Rituximab-conjugated, doxorubicin-loaded microbubbles as a theranostic modality in B-cell lymphoma

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ABSTRACT

This study evaluated rituximab-conjugated, doxorubicin-loaded microbubbles (RDMs) in combination with ultrasound as molecular imaging agents for early diagnosis of B cell lymphomas, and as a targeted drug delivery system. Rituximab, a monoclonal CD20 antibody, was attached to the surfaces of doxorubicin-loaded microbubbles. RDM binding to B cell lymphoma cells was assessed using immunofluorescence. The cytotoxic effects of RDMs in combination with ultrasound (RDMs+US) were evaluated in vitro in CD20+ and CD20− cell lines, and its antitumor activities were assessed in Raji (CD20+) and Jurkat (CD20−) lymphoma cell-grafted mice. RDMs specifically bound to CD20+ cells in vitro and in vivo. Contrast enhancement was monitored in vivo via ultrasound. RDM peak intensities and contrast enhancement durations were higher in Raji than in Jurkat cell-grafted mice (P<0.05). RDMs+US treatment resulted in improved antitumor effects and reduced systemic toxicity in Raji cell-grafted mice compared with other treatments (P<0.05). Our results showed that RDMs+US enhanced tumor targeting, reduced systemic toxicity, and inhibited CD20+ B cell lymphoma growth in vivo. Targeted RDMs could be employed as ultrasound molecular imaging agents for early diagnosis, and are an effective targeted drug delivery system in combination with ultrasound for CD20+ B cell malignancy treatment.

INTRODUCTION

Early diagnosis is pivotal for therapeutic success in many types of cancers. Ultrasound (US) molecular imaging is a novel diagnostic approach for early detection of non-hodgkin lymphoma. Recent studies suggest that targeted microbubbles as US contrast agents (TMUCA) may serve as probes for US molecular imaging. TMUCA would improve diagnostic specificity and allow for disease monitoring in real time. TMUCAs can accumulate and remain at the tumor site for long time periods, and imaging at the molecular level can be achieved using US after TMUCA venous injection [1–3]. US molecular imaging also produces quantitative data, exhibits good temporal resolution, is noninvasive, produces no ionizing radiation, and is relatively inexpensive [4]. Over the past decade, various types of TMUCA have been applied for cell-specific targeting with US molecular imaging in vivo [5–10], specifically to assess intravascular inflammation, intravascular thrombosis and tumor blood vessels. For early tumor diagnosis, TMUCAs were conjugated with antibodies specific for tumor cell surface antigens. Previous studies showed that the tumor neovasculature is distorted, with an imperfect basement membrane, and no smooth muscle layer [11]. Permeability was also increased, with wall pores approximately 380–780 nm in diameter [11]. Therefore, TMUCAs diameters were adjusted to approximately 500 nm for easy passage through vascular endothelial cells and improved molecular imaging.

Targeted microbubbles are promising tumor-targeting drug delivery systems, although their potential utility as US contrast agents has not yet been studied. Most chemotherapy drugs currently have no targeting capabilities, and act on both diseased and non-diseased sites, leading to low therapeutic indices and severe side effects. A targeted drug delivery system can increase chemotherapy drug accumulation specifically at target sites, while reducing non-target impacts. Moreover, targeted microbubbles are both chemically stable and biodegradable, and exhibit prolonged circulation in the blood, with localized drug release. Tumor-specific ligand-like peptides [12, 13], galactose-conjugated chitosan [14], transferrin [15], folic acid [16–19], and...
monoclonal antibodies [20–22] have been employed to target microbubbles to tumor cells for the treatment of many cancers. Additionally, the combination of targeted drug-loaded microbubbles with US irradiation permeabilizes cell membranes, enhancing drug uptake by tumor cells, and selectively killing tumor cells without harming normal cells. Therefore, targeted drug-loaded microbubbles have potential use in both targeted drug delivery systems and in combination with US molecular imaging.

