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Dual targeting nanoparticles for epilepsy therapy†

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For epilepsy therapy, one-third of the patients worldwide are resistant to antiepileptic drugs mainly due to the existence of the blood–brain barrier (BBB) that prevents the drugs from reaching the epileptic lesions. Here, we design a double targeting nanoparticle carrying lamotrigine (LTG) to cross the BBB and further concentrate at the neurons. We prepare the nanoparticles on a microfluidic chip by encapsulating LTG in poly(lactic-co-glycolic acid) (PLGA) to form a core (PL) and capping the core with a shell of lipids conjugated with the D-T7 peptide (targeting the BBB) and Tet1 peptide (targeting the neuron) to form D-T7/Tet1-lipids@PL nanoparticles (NPs). *In vitro* and *in vivo* experiments show that D-T7/Tet1-lipids@PL NPs have excellent neuron targeting, antiepileptic, and protecting effects. Our approach provides a new strategy for improving the therapeutic efficacy of existing antiepileptic drugs.

Introduction

Epilepsy is one of the most common neurological diseases. There are more than 70 million people worldwide suffering from epilepsy.¹ Epilepsy is usually caused by abnormally increased neuronal excitability and abnormal activation of brain neuronal circuits due to the increase of extra neuronal excitatory neurotransmitters or the decrease of inhibitory neurotransmitters.² Antiepileptic drugs (AEDs) are the main means of epilepsy treatment through various mechanisms.^{1,3} Although there are more than 25 AEDs,¹ about one-third of epilepsy patients are resistant to existing drugs and cannot completely control seizures.⁴ The main cause of drug resistance is that most of the AEDs cannot pass through the blood–brain barrier (BBB).⁵ Moreover, even if the drugs enter the brain, it is difficult to specifically target the diseased neurons. Thus, it is necessary to explore a strategy to effectively deliver AEDs to the brain and target the neurons. Nano-delivery systems have been widely used for improving the utilization of drugs.^{6–14} The use of nanocarriers to improve the BBB-

penetration efficiency of drugs has been extensively studied.^{15–19} However, there are few reported nanocarriers for delivering AEDs, and most of them are single-targeted nanocarriers, which still cause side effects. Recently, microfluidic technology has played important roles in many biomedical fields.^{20–22} In particular, microfluidic chips can easily synthesize NPs with uniform and small sizes,^{23–26} providing a platform for preparing nano-drugs.

Here, we propose a dual-targeting nanocarrier system to deliver lamotrigine (LTG) to the diseased neurons to treat epilepsy. LTG is a first line AED in clinics.^{4,27} However, it has low solubility in water and is easily metabolized in the liver. Thus, high doses or repeated administration are required to achieve therapeutic concentrations,^{28,29} which can cause side effects such as nausea, headache, blurred vision, dizziness, and ataxia. To optimize the pharmacological action of LTG and minimize its side effects, the dual-targeting nanocarrier system had two components: (i) D-form T7 (D-T7) peptide, the retro-inverse sequence of the T7 peptide, which showed a high binding affinity with transferrin (Tf) receptor (a major component of the BBB) and was effective in guiding drug delivery to the central nervous system (CNS),^{30–35} was designed to target the BBB. (ii) Tet1 peptide, which can specifically bind to sphingomyelin and ganglioside (G_{T1b} receptor) highly expressed on the surface of neurons.^{36–39} Although the ability of the T7 peptide, D-T7 peptide, and Tet1 peptide to target their respective targets and the combination of the T7 peptide and Tet1 peptide to treat Alzheimer's disease have been reported, the combined effects of the D-T7 peptide and Tet1 peptide to target the central nervous system (CNS) have not yet been attempted. The D-T7 peptide has a higher binding affinity to the Tf receptor than the T7 peptide for targeting the CNS, so the combined effects of the D-T7 peptide and Tet1 peptide to target the CNS need to be explored.^{32,33,38,39} We synthesized a dual-targeting delivery system on a two-step microfluidic chip which has been

