Sucrose Addition Improves Targeted ECO/siBeta3 Nanoparticle Stability

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The ability of siRNAs to regulate oncogenes through RNA interference makes them a promising target in cancer therapy. siRNAs are an especially beneficial treatment tactic for TNBC, a more aggressive breast cancer that currently lacks targeted treatment options. Previous studies have shown that siRNAs can treat tumor cells by regulating $\beta 3$ integrin, a protein involved in metastasis. Previously developed ECO lipid nanoparticle carrier can target $\beta 3$ integrin through successful encapsulation and cytosolic delivery of si $\beta 3$ in tumor cells. Current clinical application of ECO/si $\beta 3$ nanoparticles is tainted by the aggregation of the nanoparticles after lyophilization or freezing storage techniques. Inconsistencies in nanoparticle size result in less effective delivery of si $\beta 3$. To combat aggregation, the impact of sucrose addition on nanoparticle size was evaluated, since sucrose is used as a cryoprotectant in pharmaceuticals. Dynamic light scattering experiments show consistency of nanoparticle size after minus 80 freezing and after lyophilization, implicating sucrose's ability to maintain the stability of ECO/si $\beta 3$ nanoparticles. Furthermore, results show that nanoparticles with addition of sucrose maintained their ability to regulate $\beta 3$ -integrin compared to nanoparticles without sucrose, ensuring that the efficacy of the nanoparticle was sustained. Results suggest that sucrose addition in ECO/si $\beta 3$ nanoparticles is a simple method to significantly improve nanoparticle stability for clinical use. Looking forward, experiments involving different cryoprotectants on the nanoparticle and different targeted ECO/siRNA nanoparticles can be performed to determine the wide-ranging applications of targeted lipid nanoparticle stability.