Engineered PLGA microparticles for long-term, pulsatile release of STING agonist for cancer immunotherapy

Xueguang Lu1†, Lei Miao1†, Wenting Gao1, Ziqi Chen1, Kevin J. McHugh1,2, Yehui Sun1, Zachary Tochka1, Stephanie Tomasic1, Kaitlyn Sadler1,3, Alain Hyacinthe1, Yuxuan Huang1, Tyler Graf1, Quanyin Hu1, Morteza Sarmadi1,4, Robert Langer1,5,6,7†, Daniel G. Anderson1,5,6,7†, Ana Jaklenec1†

Activation of the stimulator of interferon gene (STING) pathway within the tumor microenvironment has been shown to generate a strong antitumor response. Although local administration of STING agonists has promise for cancer immunotherapy, the dosing regimen needed to achieve efficacy requires frequent intratumoral injections over months. Frequent dosing for cancer treatment is associated with poor patient adherence, with as high as 48% of patients failing to comply. Multiple intratumoral injections also disrupt the tumor microenvironment and vascular networks and therefore increase the risk of metastasis. Here, we developed microfabricated poly(lactic-co-glycolic acid (PLGA) particles that remain at the site of injection and release encapsulated STING agonist as a programmable sequence of pulses at predetermined time points that mimic multiple injections over days to weeks. A single intratumoral injection of STING agonist–loaded microparticles triggered potent local and systemic antitumor immune responses, inhibited tumor growth, and prolonged survival as effectively as multiple soluble doses, but with reduced metastasis in several mouse tumor models. STING agonist–loaded microparticles improved the response to immune checkpoint blockade therapy and substantially decreased the tumor recurrence rate from 100 to 25% in mouse models of melanoma when administered during surgical resection. In addition, we demonstrated the therapeutic efficacy of STING microparticles on an orthotopic pancreatic cancer model in mice that does not allow multiple intratumoral injections. These findings could directly benefit current STING agonist therapy by decreasing the number of injections, reducing risk of metastasis, and expanding its applicability to hard-to-reach cancers.

INTRODUCTION

The advent of immune checkpoint blockade (ICB) has had a profound impact on cancer treatment, with several drugs receiving approval from the U.S. Food and Drug Administration (FDA) (1). Despite great promise, the clinical benefits of ICB remain limited by a low response rate (2). Clinical studies have shown that patients who respond to ICB have higher amounts of tumor-infiltrating lymphocytes (TILs) and display a signature of type I interferon (IFN)–producing genes indicative of innate immune system activation (3–5). Therefore, strategies to improve TIL infiltration and innate immune system activation have been proposed as combination therapies to further improve the response rate of ICB. Among many innate immune pathways that are initiated through Toll-like receptors (TLRs), mitochondrial antiviral-signaling protein (MAVS), or P2X purinergic receptor 7, activation of stimulator of interferon genes (STING) shows great promise for increasing TILs and improving the antitumor efficacy of ICB (6, 7). STING pathway activation is initiated through recognition of cytoplasmic DNA. Cyclic guanosine monophosphate–adenosine monophosphate (cGAMP) synthase senses cytoplasmic DNA and produces the second messenger cGAMP, which then binds to STING to trigger a signaling cascade through tank-binding kinase 1 (TBK1)/interferon regulatory factor 3 (IRF3) for production of type I IFNs and other cytokines (8, 9). Substantial evidence has shown that intratumoral injections of STING agonist stimulate potent antitumor immunity in clinically relevant tumor models (7, 10–13). As a result, phase I clinical trials (NCT02675439, NCT03172936, and NCT03010176) using STING agonist alone or in combination with ICB are currently under way to treat patients with advanced solid tumors and lymphoma.

The dosing regimen of STING agonist in current clinical trials consists of multiple intratumoral injections administered repeatedly (for example, three injections over a 28-day period or one injection every week for 9 weeks per treatment cycle) for as long as 2 years to achieve therapeutic efficacy (14, 15). Such high dosing frequency over a long period of time can cause chronic injection pain, increase the risk of infection, and ultimately result in poor adherence, especially when every dose requires a health care visit (16–18). The adherence rates to cancer treatment are as low as ~52%, with similar rates (~50%) reported for patients with other chronic diseases (19, 20). Poor adherence can result in failed treatment and constitutes a financial burden of about $100 billion each year in the United States alone (21). In addition, multiple intratumoral injections limit the scope of STING agonist–based therapies to readily accessible tumor types and introduce the risk of disrupting the tumor microenvironment (TME) and vascular network, potentially promoting cancer cell extravasation and metastases (22–24). Therefore, a delivery system that mimics current clinical dosage regimens within a single injection is an attractive solution to...
improve patient adherence, decrease risk of metastasis and therapeutic cost, and expand the scope of current STING agonist–based therapies.

Here, we developed such a multidose drug delivery platform through engineering poly(lactic-co-glycolic acid) (PLGA), an FDA-approved and commercially available polymer, into cubic microparticles (Fig. 1A). Unlike commonly used local drug delivery materials, such as hydrogels or microparticles produced using a double emulsion solvent evaporation technique, which exhibit sustained drug release kinetics (25–28), these microfabricated PLGA particles (PLGA-MPs) released individual doses of encapsulated STING agonist in pulses for up to several months with essentially no leakage. We demonstrated that a single injection containing multiple populations of STING agonist–loaded PLGA-MPs inhibited tumor growth and improved survival as effectively as multiple injections of soluble STING agonist in several mouse tumor models. The ability to combine multiple doses into single-injection PLGA-MPs also decreased metastasis and expanded the potential applications of current STING agonist–based therapy to hard-to-reach tumors.

RESULTS

Fabrication of PLGA microparticles with different release kinetics

We used soft lithography techniques to fabricate arrays of cubic PLGA microparticles with a fully enclosed cavity for drug loading (Fig. 1B). Briefly, PLGA was heated and pressed into a poly(dimethylsiloxane) (PDMS) mold to form microparticle bases with internal cavities (200 μm by 200 μm by 100 μm, length by width by height), which corresponds to a volumetric capacity of 4 nl. We then filled aqueous drug or model drug into these bases using a piezoelectric dispenser that can dispense volumes on the order of 100 pl. The water component of the drug solutions evaporated rapidly because of the small volume, which provided space for filling additional cargo. Multiple filling and drying cycles were used to achieve maximum cargo loading. Filled microparticles were then aligned with PLGA caps embedded in a PDMS mold and sealed by heating above the glass transition temperature (about 50°C) of PLGA. Sealed microparticles have external dimensions of 400 μm by 400 μm by 300 μm (length by width by height) and wall thickness of 100 μm in each dimension. The loading capacity of each microparticle is 8.4% by volume. Scanning electron microscopy, high-resolution x-ray computed tomography (CT), and optical images showed that microparticles can be made in large arrays (over 300 per array) with high fidelity (Fig. 1, C to G, and fig. S1).

To study the release kinetics, we filled a fluorescently labeled macromolecule, Alexa Fluor 647–labeled dextran (AF647-dextran), into PLGA microparticles with different polymer properties (table S1). These microparticles were sealed with corresponding caps and incubated in phosphate-buffered saline (PBS; pH 6.84) at 37°C to mimic the acidic TME. PLGA microparticles released AF647-dextran in pulses at about 1 ± 0, 4 ± 0, 8 ± 0, 11 ± 1, 15 ± 1, 18 ± 1, and 97 ± 2 days without detectable leakage before release (fig. S2). To mimic the dosing regimen of four consecutive doses with 3 to 4 days between each dose that has been shown to be effective at inhibiting tumor growth in animal models (7), microparticles that release AF647-dextran at day 4 (PLGA-1), day 8 (PLGA-2), and day 11 (PLGA-3) were selected for further study (Fig. 2A). To validate the release kinetics in vivo, we subcutaneously injected AF647-dextran–loaded PLGA-1, PLGA-2, and PLGA-3 into hairless mice. The release of AF647-dextran was monitored by in vivo fluorescence imaging [in vivo imaging system (IVIS)].