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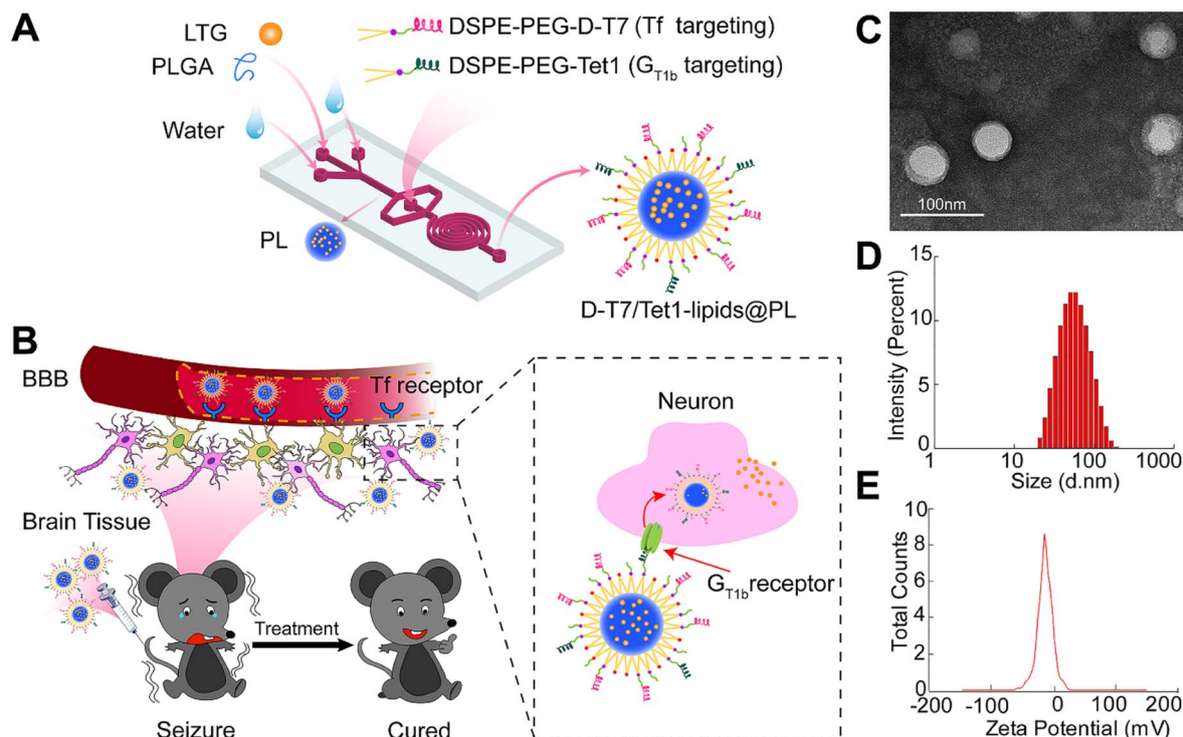


Fig. 1 Schematic diagram of the general design of the dual-targeting NPs for epilepsy therapy. (A) Preparation of D-T7/Tet1-lipids@PL using a microfluidic chip. (B) Administration of D-T7/Tet1-lipids@PL NPs to treat epilepsy. (C) TEM images of D-T7/Tet1-lipids@PL (2 : 1). (D and E) Size and zeta potential of D-T7/Tet1-lipids@PL (2 : 1) analyzed by DLS.

developed in the previous work.⁹ We encapsulated LTG in poly(lactic-co-glycolic acid, PLGA), a biodegradable and biocompatible polymer approved by the Food and Drug Administration (FDA), to form a core (PL) in the first step. We further coated a layer of lipids (DPPC, cholesterol, DSPE-PEG2k-D-T7, and DSPE-PEG2k-Tet1) as a shell to form a core-shell NP (D-T7/Tet1-lipids@PL). The co-modification of the D-T7 peptide and Tet1 peptide on the lipids@PL endowed the NPs with dual targeting capability to both the BBB and the neurons (Fig. 1B). We screened NPs with different D-T7/Tet1 concentration titrations. *In vitro* (BBB model) and *in vivo* (2 acute epilepsy models induced by kainic acid (KA) or pilocarpine and 1 chronic epilepsy model induced by KA, which are associated with human temporal lobe epilepsy.^{40,41}) experiments showed that when the molar ratio of the D-T7 peptide to Tet1 peptide was 2 : 1, D-T7/Tet1-lipids@PL NPs showed the best performance in accumulating in the brain, inhibiting the seizures and protecting the neurons. To the best of our knowledge, this is the first work on dual targeting delivery systems with multiple types of *in vivo* epilepsy models, which has important implications for the clinical treatment of human temporal lobe epilepsy.

Results and discussion

To modify D-T7 and Tet1 peptides on the lipids@PL, the peptides were coupled with DSPE-PEG-Mal through a Michael addition reaction (Fig. S1[†]), and the coupling was confirmed by mass spectrometry (Fig. S2[†]). A microfluidic chip was used to

