

## NANOROBOTS

# A microrobotic system guided by photoacoustic computed tomography for targeted navigation in intestines in vivo

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Recently, tremendous progress in synthetic micro/nanomotors in diverse environment has been made for potential biomedical applications. However, existing micro/nanomotor platforms are inefficient for deep tissue imaging and motion control in vivo. Here, we present a photoacoustic computed tomography (PACT)-guided investigation of micromotors in intestines in vivo. The micromotors enveloped in microcapsules are stable in the stomach and exhibit efficient propulsion in various biofluids once released. The migration of micromotor capsules toward the targeted regions in intestines has been visualized by PACT in real time in vivo. Near-infrared light irradiation induces disintegration of the capsules to release the cargo-loaded micromotors. The intensive propulsion of the micromotors effectively prolongs the retention in intestines. The integration of the newly developed microrobotic system and PACT enables deep imaging and precise control of the micromotors in vivo and promises practical biomedical applications, such as drug delivery.

## INTRODUCTION

Micro- and nanorobots that can be navigated into hard-to-reach tissues have drawn extensive attention for the promise to empower various biomedical applications, such as disease diagnosis, targeted drug delivery, and minimally invasive surgery (1–6). Chemically powered motors, in particular, show great potential toward in vivo application owing to their autonomous propulsion and versatile functions in bodily fluids (7–11). However, imaging and control of micromotors in vivo remain major challenges for practical medical investigations, despite the substantially advanced development of micromotors (12–15). The ability to directly visualize the dynamics of micromotors with high spatiotemporal resolution in vivo at the whole-body scale is in urgent demand to provide real-time visualization and guidance of micromotors (14). In addition to high spatiotemporal resolution, the ideal noninvasive micromotor imaging technique should offer deep penetration and molecular contrasts.

To date, optical imaging is widely used for biomedical applications owing to its high spatiotemporal resolution and molecular contrasts. However, applying conventional optical imaging to deep tissues is hampered by strong optical scattering, which inhibits high-resolution imaging beyond the optical diffusion limit (~1 to 2 mm in depth) (16). Fortunately, photoacoustic (PA) tomography (PAT), detecting photon-induced ultrasound, achieves high-resolution imaging at depths that far exceed the optical diffusion limit (17). In PAT, the energy of photons absorbed by chromophores inside the tissue is converted to acoustic waves, which are subsequently detected to yield high-resolution tomographic images with optical contrasts. Leveraging the negligible acoustic scattering in soft tissue, PAT has achieved superb spatial resolution at depths, with a depth-to-resolution ratio of ~200 (18), at high imaging rates. As a major incarnation of PAT, PA computed tomography (PACT) has attained

high spatiotemporal resolution (125- $\mu\text{m}$  in-plane resolution and 50- $\mu\text{s}$  frame<sup>-1</sup> data acquisition), deep penetration (48-mm tissue penetration in vivo), and anatomical and molecular contrasts (text S1) (19–21). With all these benefits, PACT shows promise for real-time navigation of micromotors in vivo for broad applications, particularly drug delivery.

