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Peptide conjugation: before or after nanoparticle

formation?

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ABSTRACT

We report herein a detailed study concerning the impact of different bioconjugation and nanoformulation strategies on the *in vitro* targeting ability of peptide-decorated squalenoyl gemcitabine (SQdFdC) nanoparticles (NPs). NPs have been functionalized with the CKAAKN peptide, previously identified as efficient homing device within the pancreatic pathological microenvironment. Two approaches have been followed: (*i*) either the CKAAKN peptide was directly conjugated at the surface of preformed SQdFdC nanoparticles (conjugation after NP formation) or (*ii*) it was first reacted with a maleimide squalenoyl derivative before the resulting bioconjugate was co-nanoprecipitated with SQdFdC to form the peptide-decorated NPs (conjugation before NP formation). NPs were characterized with respect to mean diameter, zeta potential and stability over time. Then, their specific interaction with the sFRP-4 protein was evaluated by surface plasmon resonance. Although both the synthetic strategies allowed to formulate NPs able to interact with the corresponding receptor, enhanced target binding and better specific avidity were observed with CKAAKN-NPs functionalized before NP formation. These NPs displayed the highest cell uptake and cytotoxicity in an *in vitro* model of human MIA Paca-2 pancreatic cancer cells.

INTRODUCTION

The ability to selectively deliver drugs to tumor cells with minimal interaction with healthy tissues is a major challenge in chemotherapy. To this aim, suitable ligands able to bind to specific receptors, overexpressed on cancer cells and relatively downregulated on healthy tissues, have been successfully employed as a valuable strategy to specifically target tumors. By means of molecular recognition, such targeted therapy has improved the balance between efficacy and toxicity of conventional systemic anticancer treatments ² · ³ as observed, for instance, in the case of antibody-conjugated chemotherapeutic agents. Several antibody-drug conjugates are currently included in clinical trials with encouraging results and two examples, Adcetris (Takeda, Seattle Genetics) and Kadcyla (Roche, Genentech), have recently been approved by FDA and EMA. Telescope and toxicity of the control of the case of antibody-trials with encouraging results and two examples, Adcetris (Takeda, Seattle Genetics) and Kadcyla (Roche, Genentech), have recently been approved by FDA and EMA.

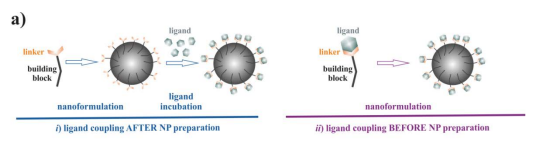
In parallel, the use of targeting ligands grafted to the surface of nanoscale drug delivery devices (*i.e.*, nanomedicines) appears as another promising approach. Compared to molecular conjugates, nanocarriers offer the advantage to protect the drug molecule from degradation and to better control drug release. Moreover, by tuning the number of ligand molecules onto the nanocarrier surface, targeting efficacy might be improved *via* a multivalent binding process. In the past 30 years, a plethora of targeted nanoparticles functionalized with various ligands belonging to the families of small molecules, polysaccharides, peptides, proteins or even antibodies has been developed. Nanomedicines are gaining more and more attention in cancer therapy with 5 examples on the market and 10 in clinical development.

However, not any ligand-decorated nanomedicine has reached the market and only 5 are currently in early clinical trials. 15-16

Several functionalization strategies and coupling methods have been developed, each of them exhibiting benefits and drawbacks (for systematic review see 11). In general, the targeting efficacy of a functionalized nanocarrier is governed by (i) the nature of the matrix and the

ligand, (*ii*) the selected coupling reaction, (*iii*) the disposition of the ligand at the surface and (*iv*) the ligand surface density.¹⁷⁻²² Two main ligation strategies have been used to display the targeting agents at the surface of the nanocarriers: the ligand can either (*i*) be directly coupled in aqueous solution to the surface of preformed nanocarriers or (*ii*) be linked to the nanocarrier components prior to nanocarrier construction (Figure 1a).

The first strategy is generally preferred because the ligand is directly grafted onto the nanoparticle surface and available for the receptor recognition, whereas the linkage of bulky ligands (such as polypeptides, proteins or antibodies) to the nanocarrier building material might alter its hydrophilic/lipophilic balance and its self-assembly properties. An undeniable advantage of the second strategy relies on the possibility to control the reaction yield, to purify and characterize the resulting conjugate. However, the solubilization in organic solvents, that is sometimes required, could cause denaturation of the proteins or antibodies compromising their affinity for the receptor. In addition, in this approach the ligand is susceptible of entrapment into the nanocarrier core during nanoparticle preparation, leading to insufficient surface coverage and lower availability for interaction with the corresponding receptor. Therefore, a comparative study about advantages/disadvantages of each strategy should be done for any new system, in order to achieve successful actively targeted nanoparticles.



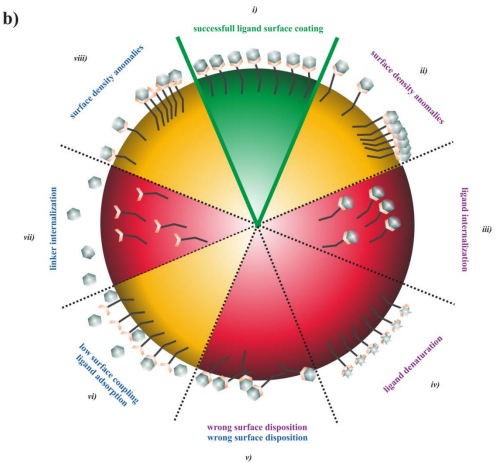


Figure 1. a) Strategies to prepare ligand-functionalized NPs: (*i*) the ligand can be coupled to the surface after nanoparticle formation or (*ii*) be directly conjugated to the nanocarrier building material prior to NP formation. **b**) Schematic representation of possible ligand disposition after both ligation strategies. Blue titles are referred to (*i*) ligand coupling after nanoparticle preparation and violet ones to (*ii*) ligand coupling before nanoparticle preparation. The green segment symbolizes the successful ligand coating, while yellow and red ones correspond to low and inefficient ligand disposition, respectively.

In this study, we report a rational approach for the design of ligand-targeted squalene-based nanoparticles for specific drug delivery to pancreatic tumor cells. The "squalenoylation" approach consists in the chemical linkage of squalene (SQ), a natural triterpene precursor in the sterol biosynthesis, to biologically active molecules. The obtained lipid bioconjugates are able to self-assemble in aqueous solution to form stable nanoparticles with high drug payloads.²³ Such innovative concept has firs been applied to gemcitabine (dFdC), which is the first-line therapeutic agent in pancreatic cancer treatment.²⁴ After intravenous administration, 4-(*N*)-trisnorsqualenoylgemcitabine (SQdFdC) nanoparticles exhibited impressively greater anticancer activity than the free drug against different tumor models (*e.g.*, experimental leukemia and pancreatic cancer).^{23, 25-27} However, the physio-pathological complexity of pancreatic cancer²⁸ urgently required the development of better targeted therapeutic approaches to enhance selective nanoparticle internalization by cancer cells, also avoiding potential side effects.

In this context, functionalized SQdFdC nanoparticles were prepared by linking the CKAAKN peptide, previously identified as efficient homing device within the pancreatic pathological microenvironment by phage display screening.²⁹ These nanoparticles were found to be more efficient than non CKAAKN-functionalized nanoparticles on a transgenic mice model of pancreatic cancer.³⁰ Herein, we performed a comparative study to establish the optimal CKAAKN peptide-SQdFdC nanoparticle ligation strategy. Briefly, CKAAKN peptide was conjugated to the nanoparticles by the well-explored thiol-maleimide Michael addition coupling strategy either (*i*) by reacting the targeting moiety with preformed nanoparticles obtained in accordance with the nanoprecipitation technique (Figure 1a – left panel) or (*ii*) by conjugating the peptide with a squalenoyl derivative before constructing nanoparticles by mixing the obtained bioconjugate with SQdFdC (Figure 1a – right panel). The resulting

CKAAKN-SQdFdC nanoparticles were characterized with respect to size, zeta potential, stability, receptor binding affinity and *in vitro* targeting ability.

