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Programmable Bispecific Nano-immunoengager That Captures T Cells and Reprograms Tumor Microenvironment

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ABSTRACT: Immune checkpoint blockade (ICB) therapy has revolutionized clinical oncology. However, the efficacy of ICB therapy is limited by the ineffective infiltration of T effector (Teff) cells to tumors and the immunosuppressive tumor microenvironment (TME). Here, we report a programmable tumor cells/Teff cells bispecific nano-immunoengager (NIE) that can circumvent these limitations to improve ICB therapy. The peptidic nanoparticles (NIE-NPs) bind tumor cell surface α3β1 integrin and undergo in situ transformation into nanofibrillar network nanofibers (NIE-NFs). The prolonged retained nanofibrillar network at the TME captures Teff cells via the activatable α4β1 integrin ligand and allows sustained release of resiquimod for immunomodulation. This bispecific NIE eliminates syngeneic 4T1 breast cancer and Lewis lung cancer models in mice, when given together with anti-PD-1 antibody. The in vivo structural transformation-based supramolecular bispecific NIE represents an innovative class of programmable receptor-mediated targeted immunotherapeutics to greatly enhance ICB therapy against cancers.

KEYWORDS: nano-immuno-engager, fibrillar transformation, T cells capture, immune checkpoint blockade (ICB) therapy

Immune checkpoint receptor pathway blockade monoclonal antibodies such as anti-PD-1, anti-PD-L1, and anti-CTLA-4 can reverse T effector (Teff) cell dysfunction and exhaustion, resulting in dramatic tumor shrinkage and sometimes complete remission in some patients, even with late stage metastatic diseases.1,2 However, the response rate varies greatly between tumor types: up to 40% in melanoma, 25% in non-small cell lung cancer, but <10% in most other tumor types. The low response rate of immune checkpoint blockade (ICB) therapy is probably ascribed to defects in Teff cell infiltration at the tumor sites.3 In addition, the tumor microenvironment (TME) can also influence tumor response to ICB therapies.4

Advancement and optimization of nano-immunotherapy lie in the development of innovative approaches to enhance the specificity and controllability of immunotherapeutic interventions and in the targeting of desired cell types. Advanced bionanomaterials or approaches in a more controlled manner could enhance immunotherapeutic potency by increasing the accumulation and prolonging the retention of the immunomodulatory agent and capturing of the immune cells at the TME while sparing the normal tissues and organs, thus reducing off-target adverse effects such as the systemic cytokine storm.5–10 Especially, in situ modulation of nanomaterial in vivo has been demonstrated to improve the performance of bioactive molecules.11–16

In order to overcome ICB resistance, here we report on a programmable tumor cells/Teff cells bispecific nano-immunoengager (NIE) that can capture Teff cells at tumor sites and sustainably release immunoagonist to reprogram the TME. This bispecific NIE, initially in nanoparticle form (NIE-NPs), is self-assembled from two transformable peptide monomers (TPMs, Figure 1a). TPM1, LXY30-KLVFFK(Pa), was comprised of three discrete functional domains: (1) the high-affinity and high-specificity LXY30 cyclic peptide (cdG-Phe(3,5-diF)-G-Hyp-NcR) ligand that targets the α3β1 integrin heterodimeric transmembrane receptor expressed by many epithelial tumors with high metastatic potential, including clinical non-small cell lung cancer;17–19 (2) the KLVFF β-sheet forming peptide domain originated from β-amyloid (Aβ) peptide;11,20–22 and

