Localized Controlled Delivery of Gemcitabine via Microsol Electrospun Fibers to Prevent Pancreatic Cancer Recurrence

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The low radical surgery rate of pancreatic cancer leads to increased local recurrence and poor prognosis. Gemcitabine (GEM) is the preferred chemotherapeutic for pancreatic cancer. However, systemic chemotherapy with GEM has reached a bottleneck due to its serious side effects after frequent injections. In this study, GEM is successfully enwrapped into electrospun fibers via microsol electrospinning technology to form a stable core–shell fibrous structure. The GEM release rate can be adjusted by altering the thickness of the hyaluronan-sol inner fiber and the quantity of loaded GEM, and the release can be sustained for as long as three weeks. In vitro assays show that these electrospun fibers effectively inhibit pancreatic cancer cells and promote apoptosis. In vivo studies show that the fibrous membranes are better for inhibiting the growth of residual tumors than that of integrated tumors. Furthermore, immunohistochemistry results show that GEM-loaded fibers promote a higher cell apoptosis rate than does systemically injected GEM in residual tumors. In addition, the local delivery of GEM with fibers significantly reduces liver toxicity. In summary, a core–shell electrospun fiber for the controlled and localized delivery of GEM, which greatly improves the treatment of residual tumors and prevents pancreatic tumor recurrence, is developed.

1. Introduction

Pancreatic ductal adenocarcinoma (PDAC) is one of the most lethal cancers worldwide; the overall five-year survival rate is less than 5%.[1] Currently, radical surgery is the most effective way to cure PDAC. Approximately 20% of PDAC patients can receive surgery when diagnosed.[2] However, due to the low rate of negative surgical margins, 80% of patients experience cancer recurrence within two years, which is a notable challenge facing the prognosis of PDAC patients.[3]

Adjuvant chemotherapy after surgery decreases the risk of recurrence and improves prognosis in only a small fraction of PDAC patients.[4] The main bottleneck is the many disadvantages of antitumor drugs, including low bioavailability and strong side effects due to their nonspecific biodistribution.[2,5] The compactness of cancerous mesenchymal cells in pancreatic cancer also acts as a physiological barrier that prevents drugs from flowing into cancerous parenchymal cells, thereby limiting the effective treatment of PDAC.[6] Gemcitabine (GEM), a chemotherapeutic, is one of the best drugs for PDAC patients.[7] Anemia, leukopenia, neutropenia, thrombocytopenia, and kidney and liver toxicity are the most common side effects of GEM.[8] As a water-soluble chemotherapeutic agent, GEM primarily relies on intravenous administration,
but only a small portion of the drug reaches to the pancreatic cancer through blood circulation. Since pancreatic cancer has the characteristics of wrapped fibrous tissue and sparse blood vessels within the tumor, drug delivery through blood circulation has low efficient, greatly reducing the ability of the drug to kill pancreatic cancer cells. Therefore, developing a more efficient way to enhance the effectiveness of chemotherapeutics and decrease the side effect of drugs to improve the prognosis of pancreatic cancer patients after surgery is urgent.

Polymer-based local drug delivery systems are promising for loading antitumor therapeutics for cancer treatment. These systems promote the bioavailability of antitumor drugs, thereby avoiding nonspecific biodistribution, increasing the accuracy of drug targeting, and decreasing the side effects of drugs. Particularly, biodegradable polymer-based drug-loaded materials can be used to control and sustain drug release over a long period of time. For example, electrospinning is a simple technique for producing drug-loaded polymers. Electrospun fibers have a larger surface area for promoting drug loading efficiency and a 3D porous structure that can prevent drug diffusion. In addition, electrospun fibers are advantageous, because they can be transformed into any shape. Therefore, electrospun fibrous membranes can be conveniently implanted into the tumor resection margins for the local delivery of chemotherapeutics and, ultimately, patient treatment.

Water-soluble drugs can be loaded into fibers through emulsion electrospinning, coaxial electrospinning, and grafting the drugs through surface functionalization of the fibers. Although emulsion electrospinning has the disadvantage of low drug loading efficiency and a large amount of surfactant addition, surface functionalized electrospun fibers can graft water-soluble drugs on the surface of fibers and prevent rapid drug release. Coaxial electrospinning can make electrospun fibers with a core–shell structure, and water-soluble drugs can be wrapped in the core of electrospun fibers. Another issue with this technique is that the water phase core is instable, therefore drug release is too fast. In addition, due to the poor stability of the core–shell structure, adjusting the drug release rate through the internal structure of the core–shell fibers is difficult. Hydrosol is a water-retaining material that fully dissolves water-soluble drugs and creates an interface to prevent drug diffusion into organic solvents, thereby acting as an excellent drug carrier. The combination of emulsion electrospinning and hydrosol for loading water-soluble drugs may achieve high drug loading capacity and controllable drug release with a sustained release profile.