RESULTS AND DISCUSSION

Preparation and characterization of non targeted nanoparticles. In order to obtain stable CKAAKN-functionalized SQdFdC nanoparticles, the formulation of the colloidal system was deeply investigated by tuning the preparation conditions. In particular, a maleimide groupbearing squalene derivative (SQMal) able to link the peptide *via* its thiol function was added to SQdFdC (1) in various molar ratios. Moreover, two different spacers were introduced between the squalene moiety and the maleimide group. In the case of the SQMal_{lipo} (3),³⁰ a 6-carbon atom lipophilic chain served as a spacer, while for the SQMal_{hydro} (4),³¹ a hydrophilic ether chain was inserted (Figure 2).

Figure 2. Structures of SQdFdC (1), CKAAKN peptide (2), SQMal_{lipo} (3), SQMal_{hydro} (4) and SQCKAAKN (5).

SQCKAAKN (5)

All NPs were prepared by nanoprecipitation, using acetone or ethanol as organic solvent and without adding any surfactant. Maleimide-bearing SQdFdC NPs were prepared by adding SQMal_{lipo} or SQMal_{hydro}.

SQdFdC and SQMal_{lipo} were co-nanoprecipitated at various SQdFdC/SQMal_{lipo} molar ratios (10:1, 5:1, 2:1) with a constant SQdFdC concentration (either in acetone or in ethanol), giving a suspension of narrow monodispersed NPs (N1). Addition of SQMal_{lipo} led to a slight size increase compared to SQdFdC NPs which displayed a mean diameter of 141 nm and 122 nm when prepared in acetone (N0_a) or ethanol (N0_e), respectively (Table 1).

SQdFdC/SQMal_{hydro} NPs (**N2**) were prepared by co-nanoprecipitation of SQdFdC and SQMal_{hydro} at two molar ratios (10:1, 5:1), with a constant SQdFdC concentration. Ethanol turned out to be the best solvent, while when acetone was employed NPs instantly aggregated. As observed for SQdFdC/SQMal_{lipo}, **N2**_e displayed a mean diameter of less than 200 nm and a narrow particle size distribution (Table 1). However, the presence of an ether chain between the maleimide group and the squalene moiety affected the nanoparticle stability, probably due to a modification of the internal inverted hexagonal phase structure of SQdFdC nanoparticles.³² On the contrary, a higher stability was observed when a lipophilic chain was used as spacer (SQMal_{lipo}).

According to these results, the SQMal_{lipo} was preferred to the SQMal_{hydro} for the preparation of peptide-targeted SQdFdC nanoparticles. Acetone was selected as solvent due to the higher colloidal stability of N1_a nanoparticles compared to those prepared in ethanol (N1_e). Using acetone as solvent and a low or medium amount of SQMal_{lipo} (SQdFdC/SQMal_{lipo} 10:1 or 5:1), no precipitation was detected after 1-week storage at 4°C.

Table 1. Nanoparticle composition and characterization.

NPs	Solvent used in the nanoprecipitation	Composition	Components molar ratio	Mean diameter (nm ± S.D.)	Polydispersity index	Zeta potential (mV ± S.D.)	Stability at 4 °C (days)
N0a	Acetone	SQdFdC	-	141 ± 42	0.087	-22 ± 4	3
N0 _e	Ethanol	SQdFdC	-	122 ± 8	0.103	-22 ± 5	3
N1a	Acetone	SQdFdC/SQMal _{lipo}	10:1	164 ± 42	0.066	-27 ± 3	7
	Acetone	SQdFdC/SQMal _{lipo}	5:1	171 ± 41	0.057	-29 ± 3	7
	Acetone	SQdFdC/SQMal _{lipo}	2:1	182 ± 20	0.047	-36 ± 3	3
N1 _a -P	Acetone	SQdFdC/SQMal _{lipo} / CKAAKN	10:1:0.5	281 ± 10	0.134	-15 ± 1	< 3
	Acetone	SQdFdC/SQMal _{lipo} / CKAAKN	10:1:0.25	187 ± 7	0.067	-11 ± 1	< 3
	Acetone	SQdFdC/SQMal _{lipo} / CKAAKN	5:1:0.5	Aggregation	/	/	/

N1 _a -P*	Acetone	SQdFdC/SQMal _{lipo} / CKAAKN	5:1:0.25	245 ± 16	0.141	-10 ± 2	3
N1e	Ethanol	SQdFdC/SQMal _{lipo}	10:1	132 ± 9	0.100	-22 ± 7	3
	Ethanol	SQdFdC/SQMal _{lipo}	5:1	167 ± 9	0.053	-22 ± 3	3
	Ethanol	SQdFdC/SQMal _{lipo}	2:1	Aggregation	/	/	/
N2 _a	Acetone	SQdFdC/SQMal _{hydro}	10:1	Aggregation	/	/	/
	Acetone	SQdFdC/SQMal _{hydro}	5:1	Aggregation	/	/	/
N2e	Ethanol	SQdFdC/SQMal _{hydro}	10:1	183 ± 5	0.055	-16 ± 1	< 3
	Ethanol	SQdFdC/SQMal _{hydro}	5:1	177 ± 3	0.053	-25 ± 3	< 3
N3	Acetone	SQMal _{lipo} 0.5 mg/mL	-	600 ± 180	0.091	-56.57 ± 3	7
	Acetone	SQMal _{lipo} 1 mg/mL	-	387 ± 101	0.068	-38.47 ± 3	7

	Acetone	SQMal _{lipo} 2 mg/mL	-	436 ± 148	0.099	-49.81 ± 4	7
N4-P	Ethanol	SQdFdC/SQCKAAKN	1:0.1	186 ± 104	0.032	11 ± 2	< 3
	Ethanol	SQdFdC/SQCKAAKN	1:0.05	897 ± 11	0.332	7 ± 23	< 3
N4-P*	Ethanol	SQdFdC/SQCKAAKN	1:0.01	168 ± 16	0.117	-6 ± 5	3

a NPs formulated using acetone as solvent

e NPs formulated using ethanol as solvent

st Formulations chosen for Surface Plasmon Resonance analysis and $in\ vitro$ studies

SQMal_{lipo} alone was able to spontaneously self-assemble in water forming nanoparticles (**N3**) which, however, displayed a large mean diameter (400-600 nm) (Table 1).

Compared to SQdFdC NPs (N0) or SQMallipo NPs (N3), mixed SQdFdC/SQMallipo NPs showed an intermediate mean diameter with higher colloidal stability, suggesting that the conanoprecipitation technique allowed an efficient incorporation of SQMallipo in the resulting composite nanoparticles probably due to the capacity of the squalene chain to insert into the nanoparticle core. SQMallipo NPs (N3) showed zeta potential values ranging from around -30 to -60 mV. Accordingly, the zeta potential value of N1a (SQdFdC/SQMallipo NPs prepared using acetone) gradually decreased (from around -20 to -40 mV) with the increase of SQMallipo in the formulation (Table 1). On the other hand, when ethanol was used as solvent (N1e), the amount of SQMallipo did not influence the zeta potential, suggesting that this experimental setting not only led to less stable NPs, but probably also did not allow the right exposition of the maleimide groups. Although for pure SQdFdC NPs the choice of the solvent did not influence the colloidal characteristics^{23, 33-34}, these results were in concordance with other experiments that highlighted the crucial role of the solvent in the nanoparticle self-organization³⁵⁻³⁸; they also supported the choice of using only the acetone-formulated SQdFdC/SQMallipo NPs for further investigations.