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(3) the pheophorbide a (Pa) moiety with fluorescence property, serving as a hydrophobic core to induce the formation of micellar nanoparticles. TPM2, proLLP2A-KLVFFK(R848), was also comprised of three discrete functional domains: (1) proLLP2A, the “pro-ligand” version of LLP2A ligand that is a high-affinity and high-specificity peptidomimetic ligand against the activated \( \alpha_3\beta_1 \) integrin (tumor cell membrane), exposure of LLP2A ligands binding \( \alpha_3\beta_1 \) integrin (T cell membrane), and release of TLRs 7/8 agonist (R848) from NIE-NFs through esterase hydrolysis. (b) Steps II–V show the processes for how a programmable bispecific NIE synergizes ICB therapy in tumor tissue: (I) NIE-NPs bind tumor cells and in situ transforms them into nanofibrils (NIE-NFs) on the surface of tumor cells. (II) NIE-NFs expose LLP2A and release R848 for (III) capturing T cells to tumors cells and (IV) re-educating TAMs from M2 to M1 phenotype. (V) Meanwhile, anti-PD-1 antibody greatly activates the NIE homed cytotoxic T cells for ICB therapy. (TPMs, transformable peptide monomers; NIE-NPs, nano-immuno-engager nanoparticles; NIE-NFs, nano-immuno-engager nanofibrils; R848, resiquimod, a TLRs 7/8 agonist; M1-TAM, M1-like tumor-associated macrophage; M2-TAM, M2-like tumor-associated macrophage.)
tumor cells, thus maintaining a prolonged retention (at least 7 days). With the abundant esterase in the TME and on the tumor cells, proLLP2A would quickly be converted to LLP2A against activated α4β1 integrin. LLP2A displayed on the fibrils would capture T\textsubscript{eff} cells (e.g., CD8\textsuperscript{+} T cells) and facilitate their long-term retention at the TME and adjacent to the tumor cells. The capturing and retention of large numbers of T\textsubscript{eff} cells would greatly enhance ICB therapy.

Besides, the sustained release of R848 from the nanofibrillar network would improve the immunosuppressive tumor microenvironment, e.g., activate antigen-presenting cells, promote immune cells to produce antitumor response factors, and re-educate the phenotype of the macrophage from M2 to M1. This in vivo structural transformation-based supramolecular bispecific NIE represents an innovative class of programmable receptor-mediated targeted immunotherapeutics against cancers through capturing of T cells and enhancement of the antitumor immune state at the TME (Scheme 1).

TPM1 and TPM2 were synthesized and characterized (Figures 1a and S1). As the proportion of water in the mixed solvent (water and DMSO) of the TPM1 and TPM2 mixture solution was increased, there was a gradual decrease in the fluorescence peak at 675 nm due to the aggregation-caused quenching properties of Pa dye (Figure 1b), reflecting the gradual formation of NIE-NPs via self-assembly. Concomitantly, there was a modest decrease in the absorption peak at both 405 and 680 nm (Figure S2a). TPM1 and TPM2 each alone were able to self-assemble to form nanoparticles (NPs\textsubscript{TPM1} and NPs\textsubscript{TPM2}) at 18 and 55 nm. NIE-NPs, assembled from a 1:1 mix of TPM1 and TPM2 into nanoparticles at around 28 nm, which
fell between the sizes of NPs\textsubscript{TPM1} and NPs\textsubscript{TPM2} (Figure S2b).

The critical aggregation concentration (CAC) of the NIE-NPs was determined to be 8 μM (Figure S2c). We have also demonstrated that NIE-NPs could maintain good serum stability and proteolytic stability over 7 days at 37 °C (Figure S2d).

To verify the receptor-mediated fibrillar transformable process of NIE-NPs in \textit{vitro}, soluble α\textsubscript{4}β\textsubscript{1} integrin protein