We hypothesized that Rituximab-conjugated, doxorubicin (DOX)-loaded microbubbles (RDMs) could serve as effective, biocompatible B cell lymphoma-targeting theranostic agents. The present work evaluated the specific binding potential of RDMs targeting CD20 antigen, a tetraspan membrane receptor overexpressed in B cell malignancies, in lymphoma Raji cells. We also assessed the cytotoxicity and antitumor activity of these RDMs in combination with US irradiation in vitro and in vivo. Finally, targeted US molecular imaging was explored in Raji and Jurkat cell-grafted mice.

RESULTS

RDM characterization

In this study, RDMs targeted the lymphoma B cell CD20 antigen (Figure 1). US triggered DOX release, which was then delivered into lymphoma B cells. RDM morphologies and size distributions were observed via scanning electron microscopy (SEM) (Figure 2A) and transmission electron microscopy (TEM) (Figure 2B), respectively. Mean RDM diameter was 411±125.1 nm (Figure 2C). Fluorescence imaging of RDMs revealed dense green (DyLight488-labeled avidin) and red (DOX) fluorescence with morphologies consistent with those observed via confocal laser scanning microscopy (CLSM) (Figure 3A). Targeting moiety quantities on RDM surfaces were evaluated by detecting RDM suspension fluorescence intensity after conjugation. RDM fluorescence intensity was 72.15% that of the DyLight488-labeled avidin samples (Figure 3B). Due to the high affinity of avidin to biotin, we presumed the same high level of adhesion of biotinylated rituximab to the avidin-conjugated DMs.

Drug loading and release

RDMs as a targeted drug delivery system were evaluated via the encapsulation efficiency of DOX. DOX encapsulation efficiency in RDMs was 51.2±2.05%. The release profiles of DOX from RDMs and DMs as triggered by US were also examined. The release profile was described as the percentage of cumulative released DOX as a function of time (Figure 3C). Total DOX released was the same for DMs + US and RDMs + US. The results indicated that DOX was about 50% unloaded after 5 h

Figure 1: Illustration of RDM structure, and depiction of antigen-specific, tumor cell targeting drug delivery. After RDM attachment to CD20 antigens on Raji cells, US irradiation triggers DOX release and delivery into cells. DOX, doxorubicin; RDMs, rituximab-conjugated, DOX-loaded microbubbles.
Figure 2: Microbubble characterization. SEM A. and TEM B. images of RDMs. RDM size distributions as determined by DLS C.

Figure 3: Microbubble fluorescence and drug release characterization. RDM imaging was performed using CLSM as follows: bright field Aa., dylight488-avidin on RDMs (green fluorescence) Ab., encapsulated DOX in RDMs (red fluorescence) Ac. (scale bar = 25μm). Fluorescence absorbance of RDMs B. DMs and free avidin were examined to assess rituximab conjugation efficiency. In vitro US-triggered DOX release from DMs and RDMs C.
with sonication and about 90% unloaded after 72 h with sonication. This implied that US could promote DOX release from RDMs and DMs through cavitation.

**Targeted properties of RDMs**

To estimate the targeted binding capability of RDMs, the affinity of RDMs to CD20 antigen on Raji cells was determined *in vitro*. RDM attachment to CD20 antigen was greater than that of DMs. CLSM imaging showed large amounts of RDMs (green and red fluorescence) aggregated on Raji cell membranes, demonstrating that rituximab enhanced RDM targeted binding to CD20 antigen. Few DyLight488-labeled avidin-conjugated biotinylated DMs were observed on Raji cell membranes. Competition experiments revealed that RDM targeted binding of Raji cells was reduced as CD20 antigen was blocked following rituximab pre-incubation, as indicated by absence of red and green fluorescent microbubbles. Few RDMs were observed on CD20− CEM cell membranes (Figure 4).

