Magnetic Nanoparticle-Based Hyperthermia Mediates Drug Delivery and Impairs the Tumorigenic Capacity of Quiescent Colorectal Cancer Stem Cells

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ABSTRACT: Cancer stem cells (CSCs) are the tumor cell subpopulation responsible for resistance to chemotherapy, tumor recurrence, and metastasis. An efficient therapy must act on low proliferating quiescent-CSCs (q-CSCs). We here investigate the effect of magnetic hyperthermia (MHT) in combination with local chemotherapy as a dual therapy to inhibit patient-derived colorectal qCR-CSCs. We apply iron oxide nanocubes as MHT heat mediators, coated with a thermoresponsive polymer (TR-Cubes) and loaded with DOXO (TR-DOXO) as a chemotherapeutic agent. The thermoresponsive polymer releases DOXO only at a temperature above 44 °C. In colony-forming assays, the cells exposed to TR-Cubes with MHT reveal that qCR-CSCs struggle to survive the heat damage and, with a due delay, restart the division of dormant cells. The eradication of qCR-CSCs with a complete stop of the colony formation was achieved only with TR-DOXO when exposed to MHT. The in vivo tumor formation study confirms the combined effects of MHT with heat-mediated drug release: only the group of animals that received the CR-CSCs pretreated, in vitro, with TR-DOXO and MHT lacked the formation of tumor even after several months. For DOXO-resistant CR-CSCs cells, the same results were shown, in vitro, when choosing the drug oxaliplatin rather than DOXO and applying MHT. These findings emphasize the potential of our nanoplatforms as an effective patient-personalized cancer treatment against qCR-CSCs.

KEYWORDS: magnetic hyperthermia, magnetic nanoparticles, doxorubicin, cancer stem cells, colorectal cancer

INTRODUCTION

Tumor heterogeneity is a major obstacle for the successful treatment of solid cancers.1 CSCs have been identified as the subpopulation of tumor mass able to self-renew through asymmetric divisions, providing fast proliferating cells and quiescent cells. The latter ones are able to repopulate and re-establish an entire tumor mass starting from one single cell under stress-related conditions. Therefore, the existence of this subset of tumor cells significantly contributes to tumor heterogeneity with consequent resistance to chemo- or radiotherapy.5,7,8 Hence, many efforts have been dedicated to identify and isolate CSCs from different tumors, including colorectal (CR) cancer, the third most common cancer worldwide, to study their survival to antitumoral treatment and to design more efficacious targeted therapies for their eradication.5,6,9 In particular, when using antiproliferating agents such as chemotherapeutic agents or radiation therapy, it was found that reduced cell cycling and increased DNA repairing capacity are typical resistance mechanisms of q-CSCs.
that make them less susceptible to therapy than the bulk cancer cells.\textsuperscript{10,11} Several studies have shown that by eradicating the fast proliferating tumor cells while sparing q-CSCs, the disease’s progression is only temporarily suppressed. In fact, killing the proliferating cells stimulates the reawakening of q-CSCs that have evaded therapy, eventually leading to cancer relapse.\textsuperscript{2,10} Among all the different therapeutic approaches for treating cancer, hyperthermia employs a moderate temperature (ca. 41–46 °C) to damage tumor cells.\textsuperscript{12} It is currently used in clinics as an adjuvant treatment to the more common types of antitumoral therapies.\textsuperscript{13} Indeed, if applied simultaneously or shortly after radiotherapy or chemotherapy, hyperthermia can interfere with the repair of therapy-induced DNA damage, thereby contributing to kill tumor cells.\textsuperscript{14–16} Novel nanoparticle-based platforms are emerging as a promising new technology to deposit heat in a more localized and selective mode enabling, simultaneously, the remote activation of the nanoparticles working as heat transducers. In MHT, the heat is produced by magnetic nanoparticles exposed to an alternating magnetic field (AMF) at a radiofrequency range (100 kHz) that has no body penetration restrictions and is not harmful for human patients. MHT has been introduced in clinics for the treatment of glioblastoma and is now under clinical trials for prostate and metastatic bone cancers.\textsuperscript{17,18} While many works have optimized the use of nanoparticles for cancer treatment, less is reported on the effect of nanoparticle-based hyperthermia on CSC eradication. In photothermal therapy, when using an infrared (IR) source to activate nanoparticles (such as carbon nanotubes\textsuperscript{19} or gold nanoparticles\textsuperscript{20}) and generate heat, clear indications of CSCs’ sensibility to heat were observed through viability studies, heat-shock protein (HSP) expression, and DNA damage analysis. However, low tissue penetration depth is a major drawback of light sources used in photothermal treatment.

Sadhuksa et al. demonstrated for the first time the in vivo potential of using superparamagnetic iron oxide nanoparticles (SPIONs) as MHT agents to reduce the tumorigenic potential of CSCs from immortalized adenocarcinoma cell lines.\textsuperscript{21} More recently, Kwon and co-workers have reported the use of magnetic nanoclusters to demonstrate the cytotoxic effect in vitro of MHT treatment on CSCs using a breast cancer cell line.\textsuperscript{22} While this study is pioneering the field of MHT on CSCs, it employs AMF field conditions (\( f = 290 \text{ kHz}; H = 60 \text{ kA/m} \)) that are not suitable for the safe application of MHT therapy in clinics (the product \( Hf \) should not exceed \( 5 \times 10^9 \text{ Am}^2 \text{ s}^{-1} \) to avoid patient discomfort due to eddy currents at the patient tissue).\textsuperscript{23} The effects of the MHT on CSCs urge to be conducted and evaluated under clinically applied conditions (100 kHz and 24 kA/m) or at an \( Hf \) factor that is clinically safe.\textsuperscript{23} This requires the use of magnetic nanoparticles with outstanding MHT heat properties.

In a more advanced study, Liu et al. exploited the CD20 antibody as a molecular target to accumulate silica-based magnetic nanoparticles on CSCs in a human lung cancer model.\textsuperscript{24} Furthermore, multiple cycles of MHT (up to 10 cycles) under clinically acceptable magnetic field conditions in combination with chemotherapy were performed to effectively reduce the tumor growth in vivo.