synthesize D-T7/Tet1-lipids@PL with different D-T7/Tet1 ratios (1 : 2, 1 : 1, and 2 : 1) (Fig. 1A). As controls, lipids@PL, D-T7-lipids@PL, and Tet1-lipids@PL were synthesized on the same chip. Transmission electron microscopy (TEM) and dynamic light scattering (DLS) showed that the D-T7/Tet1-lipids@PL NPs with various D-T7/Tet1 ratios had a typical core-shell structure and hydrodynamic diameters ranging from 58.89 ± 0.88 to 71.86 ± 0.31 nm (Fig. 1C-E, S3A and B[†]). We selected D-T7/Tet1-lipids@PL (2 : 1) to characterize the physicochemical properties of the NPs. Compared to the size of NPs synthesized by a traditional method (124.30 ± 1.18 nm) (Fig. S3C and D[†]), the size of the NPs synthesized using the microfluidic chip was much smaller (68.93 ± 0.59 nm), and the synthesis speed was faster,³³ demonstrating the advantage of the microfluidic chip. We tested the stability of various NPs, which are all stable at 4 °C for 10 days (Fig. S4[†]). D-T7/Tet1-lipids@PL (2 : 1) had a LTG encapsulation efficiency of approximately 85.85%, and a cumulative release rate of more than 50% within 72 h (Fig. S5[†]), indicating a high encapsulation capability and an appropriate drug release profile. UV-vis spectra characterization indicated that DSPE-PEG-D-T7 and DSPE-PEG-Tet1 encapsulation efficiency in D-T7/Tet1-lipids@PL (2 : 1) was 83.33% and 71.34% respectively. The content of DSPE-PEG-D-T7 and DSPE-PEG-Tet1 in the D-T7/Tet1-lipids@PL (2 : 1) solution was 57.33 and $43.05 \mu\text{g mL}^{-1}$ respectively (Fig. S6A and B[†]).

We conducted experiments on the internalization of different DiI-labelled NPs in bEnd.3 cells which is a cell line from mice BBB. After 6 h of co-cultivation with the cells, the NPs



displayed different intensities of fluorescence signals (Fig. S7A and C†). D-T7/Tet1-lipids@PL (2 : 1) showed higher uptake efficiency compared to other groups except Tet1-lipids@PL (Fig. S7B and D†).

We verified the internalization of various DiI-labelled NPs in hippocampal neurons and the human neuroblastoma (SY5Y) cell line. SY5Y cells were used as a control group to verify if the nanoparticles have a good targeting effect on neurons. According to the results of the confocal microscopy experiments, the NPs displayed different intensities of fluorescence signals. Except for Tet1-lipids@PL, the uptake efficiency of D-T7/Tet1-lipids@PL (2 : 1) was higher than those of other groups (Fig. S8†). Quantitative data by flow cytometry revealed that among the experimental groups, D-T7/Tet1-lipids@PL (2 : 1) showed a higher level of internalization by the hippocampal neurons compared to other groups except Tet1-lipids@PL (Fig. S8, S9A and C†). This means that the Tet1 peptide plays a key role in targeting the neurons. Confocal microscopy images revealed that D-T7/Tet1-lipids@PL (2 : 1) exhibited the highest fluorescence intensity in the hippocampal neurons among all the experimental groups, demonstrating its high internalization efficiency by neurons (Fig. S10A and B†). All the NPs showed a lower level of internalization in the SY5Y cells than those in the hippocampal neurons (Fig. S9B, D, S10B and D†), indicating the high specificity of the nanoparticles.

We verified the penetration ability of different NPs through an *in vitro* BBB model composed of a bEnd.3 monolayer and

a hippocampal neuron layer (Fig. 2A). The bEnd.3 cell is a mouse brain microvascular endothelial cell line which expresses the Tf receptor. D-T7/Tet1-lipids@PL (2 : 1) was supposed to transcytosis the bEnd.3 membranes by binding to the Tf receptor. Z-axis confocal imaging showed that different nanoparticles appeared in the bEnd.3 monolayer with different fluorescence intensities and penetration depth. Tet1-lipids@PL, although having the highest fluorescence inside the bEnd.3 monolayer, showed relatively low fluorescence intensity beneath the monolayer. This means that although Tet1-lipids@PL can bind to the bEnd.3 cells, they only have a weak chemotaxis tendency towards the hippocampal neurons beneath the bEnd.3 monolayer (Fig. 2B). D-T7/Tet1-lipids@PL (2 : 1) had the largest penetration depth among all the experimental groups (Fig. 2B), which means that D-T7/Tet1-lipids@PL (2 : 1) had the most apparent chemotaxis tendency towards the neurons. Flow cytometry showed that all the NPs penetrated the BBB model and were internalized by the hippocampal neurons beneath the bEnd.3 monolayer (Fig. 2C and D). Among all the NP groups, D-T7/Tet1-lipids@PL (2 : 1) showed the highest fluorescence intensity, demonstrating its excellent BBB penetration and neuron targeting performance (Fig. 2C and D).