Fig. 1. Design and fabrication of PLGA-MPs. (A) Schematics of single-injection drug delivery platform for cancer immunotherapy. Different PLGA microparticles reside in the tumor after a single intratumoral injection, release encapsulated STING agonist in pulses at discrete time points, and promote infiltration of TILs. (B) Schematics of the fabrication process of PLGA-MPs, which are prepared by filling cargo of interest into particle bases and then sealing the bases with corresponding particle caps by briefly applying heat. (C to E) Representative scanning electron microscopy images of EP bases (C) and sealed array of particles (D) or an individual particle (E). Scale bars, 500 μm (C and D) and 100 μm (E). (F) Representative high-resolution x-ray CT image of a sealed particle encapsulating 3′3′-cGAMP. Red color represents dried 3′3′-cGAMP. Scale bar, 100 μm. (G) Representative optical image of an array of sealed particles encapsulating Alexa Fluor 647–labeled dextran. Scale bar, 1 mm.

SCIENCE TRANSLATIONAL MEDICINE | RESEARCH ARTICLE

HF et al., Sci. Transl. Med. Lu defined environment (Fig. 2E). To further demonstrate that PLGA- and aggregated at the injection site due to the low mobility in a con-

microparticles in tumors, we doped 5% phosphotungstic acid (PTA) showed consistent release kinetics in both tumors and the subcuta-

dmatic environment (Fig. 2, C and D). To study the distribution of neous environment (Fig. 2, C and D). To study the distribution of

showed an over 100-fold increase in fluorescence intensity compared with encapsulated AF647-dextran in particles due to the self-quenching effects of the fluorophore when dry or at an extremely high local concentration (fig. S3). Figure 2B and fig. S4 showed that micro-

We then evaluated the influence of different TEMs on release kinetics. AF647-dextran–loaded PLGA-2 was intratumorally injected into mice bearing B16F10 melanoma or 4T1 breast tumors. Release kinetics was then monitored by daily IVIS imaging. PLGA-2 showed consistent release kinetics in both tumors and the subcuta-

nous melanoma model (n = 4). Data are shown as means ± SEM. Scale bars represent radiant efficiency. (E) MicroCT image of the B16F10 tumor, which was isolated from mice 1 hour after injection of 5% phosphotungstic acid–doped PLGA-1. Scale bar, 2 mm. (F) Treatment and sampling schedule of B16F10 tumor–bearing mice after intratumoral injection of AF647-loaded PLGA-1. (G) Cumulative in vivo release of AF647 from PLGA-1 in B16F10 tumors (n = 4). (H) AF647 concentration in serum after intratumoral injection of AF647-loaded PLGA-1 (n = 4). Data are shown as means ± SEM.

We measured the amount of unreleased AF647 in tumors every day and back calculated the amount of released AF647 (Fig. 2F). Figure 2G shows that PLGA-1 completely released AF647 from 3 to 6 days in tumors. Some released AF647 also diffused into the bloodstream, as demonstrated by increased AF647 concentration in serum from day 3 to day 6 (Fig. 2H). These data demonstrated that PLGA-MPs released all encapsulated cargos at anticipated time points in tumors.

We then fabricated STING agonist–loaded PLGA-1, PLGA-2, and PLGA-3 with a drug loading of 2 µg per particle. 3′3′-cGAMP, a linkage isomer of naturally produced 2′3′-cGAMP, is used here because of its enhanced stability against ecto-nucleotide pyrophosphatase/phosphodiesterase 1 (ENPP1), which primarily hydrolyzes cGAMP (29). 3′3′-cGAMP was released from PLGA-1, PLGA-2, and PLGA-3 in pulses at nearly the same times as the fluorescent molecules in vitro (Fig. 3A). The stability of encapsulated cGAMP in physiological conditions is critical for the retention of bioactivity upon release. To study the stability of 3′3′-cGAMP in particles, we incu-

bated cGAMP-loaded PLGA-2 particles in PBS at 37°C and analyzed the structure integrity of cGAMP in the supernatant over time by liquid chromatography–mass spectrometry (LC-MS). Released cGAMP showed identical elution time and molecular mass to those of standard 3′3′-cGAMP (Fig. 3B and fig. S7). The bioactivity of released cGAMP was also tested by an IRF reporter cell line [RAW-Lucia interferon-stimulated gene (ISG) cells]. Released cGAMP from PLGA-2

<table>
<thead>
<tr>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Days 0 2 4 6 8 10 12 14</td>
<td>Days 0 2 4 6 8 10 12 14</td>
<td>Days 0 2 4 6 8 10 12 14</td>
</tr>
<tr>
<td>% Released</td>
<td>% Released</td>
<td>% Released</td>
</tr>
<tr>
<td>PLGA-1</td>
<td>PLGA-2</td>
<td>PLGA-3</td>
</tr>
</tbody>
</table>

Fig. 2. Release kinetics of PLGA-MPs. (A and B) Cumulative in vitro (A) and in vivo (B) release kinetics of AF647-dextran from PLGA-1, PLGA-2, and PLGA-3. PLGA-MPs were administered subcutaneously (n = 6 to 8). Data are shown as means ± SEM. (C and D) Cumulative in vivo release kinetics (C) and representative fluorescence images (D) of AF647-dextran–loaded PLGA-2 that were administered subcutaneously in 5K/H1E hairless mice (n = 8) or intratumorally (IT) in the B16F10 melanoma model (n = 4) and 4T1 breast cancer model (n = 4). Data are shown as means ± SEM. Scale bars represent radiant efficiency. (E) MicroCT image of the B16F10 tumor, which was isolated from mice 1 hour after injection of 5% phosphotungstic acid–doped PLGA-1. Scale bar, 2 mm. (F) Treatment and sampling schedule of B16F10 tumor–bearing mice after intratumoral injection of AF647-loaded PLGA-1. (G) Cumulative in vivo release of AF647 from PLGA-1 in B16F10 tumors (n = 4). (H) AF647 concentration in serum after intratumoral injection of AF647-loaded PLGA-1 (n = 4). Data are shown as means ± SEM.

Downloaded from https://www.science.org at Purdue University on January 20, 2022
maintained over 95% of bioactivity (Fig. 3C). Collectively, these data demonstrated that encapsulated cGAMP remained stable and could be completely released from PLGA microparticles.

**Single injection of cGAMP-loaded particles effectively inhibited tumor growth**

To test our hypothesis that a single injection of several timed release populations of cGAMP-loaded PLGA-MPs could stimulate anti-tumor immunity comparable to multiple injections of soluble cGAMP (cGAMP-S), mice bearing poorly immunogenic B16F10 melanoma tumors were treated intratumorally with (i) a single injection of cGAMP-S (10 μg) combined with cGAMP-loaded PLGA-1, PLGA-2, and PLGA-3 microparticles (cGAMP-MPs, 10 μg of cGAMP per formulation) to mimic four doses; (ii) four injections of cGAMP-S (10 μg of cGAMP per injection) administered at multiple time points corresponding to PLGA release (4×cGAMP-S); (iii) a single intratumoral injection of cGAMP-S combined with cGAMP-loaded PLGA-1, PLGA-2, and PLGA-3 microparticles (cGAMP-MPs, 10 μg of cGAMP per formulation) to mimic four doses.