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Figure 2. \textit{In vitro} bispecific NIE binds both 4T1 breast cancer cells and CD8 T cells and re-educates tumor-associated macrophages. (a) Cellular fluorescence distribution images of interaction of NIE-NPs and CNIE-NPs for 6 h with 4T1 tumor cells to show NIE-NPs around cells and CNIE-NPs inside cells. Scale bar is 10 μm. (b) Cellular fluorescence signal retention images of 4T1 tumor cells after exposure to NIE-NPs and CNIE-NPs for 6 h followed by incubation with fresh medium without nanoparticles for 18 h to show long retention of NIE and short retention of CNIE. Scale bar is 10 μm. (c) Representative SEM images of untreated 4T1 tumor and 4T1 tumor cells treated with NIE-NPs for 6 h. The concentration of NIE-NPs was 50 μM. (d and e) (d) Cellular fluorescence distribution images and (e) representative SEM images of murine CD8 T cells (isolated from mouse spleen, CellTracker green CMFDA dye labeled, green fluorescence) after incubation with esterase-pretreated NIE-NPs to show NIE around cells. α\textsubscript{4}β\textsubscript{1}, integrins on the surface of murine CD8 T cells were preactivated by Mn\textsuperscript{2+} (1 mM). Scale bar is 10 μm. (f) Experimental scheme and cellular fluorescence distribution images of NIE-NPs (fluorescent red), after interaction with 4T1 tumor and murine CD8 T cells (fluorescent green, α\textsubscript{4}β\textsubscript{1}). It shows that nanofibrillar networks (bispecific NIE) cover 4T1 tumor cells, which in turn capture CD8 T cells. More incubation time, more bound CD8 T cells. Scale bar is 10 μm. (g) Representative SEM images of 4T1 tumor and CD8 T cells after treatment with NIE-NPs (see part f). (h) Representative images of M2-like murine macrophages and subsequent re-education by NIE-NFs, NIE-NFs plus esterase, or R848 at different time points. Scale bar is 20 μm. Statistical significance was calculated using one-way ANOVA followed by Tukey’s post hoc analysis: *P < 0.05, **P < 0.01, ***P < 0.001.
LLP2A resulted in complete loss of its binding to Esterification of the carboxyl side chain of LLP2A to form affinity and selectivity of LXY30 to integrin, and the reverse was true for LLP2A ([Figure S6a and c]).

After 24 h of incubation at room temperature, a fibrillar network (NIE-NFs, width diameter about 8 nm) with a broad size distribution was clearly detected ([Figure 1c and f]). No transformation was observed in the preparation of NIE-NPs without the addition of $\alpha_\beta_1$ integrin protein, even after 24 h ([Figure S2c]). Addition of $\alpha_\beta_1$ integrin protein triggered a gradual decrease in the fluorescence intensity of Pa over time, reflecting an increase in fluorescence quenching in NFs1, indicating that the packing of Pa in NIE-NFs was denser ([Figure 1d]). The CAC of NIE-NFs was confirmed to be lower than that of NIE-NPs ($5 \mu M$ vs $8 \mu M$, [Figure S2f]).

In addition, a fluorescence peak reversal (from 680 to 725 nm) was observed, indicating that the disordered arrangement of Pa in the core of NIE-NPs was transformed into $f$-aggregate form in NIE-NFs. We also investigated the responsiveness of proLLP2A displayed on the NIE-NPs surface to soluble $\alpha_\beta_1$ integrin protein in the presence and absence of esterase ([Figure 1e and f and Figure S3]). Successive addition of esterase, followed by soluble $\alpha_\beta_1$ integrin protein, was able to elicit conversion of NIE-NPs to fibrillar network after 24 h of incubation. This expected result confirmed that esterase was able to convert pro-lllp2a to ligand LLP2A, which in turn was able to trigger receptor-mediated transformation of NIE-NPs to NIE-NFs. We monitored the conversion of proLLP2A ligand to LLP2A ligand by HPLC. The majority of LLP2A ligand by HPLC. The majority of proLLP2A was found to be converted to LLP2A after incubation with esterase for 8 h at pH 7.4 and 37 °C ([Figure S4]).

Circular dichroism (CD) spectroscopic analysis of the transformation process of NIE-NPs showed a gradual progression of a negative signal at 216 nm and a positive signal at 195 nm upon incubation with $\alpha_\beta_1$ integrin protein or combination esterase/$\alpha_\beta_1$ integrin protein, indicative of $\beta$-sheet formation ([Figure 1g]). The in vitro release behavior of R848 from NIE-NFs was studied at pH 6.5 with addition of esterase to simulate TME conditions. About 45% of R848 was released in the first 24 h, after which the release rate gradually slowed down and about 86% cumulative release was observed by 168 h, indicating that prolonged and sustained release of R848 could occur at the TME ([Figure 1h]). To demonstrate the unique transformable property of NIE-NPs, we designed a related negative control nano-immunoengager nanoparticle (CNIE-NP) formed by assembly of two negative control TPMs without a $\beta$-sheet forming KLVFF peptide sequence, at a ratio of 1:1 (CTPM3:LXY30-KAAGGGK(Pa) and CTPM4:proLLP2A-KAAGGGK(R848), [Figures S1 and S5]).