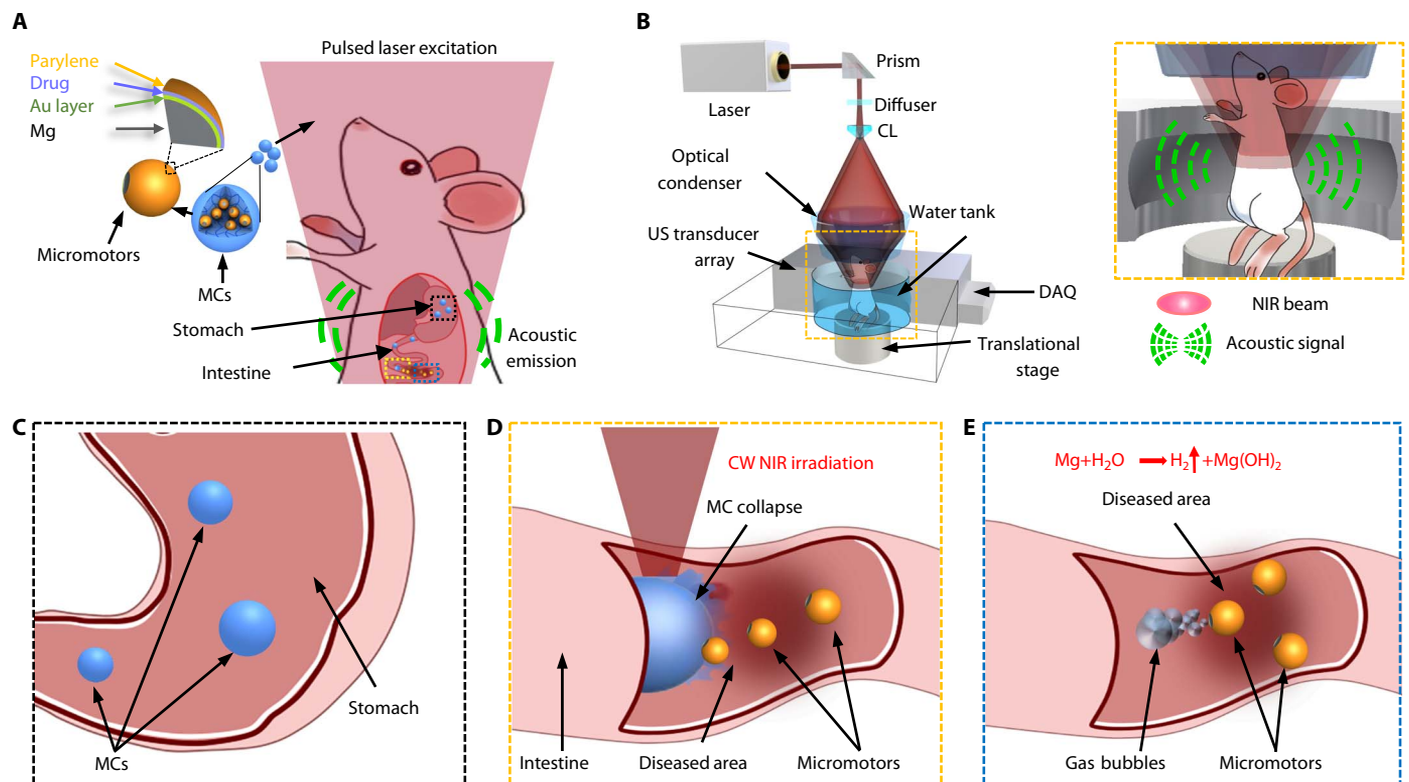
Drug delivery through the gastrointestinal (GI) tract serves as a convenient and versatile therapeutic tool, owing to its cost effectiveness, high patient compliance, lenient constraint for sterility, and ease of production (22, 23). Although oral administration of various micro/nanoparticle-based drug delivery systems has been demonstrated both to survive the acidic gastric environment and to diffuse into the intestines, drug absorption is still inefficient because of the limited intestinal retention time (24). A large number of passive diffusion-based targeting strategies have been explored to improve the delivery efficiency, but they suffer from low precision, size constraint, and specific surface chemistry (25). With precise control, microrobotic drug delivery systems can potentially achieve targeted delivery with long retention times and sustainable release profiles, which are in pressing need (26). Because of the lack of imaging-guided control, there is no report yet for precisely targeted delivery using micromotors in vivo (14). In addition, biodegradability and biocompatibility are required, and an ideal microrobotic system is expected to be cleared safely by the body after completion of the tasks (5, 27, 28).

Here, we describe a PACT-guided microrobotic system (PAMR), which has accomplished controlled propulsion and prolonged cargo retention in vivo (Fig. 1A and movie S1). Because of high spatiotemporal resolution, noninvasiveness, molecular contrast, and deep penetration, PACT provides an attractive tool to locate and navigate the micromotors in vivo (Fig. 1B) (18–20). Ingestible Mg-based micromotors were encapsulated in enteric protective capsules to prevent reactions in gastric acid and allow direct visualization by PACT (Fig. 1, A to C). PACT monitored the migration of micromotor capsules (MCs) in intestines in real time; continuous-wave (CW) near-infrared (NIR) light irradiation induced phase transition of microcapsules and triggered propulsion of the micromotors (Fig. 1D); autonomous and efficient propulsion of the micromotors enhanced the retention in