Recently, we have developed a magnetic nanoparticle-based platform consisting of cubic shape iron oxide nanoparticles (IONCs) coated with a thermoresponsive polymer (TR-Cubes) that can carry and release doxorubicin (TR-DOXO).\textsuperscript{25} Using such TR-DOXO, the release of DOXO is achieved following a heat stimulus provided by the nanocubes under MHT. Our group proved that the DOXO released from the TR-DOXO is significantly accelerated under clinically acceptable AMF conditions. The synergic therapeutic effects of the heat-mediated chemotherapy and the direct heat damage caused a complete eradication of human adenocarcinoma tumors in a xenograft mice model only when the animals were treated with the TR-DOXO and exposed to clinically MHT conditions.\textsuperscript{25} Remarkably, the dose of DOXO needed for the combination therapy was much lower than that used in other in vivo studies.\textsuperscript{26,27} This might eventually reduce the side effects of standard chemotherapy doses.

Herein, by exploiting this innovative magnetic nanoparticle platform (TR-DOXO), we aim to understand the effect of MHT in combination with heat-mediated chemotherapeutic drug release not only on the whole tumor but on a unique dormant cancer subset, the quiescent CSCs. For this purpose, we performed our experiments on CSCs, growing as spheroids, isolated from CR cancer specimens since their resistance to chemotherapeutic treatments is by far more aggressive than that of immortalized tumor cells. More specifically, we have implemented standard assays for identifying qCR-CSCs and study the effects of the MHT with TR-Cubes and TR-DOXO on those cells. Through a variety of in vitro analyses including colony-forming assay after fluorescence-activated cell sorting (FACS), confocal and transmission electron microscopy analysis of tumor cells, viability assays such as trypan blue, and apoptotic/necrotic cell quantification by flow cytometry analysis, our study proves the synergistic effects of the dual therapy (MHT and chemotherapy) to completely stop tumor growth. The use of MHT only or drug chemotherapy (DOXO) treatment alone has an effect on the overall viability of tumor spheroids, but it does not fully eradicate the qCR-CSCs as they reawake and regrow the tumor mass both in vitro and in vivo. Instead, when the cells are treated with TR-DOXO and exposed to MHT, this dual therapy enables the complete eradication of the tumor cells. Moreover, by checking over time the evolution of the number of alive PKH\textsuperscript{pos} qCR-CSCs, after MHT, we have found that the heat stress leads the qCR-CSCs to exit quiescence and start to divide to react and repopulate the tumor colony. At this stage, the DOXO released by the nanocubes is favorably internalized by the dividing CSCs, inducing an additional cytotoxic drug action on the tumor cells. These data were also supported by in vivo studies revealing the absence of initiation and relapse potential for the cells previously exposed in vitro to MHT with TR-DOXO.

\section*{RESULTS AND DISCUSSION}

Characterization and Isolation of Quiescent PKH\textsuperscript{pos} CR-CSCs. Spheroids derived from CR cancer patient tissues (hereafter referred to as CR-CSCs) were isolated as described by Todaro et al.\textsuperscript{28,29} CR-CSCs are characterized by the expression of different stemness markers, including CD133 and CD44v6, and by their in vivo tumorigenic potential.\textsuperscript{28,29} To study the effects of MHT using TR-Cubes or TR-DOXO on the qCR-CSC spheroid population, we needed to discriminate the qCR-CSCs within the spheroids. A diluting antibody as a molecular target to accumulate silica-based magnetic nanoparticles on CSCs in a human lung cancer model.\textsuperscript{24} Furthermore, multiple cycles of MHT (up to 10 cycles) under clinically acceptable magnetic field conditions in combination with chemotherapy were performed to effectively reduce the tumor growth in vivo.

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highly proliferative tumor cells were diluting PKH67 in a faster rate, leading to lower to no fluorescence (10–11 days) compared to the slow-dividing qCR-CSCs that retained a strong fluorescent signal for a prolonged period of time as observed by confocal microscopy (Figure 1a). This result was also corroborated by flow cytometry: nearly 10% of green stained cells, representing the qCR-CSCs (PKHpos), were found 11 days post-staining (Figure 1b). The self-renewal ability of the sorted PKHpos cells in respect to the nonstained fraction containing the fast proliferating tumor cells (PKHneg) was evaluated in vitro through a spheroid-forming assay. Plating a single sorted PKHpos qCR-CSC in a 96-well plate led to the formation of small spheroids by day 3, after which they continued to grow, developing a larger cell mass between day 6 and 2 weeks post-seeding, as observed under an optical microscope (Figure 1c). Contrarily, in PKHneg cells, colony formation from a single cell was not observed. In a comparative study, the colony formation assay was slightly modified: after FACS sorting, the PKHpos and PKHneg cells were separately seeded at low cell density (1000 PKHneg cells/well or 100 PKHpos cells/well) and the colony formation was monitored over time (Figure S1). The results confirmed the presence of PKHpos spheroids already after 2 days of culture, indicating a higher self-renewal capacity. On the contrary, PKHneg cells show a much slower growth rate with no signs of sphere formation even after 7 days of culture. Such outcome is in line with data from the literature reporting that fast proliferating cells eventually become exhausted after dividing.31 To further verify if the sorted PKHpos cells were enriched in CSCs, we evaluated the expression of CD44v6 as a marker of metastatic cells29 (Figure 1d). In accordance with functional studies, PKHpos CR-CSCs displayed higher levels of CD44v6 than PKHneg cells (Figure 1d). Likewise, PKHpos CR-CSCs showed a more than threefold upregulation in the expression of several genes showing a more than threefold change in PKHpos than PKHneg CR-CSCs are shown in red and green, respectively.