The aim of this study is to further explore a safe, efficient way to improve the survival of PDAC patients after surgery using as a model drug the water-soluble GEM and electrospun fibrous scaffolds comprising poly(l-lactide) (PLA) and hyaluronic acid (HA, approved by the Food and Drug Administration, FDA) as a model drug. In this study, we apply hydrosol and microsol electrospinning technology to load GEM and develop a GEM-loaded PLA electrospun fibrous membrane, which controls GEM release through changing the thickness of the inner-hydrosol in the fiber. The biocompatibility and drug release behavior of the material are evaluated in vitro, and the cytotoxicity and ability of this material to inhibit pancreatic cancer are investigated in vitro in a nude mouse xenograft tumor and residual tumor models.

2. Results

2.1. Characterization of Fibrous Scaffolds and GEM Release

First, we prepared hyaluronic acid hydrosol and used it to dissolve GEM to form GEM-HA-sol. Next, GEM-HA-sol was dispersed into PLA solution to form the electrospinning solution through emulsification during which sol particles were dispersed. Finally, sol-electrospinning was used to construct the electrospun fibrous membrane with a stable core–shell fibrous structure and loaded with GEM (Figure 1A). We prepared electrospun fibrous membranes containing various amounts of GEM (denoted GEM1@PLA-HA, GEM2@PLA-HA, and GEM3@PLA-HA) with different contents of sol (denoted GEM1@PLA-HA, GEM1@PLA-HA2, and GEM1@PLA-HA3). Figure 1B shows the morphology of the electrospun fibrous membrane, and no HA-sol particles were observed on the surface of the fiber. The diameters of PLA, GEM1@PLA-HA, GEM2@PLA-HA, GEM3@PLA-HA, GEM1@PLA-HA2, and GEM1@PLA-HA3 fibers were 1.15 ± 0.38, 1.26 ± 0.35, 1.32 ± 0.46, 1.41 ± 0.33, 1.30 ± 0.47, and 1.21 ± 0.39 µm, respectively, and none of the membranes showed significant differences with different quantities of GEM.

The fibrous morphology was further evaluated with transmission electron microscopy (TEM), and a clear core–shell structure was observed for HA-sol-PLA. The inner fibers formed by the HA-microgel had diameters of 0.37 ± 0.07, 0.48 ± 0.11, 0.39 ± 0.09, 0.30 ± 0.08, and 0.24 ± 0.06 µm for GEM1@PLA-HA, GEM2@PLA-HA, GEM3@PLA-HA, GEM1@PLA-HA2, and GEM1@PLA-HA3, respectively. Figure 1B shows that the outer diameter of GEM@PLA-HA was larger than that of PLA, indicating that GEM-sol loading slightly affected the PLA fiber diameter (Table 1). As shown in Table 1, with the same HA-sol content, increasing the amount of GEM content did not have a clear effect on the core and shell diameter of the fiber. Both core and shell diameters decreased with a decrease in HA-sol content. Therefore, HA-sol not GEM played a dominant role in affecting the core and shell diameter of the fiber.

The surface wettability of the electrospun membranes was measured by water contact angle (WCA), and the WCAs were 128.3 ± 2.7°, 130.4 ± 3.3°, 127.2 ± 3.18°, 128.57 ± 3.5°, 131.7 ± 4.3°, and 130.1 ± 4.3° for PLA, GEM1@PLA-HA, PLA-GME2, GEM3@PLA-HA, GEM1@PLA-HA2, and GEM1@PLA-HA3, respectively. The wettability of all electrospun membranes revealed a hydrophobic surface, and there were no significant differences among the different groups. Therefore, the fibrous membranes showed poor water permeability.

Subsequently, we examined the release behavior of GEM from the fibers containing different amounts of GEM and HA-sol (Figure 1C). With the increased amount of GEM in HA-sol, the release rate of GEM from the electrospun fibers increased (Figure 1C). With increased GEM loading, the fibrous membrane showed an initial burst release of GEM of 21.4, 25.7, and 29.8% within 1 d (Figure 1C). GEM3@PLA-HA released GEM over more than two weeks, whereas GEM2@PLA-HA and GEM1@PLA-HA released GEM for longer than three weeks.