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**Fig. 1. Schematic of PAMR in vivo.** (A) Schematic of the PAMR in the GI tract. The MCs are administered into the mouse. NIR illumination facilitates the real-time PA imaging of the MCs and subsequently triggers the propulsion of the micromotors in targeted areas of the GI tract. (B) Schematic of PACT of the MCs in the GI tract in vivo. The mouse was kept in the water tank surrounded by an elevationally focused ultrasound transducer array. NIR side illumination onto the mouse generated PA signals, which were subsequently received by the transducer array. (Inset) Enlarged view of the yellow dashed box region, illustrating the confocal design of light delivery and PA detection. US, ultrasound; CL, conical lens; DAQ, data acquisition system. (C) Enteric coating prevents the decomposition of MCs in the stomach. (D) External CW NIR irradiation induced the phase transition and subsequent collapse of the MCs on demand in the targeted areas and activated the movement of the micromotors upon unwrapping from the capsule. (E) Active propulsion of the micromotors promoted retention and cargo delivery efficiency in intestines.

targeted areas of the GI tract (Fig. 1E). We believe that the proposed integrated microbotic system will substantially advance GI therapies.

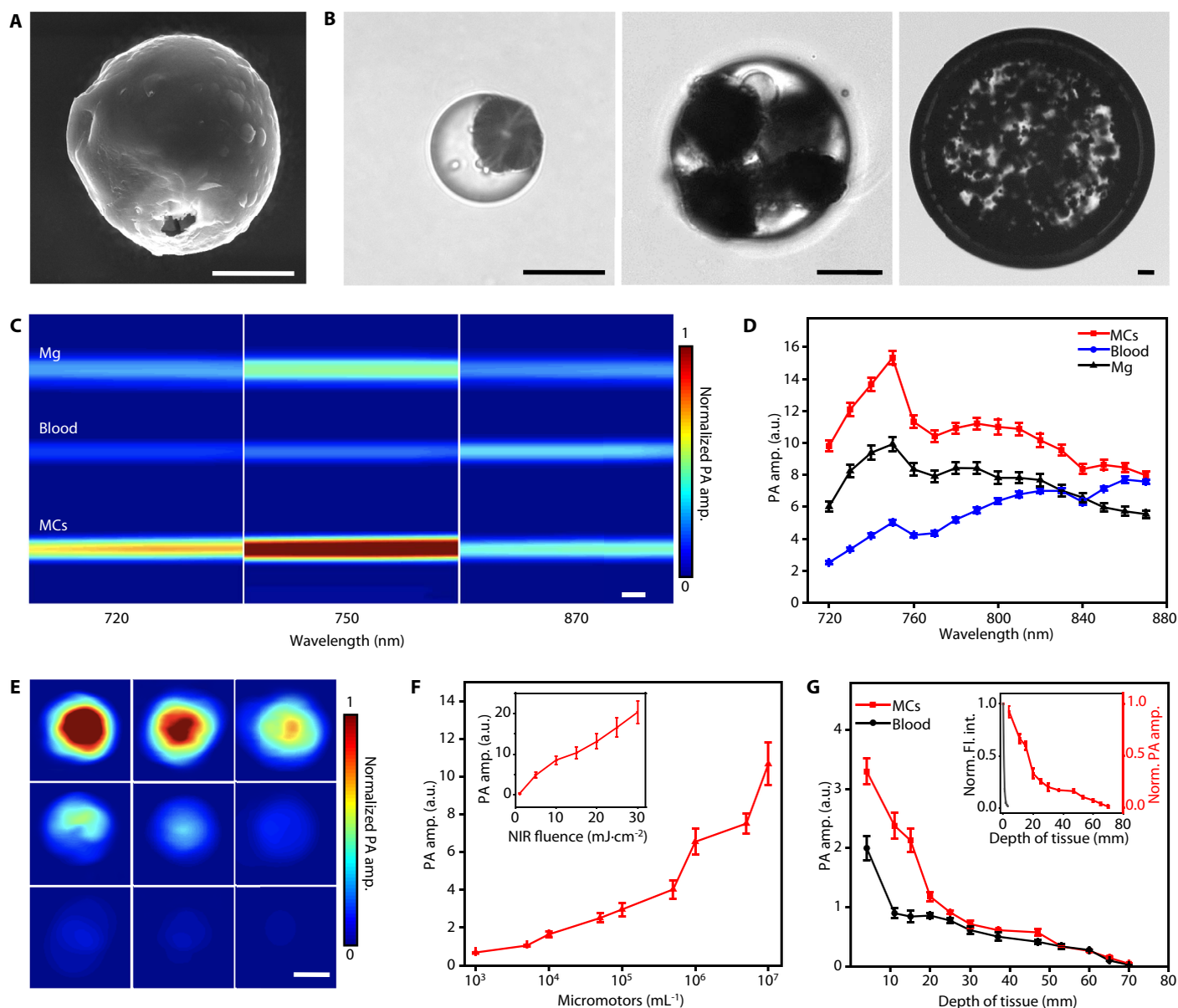
## RESULTS

### Fabrication of the MCs

The fabrication of MCs mainly consists of two steps: the fabrication of Mg-based micromotors (see fig. S1 and Materials and Methods) and the formation of MCs (see fig. S2 and Materials and Methods). In the first step, Mg microparticles with a diameter of  $\sim 20 \mu\text{m}$  were dispersed onto glass slides, followed by the deposition of a gold layer, which facilitates the autonomous chemical propulsion in GI fluids and enhances PA contrast of the micromotors. An alginate hydrogel layer was coated onto the micromotors by dropping aqueous solution containing alginate and drugs (e.g., doxorubicin) on the slides. A parylene layer, acting as a shell scaffold that ensures the stability during propulsion, was then deposited onto the micromotors. Figure 2A illustrates a fabricated spherical micromotor ( $\sim 20 \mu\text{m}$  in diameter). A small opening ( $\sim 2 \mu\text{m}$  in diameter), attributed to the surface contact of the Mg microparticles with the glass slides during various layer coating steps, acts as a catalytic interface for gas propulsion in the intestinal environment. Next, the micromotors were encapsulated into the enteric gelatin capsules by the emulsion method (fig. S2). Green fluorescence from the fluorescein isothiocyanate-labeled bovine serum