To evaluate the BBB penetration ability of the nanoparticles, we used an *in vivo* imaging system (IVIS) to monitor the fluorescence distribution of DiI-labelled NPs in mice. Compared with other NPs, D-T7-lipids@PL and D-T7/Tet1-lipids@PL (2 : 1) showed a stronger fluorescence signal in the brain at 6- and 12-h

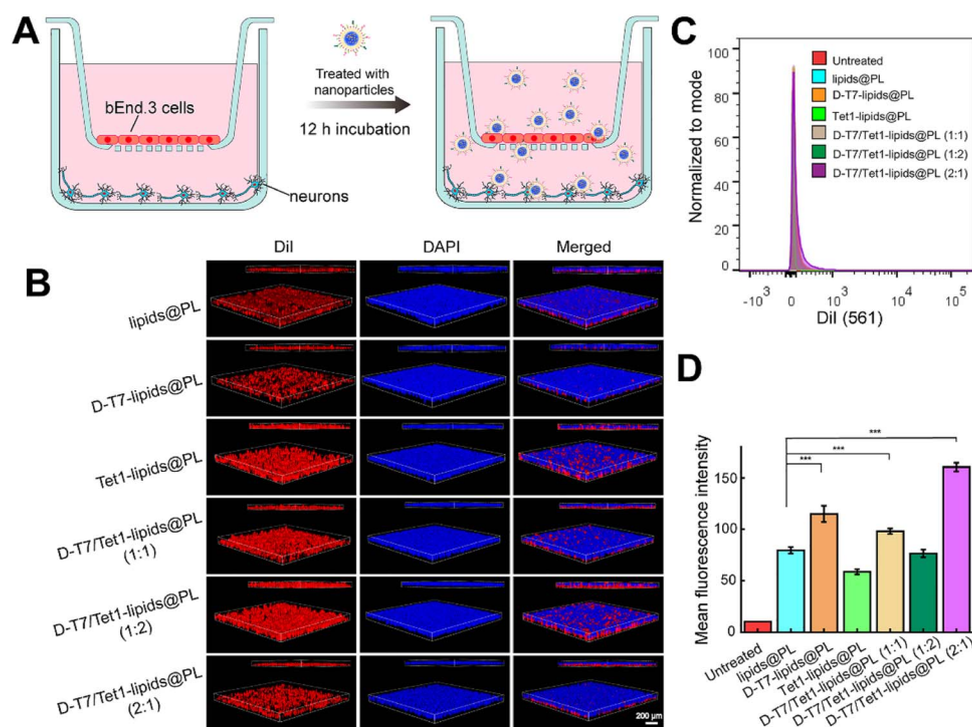


Fig. 2 The *in vitro* BBB model verifies the penetration efficiency of various NPs with different D-T7/Tet1 concentration titrations through the BBB. (A) Schematic diagram of the ability of nanoparticles to cross the BBB model *in vitro*. (B) 3D images of immunofluorescence of the bEnd.3 monolayer treated with various nanoparticles. The nanoparticles are labelled with DiI (red). The cell nuclei are stained with DAPI (blue). Scale bar, 200 μ m. (C) Flow cytometry histogram of internalization of nanoparticles by hippocampal neurons after passing through the bEnd.3 monolayer. (D) Data from the flow cytometry which show the mean fluorescence intensity of the nanoparticles in the hippocampal neurons. *** $P < 0.001$.