**Cytotoxicity *in vitro***

The therapeutic efficacies of DOX, DOX + rituximab, DMs + US, RDMs + US, and RDMs + US + rituximab were explored *in vitro*. Raji, Daudi, Jurkat, and CEM cell proliferation inhibition was assessed after treatment for 24, 48 and 72 h. Raji and Daudi cell proliferation inhibition was limited, and was the same for DOX, DOX + Rituximab, DMs + US, and RDMs + US + Rituximab. In contrast, RDM + US inhibited Raji cell proliferation after 24 (35.42±2.16%), 48 (52.32±3.42%), and 72 h (82.74±2.97%) (Figure 5A). RDM + US in Daudi cells also reduced proliferation at 24 (32.84±3.31%), 48 (48.79±2.71%), and 72 h (74.85±3.52%) (Figure 5C). Compared with other treatments, RDM + US effectively inhibited Raji and Daudi cell proliferation (P<0.05). However, in Jurkat and CEM cells, all treatments exhibited similar cytotoxicity (Figure 5B & 5D). RDM + US inhibited proliferation in Raji and Daudi cells more than in Jurkat and CEM cells (P<0.05). We also observed time-dependent cytotoxic effects in all groups. These results demonstrated that US-mediated RDM destruction efficiently inhibited CD20+ lymphoma B cell proliferation.

**Cell apoptosis *in vitro***

Raji, Daudi, Jurkat, and CEM cell apoptosis rates were detected quantitatively by flow cytometry 24, 48 and 72 h after various treatments. Raji and Daudi cell apoptosis rates were comparable following DOX, DOX+ rituximab, DMs + US, and RDMs + US + rituximab treatment,
although apoptosis was increased in all groups compared to controls. Importantly, RDMs + US induced higher apoptosis rates than other treatments (Figure 6A & 6C, \( P<0.05 \)). Jurkat and CEM cell apoptosis rates were similar for DOX, DOX + rituximab, DMs + US, and RDMs + US + rituximab, but higher compared to controls (Figure 6B & 6D). Additionally, RDM + US induced higher apoptosis rates in Raji and Daudi cells as compared to Jurkat and CEM cells (\( P<0.05 \)). Time-dependent apoptosis rates was detected in all groups (Figure 6). Apoptosis rate measurements were consistent with proliferation inhibition results.

**Cellular uptake of DOX**

Because DOX auto-fluoresces, we detected whether 48 h RDMs + US treatment improved Raji, Daudi, Jurkat, and CEM cell DOX uptake using flow cytometry. DOX intracellular RFI for Raji and Daudi cells was similar following DOX, DOX + rituximab, DMs + US, and RDM + US + rituximab treatment. However, DOX intracellular RFI following RDM + US was higher compared to all other treatments (Figure 7A & 7C, \( P<0.01 \)). DOX intracellular RFI was also similar following all treatments in Jurkat and CEM cells (Figure 7B & 7D). Additionally, DOX intracellular RFI following RDM + US treatment was higher in Raji and Daudi cells as compared to Jurkat and CEM cells (\( P<0.01 \)). These results suggest that RDM + US treatment increased DOX transfer into lymphoma B cells more than other treatment groups.

**In vivo imaging**

Arrival time, time to peak, peak intensity, and duration of contrast enhancement were compared via US imaging between non-targeted DMs and targeted RDMs in Raji and Jurkat cell-grafted mice. In Raji cell-grafted mice, there was no difference between DMs and RDMs for arrival time or time to peak, but peak intensity and duration of contrast enhancement were higher for RDMs (\( P<0.05 \)). In Jurkat cell-grafted mice, there was no difference between DMs and RDMs in any US measurement (Table 1). Additionally, arrival times and times to peak for targeted RDMs were the same in Raji and Jurkat cell-grafted mice. However, RDM peak intensities and the durations of contrast enhancement were higher in Raji as compared to Jurkat cell-grafted mice (Figure 8, \(*P<0.05\)). Targeted RDM (Figure 9C & 9F) and non-targeted DM (Figure 9B & 9E) peak intensity images are shown for Raji and Jurkat cell-grafted mice.

**In vivo antitumor activity**

This study used a lymphoma nude mouse model to investigate the antitumor effects of RDMs + US *in vivo*. RDMs + US exhibited the strongest tumor inhibition effect in Raji-cell-grafted mice. DMs + US, and RDMs + US + rituximab-treated mice exhibited similarly-reduced Raji cell tumor growth rates compared to controls, and inhibited tumor growth more than treatment with DOX and DOX + rituximab.