In addition, for the same drug loading content, GEM1, the drug release further decreased by reducing the amount of HA-sol in the fibers. With more HA-sol content in the fibers, the
GEM release rate increased (Figure 1C). The initial drug release from GEM1@PLA-HA, GEM1@PLA-HA2, and GEM1@PLA-HA3 was 21.4, 19.7, and 6.8%, respectively, within 1 d. On the 26th d, the content of GEM released from GEM1@PLA-HA was only two-thirds of that released from GEM1@PLA-HA. Electrospun fibers swell in water, and drugs and sol molecules are gradually released from the fiber by diffusion.[22] The early burst of drug release was mainly caused by the diffusion of the

Figure 1. A) Schematic illustration of the process of constructing GEM-loaded PLA electrospun fibrous membranes with sol-electrospinning technology. B) SEM images of PLA (a), GEM1@PLA-HA (b), GEM2@PLA-HA (c), GEM3@PLA-HA (d), GEM1@PLA-HA2 (e), and GEM1@PLA-HA3 (f) fibers (inset pictures are TEM images of the fiber to show the core–shell). C) GEM release profiles of fibers containing different drug content (GEM1@PLA-HA, GEM2@PLA-HA, and GEM3@PLA-HA) and different sol content (GEM1@PLA-HA, GEM1@PLA-HA2, and GEM1@PLA-HA3) (n = 3). Data represent the mean ± SD.
drug out of HA molecules that created pores in the core. Based on TEM images, GEM1@PLA-HA3 had a thicker shell than did GEM1@PLA-HA (0.97 μm vs 0.89 μm); this difference in shell thickness might account for the delay of HA and drug diffusion out of the fibers. This result is in good agreement with the findings of a previous study by Wen et al., who showed that increasing the shell thickness can delay the drug release rate.\[23\] Thus, by adjusting the GEM- and HA sol content, it is possible to control GEM release in fibers and tailor them to different kinds of biomedical applications.

### 2.2. In Vitro Compatibility of PLA Membrane

Fibroblast cells play an important role in the progression of pancreatic cancer. These cells promote the proliferation and metastasis of pancreatic cancer cells and can also induce GEM resistance in pancreatic cancer. Hence, we investigated whether a PLA-HA electrospun membrane would affect fibroblast cells and pancreatic cancer cells.\[23\] We first performed a CCK8 assay to investigate cell growth on PLA or PLA-HA electrospun membranes to evaluate whether they were biocompatible. PANC-1 and 3T3 cells (mouse fibroblast cell line) were inoculated on PLA and PLA-HA electrospun membranes in 24-well plates, and cell proliferation was evaluated on days 1 and 3. Comparing with controls, PANC-1 and 3T3 cells grown on electrospun PLA showed no change in cell viability; however, cells grown on HA-PGA showed slightly higher viability than did those grown on the control and PLA because HA promotes the adhesive capacity of cells (Figure 2A). This result indicates that HA-PLA and PLA electrospun membranes displayed no cytotoxicity to cells. Furthermore, the adhesive morphologies of 3T3 and PANC-1 cells growth on PLA or PLA-HA electrospun membranes on days 1 and 3 were detected by phalloidin staining to investigate whether the PLA-HA membrane affects fibroblasts or pancreatic cancer cells. We investigated this aspect, because changes in morphology will affect cell function in a certain respect; pancreatic cancer cells change from a polygon shape into a shuttle shape, which promotes metastasis.\[25\] Moreover, the shuttle shape of fibroblasts might affect the growth rate.\[26\] Figure 2B shows that the cell size decreased and the outline of the cells changed when inoculated on PLA or electrospun PLA-HA, which might be due to the 3D structure displayed in the 2D photo. The morphology of fibroblasts remained in a shuttle shape and pancreatic cancer cells remain in a polygonal shape. In addition, no cell morphology change was observed between PLA and PLA-HA in scanning electron microscopy (SEM) (Figure 2C). The above results demonstrate that PLA and PLA-HA electrospun fibrous membranes are biocompatible and can be used for biomedical applications.

### 2.3. GEM-Loaded PLA-HA Membranes In Vitro

The anticancer effect of GEM-loaded PLA-HA membranes in vitro was studied by a 3-[4,5-dimethylthiazol-2-yl]-2,5 diphenyl tetrazolium bromide (MTT) assay. The three GEM-loaded PLA-HA membranes contained different quantities of GEM. In Figure 1C, GEM release experiments showed that GEM3@PLA-HA was loaded with the maximum amount of GEM, followed by GEM2@PLA-HA and GEM1@PLA-HA. Figure 3A shows the cell viability of BxPC-3 and PANC-1 cells treated with PLA-control (no GEM-loaded PLA membranes), GEM1@PLA-HA, GEM2@PLA-HA, or GEM3@PLA-HA. Compared with the control group, the PLA and PLA-HA groups showed no cytotoxicity when incubated with PANC-1 and BxPC-3 cells. For the other groups, GEM3@PLA-HA showed higher antiproliferation activity than did GEM2@PLA-HA and GEM1@PLA-HA against PANC-1 and BxPC-3 cells (P < 0.001), because GEM3@PLA-HA loaded and released more GEM than did GEM2@PLA-HA and GEM1@PLA-HA. In addition, the PANC-1 cell viability decreased with an increased incubation time with PLA-HA-GEM; cell viability almost decreased to zero at day 7, the results suggested that the GEM-loaded membranes needed time to release more gemcitabine to effectively kill pancreatic cancer cells. Since GEM induces apoptosis in cancer cells through interfering with DNA replication,\[27\] we performed an Annexin V/propidium iodide (PI) double staining to determine which kind of material possessed the greatest ability to induce cell apoptosis in BxPC-3 and PANC-1 cells. GEM3@PLA-HA had the strongest effect on apoptosis induction, which was in good agreement with the MTT assay (Figure 3B). Thus, cell cytotoxicity was positively associated with the loaded amount of GEM in the fibers and the treatment time, and we selected GEM3@PLA-HA for subsequent in vivo studies.