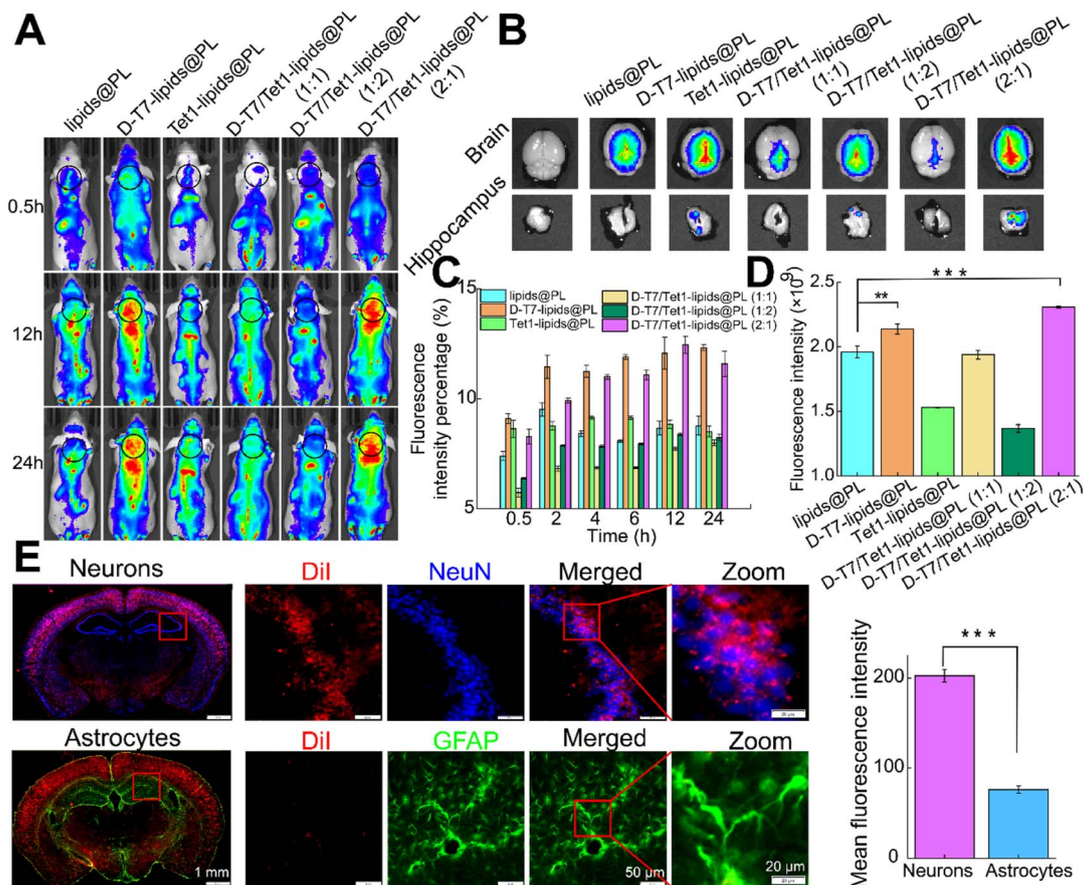


Fig. 3 Fluorescence distribution of different NPs *in vivo*. (A) *In vivo* imaging shows the distribution of different NPs with different D-T7/Tet1 concentration titrations in mice at different time points. (B) *In vivo* imaging of mouse brains in different groups after 24 h. (C) Quantification of the enrichment of different NPs in the brains from the fluorescence intensity. (D) Fluorescence retention rate of different NPs in the brains ($n = 3$) $***P < 0.001$. (E) Immunofluorescence of the mouse brain treated with D-T7/Tet1-lipids@PL (2 : 1) labelled with Dil. The neuron nuclei are stained with NeuN (blue) and the astrocytes are stained with GFAP (green). $**P < 0.01$, $***P < 0.001$.

time points (Fig. 3A, B, and S11A†). The presence of PEG in the nanoparticles can enhance the long-term circulation of the nanoparticles in mice, leading to increased fluorescence signals after 24 h. The mice were sacrificed and dissected after 24 h. The brains of the mice injected with D-T7/Tet1-lipids@PL (2 : 1) showed the brightest fluorescent signal among all the experimental groups (Fig. 3C). We extracted the hippocampus from different experimental groups and found that the hippocampus of the mice treated with D-T7/Tet1-lipids@PL (2 : 1) had the strongest fluorescent signal, indicating its excellent performance to target the hippocampus region (Fig. 3D and S12†). To evaluate the neuron-targeting capability of D-T7/Tet1-lipids@PL (2 : 1), the mouse brain tissues were dissected and NeuN (neuron marker) and GFAP (astrocyte marker) were stained specifically 6 h after the NP injection. Most of the D-T7/Tet1-lipids@PL (2 : 1) NPs co-localized with the NeuN-labelled cells (Fig. 3E, S13 and S14†), indicating that D-T7/Tet1-lipids@PL (2 : 1) can specifically target the neurons. A previous study reported that nanoparticles within 200 nm can also cross the BBB.⁵ The size of D-T7/Tet1-lipids@PL (2 : 1) was less than 100 nm (Fig. 1D), so, it is easy for them to penetrate the BBB. Also, D-T7/Tet1-lipids@PL (2 : 1) has dual targeting properties by binding

to Tf receptors *via* the D-T7 peptide and binding to gangliosides on neurons *via* the Tet1 peptide. The Tf receptors have been reported to be expressed in the cerebral cortex and hippocampus, and are also widely expressed in the neurons.⁴² Thus, D-T7/Tet1-lipids@PL (2 : 1) can not only target the BBB, but also accumulate in the neurons. Since both *in vivo* and *in vitro* experiments have proven that D-T7/Tet1-lipids@PL (2 : 1) had the best BBB penetration and neuron targeting capability among the NPs, we selected it for further study. In addition, D-T7/Tet1-lipids@PL (2 : 1) was also present in the blood vessels of the brain (Fig. S15†). Apart from the brain, the liver and the spleen showed strong fluorescence, implying that the NPs tend to accumulate in the liver and can be metabolized quickly (Fig. S11B†).

We constructed two acute epilepsy models, the KA-initiated model (Fig. 4A) and the pilocarpine-initiated model (Fig. 4B), which were used for electrical signal monitoring and behavior analysis respectively. We verified the antiepileptic effect of D-T7/Tet1-lipids@PL (2 : 1) on the KA epilepsy model. C57BL/6 mice were randomly divided into seven groups and injected with various NPs intravenously 2 h before the model construction (Fig. 4A). The PBS, LTG, lipids@PL, Tet1-lipids@PL, and D-T7-