Preparation and characterization of targeted nanoparticles by coupling CKAAKN peptide to preformed nanoparticles. As mentioned before, the CKAAKN peptide is able to react with the maleimide function of the spacer *via* the cysteine thiol group present in its sequence, leading to the formation of a stable thioether bond. Noteworthy is that this site of derivatization does not alter the ligand/receptor recognition, as confirmed by the fact that the same function was used for performing the previous phage display *in vivo* screening.²⁹

CKAAKN conjugation on preformed nanoparticles allows peptide moieties to be disposed only at the surface of the carriers, thus being more available for the receptor recognition. For this purpose, the peptide thiol group was reacted with maleimide functions of N1a (SQdFdC/SQMallipo molar ratio 10:1 or 5:1) using different Mal/peptide molar ratios (1:0.5 or 1:0.25) to obtain N1a-P. Targeted NPs showed a higher mean diameter compared to untargeted ones (Table 1). In some cases, aggregation occurred and/or the stability was very low. The zeta potential value increased with the peptide amount (from -23 to -10 mV), probably due to the presence of two protonable amino groups in the CKAAKN moiety ((2) Figure 2). SQdFdC/SQMallipo/CKAAKN molar ratio 5:1:0.25 NPs (N1a-P*) showed the highest stability and only this nanoformulation was therefore used for further studies. The analyses of the amount of conjugated peptide on these NPs revealed the presence of 106866 peptide molecules per NP.

The interaction between the thiol groups and maleimide functions onto $N1_a$ surface was investigated by isothermal titration calorimetry (ITC) analysis. This technique is widely used in nanotechnology for characterizing thermodynamics and stoichiometry of intermolecular interactions allowing the evaluation of the association constant (K), the stoichiometry, the enthalpy (ΔH) and the entropy (ΔS) from which the Gibbs free energy (ΔG) of the process can be calculated.³⁹⁻⁴¹ The heat flows were determined when peptide aliquots were added to $N0_a$ (used as control) or $N1_a$ suspension placed in the titration cell accurately thermostated (Figure 3a). ITC thermograms showed two different signatures: (i) a first one mainly related to the non-specific adsorption of the peptide to the nanoparticle surface and (ii) a second one mostly attributed to the interaction with maleimide functions, observed only in the presence of these groups onto $N1_a$. Such a hypothetical mechanism could be supported by the enthalpograms of the interaction between cysteine and maleimide groups in solution (Figure 3b). Whether the thiol-maleimide Michael addition happened on the nanoparticle surface (Figure 3b – blue line)

or in solution (Figure 3b – black line), the thermodynamics parameters, including the stoichiometry of the interaction, were comparable. Despite the maleimide group was highly specific for thiols, at basic pH some secondary nucleophilic additions from the amine groups present in the two lysines of CKAAKN sequence could also take place, ⁴²⁻⁴³ modifying the recognition sequence of the peptide. For this reason, we also verified that no interactions between lysines and the maleimide groups were established in the same conditions (Figure 3b – grey line).

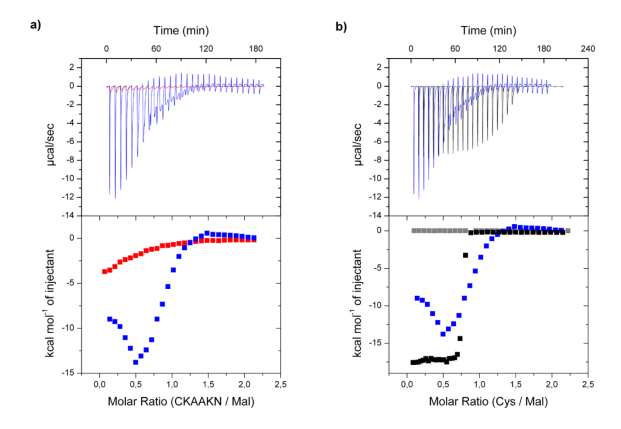


Figure 3. Isothermal titration calorimetry analysis of the interaction between CKAAKN, cysteine (Cys) and maleimide groups. The raw and the integrated binding heats are shown in the upper and lower panels, respectively. Heat flows accounting for dilution effects were further subtracted from each experimental heat flow. **a)** ITC thermograms upon injection of CKAAKN solution into either **N0**_a (red) or **N1**_a (blue) suspension. **b)** ITC thermograms upon injection of (i) a Cys solution into either maleimide solution (black) or **N1**_a suspension (blue) and (ii) injection of lysine into maleimide solution (grey).

To be noted that, according to ITC results, the Michael addition reaction occurred in a larger extent for CKAAKN/Mal molar ratios (N) higher than 0.5. On the light of these results, it appeared that the CKAAKN/Mal molar ratio (1:0.25) used to formulate the N1_a-P* led to the co-existence of both non-covalent interactions (demonstrated by ITC in other cases such as bio-

molecular recognition or inclusion complexes $^{39-41}$) and covalent conjugation of the CKAAKN onto the NP surface (Figure 1b – vi). However, higher CKAAKN/Mal molar ratios could not be used because they led to $\mathbf{N1}_a$ aggregation, (Table 1) likely due to peptide instability at the nanoparticle surface.

Preparation of targeted nanoparticles by coupling CKAAKN to SQMal_{lipo} prior to nanoparticle preparation. In this approach the CKAAKN peptide was previously conjugated to SQMal_{lipo} to give SQCKAAKN bioconjugate (5) which was then co-nanoprecipitated with SQdFdC at various SQdFdC/SQCKAAKN molar ratios (1:0.1, 1:0.05, 1:0.01), leading to the formation of peptide-functionalized NPs (N4-P). In this case only ethanol was used as organic solvent because of the limited solubility of SQCKAAKN in acetone. The main problem of this synthetic strategy is that the targeting agent could be buried in the nanoparticle core during the nanocarrier formation. Interestingly, the zeta potential gradually increased (from around -22 to +11 mV) with the increase of the percentage of SQCKAAKN in the formulations, suggesting the surface distribution of the peptide. As observed by DLS analysis, a SQdFdC/SQCKAAKN molar ratio higher than 1:0.01 led to nanoparticle destabilization and aggregation. Hence, SQdFdC/SQCKAAKN 1:0.01 molar ratio NPs (N4-P*), which showed a significant decrease of the absolute surface charge compared to untargeted NPs (N0_e) (-6 vs -22 mV) and only a slight increase of the mean diameter, were chosen for further investigations. According to the hydrophilic nature of the peptide we might assume that during the nanoprecipitation process the CKAAKN molecules localize on the NP surface. On the basis of the SQdFdC/SQCKAAKN 1:0.01 molar ratio used for N4-P* formulation, a value of 28385 molecules of CKAAKN per NP was obtained. Interestingly, N4-P* showed a marked reduction of the zeta potential value when increasing the pH of the medium from 7.4 to 9.8 (Figure 4). This probably resulted from the deprotonation of the lysine amino groups, thus supporting again the hypothesis of the presence of the peptide onto the surface of the nanoparticles, which is a crucial parameter to obtain an efficient targeting towards cancer cells. Only a slight increase of **N4-P*** mean diameter has been observed over a period of 72 h after incubation at 37 °C in water and in cell culture medium containing 10% FBS (SI Figure S4).

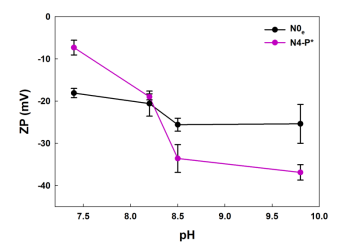


Figure 4. The surface charge of $N0_e$ and $N4-P^*$ as a function of pH was investigated by zeta potential (ZP) measurement at 25 °C after dilution with 0.02 M phosphate buffer at pH values ranging from 7.4 to 9.8. Values represent mean \pm SD, n=3.