albumin (FITC-albumin) and red fluorescence from doxorubicin (DOX) were observed from the micromotors (see fig. S3 and Materials and Methods) and the MCs (see fig. S4 and Materials and Methods), confirming a successful drug loading. The size of MCs could be varied by changing the speed of magnetic stirring (fig. S5). The microscopic images in Fig. 2B show three MCs with diameters of 68, 136, and  $750 \mu\text{m}$ .

For deep tissue imaging in vivo, it is crucial that the MCs have a higher optical absorption than the blood background. To evaluate the PA imaging performance of the MCs, we measured the PA amplitudes of the MCs, whole blood, and bare Mg particles (see Materials and Methods). NIR light experiences the least attenuation in mammalian tissues, permitting the deepest optical penetration. As shown in Fig. 2C, the MCs exhibit strong PA contrast in the NIR wavelength region, ranging from 720 to 890 nm. To assess quantitatively the optical absorption of the MCs, we extracted amplitude values from the above PA images and subsequently calibrated them with optical absorption of hemoglobin (29, 30). At the wavelength of 750 nm, the MCs display the highest PA amplitude of 15.3 (Fig. 2D). The bare Mg particles display a similar PA spectrum, with a lower PA peak with an amplitude of 10.0 at 750 nm. The difference due to the Au layer is expected to significantly improve the imaging sensitivity in the NIR wavelength region (Fig. 2D) (31, 32). In addition, the approximate threefold increase in PA amplitudes of the MCs compared with that



**Fig. 2. Characterization of the MCs.** (A) SEM image of an ingestible micromotor. Scale bar, 10  $\mu\text{m}$ . (B) Microscopic images of the MCs with different sizes. Scale bars, 50  $\mu\text{m}$ . (C) PACT images of Mg particles, blood, and MCs in silicone rubber tubes with laser wavelengths at 720, 750, and 870 nm, respectively. Scale bar, 500  $\mu\text{m}$ . (D) PACT spectra of MCs (red line), blood (blue line), and Mg particles (black line). (E and F) PACT images (E), the corresponding PA amplitude (F) of the MCs with different micro-motor loading amounts, and the dependence of the PA amplitude on the fluence of NIR light illumination [inset in (F)]. Scale bar, 500  $\mu\text{m}$  (E). (G) Dependence of PA amplitude of the MCs (red line) and blood (black line) on the depth of tissue and the normalized PA amplitude and fluorescence intensity of the MCs under tissues (inset). Norm., normalized; amp., amplitude; Fl, fluorescence; int., intensity. Error bars represent the SDs from five independent measurements.