### 2.4. In Vivo Anti-Integrated Tumor Capability

A xenograft tumor model of pancreatic cancer was used to evaluate the antitumor ability of GEM3@PLA-HA in vivo. We implanted a GEM3@PLA-HA electrospun membrane into the tumor surface as depicted in Figure 4A. The tumor volume and tumor weight of mice that received GEM3@PLA-HA...
local chemotherapy were almost the same as those of mice that received GEM chemotherapy (GEM group) (Figure 4B–D). Weight change was an easily measured indicator of the side effect of chemotherapy. We noticed that the weights of mice in the GEM group were significantly lower than those of mice in the PLA-GEM3 group (Figure 4E). Systemic administration of GEM caused side effects because of the whole biodistribution of GEM unlike local chemotherapy with PLA-GEM3, which minimized the side effects of the drug, since GEM mainly acted at the tumor site instead of being distributed throughout the whole body.

We also performed immunohistochemical (IHC) staining to evaluate the proliferation and apoptosis status in the tumor tissues of the two groups (Figure 4F). The staining levels of the proliferation index Ki-67 and the antiapoptosis index Bcl-2 were similar between the two groups, and the proapoptosis index cleaved caspase-3 also exhibited the same tendency (Figure 4G). These results demonstrate that the efficiency of local chemotherapy with a GEM3@PLA-HA electrospun membrane was as good as that of systemic chemotherapy with GEM, while extensively minimizing the side effects of the drug.

2.5. In Vivo Antiresidual Tumor Effect

Radical surgery for pancreatic cancer is difficult to achieve; therefore, the local tumor recurrence rate after surgery remains high, and compared with an integrated tumor, a residual tumor is often more aggressive and difficult to treat. In this study, we further investigated the anti-residual xenograft tumor capability of GEM3@PLA-HA. The GEM3@PLA-HA electrospun membrane (Figure 5A) was implanted into the residual tumor and compared with systemic chemotherapy with GEM. Interestingly, the anti-tumor capability of GEM3@PLA-HA was superior to that of GEM, and compared with the control, GEM also showed an antitumor effect (Figure 5B). As shown in Figure 5C,D, the tumor volume \( (P = 0.0188) \) and tumor weight \( (P = 0.0134) \) of the GEM3@PLA-HA group were significantly lower than those of the control and GEM groups.

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Figure 2. A) Cell viability of 3T3 and PANC-1 cells grown on PLA or PLA-HA electrospun membranes for 1 and 3 d \( (n = 4) \). B) Morphology of 3T3 and PANC-1 cells grown on PLA or PLA-HA electrospun membranes or culture plates for 1 and 3 d of incubation. Red scale bars indicate 100 \( \mu m \). C) Morphology of 3T3 and PANC-1 cells grown on PLA or PLA-HA electrospun membranes for 3 d were imaged by SEM. Scale bars indicate 50 \( \mu m \). Data represent the mean \( \pm SD \), and ** indicates \( p < 0.01 \).
of the GEM group and control group. The mouse weight of the GEM3@PLA-HA group was higher than that of the GEM group and control group (Figure 6E). Furthermore, we detected the expression of Ki-67, Bcl-2, and cleaved caspase-3 with IHC (Figure 6F). The proliferation index Ki-67 and the antiapoptosis index Bcl-2 were lower in the GEM3@PLA-HA group, and the apoptosis index cleaved caspase-3 was higher in the GEM3@PLA-HA group (Figure 6G). All these results demonstrate that local GEM delivery with PLA-HA was more effective than systemic drug delivery for treating residual tumors when tumor integrity was destroyed. Thus, the deficiency in the protective membrane of the tumor benefited the GEM delivery from the GEM-loaded electrospun membranes.