Surface plasmon resonance analysis. While for N4-P* the peptide was covalently conjugated to the nanoparticle surface, in the case of N1_a-P* the ITC results suggested both adsorption and conjugation of the peptide on the NP surface. Covalent conjugation, in contrast to adsorption, ensures a stronger association between the ligand and the nanoparticles, especially after *in vivo* administration.⁴⁴ However, ligand association to the nanoparticle by weak interactions or surface adsorption has also been proven as an alternative strategy to tailor targeted nanoparticles.⁴⁴⁻⁴⁵ Therefore, both coupling strategies were further investigated and the peptide targeting capability of N1_a-P* and N4-P* was evaluated by surface plasmon resonance (SPR). Since it was reported that the sequence CKA-K shared motifs with the Wnt-2 protein ²⁹,

sFRP-4, a secreted frizzled-related protein that binds Wnt-2, was immobilized onto the sensor chip surface.^{30, 46} The ability of the free CKAAKN peptide to specifically interact with the sensor chip-immobilized sFRP-4 was also verified.³⁰ Although both $N1_a$ -P* and N4-P* were able to specifically interact with the receptor (which was not the case for untargeted $N0_e$), N4-P* enabled to reach the more important plasmonic signal after PBS washing (Figure 5).³⁰

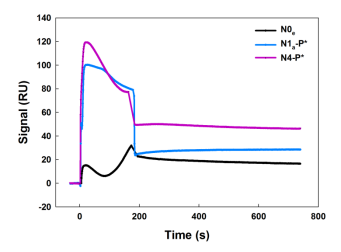


Figure 5. SPR sensorgrams (resonance units (RU) *versus* time) obtained by simultaneous injections of non functionalized SQdFdC NPs (**N0**_e) 180 μM, SQdFdC/SQMal_{lipo}/CKAAKN 5:1:0.25 NPs (**N1**_a-**P***) 180 μM, SQdFdC/SQCKAAKN 1:0.01 NPs (**N4-P***) 180 μM over sFRP-4 immobilized on two parallel channels of the same sensor chip.

The enhanced target binding and specific avidity of **N4-P*** suggested that the peptide conjugation prior to nanocarrier formation was the most efficient methodology to obtain enhanced binding and avidity, ensuring stronger association between nanoparticles and sFRP-4 receptor. This is of significance since earlier studies^{22, 47-51} have shown that the binding of a surface-functionalized nanoparticle to its target receptor is not only relied to the presence of the ligand on the nanoparticle surface.

In vitro studies. In order to confirm the on the bench SPR results, the receptor targeting ability as a function of the synthetic strategy was evaluated in *in vitro* cell culture experiments. To this purpose, a preliminary Western Blot analysis on several cell lines was performed in order to assess their frizzled-5(FZD-5) receptor expression (SI Figure S2). The MIA PaCa-2 cells were identified as high FZD-5-expressing cells and this result was further confirmed by immunocytochemistry (SI Figure S3). MIA PaCa-2 cells were then incubated for 6 h with BChol-green-labeled N0_e, BChol-green-labeled N1_a-P* or BChol-green-labeled N4-P*.

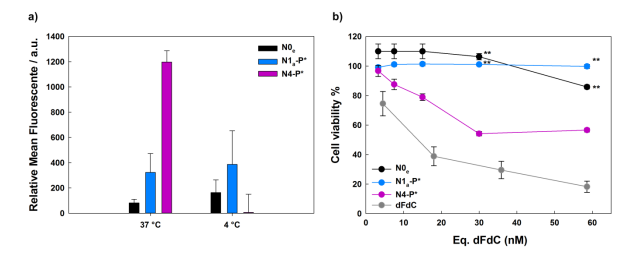


Figure 6. a) MIA PaCa-2 cell uptake quantification of SQdFdC NPs (N0_e)

SQdFdC/SQMal_{lipo}/CKAAKN 5:1:0.25 NPs (N1_a-P*) and SQdFdC/SQCKAAKN 1:0.01 NPs (N4-P*) after 6-h incubation at 37 °C and 4 °C. All NPs were labeled with BChol-green and incubated at a final concentration of 1 μ M eq. SQdFdC. b) Cell viability of MIA PaCa-2 cells treated with increasing concentrations of dFdC as free drug (dFdC), SQdFdC NPs (N0_e), SQdFdC/SQMal_{lipo}/CKAAKN 5:1:0.25 NPs (N1_a-P*) or SQdFdC/SQCKAAKN 1:0.01 NPs (N4-P*) for 72 h at 37 °C. Cytotoxicity of nanoparticles was compared to non-treated cells. Values represent mean \pm SEM. Statistical difference N0_e vs N4-P* and N1_a-P* vs N4-P* (Student's t-test with Bonferroni correction for multiple comparisons) is marked by ** (p < 0.01).

Although slight differences were detected after treatment with $N0_e$ and $N1_a$ - P^* , remarkably the cell fluorescence intensity dramatically increased (by 14–fold) after N4- P^* incubation (Figure 6a). These results clearly indicated a more efficient internalization of N4- P^* in comparison to $N1_a$ - P^* in MIA PaCa-2 cells.

When cells were incubated at 4 °C, the internalization of **N4-P*** dramatically decreased, suggesting that peptide-functionalized NPs were internalized *via* an energy dependent route. The internalization of the nanocarrier is a key issue for the drug to be efficient.⁵² Thus, in order to investigate whether the peptide functionalization enabled to achieve an increased drug efficacy compared to the untargeted NPs, the *in vitro* cytotoxicity of the different NPs and the free drug was investigated. As previously reported ²⁵, all NPs were less cytotoxic than free dFdC due to their prodrug nature. Nevertheless, **N4-P*** caused a higher cytotoxicity compared to both **N0**_e and **N1**_a-**P*** (Figure 6b). It is worth mentioning that for **N4-P*** the 50% inhibition of cell viability (IC₅₀) was already reached at 30 nM (eq. dFdC), while untargeted **N0**_e were

not efficient in killing cancer cells in these conditions. These results clearly demonstrated the ability of CKAAKN-functionalized NPs not only to keep the anticancer activity of the SQdFdC, but also to significantly enhance the cytotoxicity. As expected, according to the ITC, SPR and internalization data, N1_a-P* did not improve cytotoxicity in comparison to N0_e treatment, thus suggesting the absence of specific tumor targeting ability when the peptide was mainly adsorbed at the NP surface.

To further investigate the ability of N4-P* to specifically target cancer cells, the interaction with NIH/3T3, a non cancerous fibroblastic cell line, was studied and compared to the uptake by MIA PaCa-2 cancer cells. Western blot analysis showed that NIH/3T3 healthy cells displayed a low expression of FZD-5 receptor (Supporting Information S2). As shown in Figure 7, non-functionalized nanoparticles (N0e) were taken-up mainly by NIH/3T3 fibroblasts in a nonspecific manner, resulting in higher accumulation in this cell line. On the contrary, N4-P* were internalized into MIA PaCa-2 cells by a receptor-mediated mechanism, which allowed nanoparticles achieving highly selective tumor cell uptake and toxicity, while decreasing nonspecific accumulation into healthy cells (Figure 7a). As shown by fluorescence quantification studies after 6 h, the peptide-functionalization caused a 6-fold decrease of nanoparticles uptake by healthy NIH/3T3 cells compared to MIA Paca-2 cells (Figure 7b). Cytotoxicity studies clearly demonstrated that peptide functionalization was able to selectively increase cytoxicity toward MIA PaCa-2 pancreatic cancer cells. Indeed, the same concentration of N4-P* (i.e, 30 nM eq. dFdC) responsible of the 50% reduction of cell viability on malignant MIA PaCa-2 cells maintained more than 93% of viable healthy NIH/3T3 cells (Figure 7a). Of note, non functionalized NPs were not efficient in killing cancer cells, since 100% cell viability was still observed after incubation at the same concentration. The higher uptake of N0_e by NIH/3T3 fibroblasts was not accompanied by important cytotoxicity, as consequence of a lower sensitivity of this cell line to gemcitabine at the tested doses, as already reported in literature⁵³- ⁵⁴. The selective capture of functionalized NPs by cancer cells was further confirmed in a coculture experiment of healthy NIH/3T3 cells (stained in green) and MIA PaCa-2 cells (unstained) incubated with red fluorescently-labeled nanoparticles (Figure 7c,d). Confocal images clearly showed similar uptake of non-functionalized NPs in both tumor and healthy cells (Figure 7c), whereas **N4-P*** were selectively captured by tumor cells and healthy cells were saved from nanoparticles recognition (Figure 7d).