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**Figure 5: Raji, Daudi, Jurkat and CEM cell growth inhibition.** Cell proliferation inhibition was measured 24, 48, and 72 h after treatment with DOX, DOX + rituximab, DMs + US, RDMs + US, and RDMs + US + rituximab via CCK8 assay. Data are represented as means ± SD (n=3). \(*P<0.05\) compared with RDM + US.
(Figure 10A). DOX and DOX + rituximab only slightly inhibited tumor growth in vivo. Similarly, DMs + US, RDMs + US, and RDMs + US + rituximab treatment reduced grafted Jurkat cell tumor growth as compared with DOX and DOX + rituximab. Jurkat cell-grafted mouse treatment with DMs + US, RDMs + US and RDMs + US + rituximab resulted in comparable growth inhibition rates (Figure 10C).

**Figure 6:** Raji, Daudi, Jurkat and CEM cell apoptosis. Cell apoptosis was measured 24, 48, and 72 h after treatment with DOX, DOX + rituximab, DMs + US, RDMs + US, and RDMs + US + rituximab by flow cytometry. Data are represented as means ± SD (n=3). *P<0.05 compared with RDM + US.

**Figure 7:** DOX relative fluorescence intensity (RFI) in Raji, Daudi, Jurkat and CEM cells. Cell RFIs were analyzed 48 h after treatment with DOX, DOX + rituximab, DMs + US, RDMs + US, and RDMs + US + rituximab by flow cytometry. Data are represented as means ± SD (n=3). **P<0.01 compared with RDM + US.
To assess the potential systemic toxicity of RDMs + US in vivo, nude mouse body weights were periodically examined. Raji and Jurkat cell-grafted mice treated with DOX and DOX + rituximab exhibited slow, continuous weight loss beginning on day 8. In contrast both Raji and Jurkat cell-grafted mouse weights increased gradually with saline, DMs + US, RDMs + US, and RDMs + US + rituximab treatments (Figure 10B & 10D). This suggests that DOX treatment caused severe systemic toxicity in nude mice.

TUNEL staining was used to evaluate apoptosis in Raji and Jurkat cell tumors. Sparse apoptosis (green fluorescence) in Raji and Jurkat lymphoma tissues was observed in mice treated with DOX and DOX + rituximab. Raji cell-grafted mouse tissues treated with DMs + US, and RDMs + US + rituximab showed moderate apoptosis, while RDMs + US treatment induced the most apoptosis. Jurkat cell-grafted mouse tissues treated with DMs + US, RDMs + US, and RDMs + US + rituximab showed moderate cell

### Table 1: Four parameters (means ± SD) for DMs and RDMs in Raji and Jurkat cell-grafted mice.

| Indicators       | Arrival time (sec) | Time to peak (sec) | Peak intensity (sec) | Duration of contrast enhancement (min) |
|------------------|--------------------|--------------------|----------------------|---------------------------------------|
| Raji cell lymphoma |                    |                    |                      |                                       |
| DMs              | 2.71 ± 0.09        | 17.91 ± 0.62       | 19.98 ± 0.84*        | 20.12 ± 0.97*                         |
| RDMs             | 2.75 ± 0.10        | 17.98 ± 0.56       | 23.27 ± 1.07         | 24.27 ± 1.05                          |
| Jurkat cell lymphoma |                |                    |                      |                                       |
| DMs              | 2.79 ± 0.07        | 18.25 ± 0.97       | 20.06 ± 1.32         | 20.62 ± 1.05                          |
| RDMs             | 2.82 ± 0.08        | 18.31 ± 1.12       | 20.17 ± 1.60*        | 20.99 ± 1.57*                         |

Paired t-test in each group between DMs and RDMs, *P<0.05.

Figure 8: RDM arrival time, time to peak, peak intensity, and duration of contrast enhancement in Raji and Jurkat cell-grafted mice. RDM arrival times and times to peak were the same in Raji and Jurkat cell-grafted mice. RDM peak intensities and contrast enhancement durations were greater in Raji cell-grafted mice than in Jurkat cell-grafted mice. Data are represented as means ± SD (n=3). *P<0.05, #P>0.05.
apoptosis. We observed that RDMs + US induced greater apoptosis levels in Raji as compared to Jurkat cell-grafted mice (Figure 11). These results confirmed that RDMs + US could inhibit lymphoma cell growth by inducing apoptosis.