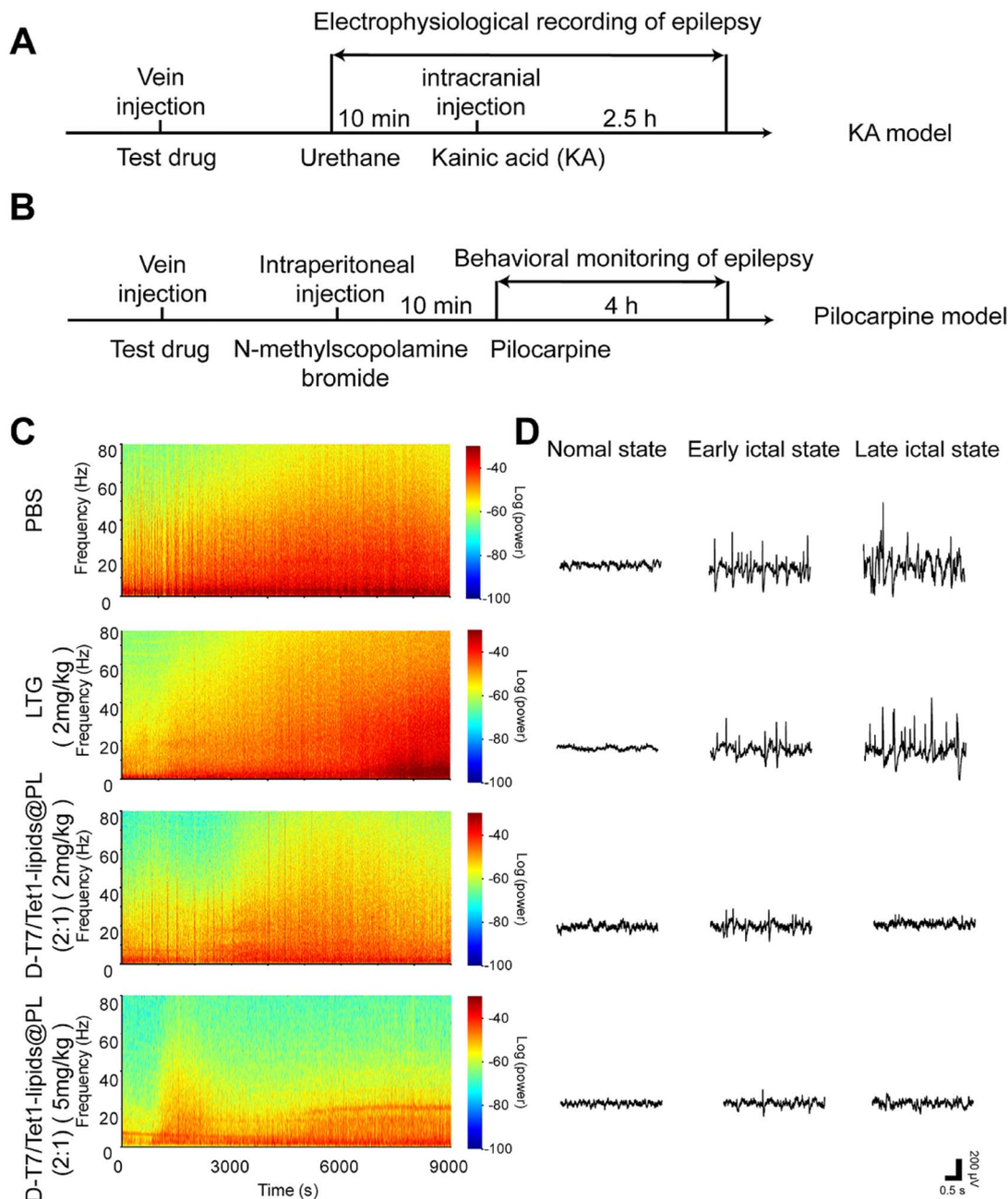


Fig. 4 Electrophysiological data of the acute epilepsy model treated with the NPs. Experimental designs of the KA model (A) and pilocarpine model (B) are shown in the diagram. (C) A 2.5-hour spectrogram of LFPs recorded in the dorsal hippocampus (dHPC) after modelling. (D) Representative LFPs of the normal state, early ictal state and late ictal state recorded in the dHPC of mice after KA injection.

lipids@PL groups showed the onset of ictal seizures at about 30 min. However, ictal discharges start at about 1 h later (early ictal state, Fig. 4C, D, and S16[†]) in the D-T7/Tet1-lipids@PL (2 : 1) group. About 2 h later (late ictal state), the local field potential (LFP) power (Fig. 4C) and amplitude (Fig. 4D) in the D-T7/Tet1-lipids@PL (2 : 1) group were obviously lower than those of other groups (Fig. 4C, D, and S16[†]). The D-T7-lipids@PL (2 : 1) group, whether with a high (5 mg kg⁻¹) or low (2 mg kg⁻¹) LTG dose, exhibited apparent inhibitory effect on the seizures (Fig. 4C, D and S16[†]). Thus, D-T7/Tet1-lipids@PL (2 : 1) can delay and shorten the time of epileptic seizures and reduce the degree of