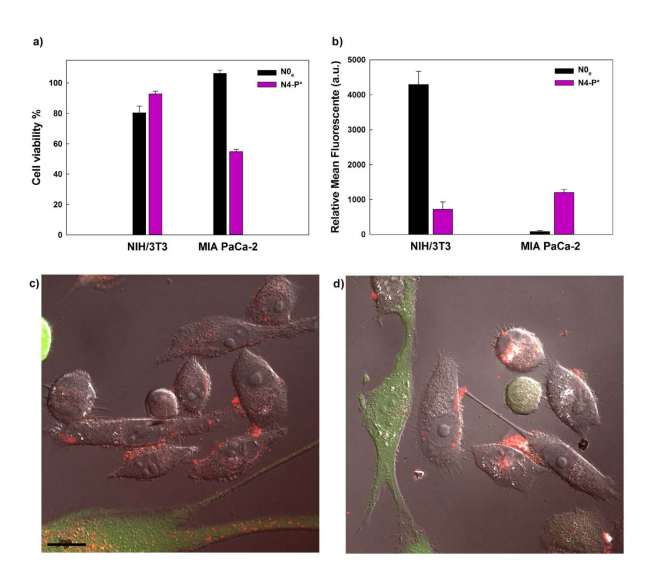


Figure 7. (a) Viability of NIH/3T3 and MIA Paca-2 cells treated for 72 h with SQdFdC NPs (N0_e) and SQdFdC/SQCKAAKN NPs (N4-P*) (30 nM eq. dFdC). Values represent mean ±

SEM. (b) Quantification of cellular uptake of BChol-green-labeled SQdFdC NPs ($N0_e$) and BChol-green-labeled SQdFdC/SQCKAAKN NPs (N4-P*) in NIH/3T3 and MIA PaCa-2 cells at 37 °C. Values represent mean \pm SEM. (c) Confocal laser-scanning microscopy images of a coculture of NIH/3T3 (green) and MIA PaCa-2 cells (unstained) after incubation with BChol-red-labeled SQdFdC NPs ($N0_e$) and (d) BChol-red-labeled SQdFdC/SQCKAAKN NPs (N4-P*). Scale bar = 20 μ m.

CONCLUSIONS

This study demonstrates that the best strategy to prepare CKAAKN-functionalized SQdFdC NPs relies in the conjugation of the peptide to the SQMal_{lipo} derivative prior nanoparticle formation. Taken together, our results clearly highlight that actively targeted nanoparticles should not be simply designed as single modular assemblies of scaffold, linker and targeting ligand, but rigorous studies must be performed in order to clarify how different structural components collaborate to reach the most efficient cell targeting ability. In a nutshell, the successful functionalization of SQdFdC nanoparticles makes this system a good example to highlight the subtle equilibrium which governs the design of actively targeted NPs able to effectively improve anticancer activity.

EXPERIMENTAL PROCEDURES

Chemicals. Gemcitabine (2',2'-difluorodeoxycitidine, dFdC) hydrochloride was purchased from Sequoia Research Products Ltd. (Pangbourne, UK). CKAAKN peptide was purchased from CASLO Laboratory Aps (Lyngby, Denmark). Squalene, dextrose, maleimide and all other reagents were obtained by Sigma-Aldrich Chemical Co. (Milan, Italy). Squalene derivatives (Figure 2) (4-(*N*)-trisnorsqualenoylgemcitabine (SQdFdC) (1)²³, 6-(maleimidocaproyl)hydrazone of squalene (SQMal_{lipo}) (3)³⁰, 2-[2-(2,5-dioxo-2,5-dihydro-1H-pyrrol-1-yl)ethoxy]ethyl (4E,8E,12E,16E)-4,8,13,17,21-pentamethyl-docosa-4,8,12,16,20-pentaenoate (SQMal_{hydro}) (4)³¹) were obtained as previously reported. All solvents were of analytical grade from Carlo Erba Reagenti (Milan, Italy) or VWR (Fontenay-sous-Bois, France).

Preparation and characterization of non targeted nanoparticles. SQdFdC, SQdFdC/SQMal_{lipo}, SQdFdC/SQMal_{lipo} and SQMal_{lipo} nanoparticles (NPs) were prepared by nanoprecipitation.^{23, 55} (Table 1) Practically, for SQdFdC NPs (**N0**), 2 mg of SQdFdC were

dissolved in acetone ($N0_a$) or ethanol ($N0_e$); the organic solution was then added dropwise under magnetic stirring into MilliQ[®] water (solvent/water 1:2 v/v). Formation of the nanoparticles occurred spontaneously without using any surfactant. After solvent evaporation under reduced pressure, an aqueous suspension of nanoparticles was obtained (final SQdFdC concentration: 2 mg/mL).

For SQdFdC/SQMallipo NPs (N1), SQdFdC and SQMallipo were co-dissolved in acetone (N1a) or ethanol (N1e) in various SQdFdC/SQMallipo molar ratios (10:1 (2 mg of SQdFdC and 0.183 mg of SQMallipo), 5:1 (2 mg of SQdFdC and 0.366 mg of SQMallipo) and 2:1 (2 mg of SQdFdC and 0.915 mg of SQMallipo)). The organic solution was then added dropwise under magnetic stirring into MilliQ® water (solvent/water 1:2 v/v). Formation of the nanoparticles occurred spontaneously without using any surfactant. After solvent evaporation under reduced pressure, an aqueous suspension of nanoparticles was obtained (final SQdFdC concentration: 2 mg/mL). For SQdFdC/SQMalhydro NPs (N2), SQdFdC and SQMalhydro were co-dissolved in ethanol in various molar ratios (10:1 (2 mg of SQdFdC and 0.164 mg of SQMalhydro) and 5:1 (2 mg of SQdFdC and 0.328 mg of SQMalhydro)). The organic solution was then added dropwise under magnetic stirring into MilliQ® water (ethanol/water 1:2 v/v). Formation of the nanoparticles occurred spontaneously without using any surfactant. After solvent evaporation under reduced pressure, an aqueous suspension of nanoparticles was obtained (final SQdFdC concentration: 2 mg/mL).

For SQMal_{lipo} NPs (**N3**), 2 mg of SQMal_{lipo} were dissolved in acetone at various concentrations. After the addition of the organic solution to MilliQ[®] water (acetone/water 1:2 v/v) and solvent evaporation, an aqueous suspension of nanoparticles was obtained (SQMal_{lipo} concentration: 0.5, 1 or 2 mg/mL).

Fluorescently-labeled SQdFdC NPs and fluorescently-labeled SQdFdC/SQMal_{lipo} NPs were obtained using the same procedure, unless 1% (w/w) of the fluorescent probe CholEsteryl

BODIPY® FL C12 (BChol-green) or CholEsteryl BODIPY® 542/563 C11 (Life Technologies, Molecular Probes, Saint Aubin, France) was dissolved in the ethanolic solution before dropwise addition into water.

The mean particle size and the polydispersity index of all formulations were measured by dynamic light scattering (DLS) with a Nano ZS from Malvern (UK) (173° scattering angle) at 25 °C. The measurements were performed after dilution of the NP suspensions (1/25 v/v) in MilliQ® water. The NP surface charge was investigated by zeta potential measurements at 25 °C after dilution with 0.05 mM KCl solution applying the Smoluchowski equation and using the same apparatus. Measurements were carried out in triplicate. The colloidal stability of the formulations was evaluated by measuring the size and the zeta potential of the nanoparticles over a storage period of 7 days at 4 °C.