**DISCUSSION**

Microbubble-based targeted drug delivery has been widely investigated as an anti-tumor therapy in
combination with US irradiation [23, 24]. Most targeted drug delivery systems exhibit high therapeutic efficacies in vitro and in vivo. However, few studies have assessed their potential roles, and the roles of microbubbles, in ultrasonic molecular imaging for diagnosis.

In the present work, we conjugated rituximab to DMs (RDMs) to construct TMUCA. We investigated the potential benefits of TMUCA in both US molecular imaging and targeted anti-lymphoma treatment. DMs encapsulated rituximab with 51.2% efficiency, similar to a previous study [25]. The results of US-mediated drug release assays implied that US induced DMs to release DOX in a controlled manner, probably due to cavitation. RDMs effectively targeted the CD20 antigen on lymphoma B cells. Very few non-targeting DMs were observed on or near Raji cells due to the absence of rituximab. Competition experiments indicated that RDM-Raji cell targeting was reduced when CD20 was blocked by rituximab pretreatment. These results indicate that rituximab conjugation greatly improved DM-lymphoma cell targeting.

In Jurkat and CEM cells in vitro, DMs + US, RDMs + US, and RDMs + US + rituximab treatment induced similar levels of cytotoxicity compared to DOX and DOX + rituximab. This verified that US-triggered drug release from the microbubbles was highly efficient. However, RDMs + US treatment inhibited Raji and Daudi cell proliferation most effectively, in agreement with cell apoptosis assay and intracellular DOX RFI results, as well as the results of a previous report [26]. Additionally, another group reported that anti-CD20 nanoparticles carrying hydroxychloroquine and chlorambucil showed promise for treatment of Burkitt’s lymphoma [27].

US molecular imaging allows for exploration of pathological processes in vivo at cellular and molecular levels [28, 29], and will improve cancer diagnosis and treatment options. B7-H3-targeted US molecular imaging can reportedly improve breast cancer diagnostic accuracy [30]. US molecular imaging also provides a tool for imaging at a molecular level to enhance B cell lymphoma early diagnosis. In this study, we performed enhanced-contrast imaging experiments with targeted RDMs in a B cell lymphoma nude mouse model. RDM peak intensities and durations of contrast enhancement were higher compared to non-targeted DMs in Raji cell-grafted mice, while there was no difference between RDMs and DMs in CD20– Jurkat cell-grafted mice. Antigen-antibody binding occurred in Raji cell lymphoma tissues after targeted RDMs passed through vascular endothelial cell gaps, leading to a considerable number of RDMs adhering to CD20-overexpressing Raji cells, and generating more stable and longer-lasting contrast enhancement in vivo compared to non-targeted DMs. Moreover, targeted RDMs remain in tissues long term, leading to higher peak intensities and persistent contrast enhancement in CD20+ Raji cell-grafted mice as compared to CD20–Jurkat cell-grafted mice. Thus, peak intensity and duration of contrast enhancement may be useful primary parameters in further investigations of targeted RDMs in CD20+ B cell malignancies.

In this study, DOX and DOX + rituximab only slightly inhibited lymphoma cell growth in vivo. This was attributed to the rapid diffusion of DOX into all mouse tissues following intravenous injection, leading to potential toxic side effects, and smaller amounts of DOX at tumor sites. The DOX molecule also has a short half-life in vivo, and may be metabolized in the circulation before reaching the tumor [31]. The lower systemic toxicity and enhanced anti-tumor activity of RDMs + US observed via TUNEL staining may result from the following factors: (1) encapsulation effectively protected DOX from quick clearance in the circulation; (2) RDMs passively and actively aggregated at the tumor site; (3) rituximab-CD20 binding contributed to RDM-Raji cell attachment, facilitating interaction between drug-loaded microbubbles and cells, and improving local drug delivery efficiency [32]; (4) sonoporation may have enhanced drug uptake by tumor cells via cell membrane pore formation [33], and; (5) rituximab as a targeted drug induced antibody-dependent, cell-mediated cytotoxicity, reducing systemic side effects. Thus, the synergistic effect of DOX and rituximab in RDMs effectively inhibited tumor growth. Our results show that RDMs + US is a promising anti-B cell lymphoma treatment strategy.