2.6. Side Effect Reduction of GEM-Loaded Electrospun Fibers

Only a small amount of GEM can enter the tumor tissue when systemic chemotherapy is performed in patients, and most of the GEM is absorbed by normal tissues. Therefore, GEM leads to serious side effects in pancreatic cancer patients, and liver toxicity is particularly obvious. Our study found that...
GEM-loaded electrospun fibrous membranes reduced the side effects of GEM on liver toxicity. As shown in Figure 6, nude mice receiving systemic chemotherapy with GEM had liver damage and hematoxylin and eosin (HE) staining of the liver showed liver cell necrosis, while no necrosis was observed for the mice receiving GEM3@PLA-HA localized chemotherapy or saline alone (Figure 6A). We also measured blood plasma aspartate aminotransferase (AST) and alanine aminotransferase (ALT), which reflect the liver function of nude mice treated with GEM or GEM3@PLA-HA. The AST and ALT levels of mice in the GEM group were significantly higher than those in the GEM3@PLA-HA group (Figure 6B). These results suggest that the localized delivery of GEM with PLA-HA electrospun membranes to pancreatic cancer could decrease the toxicity of GEM, which is clinically important for the treatment of PDAC.

2.7. Mechanism for Drug Release and Chemotherapy

Currently, the most effective treatment for pancreatic cancer is surgery, but only approximately 20% of pancreatic cancer patients can receive surgery, because most patients are in the advanced
stage when diagnosed. However, only 20% of these patients can receive radical surgery, and the remaining 80% who undergo pancreatic cancer surgery might have local tumor cell residue.[29] Systemic chemotherapy with GEM is a way to improve the survival of pancreatic cancer patients.[28,30] Due to its special pathologic characteristics, pancreatic cancer has a poor blood supply; pancreatic cancer cells are surrounded by abundant fibrous connective tissue that compresses tumor blood vessels and cause chemotherapeutics to concentrate; compared with other types of solid tumors, pancreatic tumors exhibit less blood flow into the cancerous tissue through the circulation.[10a,31]

In view of the above situation, we used GEM-loaded PLA-HA electrospun fibrous membranes with a core–shell structure using sol-electrospinning technology to provide sustained release of GEM at the site of the tumor. The release time of GEM could be adjusted by the thickness of core and sustained over three weeks. Localized drug delivery has been proven to effectively inhibit the growth of tumors in cervical,[12b] colorectal,[32] and liver cancers,[11b] it also reduces the side effects of chemotherapeutics.

As illustrated in Figure 7, one of the physiological features of PDAC is that the tumor is tightly packed and surrounded with fibrous tissue, which presents a great challenge for drug penetration into tumor.[33] Thus, in the xenograft pancreatic tumor model, GEM@PLA-HA did not show better efficiency than systemically administered GEM (Figure 7A). Localized delivery of the drug to a particular body compartment, thereby lowering the systemic drug level, suggests that localized delivery drug could penetrate to vessels.[34] However, the fibrous tissue in pancreatic cancer not only prevented the gemcitabine released by gemcitabine-loaded membranes from penetrating to tumor and tumor vessels, but also compressed the tumor vessels which reduced the concentration of gemcitabine. Therefore, after surgery, with the loss of tumor integrity, the fibrous tissue around the tumor is also destroyed. The destruction of the fibrous matrix in pancreatic cancer could promote GEM penetration into the tumor tissue.[10b,35] Thus, the gemcitabine locally released from GEM-loaded membranes was more capable of penetrating tumor tissues (Figure 7B). GEM@PLA-HA was combined with...
sol-electrospinning technology and a GEM-loaded hyaluronic acid hydrogel, which showed significantly better efficacy in the treatment of residual tumors than did systemically administered GEM. In addition, when tumor was completely resected, the GEM-loaded membranes could kill the pancreatic cancer cells which might scattered in the surgical site.

PLA-HA electrospun fibers had good biocompatibility, and another great advantage of using it to load GEM is reducing the side effects. The localized and sustained release of GEM minimized the systemic biodistribution of the drug and greatly reduced the spread of GEM to healthy tissues, such as the liver. In our study, mouse bodyweight was clearly higher in the GEM@PLA-HA group than in the GEM group, indicating the reduced side effects of the drug. GEM@PLA-HA also clearly reduced the liver injury and liver toxicity. In addition, it is very easy to attach GEM@PLA-HA to the tumor bed after surgery because the shape of the electrospun fibrous membrane could be tailored. Therefore, for patients with pancreatic cancer after

**Figure 6.** A) HE staining of the liver after treatment with GEM3@PLA-HA and GEM. Black arrows showed the necrosis of liver cells. Black scale bars indicate 200 µm. B) AST and ALT levels of mice treated with GEM3@PLA-HA or GEM. The control group included normal nude mice without tumors (n = 5). Data represent the mean ± SD. ** indicates P < 0.01 and *** indicates P < 0.001.