epileptic seizures. In the experiments on the pilocarpine epilepsy model, 40 min after the injection of pilocarpine, the seizures occurred in the PBS (level 5), LTG (level 4), and lipids@PL (level 4) groups (Video S1[†]). By comparison, only mild seizures (level 1) occurred in the D-T7/Tet1-lipids@PL (2 : 1) group after 1.5 h (Video S2[†]). 4 h later, the behavior of the D-T7/Tet1-lipids@PL (2 : 1) group returned to the normal state, while the mice in other groups were still epileptic (Video S3[†]). Thus, D-T7/Tet1-lipids@PL (2 : 1) can delay the onset of epilepsy and reduce the degree of epileptic seizures. The electrophysiological signal monitoring and behavioral observation in the two acute



epilepsy models confirmed that D-T7/Tet1-lipids@PL (2 : 1) was excellent in the treatment of epilepsy.

We further verified the antiepileptic effect of D-T7/Tet1-lipids@PL (2 : 1) on the KA-induced chronic epilepsy model (Fig. 5A). In the LFP results, spontaneous seizures in the dHPC of PBS and LTG groups were obviously observed, while the D-T7/Tet1-lipids@PL (2 : 1) group showed a relatively normal state (Fig. 5B and C). In addition, the LFP power of the D-T7/Tet1-lipids@PL (2 : 1) group was significantly lower than those of other groups (Fig. 5D and E). Neuronal loss and dispersion of the granule cell layer of the dHPC are considered as the signs of chronic epilepsy.^{43,44} As shown in dHPC images, neuronal loss

and dispersion of the granule cell layer of the dHPC in the mice treated with PBS or LTG were much more serious than those treated with D-T7/Tet1-lipids@PL (2 : 1), and the D-T7/Tet1-lipids@PL (2 : 1) group had an intact hippocampal structure (Fig. 5F). The cell density of the granule cell layer in mice treated with PBS or LTG was significantly lower than that of the D-T7/Tet1-lipids@PL (2 : 1) group (Fig. 5G). It is worth mentioning that D-T7/Tet1-lipids@PL (2 : 1) NPs were also located in cortex structures (Fig. 3E). As seizures usually propagate from the hippocampus to the cortex structures,^{45–47} our result suggested that D-T7/Tet1-lipids@PL (2 : 1) NPs may inhibit both the initiation and spreading of ictal discharges. The above data