In conclusion, this study indicated that targeted RDMs specifically bound CD20+ B cell lymphomas. RDMs combined with US irradiation enhanced tumor targeting, reduced systemic toxicity, and inhibited B cell lymphoma cell growth in vivo. Additionally, targeted RDMs increased peak intensity and contrast enhancement duration compared to non-targeted DMs in CD20+ B cell lymphoma-grafted mice. Our findings show that targeted RDMs could potentially be employed as US molecular imaging agents for early diagnosis, and are an effective targeted drug delivery system in combination with US irradiation for the treatment of CD20+ B cell malignancies.

MATERIALS AND METHODS

Materials

Poly(lactic-co-glycolic acid) (PLGA; 50% lactide, 50% glycolide, MW=10,000 Da) was purchased from Shandong Shuyuan Biotechnology Co., Ltd (Shandong, China). Poly(vinyl alcohol) (PVA, 87–89%, MW=31,000–50,000) was obtained from Sigma-Aldrich (St Louis, MO, USA). Doxorubicin (DOX) was obtained from Shenzhen Wanle Pharmaceutical Co., Ltd. (Shenzhen, China). EZ-LinkTM Sulfo-LC-Biotinylation kit was purchased from Thermo Fisher Scientific, Inc. (Rockford, USA). Amine-Peg2000-Biotin was purchased from Nanjing Ling Di Ren Chemical Technology Co., Ltd (Nanjing, China). Avidin and dyLight488-labeled avidin were obtained from Wuhan Boster Biotechnology Co., Ltd (Wuhan, China). Rituximab was obtained from Hoffmann-La Roche, Inc.
(New Jersey, USA). Cell Counting Kit-8, the Annexin V-FITC cell apoptosis detection kit and the TUNEL apoptosis detection kit were purchased from Beyotime Biotechnology Co., Ltd (Shanghai, China). All chemicals were analytical grade and used without further processing.

**DM preparation**

PLGA microbubbles incorporating DOX were fabricated via a double US emulsion evaporation procedure. 0.5 g PLGA was fully dissolved in 10 ml of liquid chloroform via agitation. The PLGA solution was then combined with a 5 mg DOX solution (dissolved in 1.0 mL superpure water), and the mixture was emulsified via US for 120 min at 100 w. 1.0 mL span-80 was then added. The vial was degassed and re-perfused with nitrogen with stirring at 23,000 rpm for 5 min to obtain primary emulsified DMs. The primary emulsion was poured into cold PVA (40 ml, 5%) containing 1.0 ml tween-80, and stirred at 21,000 rpm for 30 min at room temperature for the second emulsion. The double emulsion was poured into isopropyl alcohol (40 ml, 2.5%) and mechanically agitated for 180 min at room temperature to volatilize the chloroform. The supernatant was removed after the solution was centrifuged at 4,800 rpm for 5 min. The precipitate was centrifuged again at 1,800 rpm for 5 min, and resuspended in superpure water. The superpure water wash was repeated several times until the supernatant became transparent. Precipitates were resuspended a final time in superpure water and stored at 4°C. DMs were sterilized via cobalt 60 (60Co) irradiation.