**Fig. 3. Single injection of cGAMP-MPs inhibited tumor growth and prolonged animal survival.** (A) Cumulative in vitro release of 3′3′-cGAMP from PLGA-1, PLGA-2, and PLGA-3 (n = 6 to 8). Data are shown as means ± SEM. (B) Mass spectrum of 3′3′-cGAMP released from PLGA-2 on day 8 showed molecular ions [M + H]+ = 675.11, [M + Na]+ = 697.09, and [M + 2Na]+ = 719.07. Particles were incubated at 37°C in PBS. (C) Response of encapsulated cGAMP in PLGA-2 after sealing or released cGAMP from PLGA-2 (incubated at 37°C in PBS for 8 days) on an interferon regulatory factor (IRF) reporter cell line (n = 6). Stock solution of cGAMP was used as a positive control. Data are shown as means ± SD. Statistical significance was calculated using one-way analysis of variance (ANOVA). ns, not significant. (D) Treatment scheme of B16F10 melanoma and orthotopic 4T1 breast tumor models. Tumor-bearing mice were treated with a single intratumoral injection of EP, cGAMP-S + EP (40 μg of cGAMP), cGAMP-collagen (40 μg of cGAMP), or cGAMP-MPs (40 μg of cGAMP; 10 μg in each of cGAMP-S, PLGA-1, PLGA-2, and PLGA-3) at day 7, or four intratumoral injections of cGAMP-S at days 7, 11, 15, and 18 (4×cGAMP-S, 10 μg per injection) after tumor inoculation. (E and F) Average tumor growth (E) and Kaplan-Meier survival curves (F) of B16F10 melanoma–bearing mice treated with different therapeutic combinations (n = 8 biologically independent samples). (G and H) Average tumor growth curve (G) and survival analysis (H) of mice bearing orthotopic 4T1 breast tumors (n = 8 biologically independent samples). Statistical significance was calculated by two-way ANOVA and Tukey’s multiple comparisons test: *P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001. Data are shown as means ± SEM.
of empty PLGA-1, PLGA-2, and PLGA-3 microparticles; or (iv) a single intratumoral injection of high-dose cGAMP-S (40 μg) and EP (cGAMP-S + EP; Fig. 3D). Untreated mice were used as negative controls. To compare the therapeutic efficacy of cGAMP-MPs to other sustained-release systems, we fabricated three sustained-release formulations including dextran hydrogel, polyvinyl alcohol (PVA) hydrogel, and collagen hydrogel. The in vitro release kinetics study showed that over 99% of cGAMP was released from all three hydrogel formulations within 24 hours (fig. S8), which is consistent with previously reported sustained-release systems for cGAMP (30, 31). Collagen gel, which exhibits the slowest release rate among the tested gels, was loaded with 40 μg of cGAMP and intratumorally administered into tumor-bearing mice as controls (cGAMP-collagen). Tumors grew rapidly in untreated and EP-treated groups, and all mice died within 21 days, indicating that PLGA microparticles alone did not inhibit tumor growth (Fig. 3, E and F). A single injection of cGAMP-S + EP exhibited an antitumor effect at early time points but failed to achieve sustained tumor inhibition. The survival time was slightly extended from 21 days for untreated mice to 25 days, indicating the necessity of multiple doses for effective tumor inhibition. cGAMP-collagen did not show superior tumor inhibition or survival compared with cGAMP-S + EP. In contrast, cGAMP-MPs effectively inhibited tumor growth and prolonged animal survival with no statistical difference compared with 4×cGAMP-S at equivalent doses. Similar results were also observed in both orthotopic (Fig. 3, G and H) and subcutaneous (fig. S9) triple-negative breast cancer model (4T1) with the same treatments. The systemic interleukin-6 (IL-6) response of orthotopic 4T1 tumor-bearing mice was evaluated from day 1 to day 7. The IL-6 concentrations were increased in both cGAMP-MPs- and 4×cGAMP-S–treated groups compared with the untreated group (fig. S10), suggesting the successful release of cGAMP from PLGA-MPs into tumors and into the bloodstream.

**Single injection of cGAMP-MPs stimulated potent antitumor immunity**

Next, we investigated the activation of STING pathway and antitumor immunity within the TME of B16F10 melanoma tumors. A combination of cGAMP-S, cGAMP-loaded PLGA-1, and cGAMP-loaded PLGA-2 micro-particles (cGAMP-MPs) was injected intratumorally at day 7 after tumor inoculation to mimic a total of three doses (fig. 4A). Mice treated with a single intratumoral injection of EP or multiple intratumoral injections of cGAMP-S at days 7, 11, and 15 were used as controls. cGAMP-MPs substantially inhibited tumor growth (fig. S11), which is consistent with our tumor inhibition findings. Tumors were isolated 1 day after the third cGAMP-S injection and analyzed by flow cytometry of empty particles (EPs) did not increase TILs within the TME, confirming its inability to activate the STING pathway. We did not observe differences in regulatory T cells between any of the groups (fig. S16).

We next evaluated the changes in dendritic cells (DCs) and myeloid composition in B16F10 melanoma TME after treatments. Both 3×cGAMP-S and cGAMP-MPs promoted the influx of DCs (CD11b+CD11c+), basophils (CD11b−Gr-1−CD200R3+), monocytes (CD11b+Ly6c−Ly6g+), and macrophages (CD11b+Ly6c−Ly6g−), creating an innate inflammatory niche with the potential to prime adaptive immunity (Fig. 4F) (37). In contrast, EPs did not increase the myeloid cell population, suggesting the low immunogenicity of PLGA. In addition, surface expression of CD86, which is a maturation marker overexpressed on activated tumor-infiltrating DCs (38), was increased by 1.7- and 1.5-fold for 3×cGAMP-S– and cGAMP-MPs–treated groups, respectively (Fig. 4G). The maturation of DCs in combination with enriched TILs and increased intratumoral ISGs suggests potential activation of adaptive immunity (10, 39, 40). We next evaluated the polarization of macrophages within the TME, another key function associated with cGAMP (34, 41). After 3×cGAMP or 1×cGAMP-MPs treatment, repolarization of M2-like macrophages in the tumor to M1-like phenotype was observed, which is consistent with previous studies of STING agonist–treated tumors and suggests a less immunosuppressive TME (10, 34, 42). cGAMP-MPs downregulated the canonical M2 surface marker (CD206) and up-regulated M1 surface markers (CD86; Fig. 4H). Quantitative analysis showed that cGAMP-MPs induced about a twofold greater M1/M2 ratio than did the 3×cGAMP-S–treated group. EPs had no significant effect on the M1/M2 ratio compared with the untreated group.

**Single injection of cGAMP-MPs triggered potent systemic antitumor immunity**

To study whether activation of STING in the TME triggers systemic antitumor immunity, mouse serum was collected from the anti-tumor efficacy study of B16F10 melanoma model (Fig. 3D) 21 and 28 days after tumor inoculation and analyzed by flow cytometry. cGAMP-MPs and 4×cGAMP-S treatments increased IFN-γ CD8+ effective cancer immunotherapy (34). Flow cytometry analysis of tumors showed that 3×cGAMP-S and cGAMP-MPs increased TILs by about 23.5- and 17.6-fold compared with the untreated group (figs. S12 and S13). Among these TILs, tumor-infiltrating CD8+ and CD4+ T cells had been increased by 24.4- and 23.6-fold for the 3×cGAMP-S–treated group, and 16.2- and 22.1-fold for the cGAMP-MPs–treated group, respectively (Fig. 4D and fig. S14). The amounts of CD8+ and CD4+ T cells in the cGAMP-S–treated group were not substantially different from those in the cGAMP-MP–treated group. 3×cGAMP-S and cGAMP-MPs treatments also showed 1.2- and 1.5-fold increase in CD8+CD4+ T cell ratio compared with the untreated group, respectively (fig. S13), which is a commonly reported protective prognostic indicator of immunotherapy (10, 35). In agreement with this enriched CD8+ T cell infiltration and enhanced antitumor activity, terminal deoxynucleotidyl transferase–mediated deoxyuridine triphosphate nick end labeling (TUNEL) showed a greater abundance of apoptotic cells for the cGAMP-MPs– and 3×cGAMP-S–treated groups than for the untreated and EP-treated groups (fig. S15). In addition, 3×cGAMP-S and cGAMP-MPs increased infiltrating natural killer (NK) cells (Fig. 4E), another important group of cytotoxic lymphocytes that shape the adaptive immune response and were found to be effective for spontaneous STING-mediated protection against B16F10 tumors (36). Conversely, the group receiving EPs did not increase TILs within the TME, confirming its inability to activate the STING pathway. We did not observe differences in regulatory T cells between any of the groups (fig. S16).
T cells in serum at day 21 by 5.1- and 4.9-fold, respectively, compared with the untreated group (Fig. 5A and fig. S17). The number of IFN-γ CD8+ T cells remained unchanged at day 28 (Fig. 5A), demonstrating long-lasting systemic immune response. In addition, cGAMP-MPs also increased the number of memory CD62L−CD44+CD4+ T cells (~5.4-fold over untreated group) in TME (Fig. 5B) (43, 44). Collectively, a single injection of cGAMP-MPs generated long-lived systemic antitumor immunity and local immunological memory, which may be able to prevent tumor recurrence and metastases.