Figure 1. Identification of quiescent colorectal cancer stem cells (PKHpos qCR-CSCs #21). Proliferative PKH67 staining assay enables to identify, after 11 days, the qCR-CSCs by (a) confocal microscopy (the green cells) and (b) flow cytometry analysis. The y axis of the histograms represents the counts, while the x axis represents the fluorescent signal measured for green PKH67. (c) Self-renewal capability of PKHpos qCR-CSC sorted by FACS and cultured in an isolated environment (one cell per well, scale bars: 50 μm). (d) Flow cytometry analysis of CD44v6 in PKH67-stained CR-CSC (#21) after 11 days of PKH67 staining. (e) Stemness-related gene expression analysis of up- and downregulated genes in FACS-sorted PKHpos and PKHneg CR-CSCs following 11 days of PKH67 staining.
crucial stemness genes with respect to fast proliferating PKH\textsuperscript{neg} cells (Figure 1e and Figure S2).

**TR-Cubes and TR-DOXO Preparation and Characterization for the Treatment of CR-CSCs.** Iron oxide nanocubes (IONCs) functionalized with a thermoresponsive polymer (TR-Cubes), namely, poly(diethylene glycol methyl ether methacrylate-co-polyethylene glycol methyl ether methacrylate) (P(DEGMA-co-PEGMA)), were used as a smart nanocarrier to combine controlled drug delivery and MHT. The TR-Cubes were synthesized by the surface-initiated atom transfer radical polymerization (Photo-ATRP) following a recently published protocol of our group (see the scheme of the synthesis in Figure 2a). Initially, IONCs (18 nm) underwent ligand exchange with catechol functionalized ATRP initiators (DOPA-BiBAm) to introduce initiating sites on their surface. Afterward, the solution containing the modified ATRP initiator, the polymer monomers, and the copper catalyst was exposed to a commercial UV lamp (excitation at 365 nm) for a short period (3 h) to induce the formation of P(DEGMA-co-PEGMA) on IONCs’ surface. Recovery of TR-Cubes was obtained by precipitation in diethyl ether followed by ultracentrifugation in a sucrose gradient to remove the nongrafted polymer. Here, Photo-ATRP offers the peculiarity to grow a thick and uniform shell of polymer on the surface of IONCs in a quick manner, thus preventing the IONCs’ tendency to cluster. This feature results in the formation of individual IONCs coated with P(DEGMA-co-PEGMA) that is crucial to maintain their heating performance under clinically relevant MHT. Indeed, a SAR value as high as 250 Wg\textsuperscript{−1} (20 kA\textsuperscript{−1} m\textsuperscript{−1}, 110 kHz) was determined by calorimetric measurement. Additionally, as shown in Figure 2c, the TR-Cubes deposited from water exhibit a well-dispersed feature on TEM grid, indicating a high stability in aqueous media. In addition, the DLS trace of TR-Cubes in saline reveals a single peak at 86 nm with no sign of aggregation, thus being in good agreement with the TEM

result (Figure 2d, blue curve). Notably, a thick and uniform polymer shell surrounding IONCs can be identified at high-magnification TEM, which further confirms a successful surface-initiated polymerization step. Finally, by transmittance measurements, a transition temperature (lower critical solution temperature (LCST)) of 43 °C was determined for the TR-Cubes (Figure 2b).

Doxorubicin hydrochloride (DOXO) was loaded onto TR-Cubes by simple mixing of the drug and the nanocubes in a saline solution at room temperature for an overnight incubation. TR-DOXO were recovered and separated by free DOXO by means of magnetic collection, and the amount of loaded DOXO in the nanocubes was found to be of 40 μg/mg of Fe as determined by DMSO release, as previously described.\textsuperscript{25} Even the TR-DOXO show no sign of aggregation and very good stability in the solution (Figure 2c,d). Interestingly, 30% of loaded DOXO can be released within 1.5 h under MHT when a temperature of 45 °C is reached (110 kHz and 24 kA\textsuperscript{−1}), while at room temperature (25 °C), it took several days (8 days) to release the equivalent amount of DOXO just by diffusion. Along with the outstanding SAR value and the heat-triggered DOXO release, TR-DOXO are an interesting candidate for novel treatment against cancer stem cells.

**MHT Effect of TR-Cubes and TR-DOXO on CR-CSC Colonies.** Due to the peculiar shape of our cubic TR-Cubes and TR-DOXO that determined their outstanding SAR values and thus heat performances in MHT, the needed dose of magnetic materials to achieve a therapeutic temperature at the tumor is one order of magnitude lower than that of spherical nanoparticles employed in clinics.\textsuperscript{18,32,33} Moreover, we have also previously shown by \textit{in vivo} biodistribution studies that TR-Cubes can be renally excreted within a few months.\textsuperscript{25}

Herein, we first look at the effects caused by MHT with TR-Cubes and then the combination of MHT and chemotherapy with TR-DOXO.
To mimic the MHT treatment applied in vivo, 5 million CR-CSCs, simulating a tumor mass with a size of a few millimeters, were treated with TR-Cubes (50 μL at 4 g Fe/L or 0.2 mg Fe) or TR-DOXO (50 μL at 4 g Fe/L corresponding to 0.2 mg Fe and containing approximately 40 μg of DOXO/mg Fe) and exposed to MHT. First, we evaluated the effect of the MHT by varying the exposure time to the radiofrequency while ensuring that the therapeutic temperature reached during the MHT treatment never exceeded 45 °C. To achieve this constant temperature, the radiofrequency (f) was set at a clinically accepted 182 kHz and the field intensity was varied such that it never exceeded 27 kA/m (this also guaranteed that the H\textsubscript{f} factor was below 5 \times 10\textsuperscript{9} Am\textsuperscript{−1} s\textsuperscript{−1}) (see Figure 3 and Figure S3). MHT cycles were applied for 10, 30, 60 (2 \times 30 min), and 90 (3 \times 30 min) min. The three cycles of 30 min each of MHT correspond to MHT cycles as applied in our preclinical studies.\textsuperscript{25} The overall cell response was followed by monitoring the cell growth rate of the spheroids (Figure 3b and Figures S4–S6) and by estimating cell survival by trypan blue assay (Figure 3c–f). When the MHT treatment exposure...
was increased (from 30 to 90 min), a proportional decrease in cell survival and spheroid formation was observed (Figure 3c–e and Figures S4–S6). This was also accompanied by extensive and evident signs of cellular damage after longer MHT treatments as observed by morphological cell changes under TEM analysis of treated cells (Figures S7–S11). As also indicated by cell viability and cell growth curves (Figure 3c–f and Figure S12), the cells treated with TR-Cubes and TR-DOXO did not show a significant difference in their survival profiles at the first days after MHT treatment (up to day 3). These data suggest that in the first stage post-MHT, the cells’ suff erance is mainly due to the heat therapy.