We have previously reported the binding affinity of LLP2A against $\alpha_\beta_1$ integrin to be at the sub-nanomolar level and that of LXY30 against $\alpha_\beta_1$ integrin to be at the mid-nanomolar level. LXY30 bound strongly to $\alpha_\beta_1$ integrin but not $\alpha_\beta_1$ integrin, and the reverse was true for LLP2A ([Figure S6a and c]). Esterification of the carboxyl side chain of LLP2A to form proLLP2A resulted in complete loss of its binding to $\alpha_\beta_1$ integrin. Treatment of proLLP2A with esterase was able to completely restore the binding affinity of LLP2A toward $\alpha_\beta_1$ integrin ([Figure S6b]). We have also confirmed the high binding affinity and selectivity of LXY30 to $\alpha_\beta_1$ integrin on the surface of 4T1 cells via flow cytometry ([Figure S7]) and also found that NIE-NPs were slightly cytotoxic against 4T1 tumor cells, with 84.5% cell viability at 50 $\mu M$ ([Figure S8]).

We investigated the distribution of nanoparticles by tracking the red fluorescent signal emitted by Pa using confocal laser scanning microscopy (CLSM). Six hours after incubation of 4T1 tumor cells with NIE-NPs, a strong red fluorescence signal was observed on the cell surface but not inside the cells ([Figure 2a]). The red fluorescence signal of transformed NIE-NPs was found to merge with plasma membrane on the cell surface but not with intracellular actin ([Figures S9 and S10]). In contrast, the fluorescent signal of Pa in the CNIE-NPs-treated group was found to be concentrated primarily in the cytoplasm of the cells. To study the retention and stability of the formed nanofibrillar network on the surface of tumor cells, unbound nanoparticles were washed off after 6 h of incubation and fresh medium without nanoparticles was added to incubate cells for another 18 h. As expected, NIE-NPs-treated cells still retained strong red fluorescence signals on the cell surface at 24 h, displaying the capacity of prolonged retention of the bispecific NIE ([Figure 2b]). In sharp contrast, only a weak fluorescence signal was observed inside the cells treated with CNIE-NPs after 24 h. SEM images confirmed the presence of a nanofibrillar network on the surface of 4T1 tumor cells ([Figure 2c]). We also investigated the effect of esterase on the interactions between NIE-NPs and CD8 T cell surface $\alpha_\beta_1$ integrin (preactivated by Mn$^{2+}$), after converting pro-LLP2A to LLP2A ([Figure 2d]). SEM confirmed the presence of a fibrillar network on the surface of esterase pretreated NIE-NPs-treated CD8 T cells ([Figure 2e]).

To simulate the processes of initial fibrillar transformation of NIE-NPs on the surface of 4T1 tumor cells followed by T cell binding, we incubated 4T1 tumor cells with NIE-NPs for 6 h; unbound NIE-NPs were then washed off, followed by addition of fresh medium containing esterase but without NIE-NPs. After 1 h of incubation, murine CD8 T cells with $\alpha_\beta_1$ integrins activated by Mn$^{2+}$ were added and incubated with 4T1 tumor cells for 2 or 4 h. After that, unbound CD8 T cells were gently removed prior to CLSM imaging ([Figure 2f]). As expected, a fibrillar structure layer with red fluorescence was detected surrounding the surface of the 4T1 tumor cells, and the CD8 T cells (green fluorescence) were found to interact with the red fluorescent fibrillar network and in close proximity to the 4T1 tumor cells, after 2 h of incubation. As the incubation time was increased to 4 h, many more T cells were found around the 4T1 tumor cells, which was consistent with our notion that the nanofibrillar-based bispecific NIE would facilitate the capturing and retention of immune cells. SEM imaging provided critical evidence that the bispecific NIE had played a significant role in the direct physical contact between 4T1 tumor cells and T cells through the nanofibrillar network ([Figure 2g]). The conversion of tumor-associated macrophages (TAMs) from an immunosuppressive M2-polarized phenotype to an anti-tumor M1-polarized phenotype is one of the major immunotherapeutic strategies for reprogramming the immunosuppressive TME.