To study whether cGAMP-MPs could inhibit the growth of distant tumors, we used a contralateral B16F10 tumor model. The primary tumor was treated by a single intratumoral injection of cGAMP-MPs (cGAMP-S, cGAMP-loaded PLGA-1, and cGAMP-loaded PLGA-2) to achieve overall three doses at days 7, 11, and 15 after tumor inoculation. The distant tumor did not receive any treatment (Fig. 5C). cGAMP-MPs effectively inhibited the growth of both the primary and distant tumors compared with the untreated group, thereby demonstrating strong systemic antitumor immunity (Fig. 5, D and E). To evaluate whether cGAMP-MPs could improve the antitumor efficacy of ICB, we tested a combination therapy of...
Fig. 5. Single injection of cGAMP-MPs induced systemic antitumor immunity and inhibited metastasis. (A) Quantitative analysis of IFN-γ+CD8+ T cells in the serum collected at days 21 and 28 (n = 4 to 5, treatment scheme shown in Fig. 3D). Untreated and 1×EP-treated mice did not survive to day 28. Data are shown as means ± SEM. (B) Numbers of effective memory CL62L−CD44+CD4+CD3+ and CL62L−CD44+CD8+CD3+ T cells in the TME (treatment scheme shown in Fig. 4A). (C) Schematic of treatment regimen on a contralateral B16F10 model. Tumors were inoculated on the right (primary) and left (distant) rear flanks of mice at days 0 and 2, respectively. The mice were treated with a single intratumoral injection of cGAMP-MPs (30 μg of cGAMP: 10 μg of cGAMP in each of cGAMP-S, PLGA-1, and PLGA-2) on the primary tumor, three intraperitoneal injections of anti–PD-1 (3×ICB, 100 μg per injection), or the combination of both cGAMP-MPs and 3×ICB. The distant tumor did not receive any treatments. (D and E) Average tumor growth curves of treated (D) and distant (E) tumors (n = 8). Data are shown as means ± SEM. (F) Schematic of treatment regimen on a metastatic 4T1 model. Mice were treated with a single intratumoral injection of cGAMP-MPs (30 μg of cGAMP: 10 μg of cGAMP in each of cGAMP-S, PLGA-1, and PLGA-2) or three intratumoral injections of cGAMP-S (3×cGAMP-S). (G and H) Representative lung photographs (G) and numbers of metastatic foci (H) on lung surfaces after treatments (n = 8). Arrows point to metastatic tumors. Scale bar is 0.5 cm. (I) Representative H&E-stained lung sections and digitally processed images used for quantifying metastatic tumor cells. Scale bar, 2 mm. (J) Percentage of tumor area within total lung area after treatments (n = 4 to 5). Statistical significance was calculated by Student's t test or two-way ANOVA. *P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001.
cGAMP-MPs with anti–programmed death 1 (PD-1) in the same contralateral B16F10 tumor model. The combination of cGAMP-MPs and ICB showed greater inhibition of primary and distant tumor growth than individual therapies by themselves (Fig. 5, D and E), demonstrating the potential of combining cGAMP-MPs with ICB to enhance the therapeutic effect.

We then studied the effect of cGAMP-MPs on metastasis using an orthotopic 4T1 breast cancer model. We treated the primary tumor with a single intratumoral injection of cGAMP-MPs at day 7 or three intratumoral injections of cGAMP-S at days 7, 11, and 15 (3×cGAMP-S) after tumor inoculation. The primary tumors were surgically removed at day 18 to extend survival time, which was necessary to allow the development of metastasis. We then isolated lungs and analyzed metastasis at day 34 (Fig. 5F). cGAMP-MPs treatment significantly decreased the number of metastatic foci on lung surfaces (Fig. 5, G and H; \( P < 0.0001 \)) and reduced the relative area of tumors in the lungs compared with the untreated groups (Fig. 5, I and J; \( P < 0.01 \)). cGAMP-MPs also exhibited a greater ability to decrease the percentage of metastatic tumor cells within lungs compared with 3×cGAMP-S (Fig. 5); \( P < 0.5 \), suggesting the benefits of a single injection for decreasing metastasis.

cGAMP-MPs could expand the scope of current STING agonist–based therapies

In the clinic, patients often develop recurrent tumors after surgery because of residual microtumors and circulating tumor cells (45–47). To expand the clinical applications of cGAMP-MPs, we adopted a surgical resection melanoma model to evaluate the efficacy of cGAMP-MPs on inhibiting tumor recurrence (48, 49). Six days after tumor inoculation, about 99% of each B16F10 tumor was surgically removed. We then directly deposited cGAMP-MPs (cGAMP-S, cGAMP-loaded PLGA-1 and PLGA-2) at the fresh surgical bed to achieve three doses overall at days 0, 4, and 8. Mice treated with 3×cGAMP-S or 1×cGAMP-S + EP were used as controls (Fig. 6A). cGAMP-MPs and 3×cGAMP-S treatments improved tumor inhibition (Fig. 6B) and enhanced survival (Fig. 6C) compared with untreated and cGAMP-S + EP–treated mice. Six of eight cGAMP-MP– and 3×cGAMP–S–treated mice were tumor free and survived for over 60 days after inoculation (Fig. 6C). The tumor recurrence rates of cGAMP-MPs– and 3×cGAMP–S–treated groups were both 25% (two of eight mice), which was lower than that of the untreated group (100%, eight of eight mice) and 1×cGAMP–S + EP–treated group (87.5%, seven of eight mice; Fig. 6C). We then rechallenged these tumor-free mice through subcutaneous injection of B16F10 cells at day 60. The tumors grew significantly slower in cGAMP-MPs– and 3×cGAMP–S–treated mice than in naïve mice (Fig. 6D; \( P < 0.001 \)). Survival analysis also showed prolonged survival time for the treated groups (Fig. 6E), suggesting cGAMP-MPs and 3×cGAMP–S can provide protective immunity.

We further tested the therapeutic efficacy of cGAMP-MPs on an allograft model of pancreatic cancer (KPC model). We injected cGAMP-MPs (10 µg each of cGAMP-S, cGAMP-loaded PLGA-1 and PLGA-2) after tumor inoculation in the pancreas to achieve overall three doses at days 0, 4, and 8 (Fig. 6F). Multiple intratumoral injections of cGAMP-S are extremely difficult on such hard-to-reach tumors. Therefore, we performed a single injection of a high dose of cGAMP-S (30 µg) + EP at day 0 (Fig. 6F). Untreated mice were used as negative controls. We then analyzed the tumor growth and metastasis 25 days after treatments. cGAMP-MPs significantly inhibited primary tumor growth in the pancreas and metastasis to lungs compared with the untreated group (Fig. 6, G to I; \( P < 0.001 \)). In contrast, cGAMP-S + EP did not show benefits on tumor growth or metastasis. Collectively, these data demonstrate that cGAMP-MPs could be useful for hard-to-reach tumors and suggest the necessity of multiple doses at extended time points for effective therapy.

Toxicity analysis

PLGA has been used in many FDA-approved medical devices owing to its biodegradability and biocompatibility (50). Flow cytometry analysis of immune cells in TME indicated that intratumorally administered empty PLGA-MPs did not induce any detectable inflammation in situ (Fig. 4, D to G). In addition, we did not observe weight loss or behavior changes in any of the animals throughout the treatment period for all in vivo studies (fig. S18). Hematoxylin and eosin (H&E) staining of histological sections of major organs (heart, liver, spleen, lung, and kidney) showed no obvious change in morphology (Fig. S19). We further studied the biodegradability of PLGA-MPs by subcutaneously injecting empty PLGA-2 into immuno-competent mice and performed H&E staining of skin tissues at days 2, 8, and 30 after injection. At day 2, PLGA-MPs exhibited cubic morphology under the skin with few lymphocytes and leukocytes around the particle, suggesting the presence of minimal inflammation. The particles then deformed to an ellipse shape due to hydrolysis of PLGA with diminished immune cells at day 8, just around the release window, which is consistent with flow cytometry analysis. We did not observe any particles left in mice at day 30, which suggests complete degradation and clearance (fig. S20). Collectively, these data demonstrate that microfabricated PLGA-MPs did not show toxicity and can be completely degraded and cleared in vivo.