**Figure 7.** Schematic illustration of the GEM-loaded PLA electrospun membrane on pancreatic cancer for local chemotherapy. A) Before surgery, tumor integrity is maintained, and the fibrous tissue around the tumor could prevent GEM from penetrating tumor tissues. B) After surgery, when tumor is incompletely resected, tumor integrity is destroyed, and the fibrous tissue around the tumor is also destroyed, therefore GEM released from the GEM-loaded PLA electrospun fibrous membranes can easily penetrate tumor tissues. C) After surgery, when tumor is completely resected, and GEM released from the GEM-loaded PLA electrospun fibrous membranes can prevent the local tumor recurrence by killing the pancreatic cancer cells which may be existed in the surgical site.
surgery, the membranes could be attached to the local surgical site and continue to release GEM to kill possible residual tumor cells and prevent tumor recurrence. However, GEM-loaded fibrous membranes as a therapeutic tool for localized treatment of pancreatic cancer after surgery also had drawbacks. For example, the membranes must be placed into the surgical field only one time, resulting in limited for long-term localized treatment of pancreatic cancer after surgery. In addition, this kind of localized treatment could not significantly increase the concentration of a chemotherapeutic agent in the blood, thus it might not work well for pancreatic cancer cells that might have already metastasized into the blood circulation, combined with systemic chemotherapy of GEM might help to resolve the deficiency of localized chemotherapy, and they could collaborate to against pancreatic cancer.

3. Conclusions

GEM was dispersed in hyaluronic acid hydrogel and subsequently enraptured into PLA-HA electrospun fibers via sol-electrospinning technology to form stable core–shell fiber structures. The GEM loading content can be easily adjusted by dispersing different amounts of GEM in HA sol-gel, and the release rate is controlled by the thickness of the sol-gel core in the PLA fiber. Sustained GEM release lasted over three weeks, which facilitated its in vivo application for localized drug delivery. GEM@PLA-HA electrospun fibrous membranes were easily implanted onto the surface of tumors; however, the antitumor effect was not superior to GEM treatment in animal models with an integrated xenograft pancreatic cancer tumor, this difference might be due to the fibrous tissue around the tumor limiting GEM penetration into tumor tissues when the tumor was integrated. However, compared with intravenously administered GEM, the GEM@PLA-HA electrospun fibrous membrane was more efficient inhibiting the growth of residual tumors. Furthermore, GEM@PLA-HA significantly reduced the adverse side effects of GEM, including reduced liver toxicity. Therefore, this kind of local implanted drug-loaded scaffold has potential in the prevention and treatment of tumor recurrence after surgery. Overall, in this study, we developed a promising platform of GEM@PLA-HA electrospun fibrous membranes for local chemotherapy of pancreatic cancer.

4. Experimental Section

**Materials:** PLA ($M_w = 140$ kDa, $M_m/M_w = 1.85$) was purchased from Jinan Daigang Co. (Jinan, China). Fermentation-derived HA (sodium salt, $M_w = 0.5$ MDa) was purchased from Yuancheng Technology Co. (Wuhan, China) and used without further purification. GEM was purchased from Eli Lilly and Company (USA). All solvents were purchased from Jiangsu Chemical Instrument Co., Baoding, Hebei, China). The GEM-loaded PLA electrospun fibrous membranes obtained through microfluid electrospinning technology. PLA solution was used as the control and prepared as follows: 1.0 g PLA was added to 8.0 g DCM and stirred completely, and then 2.0 g DMF was added to the mixture. The electrospinning equipment had a high voltage station (Tianjing High Voltage Power Supply Co., Tianjing, China) set to 10–25 kV. The electrospinning solution was added by a 5 mL syringe attached to a metal needle with a 0.6 mm diameter and was electropun at the flow rate of 0.2–6 mL h$^{-1}$ controlled by a microinject pump (Lange Medical Instrument Co., Baoding, Hebei, China). The GEM-loaded PLA electrospun fibrous membranes were obtained through microfluid electrospinning technology. PLA solution was used as the control and prepared as follows: 1.0 g PLA was added to 8.0 g of DCM and stirred completely, and 2.0 g of DMF was added to the mixture. All of the fibrous scaffolds were lyophilized overnight to remove the solvent and water residue and stored at 4 °C in a desiccator until further use.

Fiber morphology were examined by field-emission scanning electron microscopy (FESEM) using a Hitachi 4800 system with an acceleration voltage of 3.0 kV. Before SEM observation, the fibers were sputter-coated with platinum. TEM images of an individual fiber’s inner structure were obtained by a Hitachi HT7700 instrument at 120 kV. The static WCAs of microfibrous membranes were measured using a contact angle analyzer (DSA255, Data Physics Corporation).