Addition of esterase to the culture medium followed by 24 h of incubation resulted in morphological change of the M2-state toward the round and flattened M1-state, a decrease in Arg1 and Mrc1, and an increase in IL-12 and Nos2 expression ([Figures 2h and S11]).

NIE-NPs were found to be nontoxic: blood counts, platelets, creatinine, and liver function tests obtained from normal Balb/c mice treated with eight consecutive q.o.d. intravenous (i.v.) doses of NIE-NPs were within normal limits ([Figures S12 and S13]). In vivo blood pharmacokinetics (PK) studies in rats showed that NIE-NPs possessed a long circulation time (T-half ($\alpha$): 2.866 h and T-half ($\beta$): 23.186 h), indicating their stability during circulation ([Figure S14]). For biodistribution studies, NIE-NPs were tail vein injected once into Balb/c mice bearing syngeneic orthotopic 4T1 breast cancer; 10, 24, 48, 72, 120, and
Figure 3. In vivo evaluation of NIE targeting tumor cells, and in situ nanofibrillar transformation to capture T_{eff} cell and facilitate the long-term retention and activate immunity. (a and b) Time-dependent ex vivo fluorescence (FL) images (a) and quantitative analysis (b) of tumor tissues and major organs (heart (H), liver (Li), spleen (Sp), lung (Lu), kidney (K), intestine (I), muscle (M), and skin (Sk)) collected at 10, 24, 48, 72, 120, and 168 h post-injection of NIE-NPs. Data are presented as mean ± s.d., n = 3 independent experiments. (c) Time-dependent ex vivo fluorescence images of tumor tissues collected at 10, 24, 48, 72, 120, and 168 h post-injection of CNIE-NPs. Data are presented as mean ± s.d., n = 3 independent experiments. (d) Representative TEM images of distribution in tumor tissue and in situ fibrillar transformation of NIE-NPs, CNIE-NPs, and the untreated control group at 72 h post-injection. “N” depicts nucleus. (f) Fluorescence distribution images of NIE-NPs in the tumor tissue region and normal skin tissue at 72 h post-injection (red, PA of NIE-NPs; blue, DAPI; scale bars, 50 μm). (g) R484 distribution retention in tumor tissues at different time points post-injection of NIE-NPs and CNIE-NPs. Injection dose of R848: 0.94 mg kg⁻¹; data were mean ± s.d., n = 3 for each time point. (h) Expression of CXCL10 chemokine within the tumor tissues after 5 days of NIE-NPs, CNIE-NPs, and saline treatment (n = 3; data were mean ± s.d.; single injection). (i) Representative flow cytometric analysis images and corresponding quantification of CD45⁺CD3⁺, CD8⁺/CD4⁺, CD4⁺Foxp3⁺, CD8⁺CD49⁺, and CD8⁺CD29⁺ T cells within the 4T1 tumors excised from mice treated with NIE-NPs, CNIE-NPs, or the saline control. (j) Immunohistochemistry (IHC) staining of T cells (CD8⁺, CD4⁺, and Foxp3⁺) and macrophage markers (CD68 and CD163). Scale bar is 100 μm. (k) The expression levels (qPCR assay) of IFN-γ, TGF-β, IL12, IL10, Nos2, and Arg-1 in 4T1 tumors excised from mice 15 days after treatment with NIE-NPs or CNIE-NPs (n = 3; data were mean ± s.d.). Statistical significance was calculated using a two-sided unpaired t test compared to the NIE-NPs group: *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001.
168 h later, tumor and main organs were excised for ex vivo fluorescent imaging (Figure 3a and b). A significant fluorescent signal of Pa was found to persist in tumor tissue for over 168 h, while the fluorescent signal in normal organs began to decline after 10 h and was almost undetectable in the main organs at 72 h. In sharp contrast, the fluorescent signal of Pa at tumor tissue treated by CNIE-NPs was found to gradually decline over time after peaking at 24 h (Figures 3c and S15). The prolonged retention of the fluorescent signal in NIE-NPs-treated mice could be attributed to in situ receptor-mediated transformation of NIE-NPs into NIE-NFs at the TME (Figure 3d).