PLGA-MPs could be a platform technology to deliver various cancer therapeutics

Loading drugs into PLGA-MPs is an independent step during particle fabrication, thus allowing us to fill other types of cancer therapeutics into PLGA-MPs. We performed preliminary studies of loading either a chemotherapeutic drug (pemetrexed) or a TLR agonist [5′-cytosine-phosphate-guanine-3′ (CpG) DNA] into PLGA-2. Pemetrexed and CpG DNA exhibited nearly the same in vitro release kinetics from PLGA-2 as cGAMP (fig. S21), indicating the potential applicability of PLGA-MPs for these cancer therapeutics. The drug-loading capacity of PLGA-MPs could also be tuned by changing the particle design. We fabricated a different particle base with a bigger cavity (300 µm by 300 µm by 200 µm, length by width by height) and wall thickness of 50 µm to increase the loading capacity. We could load ~10 µg of cGAMP into each of these particle bases (fig. S22).

DISCUSSION

Adherence to current STING agonist–based therapy is challenging because of the frequent injections over a long period of time and the need of trained health care professionals for each injection. Poor patient adherence represents a notable challenge resulting in treatment failures and large financial costs (19, 51). Frequent injections for cancer treatment also cause a substantial burden to daily lives of patients (52). Previous research efforts have mainly focused on improving the cellular uptake of STING agonist and tumor-targeting efficacy after systemic administration (10, 34, 53). Here, we describe...
Fig. 6. cGAMP-MPs prevented tumor recurrence after surgery and inhibited growth of hard-to-reach tumors. (A) Schematic of treatment regimen in a surgical removal B16F10 model. About 99% of the tumor mass was surgically removed at day 6 after tumor inoculation. cGAMP-MPs (30 μg of cGAMP: 10 μg of cGAMP in each of cGAMP-S, PLGA-1, PLGA-2) or 1×cGAMP-S + EP (10 μg of cGAMP with empty PLGA-1 and PLGA-2) was directly deposited at the surgical bed. For 3×cGAMP-S treatment, cGAMP-S was administered intratumorally at days 6, 10, and 14 (10 μg per injection). (B and C) Average tumor growth curve (B) and survival analysis (C) of treated mice (n = 8). (D and E) Tumor-free mice after cGAMP-MPs and 3×cGAMP-S treatments were rechallenged with B16F10 cells at day 60 after tumor inoculation. Naïve mice were challenged as negative controls. Tumor growth (D) and survival (E) were monitored over time (n = 6). Data are shown as means ± SEM. (F) Schematic of treatment regimen on an orthotopic pancreatic tumor model. cGAMP-MPs or 1×cGAMP-S + EP was injected into the pancreases immediately after tumor inoculation (n = 7). (G and H) Representative images (G) and weights (H) of isolated pancreatic tumors. Scale bar, 1 cm. (I) Representative photographs of lungs and H&E-stained lung sections. Red arrows point to metastatic tumors. Scale bar, 2 mm. Statistical significance was calculated by one way or two-way ANOVA and Tukey’s multiple comparisons test. *P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001.
an approach of improving the overall effectiveness of STING agonist therapy by ensuring that patients receive every required dose at the correct time.

To replace multiple intratumoral injections of soluble STING agonist, PLGA-MPs need to stay inside tumors and release biologically active cargo at anticipated times. MicroCT analysis showed that these microparticles aggregated at the tumor injection site due to low mobility. LC-MS and in vitro analysis in cells showed that cGAMP maintained >95% of bioactivity after release from PLGA-MPs. We then extensively studied the release kinetics of different cargos in vitro and in vivo. The release kinetics of PLGA-MPs were independent of the cargo released (AF647, AF647-dextran, and STING agonist) and the in vivo microenvironment (subcutaneous, B16F10, and 4T1 tumors). These observations are expected because the degradation of PLGA is predominantly driven by hydrolysis. It has been previously reported that enzymatic activity has a negligible effect on PLGA degradation (50). The slightly acidic TME did not accelerate the release of low-MW PLGAs, but may have effects on the long-term release of PLGA-MPs because of the acid-catalyzed hydrolysis of PLGA. Therefore, testing in vitro release in an acidic environment is needed to predict release kinetics in tumors.

Pulsatile release over long periods of time is usually achieved by implantable drug delivery devices, which require invasive surgery to administer and remove (54). One benefit of PLGA-MPs is that they could be injected using a regular needle and completely degraded over time, thus improving patient compliance. Other injectable long-term drug release systems, such as emulsion-based microparticles (including PLGA formulations) or hydrogels, often show an initial phase of burst release and then a second phase of sustained release for hydrophilic drugs (26, 55, 56). Such release kinetics result in an initial high dose, which could cause toxic side effects. In addition, achieving sustained release of small/hydrophilic drugs, such as STING agonists, over weeks is extremely challenging. Drug encapsulation efficacy is also relatively low for emulsion-based microparticles (57). Our engineered PLGA microparticles could achieve essentially 100% drug encapsulation efficacy and be combined to exhibit multiple burst release events at time points up to months. We tested the release kinetics of several low-MW PLGAs in this study to achieve release time within the previously reported treatment schedules using cGAMP in animal models. Expanding the release kinetics library to cover pulsatile release over months or even a year should be possible by tuning the MW, chain-end functionality, and copolymer ratio of PLGA. Therefore, we could potentially administer customizable doses by physically mixing PLGA-MPs with different release profiles within one injection.

We demonstrated that the antitumor efficacy of single-administered PLGA-MPs is comparable with multiple injections of cGAMP solutions in multiple mouse models. PLGA-MP–treated B16F10 tumors showed high amounts of ISGs and phosphorylated TBK1 and IRF3 proteins, suggesting the successful activation of the STING pathway by a sequence of pulsatile releases of cGAMP. PLGA-MPs induced an immunogenic TME, as demonstrated by increased tumor-infiltrating CD8+ T cells, NK cells, DCs, and the shift from an M2 to an M1 macrophage phenotype. Loading cGAMP in PLGA-MPs improved M1 polarization as compared with free cGAMP, possibly due to the acidic degradation products of PLGA, which have been shown to stimulate proinflammatory macrophages (58–60). Further studies on macrophage polarization kinetics are required to fully elucidate the function of PLGA-MPs on macrophage polariza-
STING agonist–based cancer immunotherapy. PLGA-MPs also expand the scope of STING agonist–based therapy to hard-to-reach tumors and as an adjuvant therapy to prevent tumor recurrence after surgery. This single injection delivery system contains only PLGA, which is widely used in FDA-approved drug products, thus holding great promise for rapid clinical translation. The modularity of this platform makes it easily compatible with loading any hydrophilic drug and many hydrophobic drugs (for example, pemetrexed and CpG DNA) or even delivering various drugs at different times for combination cancer therapies. This platform also opens up opportunities for many other diseases that require frequent or repeated local administration of therapeutics.

MATERIALS AND METHODS

Study design

The objective of this study was to develop an injectable delivery platform that could release multiple doses of STING agonist at distinct time points within one injection. We first screened the release kinetics of different PLGA-MPs in vitro to find desired releasing time points for treating tumor-bearing animal models. We further verified the release kinetics of PLGA microparticles in vivo using fluorescence imaging. The antitumor efficacy was evaluated in mouse B16F10 melanoma, 4T1 breast cancer, contralateral B16F10 melanoma, orthotopic pancreatic cancer, and incomplete surgical removal tumor models. To demonstrate the activation of STING pathway, we analyzed STING-related mRNA expression and protein abundance using qPCR and Western blots, respectively. The local and systemic immune responses of microparticles were also examined using flow cytometry and immunofluorescence staining. The toxicity and biodegradability of PLGA microparticles were studied by body weight analysis and histology. Mice were randomly divided into different treatment groups for all studies. The researchers were not blinded in this study. Power analysis was not used to predetermine sample size. Sample sizes were determined on the basis of our previous studies. The replicate numbers for each experiment and statistical analysis methods are indicated in the figure legends. Raw data are shown in data file S1.

