitized sheep erythrocytes (CH50 units) after exposure to the evaluated nanomedicines (Fig. 8).  
 559 For the same type of NM and the same PEG density, the NM-scFv adsorbed a larger amount of serum  
 560 proteins than the nanomedicine without scFv (between 25 and 150 μg of iron/mL of NHS, NM1-scFv  
 561 consumed 20 to 40% of CH50 units vs 10 to 15% for NM1). This result showed that the presence of  
 562 conjugated anti-EGFR scFv as predicted increased the absorption of protein onto the NM's surface  
 563 and may consequently prevent the long-circulating properties of PEG [34].

564 Despite the bigger size that may lead to a decline in stealthiness [26], the increase in PEG density  
 565 helped NM2-scFv and NM2 to reduce the adsorption of serum proteins onto the nanomedicine surface  
 566 and improved their stealthiness. For the whole evaluated range of NM concentrations (from 0 to 250 μg  
 567 of iron/mL of NHS), CH50 unit consumption was more important for NM1-scFv and NM1 than that of  
 568 NM2-scFv and NM2. Especially, in the zone of interest from 125 to 188 μg of iron/mL of NHS, the  
 569 CH50 unit consumption was dramatically increased from 30 to 100% for NM1-scFv while that of NM2-  
 570 scFv varied from 10 to 50% and only reached 100% at 250 μg of iron/mL of NHS.

571 Though the confirmation of NM's stealthiness *in vivo* is unavoidable, these results revealed that by  
 572 increasing the PEG density, there was a great perspective to improve our NM-scFv stealthiness for the  
 573 further *in vivo* application.



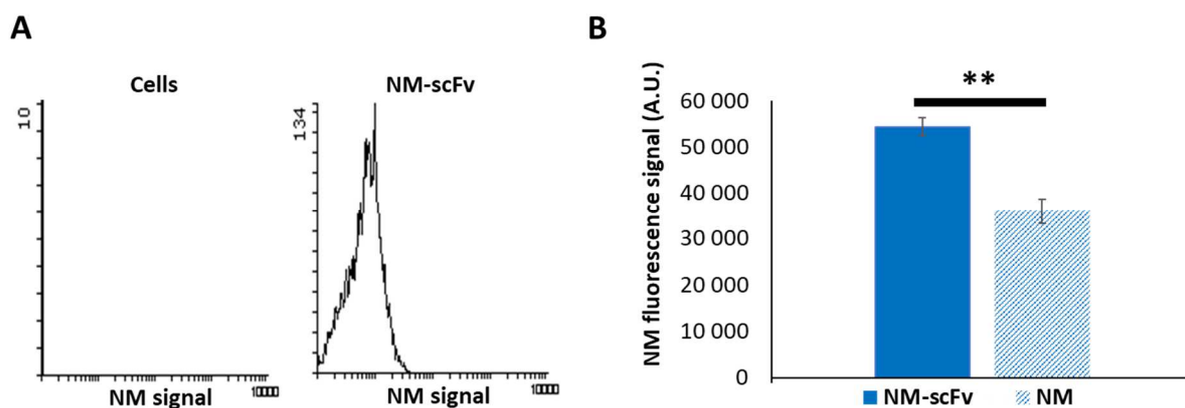
574

575 Fig. 8. Complement consumption by different NM according to iron concentration ( $\mu\text{g}$  of iron/mL of  
 576 normal human serum). Dotted line zone presents the zone of interest concentrations for *in vivo*  
 577 injection (125-188 $\mu\text{g}$  of iron/mL of NHS).

### 578 3.3.2 Cellular internalization

579 To confirm the active targeting properties of NM-scFv, a cellular internalization experiment was  
 580 performed with the optimal NM-scFv into MDA-MB-231 cells. The NM-scFv was prepared with control  
 581 siRNA at the optimal ratios of MS=10; CR=10; and CS=10. The fluorescence signal of Dylight<sup>TM</sup>680  
 582 was followed and compared. If the NMs internalize into the cancer cells, Dylight<sup>TM</sup>680 fluorescence  
 583 signal will be detected. Otherwise, there is no fluorescence detected for non-treated cells (Fig. 9A).

584 As shown in Fig. 9B, the Dylight<sup>TM</sup>680 fluorescence intensity of internalized NM-scFv was higher than  
 585 the non-targeted NM by a factor of 1.5 ( $p < 0.01$ ). This increase in fluorescence intensity demonstrated  
 586 a better cellular internalization of targeted NM compared to non-targeted NM. This better  
 587 internalization could be explained by the increased uptake via receptor-mediated endocytosis due to  
 588 anti-EGFR scFv. In comparison with the NV-scFv, the NM-scFv internalized better into the MDA-MB-  
 589 231 cells. This phenomenon could be explained by the charge of the nanoparticles. The positively  
 590 charged nanoparticles were more favorable in internalization than negatively charged nanoparticles  
 591 due to the interaction between positive charges of nanoparticles and negatively charged cell  
 592 membrane [30].