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**Figure 4.** Intrinsic antitumor immune efficacy of bispecific NIE in Balb/c mice bearing 4T1 breast tumor. (a) Expression of CXCL10 and CXCL9 chemokine on day 7 within the excised tumor tissues of mice after different treatments (13 mg/kg each dose, every other day; n = 3; data were mean ± s.d.). (b) Representative flow cytometric analysis images and relative quantification of CD8+/CD4+ and CD4+ Foxp3+ T cells within the 4T1 tumors excised from mice after different treatments at day 15. All treatment regimens were tail vein injected consecutively three times q.o.d. (13 mg/kg each dose, every other day, total three times; n = 5; data were mean ± s.d.). In parts a and b, (1) saline; (2) LXY30-KLVFFK(Pa)/(EK)3-KLVFFK(R848) (fibrillar-transformation but absence of LLP2A ligand); (3) LXY30-KLVFFK(Pa)/proLLP2A-KLVFFK(Pa) (fibrillar-transformation but absence of R848); and (4) LXY30-KLVFFK(Pa)/proLLP2A-KLVFFK(R848) (complete NIE-NPs). (c) Experimental design: orthotopic tumor inoculation and treatment protocol; regimen 6 is NIE-NPs with all four critical components: (1) saline; (2) (EK)3-KLVFFK(Pa)/(EK)3-KLVFFK(R848) (“R848 only” in micellar formulation); (3) proLLP2A-KLVFFK(R848) (single monomer); (4) LXY30-KAAGGK(Pa)/proLLP2A-KAAGGK(R848) (untransformable negative control CNIE-NPs); (5) LXY30-KLVFFK(Pa)/proLLP2A-KLVFFK(Pa) (fibrillar-transformation but absence of R848); and (6) LXY30-KLVFFK(Pa)/proLLP2A-KLVFFK(R848). (d and e) Observation of the tumor inhibitory effect (d) and weight change (e) of mice bearing orthotopic 4T1 tumor over 21 d after initiation of treatment (n = 8 per group). Data are presented as mean ± s.d. (f) Cumulative survival of different treatment groups of mice bearing 4T1 breast tumors. Data are presented as mean ± s.d. Statistical significance was calculated using one-way ANOVA followed by Tukey’s post hoc analysis: *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001.
the tumors and overlying skin revealed an intense fluorescent signal in the tumor region but negligible signal in the normal skin (Figure 3f). The tumor tissue distribution of R848 over time was also determined with high-pressure liquid chromatography–mass spectroscopy (HPLC-MS, Figure 3g). We found that the R848 uptake by the tumor was quite high at 24 h (3.62 μg per g tissue) and that about one-third of the R848 was found to be retained at the tumor site (1.14 μg per g tissue) even at 7 days.

We evaluated whether the bispecific NIE (single injection) could capture T effector cells and facilitate their long-term retention at the tumor sites and re-educate TAMs in vivo. First, NIE-NPs were found to significantly stimulate the production of chemokine CXCL10 at the tumor site (Figure 3h), which was known to facilitate recruitment of T effector cells.32,33 This should be attributed to sustained release of R848, which is known to induce the production of chemokines.34−36 We confirmed that the proportion of CD45+CD3+ and CD45+CD3+CD8+ T cells in the NIE-NPs-treated tumor tissue was substantially higher than those from mice treated with endocytic CNIE-NPs or saline (Figures 3i and S17). Second, the relative abundance of CD45+Foxp3+ T regs at the tumor site was substantially lower in mice that received the NIE-NPs treatment than that in mice treated with CNIE-NPs, i.e., 4.97% vs 13.0%. We also confirmed the higher expression (CD49β) and activation (CD29α) of αβ integrin in tumor-infiltrating CD8+ T cells after NIE-NPs treatment, which were conducive to capturing of CD8+ T cells by LLP2A displayed at the TME. Third, IHC staining of tumor sections demonstrated an increase in M1-polarized macrophage marker CD68 and a decrease in M2-polarized macrophage marker CD163 in the NIE-NPs-treated group (Figure 3j).
Fourth, the high expression levels of IFN-γ, IL-12, and Nos2 and low expression levels of TGF-β, IL-10, and Arg-1 in the tumor tissue confirmed that a strong tumor-specific immune response had been elicited (Figure 3k).