Fabrication of PLGA microparticles

PLGA were purchased from Evonik and PolySciTech. PLGA microparticles were fabricated through the Stamp Assembly of Polymer Layers process (64). Photomasks with microscale patterns of bases and caps were made by Front Range Photomask. Positive master molds of microparticle base and cap were created by SU-8 lithography on silicon wafers. The mixture of PDMS base and curing agent (Sylgard 184, Dow Corning) was poured onto the silicon master mold and then exposed to high vacuum for 1 hour. A glass slide with two coveslips at each end was then pushed against the silicon mold while curing in the oven for 2 hours at 150°C to yield a thin PDMS mold. The obtained PDMS molds were then used as negative molds to press desired microparticles. PLGA films were prepared by solvent casting 60% (w/v) PLGA in acetone solution. The thickness of PLGA films was ~1650 to 1750 μm. For molding the caps of the microparticles, a small piece of PLGA film was placed between the PDMS cap mold and a Teflon film and covered with a glass slide. A pair of spring-loaded clamps was then used to fix and compress the microparticles in a 120°C vacuum oven for 2 hours. PLGA film melted and flowed in the cap PDMS mold during heating. The setup was then allowed to cool down to room temperature and separated to yield the PLGA caps in the PDMS mold. For molding the base of the microparticles, the above steps were repeated without using the Teflon film. Therefore, the PLGA base would separate from the PDMS base mold and stick to the covering glass slide. The cargos of interest were filled into the PLGA microparticles using a BioJet Ultra picoliter dispensing instrument (BioDot). The aqueous solutions of cargos were dispensed for multiple 15-drop cycles of 180- to 200-pl drops. Filled particles were then aligned and sintered with corresponding PLGA caps using a photomask aligner (MA4, Karl SUSS) retrofitted with a Peltier heater to enable simultaneous alignment and sealing. Sealed particles were then separated from glass slides using a razor blade. Scanning electron microscopy images were collected using a JSM-5600LV scanning electron microscope (JEOL) with an acceleration voltage of 5 kV. High-resolution x-ray CT images were collected at the Biotechnology Resource Center of Cornell University.

In vitro release kinetics

To study the in vitro release kinetics of AF647-dextran (Life Technologies), 3′3′-cGAMP (InvivoGen), Cy5-CpG DNA (5′-TCC ATG ACG TTC CTG ACG TT-Cy5–3′, IDT Inc.), and pemetrexed (Sigma-Aldrich), PLGA microparticles were separately filled with 1 μg of AF647-dextran, 2 μg of 3′3′-cGMAP, 1 μg of Cy5-CpG, or 2 μg of pemetrexed, respectively. To determine the cargo loading in particles, filled particles were suspended in 200 μl of PBS buffer individually, vortexed for 15 s, and centrifuged at 14,000 rcf for 1 min. The supernatants were then analyzed by a microplate reader (AF647, AF647-dextran, Cy5-CpG), high-performance liquid chromatography (HPLC) (pemetrexed), and NanoDrop (3′3′-cGAMP, absorbance at 260 nm). The results were quantified using a standard curve of a serial dilution of stock solutions. Filled particles were then sealed with corresponding PLGA caps. Each particle was placed into 200 μl of PBS (pH 6.84) in a 0.5-ml microcentrifuge tube (Eppendorf) and incubated on an orbital shaker at 37°C. The supernatant of each centrifuge tube was collected at predetermined time points. The supernatants of the AF647-dextran and Cy5-CpG groups were analyzed by a microplate reader (Tecan Infinite M200 spectrophotometer; excitation/emission, 640/680 nm). The supernatants of 3′3′-cGAMP and pemetrexed were analyzed via HPLC (Alliance HPLC systems, Waters Co.) using a C-18 column (Acclaim PolarAdvantage II, 3 μm, 4.6 mm by 150 mm) and a photodiode detector at 260 nm for 3′3′-cGAMP and 254 nm for pemetrexed. Water and acetonitrile were used as mobile phases. The results were quantified using a standard curve of a serial dilution of stock solutions and normalized to total cumulative release (n = 6 to 8). The actual release day of each PLGA was determined as the day at which more than 50% of cargo was released.

For the sustained-release systems, fast-dextran hydrogel kits (part no. TURE2-1KT), fast-PVA hydrogel kits (part no. TRUE4-1KT), and 3D collagen Kit (part no. ECM675) were purchased from Millipore-Sigma. A total of 40 μg of cGAMP was loaded into 30-μl hydrogels according to the manufacturers’ instructions. To study the in vitro release kinetics, cGAMP-loaded hydrogels were incubated on an orbital shaker at 37°C. The supernatant of each centrifuge tube was collected at predetermined time points and analyzed by NanoDrop.

Animals and cell lines

All animal procedures were approved by the Massachusetts Institute of Technology Committee on Animal Care. Six- to 8-week-old SKH1-E, C57BL/6, and BALB/c female mice were purchased from Charles
of cGAMP stock solutions, dissolved cGAMP after particle fabrication. The supernatant containing unreleased AF647 and serum was physically broken to release AF647 during homogenization. The tumors were homogenized in PBS buffer. Unreleased AF647–PLGA-1 was physically broken to release encapsulated cargo. To evaluate the activity of 3′3′-cGAMP after particle fabrication, cGAMP-loaded PLGA-2 was placed in PBS buffer and mechanically broken by a scalpel to release encapsulated cargo. To evaluate the activity of cGAMP after release, cGAMP-loaded PLGA-2 was incubated on an orbital shaker in PBS buffer at 37°C. Supernatant was collected at the release window and quantified by NanoDrop. RAW-Lucia ISG cells (5 × 10⁶) were plated in a 96-well plate. A serial dilution of cGAMP stock solutions, dissolved cGAMP after particle fabrication, and released cGAMP were incubated with cells for 24 hours before adding QUANTI-Luc solution. The plate was then analyzed by a microplate reader. The results were quantified using a standard curve of a serial dilution of stock solutions.

**Treatment of B16F10 and 4T1 tumors**

4T1 (2 × 10⁵) or B16F10 (2 × 10⁵) cells were subcutaneously injected into the right rear flank of BALB/c or C57BL/6 female mice, respectively. For orthotopic 4T1 model, 2 × 10⁵ 4T1 cells were injected into the mammary fat pad of BALB/c female mice. Seven days after tumor injection, B16F10 tumor-bearing mice were divided into six experimental groups (n = 8 for each group): untreated, 1×EPs, 1×cGAMP-S + EP, 1×cGAMP-collagen, 4×cGAMP-S, and cGAMP-MPs. For the 4×cGAMP-S group, mice were intratumorally administered 10 μg of cGAMP in 50 μl of MC solution via an 18-gauge filter needle. The overall dose of cGAMP was 40 μg per mouse throughout the treatment period. For the cGAMP-MP group, mice were given a single intratumoral injection of a mixture of 10 μg of cGAMP-S, five PLGA-1 particles containing a total of 10 μg of cGAMP, five PLGA-2 particles containing a total of 10 μg of cGAMP, and five PLGA-3 particles containing a total of 10 μg of cGAMP in 50 μl of MC solution via an 18-gauge filter needle. The overall dose of cGAMP is 40 μg. For the 1×EP and 1×cGAMP-S + EP groups, five of each of empty PLGA-1, PLGA-2, and PLGA-3 particles with or without 40 μg of cGAMP-S were injected intratumorally in 50 μl of MC solution. Mice from the untreated groups were injected intratumorally with 50 μl of MC solution at day 7 after tumor inoculation. In the subcutaneous and orthotopic 4T1 models, tumor-bearing mice were subjected to MC solution (untreated group), 4×cGAMP-S, or 1 × cGAMP-MPs treatments at day 7 after tumor inoculation (n = 8 for each group). Tumor size was measured every other day, starting at day 7 after tumor inoculation with a digital caliper. Tumor volume was calculated using the following formula: length (mm) × width² (mm) × 0.5. Animals were euthanized when they showed signs of poor health or when the tumor size exceeded 1500 mm³.