**In Vitro Drug Release Tests:** The test specimens (20 × 20 mm, ~350 μm thick; total mass = 50 mg) of PLA fibrous membranes were immersed in 10 mL phosphate buffered saline (PBS, 154 × 10$^{-3}$ M, pH 7.4). The suspension was placed in a 50 mL centrifugal tube and maintained in a thermostat shaker (Thermo, USA) at 37 °C with 100 cycles min$^{-1}$. Next, 2.0 mL release buffer was removed from the centrifugal tube and 2.0 mL fresh PBS was replaced at different time points. The quantity of released GEM from PLA fibrous membranes in release buffer was measured by high-performance liquid chromatography (HPLC, Waters) with a mobile phase of 0.05 M ammonium acetate buffer and methanol (90:10%) at a flow rate of 1 mL min$^{-1}$, a detection wavelength of 268 nm, and a column (C18, 250 mm × 4.6 mm length, 5 μm for filler size) temperature of 30 °C. Based on the initial weight of GEM incorporated into the electrospun fibrous membrane, the percentage of released drug from the membranes was calculated.

**Cell Culture and Reagents:** Human PDAC cell lines (PANC-1 and BxPC-3) and mouse fibroblast 3T3 cells were purchased from the Chinese Academy of Sciences. PANC-1 and 3T3 cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM), and BxPC-3 cells were cultured in Roswell Park Memorial Institute-1640 (RPMI-1640) medium. All culture media contained 10% fetal bovine serum (Gibco), 100 U mL$^{-1}$ penicillin, and 100 μg mL$^{-1}$ streptomycin (Sigma). Cells were cultured at 37 °C in a humidified atmosphere containing 5% CO$_2$.

**Phalloidin Staining:** Approximately 2 × 10$^4$ cells were seeded on PLA membranes in 48-well plates for 1 and 3 d. For fluorescence microscope observations, cells were washed with PBS, fixed in 4% formaldehyde solution for 30 min, and washed twice with PBS. subsequently, the cells were incubated with 0.1% Triton X-100 for 5 min and phalloidin solution (Yearsen, China) for 30 min, washed twice with PBS, and finally incubated in 4′,6-diamidino-2-phenylindole (DAPI) solution for 10 min in the dark.

**Cell Proliferation:** Cell viability was evaluated with a cell counting kit (CCK-8, Dojindo, Japan). Fibrous materials were cut into the proper size to cover the wells in 24-well plates and placed into the bottom of the wells with an iron ring to prevent floating in the culture medium. All materials were decontaminated under ultraviolet (UV) irradiation overnight. Approximately 4 × 10$^4$ 3T3 or PANC-1 cells were seeded on the PLA electrospun membranes in 24-well plates for culturing for 1 and 3 d and cells cultured without membranes were used as the control. After incubation, the wells were refreshed with new culture medium with 10% CCK8 solution and cultured at 37 °C for another 2 h to allow the reaction to proceed. Finally, 100 μL reaction medium was transferred into 96-well plates for absorbance measurements at a wavelength of 450 nm with a microplate reader.

**Cell Viability Assay:** The cytotoxicity of a GEM-loaded electrospun membrane against PANC-1 and BxPC-3 cells was assessed by a MTT
assay (Yearsen, China). Approximately $4 \times 10^5$ PANC-1 and BxPC-3 cells were cultured in a 24-well plate with culture medium overnight. Wells were divided into the following groups: (1) control group, cells cultured in 24-well plates; (2) PLA control group, 5 × 5 mm PLA membranes were added in the bottom of the wells; (3) PLA-HA control group, 5 × 5 mm PLA-HA membranes were added; (4) GEM1@PLA-HA group, 5 × 5 mm GEM1@PLA-HA membrane were added; (5) GEM2@PLA-HA, 5 × 5 mm GEM2@PLA-HA membranes were added; and (6) GEM3@PLA-HA, 5 × 5 mm GEM3@PLA-HA membranes were added. All materials were decontaminated under UV irradiation overnight before the experiments. After culturing for 3, 5, and 7 d, MTT assays were performed for cell viability as measured by a microplate reader (MULTISKAN MK3, Thermo Scientific) at the wavelength of 490 nm.