**RDM preparation**

Covalent bonding of the activated carboxyl groups on DM surfaces was performed using the 1-ethyl-3-[3-dimethylaminopropyl] carbodiimide hydrochloride (EDC) method in the presence of N-hydroxysuccinimide (NHS). Prepared DMs were resuspended in phosphate-buffered saline (PBS; pH 4.7), and EDC and NHS in an equimolar ratio were added into the suspension. The carboxyl groups were activated for 60 min at room temperature. The supernatant was removed after centrifugation, and the precipitate was resuspended in PBS. Amine-Peg2000-Biotin in MES buffer was added, and the mixture was incubated for 120 min at room temperature to obtain biotinylated DMs. Biotinylated DMs were incubated with avidin or dyLight488-labeled avidin (1mg/mL) for 10 min at room temperature. The mixture was then centrifuged three times and resuspended in PBS to remove surplus dyLight488-labeled avidin/avidin. Rituximab was biotinylated using the EZ-LinkTM Sulfo-LC-Biotinylation kit according to the manufacturer’s instructions, and was added to the avidin-biotin conjugated DMs and incubated for 10 min. The RDM suspension was rinsed three times and centrifuged to remove surplus biotinylated rituximab.

**RDM characterization**

We explored RDM morphologies using SEM (Hitachi S-3400N, Japan) and TEM (Hitachi H-7600, Japan), and determined RDM mean diameters and size distributions via dynamic light scattering (DLS) (Nanosizer-S, Malvern, London, UK). RDMs were also observed using a CLSM (Olympus, FV1000, JAPAN). Rituximab coupling efficiency was determined by measuring dyLight488-labeled avidin solution and biotinylated DMs suspension absorbances with a fluorescence spectrophotometer (Jasco, FP-6500, JAPAN) at a maximum excitation wavelength of 493 nm and maximum emission wavelength of 518 nm. Rituximab quantities on biotinylated DMs (binding efficiency (%)) were calculated as the ratio of the intensity of biotinylated DM to the intensity of the dyLight488-labeled avidin samples.

**Assessment of DOX loading**

Drug encapsulation efficiency was assessed by ultraviolet-visible spectrophotometry (Eppendorf, BioSpectrometer, Germany). A DOX solution standard curve was measured. Then, fresh RDMs were centrifuged and collected. The RDMs were destroyed using a 5% hydrochloric acid ethanol solution, and the mixture was centrifuged at 3000 rpm for 5 min. The optical density of the supernatant was determined at an excitation wavelength of 495 nm. Drug encapsulation efficiency was calculated using the following equation:

\[
\text{Encapsulation efficiency} = \frac{W_a}{W_b} \times 100\%
\]

where Wa represents the total amount of drug in the RDMs, and Wb represents the total weight of DOX used in the RDM preparation.

**Drug release assay**

To estimate DOX release, RDM suspensions were enclosed in dialysis bags (MWCO; 10,000 Da), which were placed in 50 mL of PBS with shaking at 100 rpm at 37°C. The suspension was then sonicated with ultrasound (power density = 1.2 W/cm², frequency = 1 MHz, duty cycle = 50%) for 60 s. At 0, 2, 4, 8, 10, 20, 30, 48, 60, and 72h, 1 ml of dialysate was extracted and stored at 4°C for analysis. An equal volume of PBS was added to the container to insure a constant volume. The concentration of DOX in the sample was determined using an ultraviolet spectrophotometer. DOX release was depicted as a function of time. The DM suspension was assessed using the same method.

**Cell cultures**

Human lymphoma B cell lines Raji (CD20+) and Daudi (CD20+), human lymphoma T cell line Jurkat (CD20−), and human T-acute lymphoblastic leukemia cell
line CEM (CD20–), were grown in RPMI-1640 medium with 10% (v/v) fetal bovine serum (FBS, Gibco, Australian origin) and 1% penicillin-streptomycin, and incubated in a humidified atmosphere at 37°C with 5% CO₂. For all experiments, cells growing in suspension were subcultured by centrifugation at a ratio of 1:4.

**Cell targeting assay**

After Raji and CEM cells were anchored in culture dishes with Poly L lysine solution, 50μl targeted RDMs and non-targeted DMs were added into the dishes. Shaking was used to encourage interactions in Raji cell cultures. After 30 min at room temperature, dishes were washed twice with PBS and observed via CLSM. Five dishes were used for each experiment group. Blocking tests were performed by pre-incubating Raji cells with rituximab for 30 min followed by washing to removing excessive rituximab. CD20– CEM cells were employed as a control, and non-specific uptake of RDMs by CEM cells was examined using the same methods.