of the whole blood provides sufficient contrast for PACT to detect the MCs *in vivo* using 750-nm illumination. To evaluate the stability of the MCs under pulsed NIR PA excitation, we measured the PA signal fluctuation of the MCs during PA imaging (fig. S6). The negligible changes in the PA signal amplitude during the operation suggest a remarkably high photostability of the MCs. Figure 2 (E and F) shows the PA images and the corresponding PA amplitudes of single MCs with different concentrations of micromotors. The dependence of the PA amplitude on the NIR light fluence (i.e., energy per area) was also investigated (see Materials and Methods). As expected, the

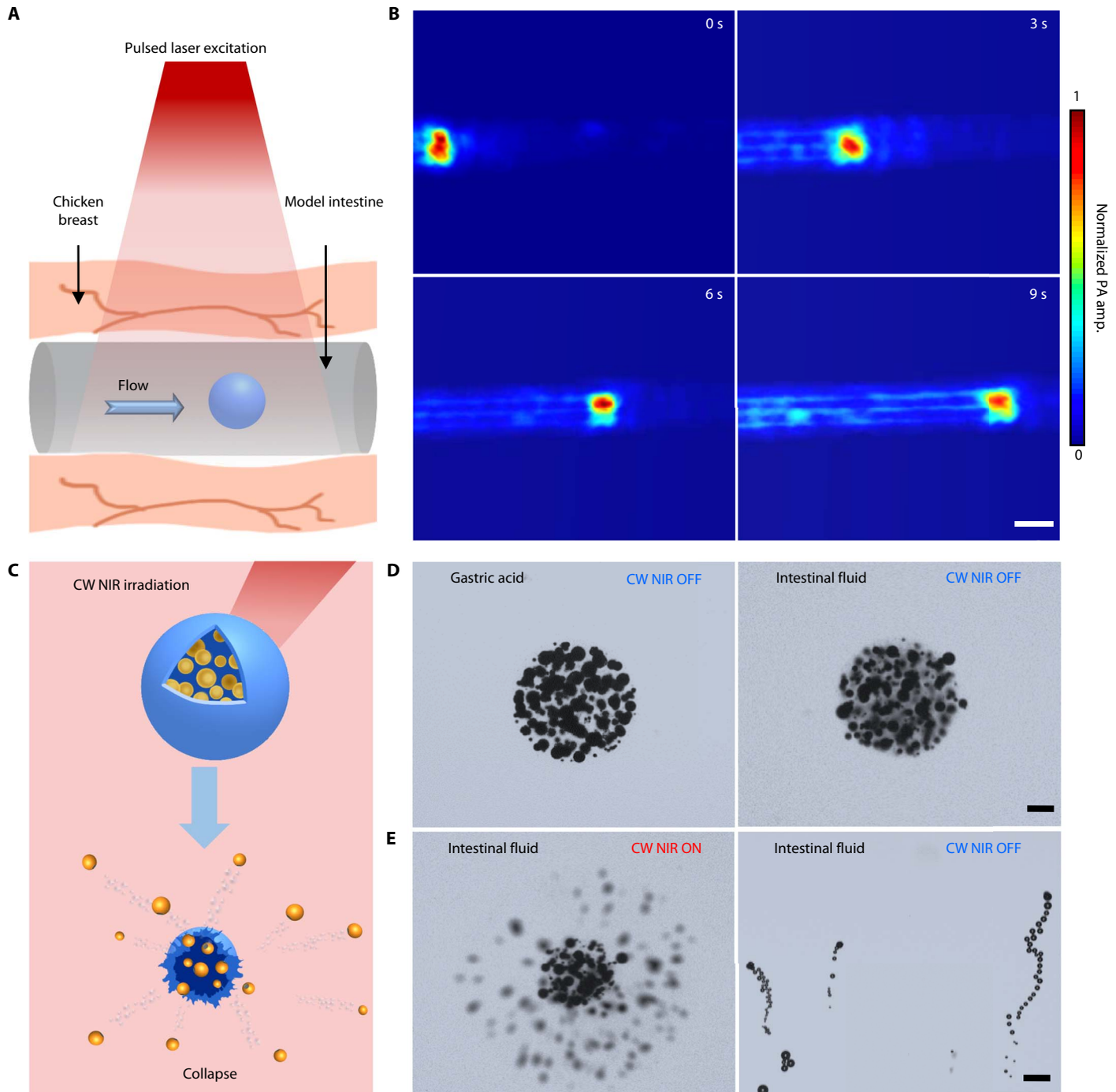
PA amplitude of the micromotors almost linearly increases with the NIR light fluence (Fig. 2F, inset). We also studied the maximum detectable depth of MCs using PACT (see Fig. 2G and Materials and Methods). The micromotors showed markedly decreased fluorescence intensity when covered by thin tissues (0.7 to 2.4 mm in thickness) and became undetectable quickly [Fig. 2G (inset) and fig. S7]. By contrast, PACT could image the micromotors inside tissue as deep as  $\sim 7$  cm (Fig. 2G), which reveals that the key advantage of PACT lies in high spatial resolution and high molecular contrast for deep imaging in tissues (19).