However, between days 3 and 7, a remarkable cell viability difference occurs when comparing the cells treated with TR-DOXO or TR-Cubes exposed to MHT. For the TR-DOXO treated CR-CSCs and qCR-CSCs, a significant decrease in cell survival contrasting that of the cells treated with TR-Cubes was clearly visible, suggesting a synergistic therapeutic effect of DOXO in addition to the heat generated via MHT. Figure 3c–e shows that depending on the heat stress provided by the MHT exposure, the cellular survival response is tuned and this response is delayed if stronger stress stimuli (longer MHT treatments) are provided. For the cells treated with 10 or 30 min of MHT, this effect was observed between days 2 and 5, while for the cells exposed to 60 or 90 min of MHT, a later cellular response is observed around days 7–14 post-treatment. Noteworthily, in all cases in which sole MHT therapy was applied, we observed a critical period, post-MHT, after which the cells were able to recover from the damage and start regrowing the spheroids. Differently, for the TR-DOXO at all MHT durations, a lower cell resistance (higher cell death) was observed, with complete cell death only when applying 90 min of MHT (Figure 3e). It is worth noting that for the cells treated with TR-DOXO but not exposed to MHT, the nonspecifi c drug release provides a reduction of cell viability, but after a week of culture, the remaining alive cells were able to restart the spheroid growth (Figure 3f).

Overall, these data suggest that the samples treated with TR-DOXO are preferred to those treated with TR-Cubes and an overall MHT duration of 90 min is the most effi cient treatment to reach the best therapeutic eff ect and completely suppress the spheroid growth. These cytotoxicity data and MHT conditions are also in agreement with the magnetic fi eld conditions used by us in the in vivo efficacy study on a xenograft tumor model of adenocarcinoma when employing the same TR-DOXO.

To better understand the cytotoxic effects caused by the heat generated during MHT on the CR-CSCs, flow cytometry analysis was used to count the number of cells in apoptotic or necrotic stages (Figure 4a). Annexin/propidium iodide staining was conducted on the cells exposed to TR-Cubes for 90 min of MHT. Most of the CR cancer cells (>80%) died mainly by necrosis during MHT, while a small fraction was detected in the apoptotic stage (ca. 10%) and an even smaller
cellular fraction was found alive (4%), which may be responsible for regrowing the spheroids (Figure 3e).

To verify if, after 90 min of MHT, the quiescent PKHpos green cells were a fraction of the dying population, a confocal microscopy study was conducted after co-staining the cells with Sytox blue, a marker for dead cells. The number of PKHpos (green signal) cells colocalized with Sytox blue was counted at different time points post-MHT (Figure 4b and Figure S13).

Quantitatively, out of the PKHpos green-stained cells that represented the qCR-CSCs, we could count a progressively higher number of dying PKHpos cells over time, reaching 50% of dead PKHpos cells 5 days after MHT. This was followed by a gradual increase in the percentage of live PKHpos cells between days 5 and 15 from 43 to 67% indicating that the surviving PKHpos fraction started to divide and repopulate the culture.

Instead, when looking at the cells exposed to the same MHT treatment using TR-DOXO, remarkable differences were observed (Figure 5). One day post-MHT treatment, a clear red signal due to DOXO internalization was already distinctly present within cells. Colocalization of DOXO (red) and PKHpos cells (green) is clearly evident since each of these components occupies different subcellular compartments of the cells (DOXO in nucleus and PKH in cell membrane, Figure 5a and also Figures S14 and S15).

Figure 5. TR-DOXO dual-therapy effect on qCR-CSCs #21. (a) Confocal microscope analysis of CR-CSCs treated with TR-DOXO and exposed to MHT (90 min; three cycles of 30 min). PKHpos cell (green signal), Sytox blue (blue signal) for dead cells, and uptake of DOXO (red signal). The colocalization of green, blue, and red signals at day 5 confirms the death of quiescent PKHpos cells that have also internalized the released DOXO following MHT. (b) Spheroid growth monitoring after MHT (f = 182 kHz reaching a fixed temperature of 45 °C for 90 min; three cycles of 30 min) using TR-DOXO compared to control cells. The lack of spheroid formation on the figures on the right panel, at day 5, confirms the dual cytotoxic effects of the treatment. All scale bars are 50 μm. (c) Estimation of PKHpos cell death based on confocal images acquired using Sytox blue for 4 weeks after MHT and TR-DOXO administration. (d) Estimation of PKHpos cell death based on confocal images acquired using Sytox blue for 4 weeks after treatment with DOXO. For panels (c) and (d), at least 500 PKHpos cells were counted per each time point and per each experimental condition in n = 3 independent experiments. The mean data are reported, and error bars represent SD. Statistical analysis was done using ANOVA followed by Dunn’s multiple comparison test. The statistical difference ***p < 0.001 was calculated for each time point.
Additionally, after adding Sytox blue representing dead cells, at day 1, we observed lower incidences of PKH<sup>pos</sup> cell death, represented by no blue co-staining with PKH<sup>pos</sup> green cells. On the contrary, later by day 5, these PKH<sup>pos</sup> cells showed colocalization of the blue signal for the cell death with green and red signals. This represents dead PKH<sup>pos</sup> cells that had internalized DOXO after MHT (Figure S16).

The presence of DOXO in PKH<sup>pos</sup> cells and the sign of cell suff erance that are more evident at a prolonged time post-MHT may explain why the spheroid growth is compromised when the cells are exposed to MHT with TR-DOXO (Figure 5b).

Further quantification of PKH<sup>pos</sup> cells (green) and Sytox blue cells, based on the confocal images, confirmed the complete death of the qCR-CSCs after MHT with TR-DOXO (Figure 5c), contrarily to the cells treated with TR-Cubes + MHT (Figure 4b) or the cells treated only with DOXO (Figure 5d). These findings demonstrate that MHT and chemotherapy are capable of eradicating the tumor cell completely by directly affecting qCR-CSCs (PKH<sup>pos</sup> cells). To test the effectiveness of our nanoplatform for a future personalized treatment, the DOXO-sensitive CR-CSC<sub>21</sub> used for all the above-mentioned studies was compared to another patient-derived DOXO-resistant CR-cell line, CR-CSC<sub>27</sub>, in a long-term study conducted for at least 28 days.