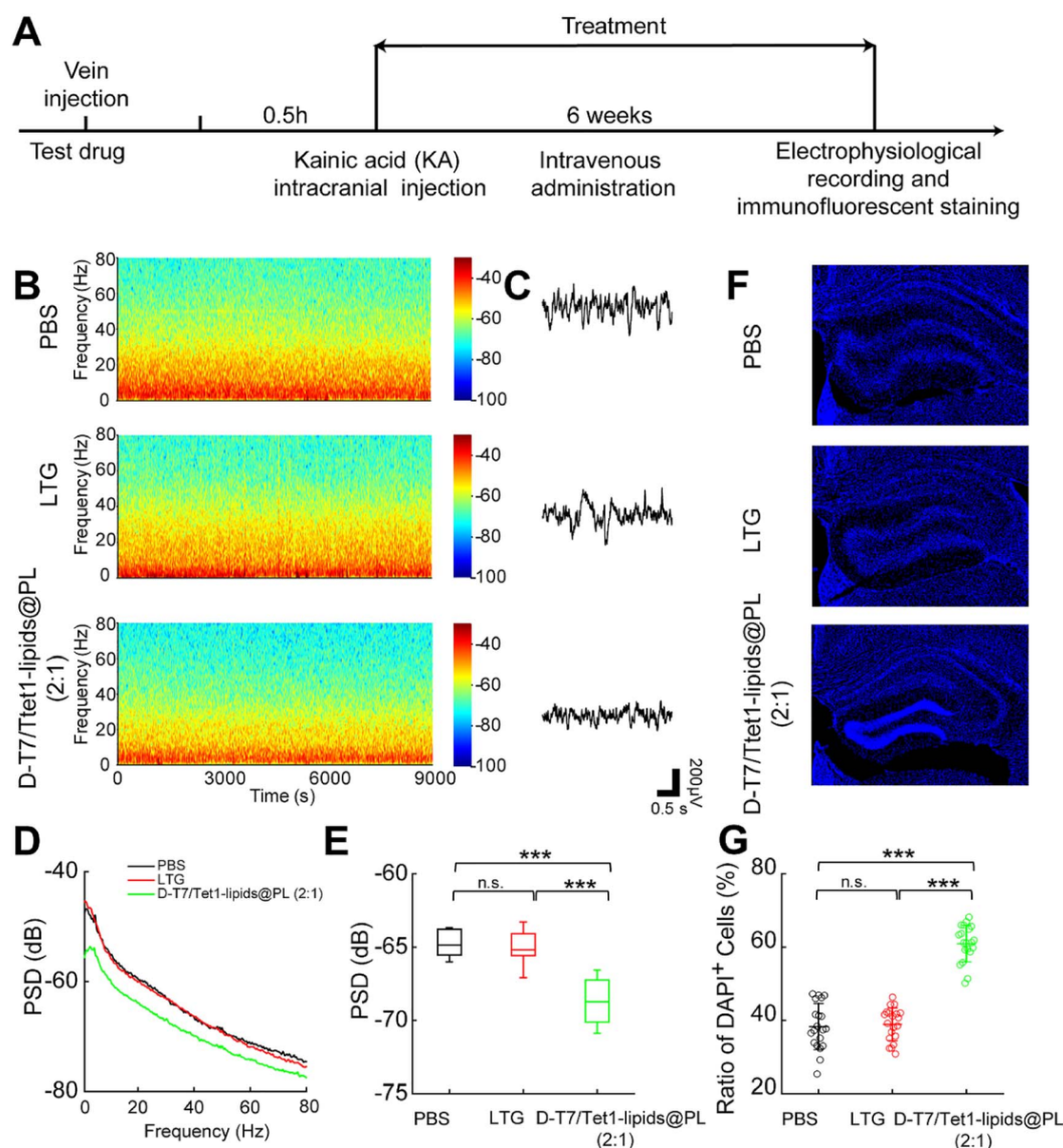


Fig. 5 Electrophysiological data of the KA-induced chronic epilepsy model treated with various NPs. (A) Experimental design of the chronic KA model. (B) LFPs recorded in the dHPC in different treatment groups. (C) LFPs recorded from representative channels in the dHPC in different treatment groups. (D–E) Average power spectra of LFPs recorded from mice injected with PBS, LTG and D-T7/Tet1-lipids@PL (2 : 1). (F) Representative dHPC images from mice injected with PBS, LTG or D-T7/Tet1-lipids@PL (2 : 1). (G) Ratio of DAPI-positive cells of dHPC images from mice injected with PBS, LTG and D-T7/Tet1-lipids@PL (2 : 1). ($n = 20$ slices from 5 mice), *** $P < 0.001$.



indicated that D-T7/Tet1-lipids@PL (2:1) had an excellent therapeutic effect in treating epilepsy and had a long-term protective effect on the brain tissues.

We evaluated the biocompatibility of D-T7/Tet1-lipids@PL (2:1) *in vitro*. Confocal microscopy showed that hippocampal neurons and bEnd.3 cells were viable after co-culturing with various NPs for 12 h (Fig. S17A and B†). CCK assays showed that after 24 h of incubation with an extremely high concentration of D-T7/Tet1-lipids@PL (2:1) ($1000 \mu\text{g mL}^{-1}$), more than 90% of the cells were alive (Fig. S17C and D†). Hemolysis tests indicated that even at an extremely high concentration ($1000 \mu\text{g mL}^{-1}$), D-T7/Tet1-lipids@PL (2:1) did not cause hemolysis (Fig. S18†). Hematological markers and biochemical parameters in mice blood showed that all the indexes of the D-T7/Tet1-lipids@PL (2:1)-treated mice were within the normal ranges in the acute (Fig. S19A†) and chronic (Fig. S20†) epilepsy models. Hematoxylin-eosin (H&E) staining showed no obvious pathological changes in the tissues of the mice treated with D-T7/Tet1-lipids@PL (2:1) in the acute (Fig. S19B†) and chronic (Fig. S21†) epilepsy models. These results indicated that the series of lipids@PL NPs had good biocompatibility.

Conclusions

In this study, we used a microfluidic chip to synthesize a dual-targeting system (D-T7/Tet1-lipids@PL) with a core-shell structure to deliver lamotrigine for treating epileptic seizures. When the ratio of D-T7 to Tet1 was 2:1, the D-T7/Tet1-lipids@PL NPs showed a maximum BBB penetration ability and the best targeting efficiency to the neurons both *in vitro* and *in vivo*. D-T7/Tet1-lipids@PL (2:1) realized excellent therapeutic effects on the seizures in both acute and chronic epilepsy models with high biosafety. Therefore, this nanocarrier may not only be used in treating epilepsy but its applications may also be extended to treating other disorders in the central nervous system such as glioma, Alzheimer's disease, and other psychiatric disorders.

Data availability

All other experimental data are available in the ESI.†

Author contributions

Qinghong Hou, Wenfu Zheng and Xingyu Jiang conceived the study; Qinghong Hou and Lulu Wang performed the experiments; Feng Xiao and Le Wang helped in the construction of the epileptic model; Xiaoyan Liu assisted in extracting hippocampal neurons. Qinghong Hou, Wenfu Zheng and Xingyu Jiang wrote the manuscript. All authors have given approval to the final version of the manuscript. Yi Lu, Lina Zhu, Wenfu Zheng, and Xingyu Jiang supervised the whole project.

Conflicts of interest

The authors declare no competing financial interest.

Acknowledgements

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