Preparation and characterization of targeted nanoparticles by coupling CKAAKN peptide to preformed nanoparticles. CKAAKN stock solution (100X) was prepared by dissolving the peptide in 1 mM freshly degased phosphate buffer at pH 8.2. To obtain CKAAKN-conjugated SQdFdC/SQMal_{lipo} NPs (N1-P), 100 μL of N1_a were added dropwise under gentle magnetic stirring to 1 mL of CKAAKN working solution (1X) in various molar ratios (SQdFdC/SQMal_{lipo}/CKAAKN molar ratio 10:1:0.5 (0.2 mg of SQdFdC, 0.0183 mg of SQMal_{lipo} and 0.0098 mg of CKAAKN) or 10:1:0.25 (0.2 mg of SQdFdC, 0.0183 mg of SQMal_{lipo} and 0.0049 mg of CKAAKN) or 5:1:0.5 (0.2 mg of SQdFdC, 0.036 mg of SQMal_{lipo} and 0.0196 mg of CKAAKN) or 5:1:0.25 (0.2 mg of SQdFdC, 0.036 mg of SQMal_{lipo} and 0.0098 mg of CKAAKN); final SQdFdC concentration: 0.18 mg/mL) (Table 1). The reaction was carried out for 1 h at room temperature under stirring. The CKAAKN-functionalized nanoparticles (N1_a-P) were then purified from unreacted peptide by centrifugal filter units (30 kDa) at 14000 g for 1 minute (Amicon Ultra-0.5, Millipore). The mean size, the polydispersity index and the zeta potential of purified N1_a-P were determined at 25 °C by DLS as described

above. The stability of the formulations was evaluated by measuring the size and the zeta potential of the nanoparticles over a storage period of 7 days at 4 °C. The amount of peptide molecules at the surface of the NPs has been evaluated by: i) titration of unreacted peptide in an unpurified aliquot of conjugated nanoassemblies by reaction with the Ellman's reagent (5,5'-dithio-bis-(2-nitrobenzoic acid), DTNB) by quantitative UV spectrophotometric analysis (absorbance at 412 nm)⁵⁶; ii) a retro-titration of the available maleimide groups present on the nanoparticle surface by quantitative fluorimetric analysis (Maleimide Quantification Assay Kit (Fluorometric) Abcam, ab112141) according to manufacturer's instructions.

Isothermal titration calorimetry (ITC) analysis of N1_a nanoparticles. The interaction between the maleimide (Mal) function and the CKAAKN peptide was investigated by isothermal titration calorimetry. The ITC instrument (VP-ITC, MicroCal, GE Healthcare Life Sciences, Velizy-Villacoublay, France) was periodically calibrated either electrically, using an internal electric heater, or chemically by measuring the dilution enthalpy of methanol in water. This standard reaction was in excellent agreement (1-2%) with the MicroCal constructor data. ⁵⁷⁻⁵⁸ In a typical experiment, aliquots of 10 μL of CKAAKN peptide, cysteine (Cys) or lysine in 0.03 M phosphate buffer solution at pH 8.2 (3.1 mM eq. Cys or 12.4 mM eq. lysine) filled into 283 μL syringe were used to titrate a suspension of either N1_a or maleimide in 0.03 M phosphate buffer solution at pH 8.2 (0.31 mM eq. Mal) into the calorimetric cell accurately thermostated at 25 °C. The corresponding heat flows were recorded as a function of time. Intervals between injections were 600 s and agitation speed was 220 rpm. Control experiments with N0_a into the calorimetric cell were carried out using the equivalent total squalene amount present in N1_a. Background of titration consisted on injecting the CKAAKN peptide or Cys in solely phosphate buffer solution placed in the sample cell.

The interaction process was analyzed by employing the models proposed in the Windows-based Origin 7 software package supplied by MicroCal. Based on the concentrations of the two

species, the software used a nonlinear least-squares algorithm to fit the series of heat flows to an equilibrium binding equation, providing the best-fit values of the stoichiometry, binding constant (K) and change in enthalpy (ΔH). From these results, the differences in free energy (ΔG) and the entropy (ΔS) were deducted according to the equation:

$$\Delta G = -RT \ln K = \Delta H - T\Delta S$$

Preparation of targeted nanoparticles by coupling CKAAKN to SQMallipo prior to nanoparticle preparation. To obtain SQdFdC/SQCKAAKN nanoparticles (N4-P), CKAAKN peptide was first conjugated to SQMallipo to give SQCKAAKN (5) (SI Figure S1). To obtain SQMallipo (3), a solution of 1,1',2-trisnorsqualenic aldehyde (8) (0.334 g, 0.868 mmol) in CH_2Cl_2 was added to dry methanol (15 mL). The resulting mixture was sonicated few minutes until complete dissolution. [6-(maleimido)hexanamido]azanium trifluoroacetate (7) (0.306 g, 0.868 mmol) and 4 Å molecular sieves (200 mg) were then added and the reaction mixture was stirred for 1 h at room temperature under nitrogen. The formation of the desired product (3) was monitored by TLC (petroleum ether/ethyl acetate 1/1 v/v, R_f : 0.65). The mixture was filtered and concentrated under reduced pressure. The residue was taken into water (5 mL) and extracted with CH_2Cl_2 (3 x 15 mL). The combined organic phases were dried over anhydrous MgSO₄ and concentrated *in vacuo*. Purification by flash-chromatography on silica column, eluting with a gradient of petroleum ether to petroleum ether/ethyl acetate 60/40 v/v, gave the product as a light yellow waxy material (0.211 g, 63% yield) (SI Figure S1).

 m/z(%) 81 (70), 110 (100), 192 (55), 591 (3). HPLC analysis: Symmetry C18 column, 5 μ m (Merck, Italy) equipped with a C18 column guard, elution with 100% methanol, detection by UV adsorption measurement at 237 nm (flow rate 1 mL/min, t_r =5.79 min). Peak heights were recorded and processed on a CBM-10A Shimadzu interface.

To synthesize the Michael adduct of CKAAKN (2) and 6-(maleimidyl)-hexanoic acid (trisnorsqualenylidene)-hydrazide (3) (SQCKAAKN, (5)), a mixture of SQMal_{lipo} (3) (13.5 mg, 0.0229 mmol) and CKAAKN peptide (2) (7.25 mg, 0.0114 mmol) in dimethylformamide (DMF)/H₂O 3/1 v/v (2 mL) was stirred for 3 h at 40 °C. The reaction mixture was then concentrated under reduced pressure and the crude product was taken into diethyl ether to remove unreacted maleimide derivative. The supernatant was withdrawn after decantation (3 times). In order to eliminate unreacted peptide, the solid was dissolved in methanol and filtered through a sintered glass funnel. The product was obtained as a translucent waxy material (10 mg, 0.0082 mmol, 70% yield).³⁰ (SI Figure S1).

¹H NMR (CD₃OD, 400 MHz) due to the presence of a 7:3 E/Z mixture at the hydrazone bond and to the asymmetric center on the maleimide moiety some signals are split, δ : 7.43 (t, J = 5.4Hz, 0.7H, CH=NNHCO), 7.28 (m, 0.3H, CH=NNHCO), 5.20 (t, J = 6.6 Hz, 1H, $HC = C(CH_3)$), 5.15-5.06 (m, 4H, $HC=C(CH_3)$), 4.71 (t, J=5.6 Hz, 1H, $NOCCH(CH_2CONH_2)NH$), 4.42-4.25(m, 5H, NOCCH(CH₃)NH, NOCCH(CH₂CH₂CH₂CH₂NH₃)NH, NOCCH(CH₂S)NH₂,), 3.70-3.60 (m, 1H, CH₂S(CHCH₂CONCO), 3.52 (t, J = 7.0 Hz, 2H, CH₂N-maleimide), 3.50-3.43 (m, 1H, $CH_2SCHCH_2CO)$, 3.30-3.10 (m, 3H, $CH_2S(CHCH_2CONCO)$, $CH_2S(CHCH_2CONCO)$, 3.02-2.92 (m, 4H, $CH_2CH_2CH_2CH_2NH_3$), 2.83 (dd, J = 16.0 Hz, J = 16.0 Hz6.4 Hz, 1H, NOCCH(CH_2CONH_2)NH), 2.76 (dd, J = 16.0 Hz, J = 4.8 Hz, 1H, NOCCH(CH₂CONH₂)NH), 2.62-2.30 (m, 4H, CH₂CH₂CON, CH₂CH₂C=NNH), 2.25-2.15 (m, 4H, $CH_2CH_2NH_3$), 2.17-1.85 (m, 22H, $=C(CH_3)CH_2CH_2$ $NOCCH(CH_2CH_2CH_2CH_2NH_3)NH)$, 1.75-1.65 (m, 4H.