Besides cell viability, we have also monitored the presence of DOXO signal over time in PKH<sup>pos</sup> cells. Together with TR-DOXO (see Figure 6c), free DOXO, at a dose that corresponded to the DOXO amount loaded on TR-DOXO, was also tested (it is worth noting that the amount of DOXO released by the TR-DOXO is always lower than the one loaded) (Figure S17).

As shown in Figure 6, even after 28 days, for CR-CSC<sub>21</sub>, the complete cell growth suppression was achieved only with TR-DOXO + MHT treatment (Figure 6a). On the contrary, for DOXO-resistant CR-CSC<sub>27</sub>, the survival rate suggests...
that no treatment is effective, as colonies were able to regrow after day 7 (Figure 6b). Notably, even the response to dual therapy when employing either TR-DOXO + MHT or DOXO + MHT is not enough to block the cell regrowth (Figure 6b and Figure S18).

To better understand why the dual therapy was effective for DOXO-sensitive CR-CSC #21 but not for DOXO-resistant CR-CSC #27, we focused on the effect of DOXO on the cells. On both patient cells, quantitative fluorescent analysis based on images acquired by confocal microscopy was used to estimate the number of PKH²⁰⁺ cells (green signal) that were also DOXO positive (red signal). We found that DOXO was present from the beginning of the drug treatment on both cell types and it was still present after a month on a small fraction of CR-CSC #21-PKH²⁰⁺ cells (ca. 17% of the cells) (Figure 6d). For CR-CSC #21, no regrowth of colony appeared up to 28 days (Figure 6a). Instead, on PKH²⁰⁺ CR-CSC #27, where a regrowth of cell colonies appeared 7 days post-treatment (Figure 6b), by analyzing confocal images, a remarkable decrease in DOXO red signal is clearly evident between days 1 and 7 post-MHT (Figure 6c,d).

These qualitative and quantitative data confirm that for these DOXO-resistant cells, the rate of expulsion of DOXO occurs very quickly between 72 h and day 7 (less % of red signal out of the PKH²⁰⁺ CR-CSC #27), in accordance with published data regarding the overexpression of drug-efflux transporters in CSCs. It is worth noting that even in this case to evaluate the toxicity of the treatment the main difference occurs between 3 and 7 days post-treatment. The DOXO elimination could suggest a possible mechanism of recovery of this DOXO-resistant cell line even if it went through MHT.

Next, to validate the efficiency of the dual treatment on the DOXO-resistant CR-CSC #27, we designed an experiment in which we co-administered the TR-Cubes as MHT heating agents together with oxaplatin (OXA), a drug that can overcome drug resistance shown by CR-CSC #27 (MHT + OXA, single step). The clinically sensitive dose range of OXA for CR-CSCs as reported by Todaro et al. is between 5 and 100 μM.28

We also aimed by combining the therapeutic effect of OXA and MHT to reduce the OXA dose required to achieve the elimination of CR-CSC #27. In a first set of experiments, CR-CSC #27 was first treated with TR-Cubes and exposed to MHT and then seeded in a fresh medium containing OXA at 10 or 100 μM (Figure 7a, treatment 1). Cells treated with MHT and 100 μM OXA were completely eradicated by day 7 post-treatment (Figure 7b, purple line). Contrarily, if a 10 μM OXA dose was administered to the cell media, a 70% cell mortality was reached by day 7 after treatment, after which the survival of the cells started to increase again (Figure 7b, green dashed line). Please note that cell viability was also tested for the free OXA drug in the solution at 50 μM (Figure S19). The viability data obtained are in accordance with those reported by Todaro et al.: there is about 60% cell death after 48 h. However, while the cells treated with 100 μM OXA were not able to recover from the treatment, the ones treated with 50 μM OXA started recovering very quickly as shown by the increase in cell viability after only 4 days post-treatment (Figure S19).

In a second set of experiments, TR-Cubes and OXA were co-administered directly to the cells before performing MHT (Figure 7a, treatment 2). Interestingly, the presence of OXA during the MHT treatment (Figure 7a, orange dashed line) brought to the complete cell elimination at a much lower OXA dose (10 μM). In fact, after 7 days, the cells do not show any sign of recovery, with a complete elimination achieved by day 12 post-treatment (Figure 7b). This result indicates the importance of having the drug during the MHT as, in this case, the heat facilitates its cell internalization and enables a more effective therapeutic effect at reduced drug dosage. Our TR-DOXO nanoplatfoemns are indeed able to encapsulate and release the drug only upon a heat stimulus, thus enabling a significant reduction in the drug dose.

**In Vivo Tumorigenicity Study.** To confirm the effective combinatorial therapeutic effects of our TR-DOXO nanoplatforaen, an in vivo tumor formation and relapse evaluation study was performed after the cells were exposed to the treatments in vitro. Nanoplatforaen were added to CR-CSC #21 (50 μL at 4 g Fe/L of TR-Cubes and TR-DOXO containing 50 mg of DOXO per gram of Fe) and exposed to MHT treatment at 45 °C in an ex vivo setup (three cycles at 182 kHz and the field intensity never exceeded 21.87 kA/m to maintain the 45 °C during the 30 min cycle). These treated cells were then injected subcutaneously in the flank of animals. With this

![Figure 7](https://doi.org/10.1021/acsaami.0c21349)