593  
 594 Fig. 9. Internalization of NMs into MDA-MB-231 cells by following Dylight™680 fluorescence signal. A)  
 595 Fluorescence profiles obtained with flow cytometer of cells treated (right figure) or not (left figure) with  
 596 NM-scFv. B) Fluorescence signal of internalized NM-scFv and NM. Double asterisk (\*\*) indicates  
 597 statistically significant factors with  $p < 0.01$ .

### 598 3.3.3 Transfection efficiency

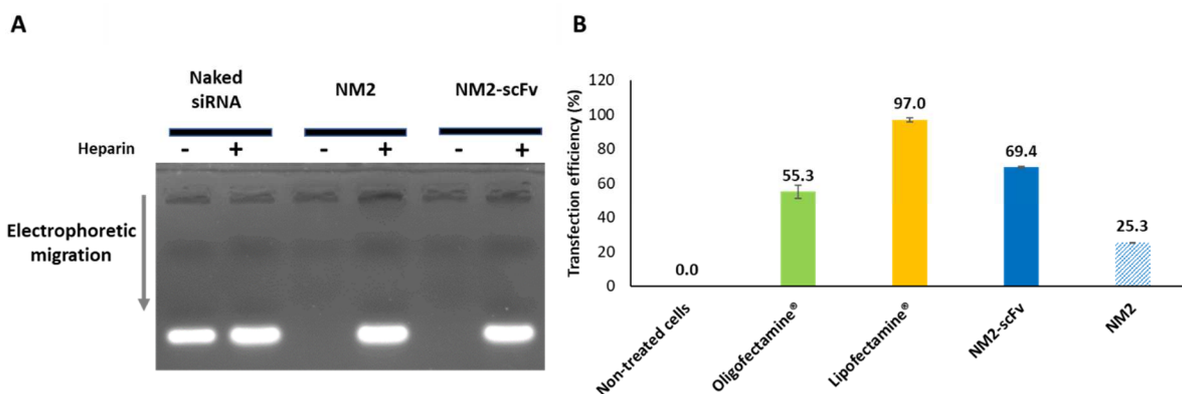
599 With the optimal ratios at CR=10 and CS=10, our targeted nanomedicine was compared to the non-  
 600 targeted NM and two other commercialized transfection agents: Oligofectamine® and Lipofectamine®  
 601 in terms of transfection efficiency. Oligofectamine® and Lipofectamine® are two liposomal agents  
 602 commonly used for *in vitro* siRNA transfection. The cationic lipids in the lipid bilayer structure can  
 603 interact with the negative siRNA to form an ionic complex to immobilize and deliver siRNA across the  
 604 cell membrane [35,36].

605 The tested formulations were firstly re-controlled in terms of physicochemical characteristics with DLS  
 606 and TEM analysis. On the one hand, the size of NM and NM-scFv evaluated with DLS was found to be  
 607 suitable for IV injection, which was inferior to 200nm with a narrow size distribution (Table SI 1). On  
 608 the other hand, the TEM analysis revealed that the state of SPIONs in different steps of synthesis or  
 609 formulation remained unchanged and not impacted by the other components. This observation may  
 610 confirm its stability and its integrity in the formulation with siRNA (Table SI 1; Fig. SI 1). As expected,  
 611 the size of SPION evaluated with TEM was smaller than that measured with DLS. The presence of  
 612 hydration layer around SPIONs' core can explain this difference when the sample was observed in the  
 613 solvated state with DLS. On the contrary, the TEM analysis allowed us to observe the SPION core in  
 614 the dry state.

615 In the next step, the siRNA protection capacity of the final formulation was evaluated on agarose gel  
 616 by electrophoresis method. Without added heparin, no fluorescent band was observed for the  
 617 complexed siRNA. On the contrary, with added heparin, the fluorescent band of complexed siRNA  
 618 was found with the same intensity as naked siRNA's (Fig. 10A). This experiment confirmed that siRNA  
 619 was completely complexed and protected.