**Characterization of the dynamics of the PAMR in vitro**

The high optical absorption of the MCs empowers the PAMR as a promising in vivo imaging contrast agent. To evaluate the dynamics of the PAMR, we conducted the PA imaging experiments initially in vitro, where silicone rubber tubes modeled intestines (see Materials and Methods). The tubular model intestine was sandwiched in chicken

breast tissues (Fig. 3A). The PA time-lapse images in Fig. 3B and movie S2 illustrate real-time tracking of the migration of an injected MC in the model intestine.

In addition to tracking and locating the MCs, propulsion of the micromotors upon unwrapping from the microcapsules could be activated on demand with high-power CW NIR irradiation (see Fig. 3C



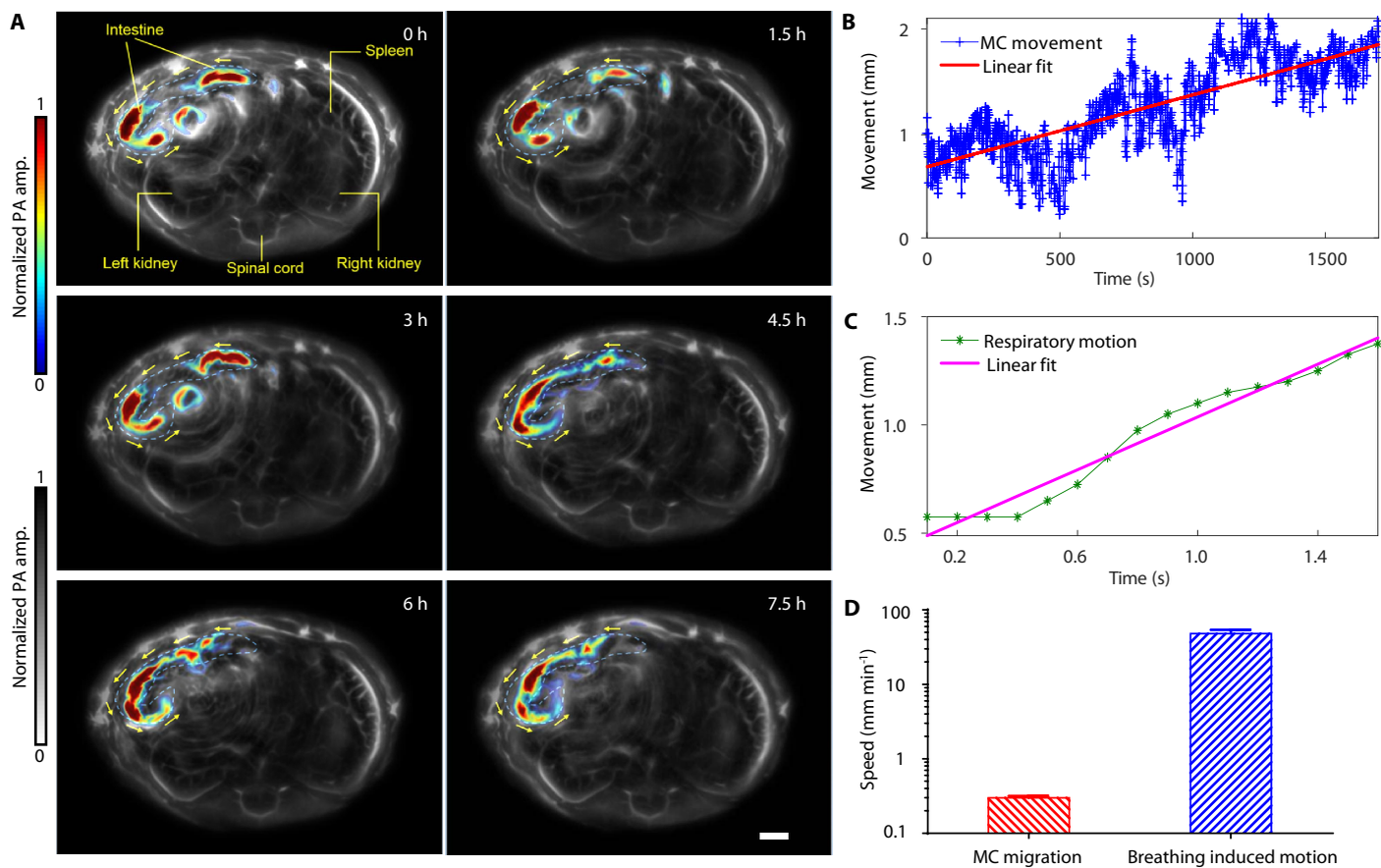
**Fig. 3. Characterization of the dynamics of the PAMR.** (A and B) Schematic (A) and time-lapse PACT images in deep tissues (B) illustrating the migration of an MC in the model intestine. Scale bar, 500  $\mu\text{m}$ . The thickness of the tissue above the MC is 10 mm. (C to E) Schematic (C) and time-lapse microscopic images (D and E) showing the stability of the MCs in gastric acid and intestinal fluid (D) without CW NIR irradiation and the use of CW NIR irradiation to trigger the collapse of an MC and the activation of the micromotors (E). Scale bars, 50  $\mu\text{m}$  (D and E).