**Western blot and qPCR**

Mice bearing B16F10 tumors were divided into the following four experimental groups (n = 5): untreated, 1×EP (five each of empty PLGA-1 and PLGA-2), 3×cGAMP-S, and 1×cGAMP-MPs (10 μg of cGAMP-S, five PLGA-1 particles containing a total of 10 μg of cGAMP, and five PLGA-2 particles containing a total of 10 μg of cGAMP). For the 3×cGAMP-S group, mice were intratumorally administered 10 μg of cGAMP in 50 μl of MC solution at days 7, 11, and 15 after tumor inoculation. Tumors were collected at day 16 after tumor inoculation and cut into 50- to 100-mg pieces in a 1.5-ml microcentrifuge tube. The tumors were lysed in radioimmunoprecipitation assay buffer (Sigma–Aldrich), homogenized, and centrifuged at 20,130 rcf for 10 min. The protein content in the supernatant was quantified using a bicinchoninic acid protein assay kit (Thermo Fisher Scientific). Equal amounts of proteins (20 μg) were separated on a 4 to 15% gradient SDS–polyacrylamide gels (Bio–Rad) and electrotransferred to nitrocellulose membranes. The membranes were then blocked with 5% milk in tris-buffered saline supplemented with 0.05% Tween 20 and further incubated with glyceraldehyde-3-phosphate dehydrogenase (GAPDH) monoclonal antibody (mAb; Invitrogen, catalog no. MA5-27912), phospho-TBK1/NAK (Ser 172) (D52C2) rabbit mAb (Cell Signaling Technology, catalog no. 5483S), or phospho-IRF3 (S396) rabbit mAb (Cell Signaling Technology, catalog no. 4947S).
at 4°C overnight. The membranes were then incubated with goat anti-rabbit immunoglobulin G (IgG) (H + L) secondary antibody, horseradish peroxidase (HRP) (Invitrogen, catalog no. TG266717) for 1 hour at room temperature. Protein bands were visualized by chemiluminescence using the ECL Western Blotting Substrate (Thermo Fisher Scientific).

For qPCR experiments, total RNAs were extracted from tumors by the RNeasy Kit (Qiagen Inc.) according to the manufacturer’s protocol. Total RNA was then reverse transcribed to complementary DNA (cDNA) using the high-capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific). The obtained cDNA was amplified with TaqMan Gene Expression Master Mix (Thermo Fisher Scientific) using a 384-well LightCycler 480 (Roche). We used primers for Irf7 (Thermo Fisher Scientific, assay Id. Mm00516793_g1), Cxcl10 (Thermo Fisher Scientific, assay Id. Mm00445235_m1), and Gapdh (Thermo Fisher Scientific, assay Id. Mm99999915_g1). Samples were analyzed in triplicates.

Flow cytometry
To stain the cell surface markers for flow cytometry analysis, cells were pretreated with anti–CD16/32-Fc blocker (BioLegend, catalog no. 101319) and stained with fluorochrome-conjugated antibody solution according to the manufacturer-suggested dilutions on ice for 1 hour. To stain each intracellular marker, for example, IFN-γ, cells were prestimulated with cell stimulation cocktail (eBioscience, catalog no. 00-4970-93) for 4 to 6 hours, fixed, and permeabilized using fixation/permeabilization solution kit (BD Biosciences, catalog no. 554714), and then stained with both anti–IFN-γ and other surface antibodies. Antibodies used for flow cytometry studies were anti–CD86-BUV395 (BD Biosciences, catalog no. 564199), anti–CD45-BUV737 (BD Biosciences, catalog no. 564880), anti–TCRβ-BV421 (BioLegend, catalog no. 109229), anti–NK.1.1-BV605 (BioLegend, catalog no. 108739), anti–NK1.1-BV605 (BioLegend, catalog no. 108739), anti–CD8α-FITC (BD Biosciences, catalog no. 553030), anti–CD4-PerCP/Cy5.5 (BD Biosciences, catalog no. 550954), anti–CD62L–phycoerythrin (PE) (BioLegend, catalog no. 104407), anti–CD19-PE/Cy7 (eBioscience, catalog no. 25-0193-81), anti–CD3-PE/P94 (BioLegend, catalog no. 100245), anti–FOXP3-APC (eBioscience, catalog no. 17-5773-80), anti–CD11b–AF700 (BioLegend, catalog no. 201222), anti–CD8α-BV421 (BioLegend, catalog no. 100737), anti–Ly6c–BV510 (BioLegend, catalog no. 127633), anti–Siglec F–BV605 (BD Biosciences, catalog no. 740388), anti–MHC II–BV786 (BD Biosciences, catalog no. 743875), anti–Ly6c–AF488 (BioLegend, catalog no. 128021), anti–CD11c–PerCP/Cy5.5 (BioLegend, catalog no. 117327), anti–CD206–PE (BioLegend, catalog no. 141705), anti–CD197–PE/P94 (BioLegend, catalog no. 120121), anti–F4/80–PE/Cy7 (BioLegend, catalog no. 123113), anti–CD200R3–APC (BioLegend, catalog no. 142207), and anti–CD11b–AF700 (BioLegend, catalog no. 101222), as well as viability dye eFluor 780 (eBioscience, catalog no. 65-0865-14). Flow cytometry data were acquired on an LSRFortessa cell analyzer (BD Biosciences) and analyzed using FlowJo software.

Immunofluorescence staining
Tumor sections (5 μm) were fixed with 4% paraformaldehyde, blocked with 3% bovine serum albumin, and permeabilized with 0.1% Triton X-100 in PBS. The tumor sections were then incubated with anti–CD8 alpha antibody (1:200; Abcam, catalog no. ab217344) at 4°C overnight and goat anti-rabbit IgG H&L (Alexa Fluor 488) (1:1000; Abcam, catalog no. ab150077) secondary antibody at room temperature for 1 hour. Apoptotic tumor cells were stained using an in situ cell death detection kit (Roche) according to the manufacturer’s instructions. Images were acquired on a Nikon A1R Ultra-Fast Spectral Scanning Confocal Microscope.

Treatment of contralateral B16F10 tumors
B16F10 cells (2 × 10^5) were subcutaneously injected into the right rear flank of C57BL/6 female mice at day 0. Another 2 × 10^5 B16F10 cells were subcutaneously injected into the left rear flank at day 2 to mimic metastatic tumor. Seven days after primary tumor inoculation, B16F10 tumor-bearing mice were divided into four experimental groups (n = 8 for each group): untreated, 1×cGAMP-MPs, 3×ICB (anti–PD-1), and 1×cGAMP-MPs + 3×ICB. cGAMP-MPs (10 μg of cGAMP-S, five PLGA-1 particles containing a total of 10 μg of cGAMP, and five PLGA-2 particles containing a total of 10 μg of cGAMP) in 50 μl of MC solution was intratumorally injected into the primary tumor (on the right side). For 3×ICB– and 1×cGAMP-MPs + 3×ICB–treated groups, 100 μg of anti–PD-1 antibody (BioLegend, catalog no. 114114) was intraperitoneally injected at days 7, 11, and 15 after primary tumor inoculation. The distant tumor (left side) did not receive any treatments. Tumor size was measured every other day starting at day 7 after tumor inoculation with a digital caliper. Tumor volume was calculated using the following formula: length (mm) × width^2 (mm) × 0.5. Animals were euthanized when they showed signs of poor health or when the tumor size on either side exceeded 1500 mm^3.