620 In Fig. 10B, the transfection efficiency of the targeted nanomedicine was higher than that of the non-  
 621 targeted NM by a factor of 3.0 ( $69.4 \pm 0.5\%$  vs  $25.3 \pm 0.2\%$ ). This result demonstrated a targeted siRNA

622 delivery into TNBC cells of our NM-scFv. This targeting delivery could be explained by i) the specific  
 623 interaction between the conjugated anti-EGFR scFv and EGFR receptors overexpressed on MDA-MB-  
 624 231 cells that resulted in a better cellular internalization [34] and ii) the successful endosomal escape  
 625 of siRNA with the help of polymers. The transfection efficiency of Oligofectamine® and Lipofectamine®  
 626 in the same conditions was  $55.3 \pm 3.9\%$  and  $97.0 \pm 1.2\%$  respectively. Though Lipofectamine® displayed  
 627 a better transfection efficiency than our NM-scFv, these cationic lipids may cause a susceptible toxicity  
 628 due to their ability to disrupt the cellular and mitochondrial membrane. Moreover, these commercial  
 629 transfection agents are not suitable for *in vivo* application [35,36]. Thus, an efficient alternative for  
 630 siRNA delivery can be achieved both *in vitro* and *in vivo* with our NM-scFv.



631  
 632 Fig. 10. Targeted nanomedicine compared to non-targeted nanomedicine and commercialized  
 633 transfection agents. A: Gel retardation assay to detect free siRNA in nanomedicines with (+) or without  
 634 (-) heparin. B: Anti-GFP siRNA transfection efficiency of optimized targeted nanomedicine (NM2-scFv)  
 635 compared to non-targeted nanomedicines, Oligofectamine®, and Lipofectamine®.

## 636 4 Conclusion

637 In this study, a targeted nanomedicine for siRNA delivery into triple negative breast cancer cells was  
 638 developed and optimized. This targeted nanomedicine (NM-scFv) was based on the targeted  
 639 nanovector (NV-scFv) functionalized with a humanized anti-EGFR scFv antibody fragment as targeting  
 640 ligand and subsequently associated with siRNA, chitosan, and PLR.

641 The NV-scFv was firstly optimized in terms of PEG layer density. By changing the PEG/iron molar ratio  
 642 from 0.3 to 0.6, an increase by a factor of 3.7 in PEG layer density was achieved for our optimized NV-  
 643 scFv ( $873 \pm 4$  vs  $236 \pm 3$  of PEGs/NV-scFv). This increase showed an important impact on: i) the  
 644 number of conjugated active targeting ligands (13 vs 9 of scFv/NV-scFv) resulting in an increase in  
 645 cellular internalization by a factor of 2 for 24h of incubation time and ii) the improvement in stealthiness  
 646 of the final NM-scFv confirmed by CH50 test. In fact, in the zone of interest for *in vivo* injection ( $125$ -  
 647  $188 \mu\text{g}$  of iron/mL of NHS), there was a significant decrease in protein adsorption for our final NM-scFv  
 648 ( $10$ - $50\%$  vs  $30$ - $100\%$  of CH50 unit consumption).

649 In the second step, the formulation between NV-scFv and siRNA was performed with the addition of  
 650 chitosan and poly-L-arginine and optimized in terms of the ratios of components. Two necessary ratios  
 651 for the formulation were evaluated including: i) Chitosan/siRNA charge ratio (CS) and ii) PLR/siRNA

652 charge ratio (CR). The optimal parameters for our NM-scFv were determined at CS=10 and CR=10.  
653 Among these parameters, CR was shown to play the most important role on the transfection efficiency  
654 with a positive effect. CS and the interaction CR&CS presented also a significant impact with a  
655 negative effect but less important than that of CR. The optimized NM-scFv presented suitable  
656 physicochemical properties for IV injection with a size of  $100.0\pm 2.3\text{nm}$ , a narrow size distribution  
657 ( $\text{PDI}=0.25\pm 0.02$ ) and an almost neutral surface charge ( $+9.2\pm 0.7\text{mV}$ ). Moreover, the benefit of the  
658 active targeting with the anti-EGFR scFv was confirmed with a higher cellular internalization into MDA-  
659 MB-231 by a factor of 1.5. Finally, NM-scFv exhibited an efficient siRNA protection capacity that was  
660 confirmed with a gel electrophoresis experiment and a promising *in vitro* transfection efficiency  
661 ( $69.4\pm 0.5\%$ ) into TNBC cancer cells with anti-GFP siRNA.

662 All the results demonstrated the promising application of our targeted nanomedicine functionalized  
663 with humanized anti-EGFR antibody fragment for efficient targeted siRNA delivery into TNBC tumors.

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## 673 **Conflict of interest**

674 The authors declare that they have no conflict of interest.

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