Apoptosis Assay: PANC-1 and BxPC-3 cell apoptosis induced by GEM-loaded PLA electrospun membranes was evaluated by flow cytometry (FCM) with Annexin V-fluorescein isothiocyanate (FITC)/PI double staining. Approximately $4 \times 10^5$ PANC-1 or BxPC-3 cells were seeded on six-well plates and cultured overnight, and GEM-loaded PLA electrospun membranes containing different amounts of GEM (GEM1@PLA-HA, GEM2@PLA-HA, and GEM3@PLA-HA) or PLA electrospun membranes (PLA-control) were placed into culture medium for 3 d of cell culture; all materials were cut into 1 × 1 cm pieces and decontaminated under UV irradiation for 24 h. Cells were collected, washed with cold PBS and centrifuged at 1500 × g for 5 min. Next, the cells were incubated with Annexin V-FITC and PI solution (Yearsen, China) for 10 min and analyzed by FCM.

Xenograft Tumor Model: Animal experiments were permitted by the Ethical Review Committee of the Sixth Affiliated Hospital of Shanghai Jiaotong University. Six-to-eight-week-old male BALB/c athymic nude mice were purchased from Shanghai Si Lai Ke Laboratory Animal Co., Ltd. China. Approximately $1 \times 10^5$ PANC-1 cells were subcutaneously inoculated into the right thigh of nude mice (n = 5). After the tumors grew to 70 mm$^3$, the mice were divided into two groups. One group received peritoneal administration of GEM (100 mg kg$^{-1}$) and was denoted the GEM group. For the other group, a small incision was made on the skin to expose the tumor, and GEM3@PLA-HA membranes (cut as 2 × 2 cm$^2$) were placed on the tumor surface; this group was denoted the GEM3@PLA-HA group. Mice were weighed, and the tumor volume was calculated by caliper every 3 d as follows: $V = 0.5 \times (length \times width)^2$. After the mice were euthanized, tumors were collected for further studies; tumors were embedded in paraffin and cut into 5-μm-thick sections for IHC analysis.

Residual Tumor Model: Approximately $1 \times 10^5$ PANC-1 cells were subcutaneously inoculated into the right flank of nude mice (n = 5). After the tumor grew to 100 mm$^3$, a small incision was made on the mouse skin to expose the tumor, and three-quarters of the tumor was removed (~4 mm × 4 mm of tumor tissue remained), as in a previous study.[26] Mice were divided into three groups, including the control, GEM, and GEM3@PLA-HA groups. Mice in the control group were intraperitoneally injected with saline; mice in the GEM3@PLA-HA group were locally implanted with GEM3@PLA-HA membrane; the mice in GEM group were intraperitoneally injected with GEM (100 mg kg$^{-1}$) twice a week. Before the materials were implanted, they were cut into 2 × 2 cm in size. Mice were weighed, and tumor volume was calculated as follows: $V = 0.5 \times (length \times width)^2$.[26] After the mice were euthanized, tumors and blood were collected for further analysis; tumors were embedded in paraffin and cut into 5-μm-thick sections for IHC analysis. Plasma was harvested from blood by centrifugation at 3000 rpm for 10 min. Next, liver function was tested by measuring AST and ALT.

IHC Staining Analysis: Tumor tissues were fixed in 4% paraformaldehyde, embedded in paraffin, cut into 5-μm-thick sections, deparaffinized with xylene, gradient-rehydrated with ethanol, and subjected to antigen-retrieval with citrate buffer at 95 °C for 15 min. Sections were later blocked with goat serum for 10 min and incubated with primary antibodies (Ki-67, caspase-3, and Bcl-2) for 30 min at room temperature. Next, the membrane was incubated with a goat anti-rabbit secondary antibody conjugated to horse radish peroxidase (HRP) for 10 min at 37 °C. The target marker was detected with 3,3′-diaminobenzidine (DAB) over 1–5 min. The nucleus was stained with hematoxylin. All staining was observed and photographed using a microscope (Nikon ECLIPSE 80i).

IHC Quantification: IHC staining for Bcl-2 was evaluated on the basis of staining intensity and staining proportion; at least five fields of 400× magnification were counted. The staining intensity was scored from grades 0 to 3 (0, absent; 1, weak; 2, moderate; and 3, intense). The staining proportion was scored from grades 1–3 (1, <10% of positive cells; 2, 10–49%; 3, >50%).[27] The IHC score was calculated by multiplying the scores of staining intensity by proportion. IHC staining for Ki-67 and cleaved caspase-3 was evaluated on the basis of the number of stained cells.

Statistical Analysis: Statistical Product and Service Solutions (SPSS) 18.0 software was applied for all statistical analysis. The results are presented as the mean ± standard deviations (SD). For samples that were used for MTT analysis, an n = 4 or 5 was used, for samples that were used for in vivo experiment, an n = 5 was applicable. Multiple group comparisons were performed by applying one-way ANOVA, and post hoc Bonferroni was performed to test the differences. Two-group comparisons were performed by using Student’s t-test. * p-value < 0.05, ** p-value < 0.01, and *** p-value < 0.001 represented statistically significant differences.

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Conflict of Interest

The authors declare no conflict of interest.

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