Treatment of metastatic 4T1 model
4T1 cells (2 × 10^7) were injected into the mammary fat pad. Seven days after injection, tumor-bearing mice were divided into three experimental groups: untreated, 3×cGAMP-S, and 1×cGAMP-MPs (10 μg of cGAMP-S, five PLGA-1 particles containing a total of 10 μg of cGAMP, and five PLGA-2 particles containing a total of 10 μg of cGAMP). For the 3×cGAMP-S group, mice were intratumorally administered 10 μg of cGAMP in 50 μl of MC solution at days 7, 11, and 15 after tumor inoculation. The primary tumor was surgically removed at day 18 to extend survival. Mice were euthanized at day 34. Lung tissues were stained with India ink and fixed in Fekete’s solution. Metastatic foci on the lung were counted under a microscope. Unstained lung tissues were fixed in formalin and stained by H&E. Quantitation of metastatic tumor cells in H&E-stained sections was performed using an Aperio ImageScope using tuned positive pixel count algorithm. Briefly, we tuned input hue value in the positive pixel count algorithm to positively select normal lung tissues in the red to orange range, whereas the tumors were negatively selected in purple. Percentage area of tumor metastasis per total lung area was calculated as the number of negative counts (purple)/the number of total counts (purple, orange, and red) × 100%. Three H&E sections per lung at different depths were analyzed and averaged to obtain the percentage of tumor over the lung for one mouse. Four or five mice were analyzed for each group.

Treatment of surgically removed B16F10 tumor
B16F10 cells (2 × 10^5) were subcutaneously injected into the right rear flank of C57BL/6 female mice. Six days after tumor inoculation, B16F10 tumor-bearing mice were randomly divided into four experimental groups (n = 8 for each group): untreated, 1×cGAMP-S + EP, 3×cGAMP-S, and 1×cGAMP-MPs. About 99% of tumor volume was...
surgically removed, leaving ~1% residual tumor to mimic residual microtumors. Upon the removal of tumors, cGAMP-MPs (10 μg of cGAMP-S, five PLGA-1 particles containing a total of 10 μg of cGAMP, and five PLGA-2 particles containing a total of 10 μg of cGAMP) or 1×cGAMP-S + EPs (10 μg of cGAMP-S, five each of empty PLGA-1 and PLGA-2 particles) in 50 μl of MC solution was directly applied to the surgical bed through a micropipette. For the 3×cGAMP-S–treated group, 10 μg of cGAMP-S was applied to the surgical bed after surgery followed by intratumoral injections of 10 μg of cGAMP at days 4 and 8 postsurgery. The wound was closed by an autoclip wound clip system. Tumor size was measured every other day starting at day 7 after tumor inoculation with a digital caliper. For the challenge experiment, 2 × 10^5 B16F10 cells were subcutaneously injected into the left rear flank of treated mice with complete responses or naïve mice. Tumor size was measured every other day with a digital caliper.

**Treatment of an orthotopic pancreatic tumor model**

A small incision was made to exteriorize the spleen and pancreas of C57BL/6 female mice. KPC cells (5 × 10^5) in 50 μl of PBS and Matrigel (1:1 mixture by volume) were injected into the tail of the pancreas. cGAMP-MPs (10 μg of cGAMP-S, five PLGA-1 particles containing a total of 10 μg of cGAMP, and five PLGA-2 particles containing a total of 10 μg of cGAMP) or 1×cGAMP-S + EPs (30 μg of cGAMP-S and five each of empty PLGA-1 and PLGA-2 particles) in 50 μl of MC solution was also injected into the tail of the pancreas. The wound was then closed by an autoclip wound clip system. Mice were euthanized 25 days after tumor inoculation. The tumors were isolated and weighed by a balance. Metastasis to lungs was evaluated by H&E stained with H&E, and imaged using an Aperio AT2 Slide Scanner (Leica Biosystems).

**Biodegradation of PLGA-MPs**

Five empty PLGA-2 microparticles were subcutaneously injected into the rear flank of SKH1-E mouse. Mice were euthanized at days 2, 8, and 30 after injection. The skin and subdermal tissue were collected and fixed in formalin-free fixative (Sigma-Aldrich) for 24 hours. Tissues were then embedded in paraffin, cut into 5-μm sections, stained with H&E, and imaged using an Aperio AT2 Slide Scanner system. Tumor size was measured every other day with a digital caliper.

**Supplementary materials**

stm.sciencemag.org/cgi/content/full/12/556/eaaz6606/DC1

**Statistical analysis**

All statistical analyses were performed using the GraphPad Prism software package (PRISM 8.0.2; GraphPad Software). Biological replicates were used in all experiments unless otherwise stated. Survival benefit was determined using a log-rank test. All experimental results were indicated as the means ± SD or the means ± SEM. One-way or two-way analysis of variance (ANOVA) was used when there were multiple comparisons. Student’s t test was used for single comparisons. The specific statistical methods are indicated in the figure legends.

**REFERENCES AND NOTES**


**Acknowledgments:** We thank L. Milling from D. Irvine laboratory for advice on orthotopic 4T1 tumor models. We thank S. Kozlov for providing KPC cells. We thank the Koch Institute Swanson Biotechnology Center for technical support, specifically microscopy, histology, flow cytometry, and animal imaging and preclinical testing facilities. **Funding:** X.L. was supported by a Ludwig Postdoctoral Fellowship. L.M. was supported by a Misrock Postdoctoral Fellowship. K.M. was supported by an NIH Ruth L. Kirschstein National Research Service Award (F32EB022416). K.S. was supported by a Ruth L. Kirschstein NRSA Postdoctoral Fellowship (1F32EB025688-0A1). This work is supported in part by the Cancer Center Support (core) (grant no. P30-CA14051) from the NIH. **Author contributions:** X.L., L.M., and A.J. conceived and designed the experiments. X.L., W.G., Z.C., K.J.M., Y.S., Z.T., S.T., A.H., and M.S. fabricated microparticles and studied release kinetics. X.L., L.M., W.G., Z.C., Y.S., Y.H., T.G., and Q.H. performed animal experiments. X.L., L.M., and K.S. performed flow cytometry experiments and analyzed the data. X.L., L.M., and A.J. analyzed the data and wrote the manuscript. R.L., D.G.A., and A.J. supervised the study. All authors discussed the results and edited the manuscript.

**Competing interests:** D.G.A. is a consultant with Translate Bio, Sigilon Therapeutics, and Verseau Therapeutics. X.L., R.L., and A.J. are inventors on patent application (Micromolded particles for cancer treatment, 21543) held by the Massachusetts Institute of Technology that covers the use of core-shell technology for treating cancer. K.M., A.J., and R.L. are inventors on patent applications (US 62/558,172 and US 14/572,631) submitted by the Massachusetts Institute of Technology that cover the core-shell particle fabrication technology. For a list of entities with which R.L. is involved, compensated, or uncompensated, see www.dropbox.com/s/yc3xb5s2s9v7x/Rev%20Langer%20COI.pdf?dl=0. All other authors declare that they have no competing interests. **Data and materials availability:** All data associated with this study are present in the paper or the Supplementary Materials.

Submitted 27 September 2019
Resubmitted 6 March 2020
Accepted 29 June 2020
Published 12 August 2020
10.1126/scitranslmed.aaz6606

Engineered PLGA microparticles for long-term, pulsatile release of STING agonist for cancer immunotherapy

Xueguang Lu, Lei Mia, Wenting Gao, Ziqi Chen, Kevin J. McHugh, Yehui Sun, Zachary Tochka, Stephanie Tomasic, Kaitlyn Sattler, Alain Hyacinthe, Yuxuan Huang, Tyler Graf, Quanyin Hu, Morteza Sarmadi, Robert Langer, Daniel G. Anderson, Ana Jaklenec

Sci. Transl. Med., 12 (556), eaaz6606. • DOI: 10.1126/scitranslmed.aaz6606

Squaring away tumors
Cancer immunotherapy has been achieving increasing prominence in recent years, but many patients' tumors still do not respond to existing immunotherapy modalities. One approach that offers promise is activation of the stimulator of interferon gene (STING) pathway, which can promote immune responses within the tumor microenvironment. STING agonists have shown promising results in mice and people, but they require multiple intratumoral injections, which are impractical for many tumors. Lu et al. designed box-shaped microparticles filled with STING agonist and optimized them to release the agonist at appropriate time intervals after being injected into a tumor just once, showing promising results in multiple mouse models of cancer.

View the article online
https://www.science.org/doi/10.1126/scitranslmed.aaz6606
Permissions
https://www.science.org/help/reprints-and-permissions