**Cytotoxicity in vitro**

Raji, Daudi, Jurkat, and CEM cells were seeded in 96-well plates at 1x10⁵ cells/well in 100 μl of RPMI-1640 medium. Microbubble samples were adjusted to contain equal amounts of DOX. Cells were treated with DOX (0.5 μg/mL final concentration), DOX + rituximab, DMs combined with ultrasound (DMs + US), RDMs combined with ultrasound (RDMs + US) and RDMs + US. Raji cells were pre-incubated with excessive rituximab for 30 min (RDMs + US + rituximab). After 24, 48 and 72 h, the Cell Counting Kit-8 was used to detect viable cells in each treatment group relative to controls.

**Cell apoptosis in vitro**

Apoptosis (early and late stage) was determined using an Annexin V/propidium iodide apoptosis kit and flow cytometry (BD Biosciences, USA) with Cell Quest software. Raji, Daudi, Jurkat and CEM cells, after various treatments for 24, 48 and 72 h, were rinsed twice with cold PBS and resuspended in 195 μl of binding buffer solution. Cells were stained with 5 μl FITC-labeled Annexin V and 10 μl propidium iodide for 20 min at room temperature in the dark. Cells treated with medium alone were used as a control.

**DOX fluorescence intensity**

Treatments were as follows: DOX, DOX + rituximab, DMs + US, RDMs + US, and RDMs + US + rituximab at final DOX concentrations of 0.5 μg/ml. Raji, Daudi, Jurkat, and CEM cells were washed three times with cold PBS after 48 h treatment, centrifuged, and resuspended in 500 μl PBS. Cells treated with medium alone were used as controls. Intracellular DOX retention (red fluorescence) was examined using flow cytometry. Relative fluorescence intensity (RFI) was calculated as: \( \frac{FI_{\text{experiment}}}{FI_{\text{control}}} \).

**RDM-enhanced contrast ultrasound imaging in vivo**

Cells were inoculated subcutaneously into the backs of five nude mice per cell type, with 6×10⁷ Raji cells per mouse and 2×10⁷ Jurkat per mouse. Imaging was performed using an iU22 ultrasound system (Phillips, Netherland) with a 12 MHz ultrasound probe, 0.1 mechanical index and 54% gain. Mice were anesthetized by injecting 10% hydral and fixed to entirely expose the tumor under the ultrasound probe. Non-targeted DMs were injected first for imaging studies in grafted mice. After the expurgation of non-targeted DMs, the same amount of targeted RDMs was injected. The process was monitored continuously by ultrasonography. US contrast data were quantified with PHILIPS QLab 8.1 software. The arrival time, time to peak, peak intensity, and duration of contrast enhancement were determined. The Animal Ethics Commitment of the Southeast University approved all animal experiments.

**In vivo antitumor activity**

Raji and Jurkat cell-grafted mice were established additionally as described above. When lymphoma volume reached approximately 100 mm³, Raji and Jurkat cell-grafted mice were randomly divided into 6 groups (five mice per group), respectively : control group (saline), DOX, DOX + rituximab, DMs + US, RDMs + US, and RDMs + US + rituximab. Each mouse was treated with the appropriate formulation (3 mg/kg) three times per week. Lymphoma sizes in all mice were examined via digital caliper, and calculated using the equation: \( V_{\text{tumor}} = LW^2/2 \) (L: tumor length, W: tumor width). Lymphoma volumes and mouse body weights were determined before every injection. After 21 d of treatment, all mice were sacrificed. Lymphomas were extracted and fixed with 4% paraformaldehyde. To detect cell apoptosis in lymphoma tissues, tissues were sliced into thin sections and stained with a TUNEL apoptosis detection kit. Samples were then stained with DAPI to visualize cell nuclei under a CLSM.

**Statistical analysis**

All experiments were performed in triplicate. Data were presented as means ± standard deviation and analyzed using SPSS 16.0 software (SPSS, Chicago, IL, USA). Comparisons were performed using Student's t-test. P<0.05 was considered a significant difference.
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CONFLICTS OF INTEREST

The authors disclose no conflicts of interest.

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