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In panels (a) and (b) we show the Schematic diagrams of experimental treatment for MHT combined with OXA administration. (a) Viability profiles, analyzed by trypan blue assay, for the DOXO-resistant CR-CSC #27 treated with MHT using TR-Cubes (red dashed line), TR-Cubes followed by OXA administration in the medium (100 μM, purple dashed line), TR-Cubes plus OXA administration in the medium (10 μM, green dashed line), and TR-Cubes plus OXA 10 μM added during hyperthermia (orange dashed line). All MHT treatments were performed for three cycles of 30 min at 45 °C (f = 182 kHz, H ~ 13). The blue dashed line indicates the control cells incubated at 37 °C. All values were calculated and normalized to the viability of the control cells at day 0.
study, the ability of CR-CSC #21 to reinitiate tumor growth was evaluated in a xenograft murine model (Figure 8a), after the cells were treated in vitro with the TR-Cubes or TR-DOXO with or without the exposure to MHT (three cycles of 30 min, at 45 °C). For the animal groups where no MHT exposure was foreseen, the nanoparticles were added to the

cells and incubated at 37 °C for the equivalent time of MHT before proceeding with the subcutaneous injection in the right flank of the 8-week-old female animals. We then monitored the tumor formation ability, tumor volume growth, animal survival, and general animal health state. As expected only for the animal group injected with CR-CSC #21 treated with TR-DOXO and exposed to MHT (TR-DOXO + MHT), in vivo tumor formation results showed a complete inhibition of tumor growth for the whole duration of the experiment (250 days). Indeed, none of the animals in the group exposed to the dual therapy developed tumors throughout the whole experimental duration. This was not the case for control animals injected with nontreated CR-CSC #21, the CR-CSC #21 treated with nonspecific DOXO release (TR-DOXO with no MHT exposure), or free DOXO (the latter administered at the same dose amount that is loaded within the TR-Cubes) or for the MHT heat damage alone (TR-Cubes + MHT). Animals belonging to these groups gradually developed tumors along the experiment, with different rates of tumor growth (Figure 8b and Figure S20). Instead, only in the group of the combined treatment of TR-DOXO + MHT was the complete tumor growth suppression induced, with no detectable tumor appearing in any of the animals of this group (Figure 8c). It is worth noting that, once more, the main viability difference between treatments occurs between days 3–7 post-treatment. Surprisingly, after 150 days of cell injection, two animals in the TR-Cubes + MHT group with absolutely no visible external tumor mass demonstrated physical signs of swallowed abdomen, and thus, they were sacrificed, respecting the humane end point of the study. Post-mortem analysis of these two animals showed the presence of bloody ascites in the peritoneal cavity, and few tumor nodules were found in the mesenterium and retroperitoneum, indicating the presence of metastasis (Figure S21e). These animals however had no evidence of cachexia and weight loss. These results are indicative of the inefficiency of hyperthermia alone (TR-Cubes + MHT) to fully eradicate the qCR-CSCs: the heat can indeed delay the tumor growth, but it cannot avoid resistant tumor cells to restart growing and cause cancer relapse or even more dangerous metastasis (Figure S21e). The presence of a low percentage of animals without tumor in DOXO, TR-DOXO, and TR-Cubes + MHT groups may be due to the random presence of quiescent CSCs in the cell population initially injected in the animal.37

Recent reports have suggested that the percentage of CSCs within a tumor cell population of patient cells can vary from 0.02 to 25% depending on the tumor type. However, this CSC fraction resisting treatments is sufficient to restart the tumor growth by an asymmetric cell division generating the growth of self-renewing cells (carrying stemness) and general progenitor tumor cells that can further differentiate. Survival rate was also monitored, and mice were sacrificed when the animal reached the human endpoint (the tumor reached 2 cm in one of its side). Control and TR-DOXO animal groups presented a higher probability of mortality due to the fast tumor growth before 100 days post-treatment. For DOXO and TR-Cubes + MHT groups, the probability of mortality was delayed to 150 days compared to control and TR-DOXO groups. This might be explained by the slower nonspecific DOXO release from the TR-DOXO compared to administering free DOXO directly. In contrast, the CR-CSC #21 treated with TR-DOXO + MHT led to significantly enhanced overall survival (probability = 1) relative to the control groups (p < 0.05, p = 0.030). Eight...
months after study initiation, all the animals of this group were still alive without tumor recurrence. (Figure 8d, Figure S20). All the animals in the study groups showed an increase in body weight over time. No significant differences in the weight of the animals among different treatment groups were observed (Figure 8e).

CONCLUSIONS

Our study demonstrates that patient-derived CR-CSCs are susceptible to the heat produced by TR-Cubes under multiple MHT exposures (3 × 30 min), at a therapeutic temperature of 45 °C, showing necrosis as the major cause of death. However, even after MHT, a small fraction of surviving qCR-CSCs can regrow the whole tumor mass. Interestingly, the longer the heat exposure time was, the longer was the delay of qCR-CSCs to reawake and restart the tumor growth (Figures 3 and 4). To inhibit the self-renewal capacity of this subcellular population, a dual therapeutic approach was required. Here, TR-DOXO, which combine the heat generated by the magnetic nanocubes under MHT with the triggered heat-mediated DOXO release, were proven to successfully eradicate qCR-CSCs (PKHpos stained cells) and avoid tumor relapse (Figures 3 and 6). Under MHT, the heat produced by TR-DOXO generates a stress on the qCR-CSCs cells, inducing them to divide. At the same time, the heat facilitates DOXO diffusion and prompts drug internalization by qCR-CSCs cells, thus making them more sensitive to DOXO (Figures 5 and 6). These in vitro findings were also supported by tumor initiation and relapse in vivo studies when using patient-derived DOXO-sensitive cells (CR-CSC #21). Notably, tumor growth was completely absent only in the animals injected with the cells pretreated with TR-DOXO and exposed to MHT (TR-DOXO + MHT, Figure 8).

The heat effect of TR-Cubes under MHT (TR-Cubes + MHT) was instead not able, by itself, to completely stop the tumor formation. Remarkably, in this study, the complete tumor inhibition with TR-DOXO was observed when applying MHT parameters that are used in clinics.