We evaluated the roles of T cell capturing ligand (LLP2A) and TLR7/8 immunoagonist (R848), the two components of NIE-NPs, in inducing chemokine production and T cell capturing at the TME in vivo. As shown in Figure 4a and b, R848 (group 2) was more effective in inducing chemokine secretion than the LLP2A ligand (group 3), but the population of CD8 T cells in group 3 was slightly higher than that in group 2. This suggested that LLP2A ligands should have played a significant role in capturing and retention of CD8 T cells at the TME, more so than R848. However, LLP2A or R848, each alone, was found to be insufficient to induce a robust antitumor immune response. It is clear that group 4 mice treated with both LLP2A and R848 were able to mount a robust antitumor immune response, resulting in a higher level of chemokine expression, more CD8 T cells, and fewer T reg cells.

A therapeutic efficacy study of a programmable bispecific NIE was performed in syngeneic orthotopic 4T1 breast cancer-bearing mice (Figure 4c–f). Regimen 6 is the complete NIE-NPs, containing all four critical components: LYY30, proLLP2A, R848, and KLH, whereas regimen 2, 3, 4, or 5 each lacks some components of NIE-NPs. Regimen 6 (NIE-NPs) was found to be the most efficacious with significant tumor growth suppression and prolonged survival, indicating the importance of the combination of the homing strategy of T cells and sustained release of TLR7/8 agonist. Regimen 5 (fibrillar transformation but no R848) demonstrated significant tumor suppression compared to groups 2, 3, and 4. None of the mice in this therapeutic study showed any symptoms of dehydration or significant body weight loss during the entire treatment period. Regimen 6 showed a great increase in the frequency of CD3+ and CD8+ T cells within the tumors and was the most efficacious in restoring the immunoreactive state of the TME (Figure S18).

Tumor sections (H&E) obtained from mice treated with NIE-NPs revealed a marked decrease in K-67 expression, an increase in CD8+ T cells, and a decrease in Foxp3 (T reg cells), compared with other control groups. There was an increase in CD68 and a decrease in CD163, and the percentage of M1-phenotype macrophage in the total macrophages was 60%, which was much higher than that of the other control groups (Figures S19 and S20). To better clarify the contribution of the recruitment of CD8 T cells to tumors in NIE-NPs’ immunotherapy, we performed a tumor protection experiment with depleting the T eff cells to the tumor sites are probably one of the main reasons why many patients remain refractory to such treatment. We believe the receptor-mediated bispecific NIE reported here can provide a relatively simple solution to this challenge. By incorporating pro-ligands LLP2A and R848 to the in vivo transformable nanofibrillar networks, we have already demonstrated in syngeneic 4T1 breast cancer and Lewis lung cancer models that this nontoxic treatment can (1) significantly enhance the capturing of T eff cells (CD8+ T cells) in the tumor region, (2) promote retention of T cells at close proximity to the tumor cells, and (3) provide sustained release of R848 at the TME, resulting in the significant enhancement of the efficacy of the anti-PD1 antibody-based ICB. Since the programmable bispecific NIE is modular, we have the options of combinatorially incorporating various ligands, pro-ligands, or immunomodulators to generate a series of nanoengagers that may be applied for capturing other beneficial antitumor immune cells.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.nanolett.2c00582.

Additional details on experimental materials and methods. Figures showing chemical characterization of peptide monomers, NIE-NPs and NIE-NFs, interaction between NPs and living cells, NPs’ treatment, and immunity index. (PDF)

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Notes

The authors declare the following competing financial interest(s): K.S.L. and L.Z. are the coinventors of a patent on fibrillar transformable nanoparticles (PCT/US2020/046495). K.S.L. is the founding scientist of LammoTherapeutics Inc., which plans to develop the nanotherapeutics described in the manuscript. The remaining authors declare no competing interests.

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