NOCCH(CH₂CH₂CH₂CH₂NH₃)NH), 3H, 1.67 (s, $=C(CH_3)_2),$ 1.64 (s, 3H, $=C(CH_3)CH_2CH_2),$ 1.60 (s. 12H, $=C(CH_3)CH_2CH_2),$ 1.60-1.45 (m, 4H, =NHNCOCH₂CH₂CH₂CH₂CH₂N-maleimide), 1.40 (d, J = 7.2 Hz, 6H, COCH(CH₃)NH), HNCOCH₂CH₂CH₂CH₂CH₂CH₂N-maleimide); MS (+ESI), m/z (%), 1226.0 (100) [M+H]⁺, 614.5 (3) $[M+2H]^{2+}$; HRMS (+ESI) calcd for $C_{62}H_{105}N_{12}O_5S$: 1225.7741; found 1225.7690. Nanoparticles were then prepared by nanoprecipitation. Practically, SQdFdC and SQCKAAKN were dissolved in 0.1 mL of ethanol in various molar ratios (SQdFdC/SQCKAAKN 1:0.1 (3.42 mg of SQdFdC and 0.64 mg of SQCKAAKN), 1:0.05 (3.42 mg of SQdFdC and 0.32 mg of SQCKAAKN) and 1:0.01 (3.42 mg of SQdFdC and 0.064 mg of SQCKAAKN)) and this organic solution was then added dropwise under magnetic stirring into 1 mL of MilliQ® water (ethanol/water 0.1:1 v/v). After solvent evaporation under reduced pressure, an aqueous suspension of nanoparticles (N4-P) was obtained (final SQdFdC concentration: 3.4 mg/mL) (Table 1). The mean size and the polydispersity index of the formulations were determined as described above. The NP surface charge was investigated by zeta potential measurements at 25 °C after dilution with 0.05 mM KCl solution or 0.02 M phosphate buffer at pH values ranging from 7.4 to 9.8, applying the Smoluchowski equation and using the same apparatus. Fluorescently-labeled nanoparticles were prepared as described above. The stability of N4-P* has been investigated over a period of 72 h after incubation at 37 °C in water and in cell culture medium containing 10% FBS.

Surface plasmon resonance (SPR) analysis. Interaction analyses were performed on a BIAcore T100 instrument using CM5 Series S sensor chip from GE Healthcare Life Sciences Europe. The activation of the carboxymethylated dextran matrix was performed with a mixture of 0.1 M *N*-ethyl-*N*'-(3-diethylaminopropyl)-carbodiimide (EDC) / 0.4 M *N*-hydroxysuccinimide (NHS) (1:1 v/v) (GE Healthcare) for 7 min. Running buffer for the immobilization process was 150 mM phosphate buffer, pH 7.4 at a flow rate of 10 μL/min. The

sFRP-4 receptor was then injected over the activated surface at 10 μg/mL in 10 mM sodium acetate buffer at pH 4.4 during 7 min, giving an average immobilization level of 2800 resonance units (RU). A solution of 1 M ethanolamine hydrochloride at pH 8.5 was then injected during 7 min to block the unreacted sites of the matrix. For reference, an additional blank flow channel was prepared in accordance with the same process without injecting the receptor over the surface. Binding capacities of the functional surfaces were tested with 180 s-injections of a peptide solution, N0ε, N1a-P* (SQdFdC/SQMal/CKAAKN molar ratio 5:1:0.25) or N4-P* (SQdFdC/SQCKAAKN molar ratio 1:0.01) at different concentrations. All experiments were carried out in duplicate at 25 °C in Dulbecco-PBS (D-PBS) running buffer on the InterMol-IPSIT platform (Châtenay-Malabry, France).

Cell culture. Human pancreatic carcinoma cell lines MIA PaCa-2, BxPC3, PANC-1, breast cancer cell line MCF-7 and embryonic murine fibroblasts NIH/3T3 were obtained from the American Type Culture Collection (Molsheim, France) and maintained as recommended. Briefly, MIA PaCa-2, PANC-1, MCF-7 and NIH/3T3 cells were grown in Dulbecco's Modified Eagle Medium (DMEM) supplemented with 10% heat-inactivated (56 °C, 30 min) fetal bovine serum (FBS). In addition, MIA Paca-2 cell medium was supplemented with 2.5% heat-inactivated horse serum (Gibco, Saint Aubin, France). BxPC3 cells were maintained in Roswell Park Memorial Institute medium (RPMI 1640 Lonza, Braine-l'Alleud, Belgium) supplemented with 10% heat-inactivated FBS. All media were further supplemented with 50 U.mL⁻¹ penicillin and 50 U.mL⁻¹ streptomycin (Lonza, Levallois, France). Cells were maintained in a humid atmosphere at 37 °C with 5% CO₂. Cells were used between passage 3 and 12 after thawing.

Western Blot. Pancreatic adenocarcinoma (BxPC3, MIA PaCa-2, PANC-1), breast cancer (MCF-7) and embryonic fibroblast cells (NIH/3T3) were lysed in RIPA buffer (Sigma-Aldrich) supplemented with PierceTM phosphatase and protease inhibitor cocktail (Thermo Fisher Scientific, Perbio Science, Brebières, France), vortexed and then centrifuged for 5 min

at 3000 g. The protein concentration in the supernatant was then measured using a colorimetric assay (Bio-Rad Protein Assay Dye, Bio-Rad Laboratories, Marnes-la-Coquette, France). Equal amount of proteins (30 µg) were boiled for 10 min with the same volume of Laemmli Sample Buffer (Bio-Rad Laboratories, Marnes-la-Coquette, France), supplemented with 5% betamercaptoethanol and then separated in sodium dodecylsulphate (SDS)-polyacrylamide gels (Mini-Protean - TGX 4-15% - Precast Gels Bio-Rad) and electrically transferred to polyvinylidene difluoride (PVDF) membrane using a semi-dry transfer system (20 V, 1 h). The membrane was blocked with 5% dry milk suspension in 0.1% Tween 20 in PBS (blocking buffer), then incubated for 2 h at room temperature with the primary antibody solution in blocking buffer and, after 30 min washing in PBS-Tween 20 0.1% buffer, incubated for 1 h at room temperature with the secondary antibody solution in blocking buffer. Antibodies were used at the following dilutions: polyclonal rabbit anti-FZD-5 diluted 1/1000 (Millipore), mouse anti-β-actin diluted 1/3000 (Sigma-Aldrich), goat anti-mouse secondary antibody conjugated to horse radish-peroxydase diluted 1/2000 (SantaCruz Biotechnology, Clinisciences, Nanterre, France) and goat anti-rabbit secondary antibody conjugated to horse radish-peroxydase diluted 1/5000 (SantaCruz Biotechnology, Clinisciences). Detection of chemiluminescence was performed using the ClarityTM Western ECL substrate (Bio-Rad Laboratories) and images were captured by the ChemiBIS system from DNR Bioimaging Systems (Thoiry, France). MCF-7 cell line was used as a positive control.⁵⁹

Immunocytochemistry. MIA PaCa-2 cells (2.5 x 10⁴ cells/mL) were plated in growing medium onto microscopic glass slides in 24-well plate and incubated for 24 h at 37 °C with 5% CO₂. Nuclei were stained with 10 μM Syto Red Fluorescent dye (SYTO61 - S11343, Life Technologies, Molecular Probes, Saint Aubin, France) according to the manufacturer's protocol. After staining, cells were fixed with 4% paraformaldehyde in PBS, incubated for 30 min at room temperature in blocking solution (1% bovine serum albumin, 0.3 M glycine, 0.1%

Tween 20 in PBS) and then with the polyclonal rabbit anti-FZD-5 diluted 1/500 (Abcam, UK) for 1 h. Then, the cells were washed with PBS-Tween 20 0.1% for 30 min and stained for 1 h at room temperature in dark with an Alexa Fuor®555-conjugated secondary antibody diluted 1/500 (Abcam, UK). Cells were then rinsed three times with PBS and analyzed by confocal laser scanning microscopy. Cells incubated with the Alexa Fuor®555-conjugated secondary antibody diluted 1/500 only were used as negative control.