Importantly, we have also proved that the dual therapy can be adjusted to provide personalized therapy by choosing the most effective chemotherapeutic drug to be combined with TR-Cubes for MHT depending on the drug resistance of patient tumor cells. Indeed, for DOXO-resistant patient-derived CR-CSCs #27, the combination of MHT and DOXO did not work but the combination of MHT using TR-Cubes with OXA, another chemotherapeutic agents, could stop in vitro tumor cell growth (Figure 7). Interestingly, this could be achieved using a much lower dose of OXA (10 μM) than the standard OXA treatment (100 μM) only when the drug and MHT were co-administered. This outcome highlights the importance of using a smart multifunctional nanoplatform, like our TR-DOXO, to provide efficient hyperthermia and trigger chemotherapy in a single step to eliminate qCSCs. Finally, our data suggest that it is reasonable to predict in vitro the cellular response of patient cells to MHT in combination with chemotherapy in a short time, with a critical time window set around 5−7 days. We consider the results obtained in this work relevant for the clinical translation of MHT therapeutic effects in combination with drug release to develop a patient-specific personalized therapy. In perspective, efforts should be made to encapsulate different chemotherapeutic drugs in such types of magnetic nanoplatform, thus providing more efficient and patient-specific therapeutic tools.

MATERIALS AND METHODS

In Vitro Cell Culture. Colorectal (CR) cancer specimens were supplied by the “P. Giaccone” University Hospital following the ethical guidelines for human experimentation of the Institutional Ethics Committee. The isolation and propagation of CR-CSCs #21 and CR-CSCs #27 were performed as previously described.15 CR-CSCs were checked routinely for mycoplasma contamination by using the MycoAlert Plus Mycoplasma Detection Kit (Lonza). The authentication of CR-CSCs was assessed by performing a short tandem repeat (STR) analysis (GlobalFiler STR Kit, Applied Biosystems) followed by ABI Prism 3130x Genetic Analyzer (Applied Biosystems) sequencing. CR-CSCs (both CR-CSC #21 and CR-CSC #27) were cultured as spheroids in Corning ultra-low attachment flasks at 37 °C and 5% CO2. The cells were plated at an optimal density of 1 × 10^3 cells/mL in an Advanced DMEM/F-12 medium (Gibco) freshly supplemented with HEPES (10 mM), l-glutamine (2 mM), penicillin-streptomycin (100 U/mL), N-acetylcyteine (1 mM), N-2 supplement (1x), bFGF (100 ng/mL), EGF (50 ng/mL), B-27 (1x), Gastrin I (human) (10 mM), and nicotinamide (10 mM).

Identification of Quiescent CR-CSCs: Cellular Membrane PKH Labeling. Typically, the staining was performed after the cells were in culture for at least a week. The cells were labeled with the lipophilic fluorescent dye PKH67 (Sigma) according to the manufacturer’s protocol. Briefly, cells were washed and incubated with the dye for 4 min. The staining was blocked with fetal bovine serum (FBS), and the cells were seeded in the supplemented medium as described above. The medium was changed every 2 days, the cells were examined daily by an optical microscope (Motic AE31) to confirm the spheroid formation, and pictures were acquired by the confocal microscope (AIR Resonant Confocal System on TiE inverted microscope, NIS Elements software) to follow the fluorescent signal dilution.

Flow Cytometry of CD44v6 in PKH67-Stained CR-CSC (#21) and Stemness-Related Gene Expression Analysis. After being washed with PKH67, PKH67-stained CR-CSC (#21) cells were washed in PBS and stained with the CD44v6/APC antibody (2F10, R&D systems) or isotype matched control (IMC) (IC002A, R&D systems). Samples were analyzed using the BD FACs Melody cell sorter.

Total RNA extraction was performed by using the Trizol Reagent (Thermo Fisher). RNA samples were retrotranscribed with the iSCRIPT reverse transcription supermix kit (Bio-Rad). Real-time PCR for genes related to stemness was performed using a PrimePCR custom array (Bio-Rad). Data were analyzed with the Bio-Rad PRIME PCR analysis software.

Fluorescent-Activated Cell Sorting (FACS) of PKH-Positive (PKHpos) and PKH-Negative (PKHneg) Cells. For PKHpos and PKHneg discrimination, FACS sorting was performed on a single-cell suspension 11 days after labeling with PKH using a BD FACSAriaII to obtain PKHpos and PKHneg cells. After gating to eliminate debris and dead cells, thresholds were set based on the signal of a control unlabeled population. All cells that were found above the established threshold were considered PKHpos.

Spheroid Forming Ability of PKH-Sorted CR-CSCs. The bright fluorescent cells, based on PKH intensity (7−10%), representing the quiescent PKHpos were sorted and plated as a single cell per well in a 96-well plate with an ultra-low attachment surface. Spheroid formation was observed by using the optical microscope for acquiring pictures with the Motic Images Plus 2.0 digital software and the confocal microscope.

MHT In Vitro Experiments. A total of 5 × 10^6 cells were suspended in 50 μL of the complete medium containing 4 g Fe L−1 of TR-Cubes or TR-DOXO. The cell suspension was exposed to an alternating magnetic field (AMF) with a fixed frequency at 182 kHz generated by a NanoScale DM100 Series (Biomagnetics Corp.). The temperature was kept at 45 °C (see Figure S3). Following the treatment, the cells were seeded in ultra-low attachment flasks at the optimal density of 6 × 10^3 cells per mL and the viability plus spheroid growth was analyzed. Every 2 days, half of the medium was replaced with a fresh one and the cell density was diluted to half.
Spheroid Growth Assay. After MHT, the spheroids were monitored by taking pictures using the optical microscope and by counting the total number of cells using an automated cell counter (Nucleocounter NC-100, ChemoMetec).

Cell Viability. All methods were applied following the manufacturers’ procedures. The resistance of the CR-CSC population after exposure to MHT was evaluated by trypsin blue and Sytox blue analysis. Cells were analyzed immediately after treatment and then followed for at least a week. Images were acquired using a confocal microscope (A1R Resonant Confocal System on a TII inverted microscope; Nikon, Amsterdam, the Netherlands) for Sytox blue staining specific for dead cells (S34857, Thermo Fisher). To evaluate the induction of apoptosis/necrosis by flow cytometry dead cell apoptosis kit with annexin V–FITC and PI from InVitrogen assay was performed on triplicate samples.

TEM Analysis of Cellular Integrity and Nanoparticle Internalization after MHT. After treatment, the cell suspension was fixed and analyzed for transmission electron microscopy (TEM). At least 500,000 cells were incubated in a growth medium supplemented with glutaraldehyde (2%) for 45 min at room temperature (RT). Cells were then centrifuged at 14,000 rpm for 10 min. The obtained pellet was then dispersed in a Na-cacodylate buffer (0.1 M, pH 7.4) supplemented with glutaraldehyde (2%) and mixed for 1 h at RT. Afterward, cells were centrifuged at 14,000 rpm for 10 min. Three washes of 10 min each were repeated in the Na-cacodylate buffer (0.1 M). Subsequently, the pellet was incubated with the Na-cacodylate buffer (0.1 M) supplemented with OsO4 (1%) for 1 h at RT. Three washes of 10 min each in the Na-cacodylate buffer (0.1 M) were performed. Then, the pellet was washed three times for 5 min with mQ water and incubated overnight (ON) in a uranyl acetate buffer 1% (in water). Next, the sample was gradually dehydrated in ethanol (EtOH) at increasing concentrations of 70, 90, 96, and 100%. Then, the pellet was washed three times for 15 min with propylene oxide. Afterward, samples were incubated in a solution of Spurr and propylene oxide 1% (in water). Next, the sample was gradually dehydrated in ethanol (EtOH) at increasing concentrations of 70, 90, 96, and 100%. Then, the pellet was washed three times for 15 min with propylene oxide. Afterward, samples were incubated in a solution of Spurr and propylene oxide 1% (in water). Next, the sample was gradually dehydrated in ethanol (EtOH) at increasing concentrations of 70, 90, 96, and 100%. Then, the pellet was washed three times for 15 min with propylene oxide. Afterward, samples were incubated in a solution of Spurr and propylene oxide 1% (in water). Finally, the pellet was incubated for 3 h in Spurr and included into it by curing at 70 °C ON. A thin section of 70 nm of selected zones was observed with the JEOL Jem1011 electron microscope operated at 100 kV.

In Vivo Studies. The study was carried out in compliance with the protocol approved by the Italian Ministry of Health Protocol No. 059 according to the DLgs 116/1992. Pathogen-free 8-week-old female immunodeficient athymic NMRI nude mice were used for all the procedures. They were housed in IVC cages in a temperature-controlled room with a 12/12-h dark/light cycle, with ad libitum access to water and food. A total of 1 × 106 CR-CSC #21 colorectal cancer patient cells, which were previously treated with TR-DOXO (50 μL at 4 g Fe/L that corresponds to 0.2 mg Fe and containing approximately 8 μg of DOXO) and exposed to an alternating magnetic field of 22 kA/m and to a frequency of 182 kHz for 90 min (three cycles of 30 min each) (TR-DOXO + MHT experimental group), were injected subcutaneously in 150 μL of the complete cell culture medium (DMEM-F12 and Matrigel (1:1)) in the flank of each animal to induce the xenograft model. Animals that received a similar number of untreated cells (Control) or cells treated previously with TR-DOXO cubes only (50 μL at 4 g Fe/L that corresponds to 0.2 mg Fe and containing approximately 8 μg of DOXO), the TR-DOXO experimental group, DOXO free-drug (10 μg DOXO) (DOXO experimental group), or previously treated with TR-Cubes (50 μL at 4 g Fe/L that corresponds to 0.2 mg Fe) and exposed for 90 min to an alternating magnetic field (TR-Cubes + MHT experimental group), served as controls (n = 6 mice/group). Animals were observed every 3 days for the appearance of palpable tumors. Tumor dimensions were measured using a digital caliper once a week, and they were calculated using the following formula: V = (a × b2)/2, where a = length and b = width. The following marked the end of the study for each animal: when the tumor size reached 1.5–2 cm/side or when the tumors became ulcerated. Weight values of the animals were also recorded at the same time as the tumoral volume measurement. It was considered by veterinary advice that all the animals with a decrease of 20% in weight should be sacrificed in compliance with animal welfare rules.

Survival Study. Kaplan–Meier survival analysis was performed to plot the time vs the fraction of all individuals surviving as of that time. Each step in the curve marks an event in the study, which, in this case, is when the animal was sacrificed as a result of the tumor lengths reaching 2 cm/side. This curve was generated in the SigmaPlot software with a 95% confidence interval for fractional survival at any particular time.

Statistics. For in vitro studies, n = 3 and the statistical calculations within multiple groups were done using ANOVA and Tukey’s, or ANOVA and Dunnett’s multiple comparison test, as specified in the caption of each figure, with 95% confidence interval using the GraphPad or SigmaPlot software. Significant differences were reported as ***p < 0.001 unless reported otherwise in the figures.

For in vivo experiments, n = 6 animals per experimental condition were used. Randomization was used to allocate animals in different groups for the nude mice xenograft of CR-CSC #21 using the EXCEL software method. For the tumor growth and weight curve, the mean and SD of the tumor volume/animal weights at the end point of each group were analyzed using two-way ANOVA and pairwise multiple comparison procedures (Dunn’s method) post hoc nonparametric test. The survival analysis of the animals was done using a Kaplan–Meier survival plot in the SigmaPlot software. p value was calculated with the log-rank test.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.0c21349.

The Supporting Information is available free of charge on the ACS Publications Web site at DOI: Additional figures, as described in the text, include results from colony-forming assays, CSC gene expression studies, nanocube heating profiles, TEM image analysis of the cells after MHT, cellular growth curves, confocal images, cell death studies, and in vivo qualitative studies (Figures S1–S21) (PDF).

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Author Contributions
S.F and T.F. contributed equally to this work.

Author Contributions
T.P., S.F., and T.F. conceived and designed the study. S.F., T.F., S.M., and P.B.B. performed in vitro experiments. S.F., T.F., and C.D.M. performed in vivo experiments. A.T., S.D.F., G.S., and M.T. provided the patient-derived cells isolated in their laboratory as well as the in vitro and in vivo procedures established in their laboratory and performed the genetic characterization of CR-CSCs. B.T.M. and J.C. produced and characterized the materials. T.P., S.F., and T.F. wrote the paper with contributions obtained from all authors.

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Notes
The authors declare no competing financial interest.

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