Cell internalization of NPs. MIA PaCa-2 or NIH/3T3 cells (5 x 10⁴ cells/mL) were cultured in 12-well plates for 24 h to achieve approximately 40% confluence. Cells were then incubated at 37 °C with 5% CO₂ for 6 h with 1 μM of freshly prepared BChol-green-labeled SQdFdC NPs (N0_e), SQdFdC/SQMal_{lipo}/CKAAKN 5:1:0.25 NPs (N1a-P*) or SQdFdC/SQCKAAKN 1:0.01 NPs (N4-P*). Free BChol-green was used as control. After treatment, supernatants were discarded, cells rinsed twice with PBS and harvested for measurement. Cell suspensions were analyzed by flow cytometry (Accuri C6, BD Biosciences, Le Pont de Claix, France) and mean fluorescence intensities were collected between 515 and 545 nm, using the 488-nm line of an argon laser for excitation. For all experiments 10000 cells were measured from each sample.

For low temperature experiments, the cells were incubated in the cold room at 4 °C for 5 h. Results were expressed as the ratio of the mean fluorescence intensity of each sample to the mean fluorescence intensity of non treated cells. This value was then corrected by the fluorescence factor of each NP suspension. All measurements were performed in triplicate or more to determine means and SD.

Cytotoxicity studies. The *in vitro* cytotoxicity of NPs was investigated on MIA PaCA-2 and NIH/3T3 cell lines through the determination of the mitochondrial activity, using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyl tetrazolium bromide test (MTT, Sigma-Aldrich). Briefly, cells were seeded in 100 μL of growth medium (MIA PaCa-2 (1 x 10⁴ cells/mL),

NIH/3T3 (0.75 x 10⁴ cells/mL)) in 96-well plates and preincubated for 24 h. Cells were then treated with 100 µL of a series of concentrations of different nanoparticle suspensions for 72 h. Initial cell density and incubation time were determined to allow cells to remain in exponential growth and to undergo two cell-doubling times during the assay. At the end of the incubation period, 20 µL of a 5 mg/mL MTT solution in phosphate buffered saline were added to each well. After 2 h of incubation, the culture medium was removed and replaced by 200 µL of dimethyl sulfoxide (DMSO), in order to dissolve the formazan crystals. The absorbance of the solubilized dye was measured spectrophotometrically with a microplate reader (LAB System Original Multiscan MS) at 570 nm. The percentage of viable cells for each treatment was calculated from the ratio of the absorbance of the well containing the treated cells *versus* the average absorbance of the control wells (*i.e.*, untreated cells). All experiments were set up in triplicate.

Coculture. NIH/3T3 fibroblasts (7.5 x 10^4 cells) were plated onto microscopic glass slides and incubated for 18 h at 37 °C with 5% CO₂ in complete medium. To visualize and distinguish between NIH/3T3 fibroblasts and MIA PaCa-2 cells in coculture, NIH/3T3 cells were stained with 5 μ M CellTracker green 5-chloromethylfluorescein diacetate (CMFDA, Life Technology, Molecular Probes) according to the manufacturer's protocol. After staining, cells were left to rest for 2 h at 37 °C with 5% CO₂ in cell culture medium, and then unstained MIA PaCa-2 cells (15 x 10^4 cells) were co-incubated. The seeding ratio between MIA PaCa-2 and NIH/3T3 cells was 2:1 due to the faster doubling time of the latter. After 18 h, the coculture was incubated with BChol-red-labeled SQdFdC NPs and BChol-red-labeled SQdFdC/SQCKAAKN NPs diluted in fresh cell culture medium at 10 μ M. At different time points (2, 6, and 24 h) nanoparticles were withdrawn and cells washed with pre-warmed culture medium before imaging. Free BChol-red was used as control.

a Zeiss LSM-510 confocal laser microscope equipped with a 1 mW argon laser and 1 mW helium neon laser, using a Plan-Apochromat 20X or 63X objective lens (NA 1.40, oil immersion). Green fluorescence (CMFDA) was observed with a band-pass 505 and 550 nm emission filter and under a 488 nm laser illumination. Red fluorescence (BChol-red) was observed with a long-pass 560 nm emission filter and under a 543 nm laser illumination. The

Confocal laser scanning microscopy. Observations were made by sequential acquisition with

pinhole diameter was set at 98 μm for CMFDA and 106 μm for BChol-red and an optical section

thickness of 0.4 µm was fixed. Twelve bit numerical images were acquired with Carlo Zeiss

ZEN 2011 software version 7.1.

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SUPPORTING INFORMATION AVAILABLE

Supporting Information

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Synthesis of SQCKAAKN, western blot analysis, immunocytochemistry analysis and stability studies are provided. This information is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

ABBREVIATIONS

dFdC, Gemcitabine; DLS, Dynamic light scattering; DMEM, Dulbecco's modified eagle medium; EDC, N-ethyl-N'-(3-diethylaminopropyl)-carbodiimide; ESI, electron spray ionization; FBS, Fetal bovine serum; FZD, Frizzled; IC50, half maximal inhibitory concentration; ITC, Isothermal titration calorimetry; MTT, 3-(4,5-dimethylthiazol-2-yl)-2,5diphenyltetrazoliumbromide; NP, Nanoparticle; NHS, N-hydroxysuccinimide; NO_a, SQdFdC nanoparticles, acetone as organic solvent; NOe, SQdFdC nanoparticles, ethanol as organic solvent; N1, SQdFdC/SQMal_{lipo} nanoparticles; N1_a, SQdFdC/SQMal_{lipo} nanoparticles, acetone as organic solvent; N1e, SQdFdC/SQMallipo nanoparticles, ethanol as organic solvent; N2, SQdFdC/SQMal_{hvdro} nanoparticles; N3. **SQMal**_{lipo} nanoparticles; N4-P. SQdFdC/SQCKAAKN nanoparticles; PDI, Polydispersity index; SD, standard deviation; SEM, standard error of the mean; SPR, Surface plasmon resonance; SQ, Squalene; SQdFdC, 4-(N)-1,1',2-Trisnorsqualenoylgemcitabine/4-(N)-1,1',2-trisnorsqualenoyl-2',2'-difluoro-2'deoxycytidine; SQdFdC/SQCKAAKN NPs, CKAAKN-functionalized squalenoylgemcitabine nanoparticles; SQCKAAKN, Michael adduct of CKAAKN and 6-(2,5-dioxo-2,5-dihydro-1H-

pyrrol-1-yl)-N'-[(4E,8E,12E,16E)-4,8,13,17,21-pentamethyldocosa-4,8,12,16,20-pentaen-1-ylidene]hexanehydrazide; SQMal $_{hydro}$, 2-[2-(2,5-Dioxo-2,5-dihydro-1H-pyrrol-1-yl)ethoxy]ethyl (4E,8E,12E,16E)-4,8,13,17,21-pentamethyldocosa-4,8,12,16,20-pentaenoate; SQMal $_{hydro}$, 6-(2,5-Dioxo-2,5-dihydro-1H-pyrrol-1-yl)-N'-[(4E,8E,12E,16E)-4,8,13,17,21-pentamethyldocosa-4,8,12,16,20-pentaen-1-ylidene]hexanehydrazide; ZP, Zeta potential.

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