and Materials and Methods). Because of the enteric coating and gelatin encapsulation, the MCs showed long-term stability in both gastric acid and intestinal fluid (Fig. 3D and fig. S8). The Au layer of the micromotors could effectively convert NIR light to heat, resulting in a gel-sol phase transition of the gelatin-based capsule followed by the release of the micromotors. Such CW NIR-triggered disintegration of the MCs usually occurred within 0.1 s. Therefore, CW NIR irradiation could activate autonomous propulsion of the micromotors (Fig. 3E and movie S3). Such a photothermal effect also significantly accelerated the Mg-water chemical reactions and thus enhanced the chemical propulsion of the micromotors. As shown in fig. S9 and movie S4, the micromotors exhibited efficient bubble propulsion in various biofluids. Further quantitative analysis indicates that the velocities of the micromotors were 45 and 43  $\mu\text{m s}^{-1}$  in phosphate-buffered saline (PBS) solution and the model intestinal fluid, respectively. Note that bare Mg particles have negligible propulsion in neutral media (i.e., intestinal fluid) and disordered propulsion in acidic condition (see fig. S10 and Materials and Methods). The highly efficient propulsion in the targeted areas in intestines provides a mechanical driving force to enhance retention and delivery. The required NIR power can be potentially adjusted by controlling the synthesis process and composition of the MCs. Other triggering mechanisms in biomedicine, such as magnetic

or ultrasonic fields, can also be used to activate propulsion of the micromotors (33).

### Dynamic imaging of the PAMR in vivo

The movement of a swarm of MCs was monitored in vivo by PACT (see Materials and Methods). The MCs were dispersed in pure water and then orally administered into 5- to 6-week-old nude mice. The mice were subsequently anesthetized, and the lower abdominal cavity was aligned with the imaging plane of the ultrasonic transducer array for longitudinal imaging (Fig. 1B). PACT images were captured at a frame rate of 2 Hz for ~8 hours (in Fig. 4A and movie S5, the blood vessels and background tissues are shown in gray, and MCs in intestines are highlighted in color). During the imaging period of the first 6 hours, the MCs migrated ~1.2 cm, roughly 15% of the length of the entire small intestine. After 5 hours, the PA signals of some MCs faded away as they moved downstream in intestines that were outside of the imaging plane. The moving speed of the swarm MCs in the intestines and the movements induced by respiratory motion were quantified (Fig. 4, B to D, and fig. S11). As shown in Fig. 4 (B to D), the abrupt motion caused by respiration was much faster than actual migration of the MCs. Despite the respiration-induced movement, PACT could distinguish the signals from the slowly migrating MCs in the



**Fig. 4. PACT evaluation of the PAMR dynamics in vivo.** (A) Time-lapse PACT images of the MCs in intestines for 7.5 hours. The MCs migrating in the intestine are shown in color; the mouse tissues are shown in gray. Scale bar, 2 mm. (B and C) Movement displacement caused by the migration of the MCs in the intestine (B) and by the respiration motion of the mouse (C). (D) Comparison of the speeds of the MC migration and the respiration-induced movement. Error bars represent the SDs from three independent measurements.

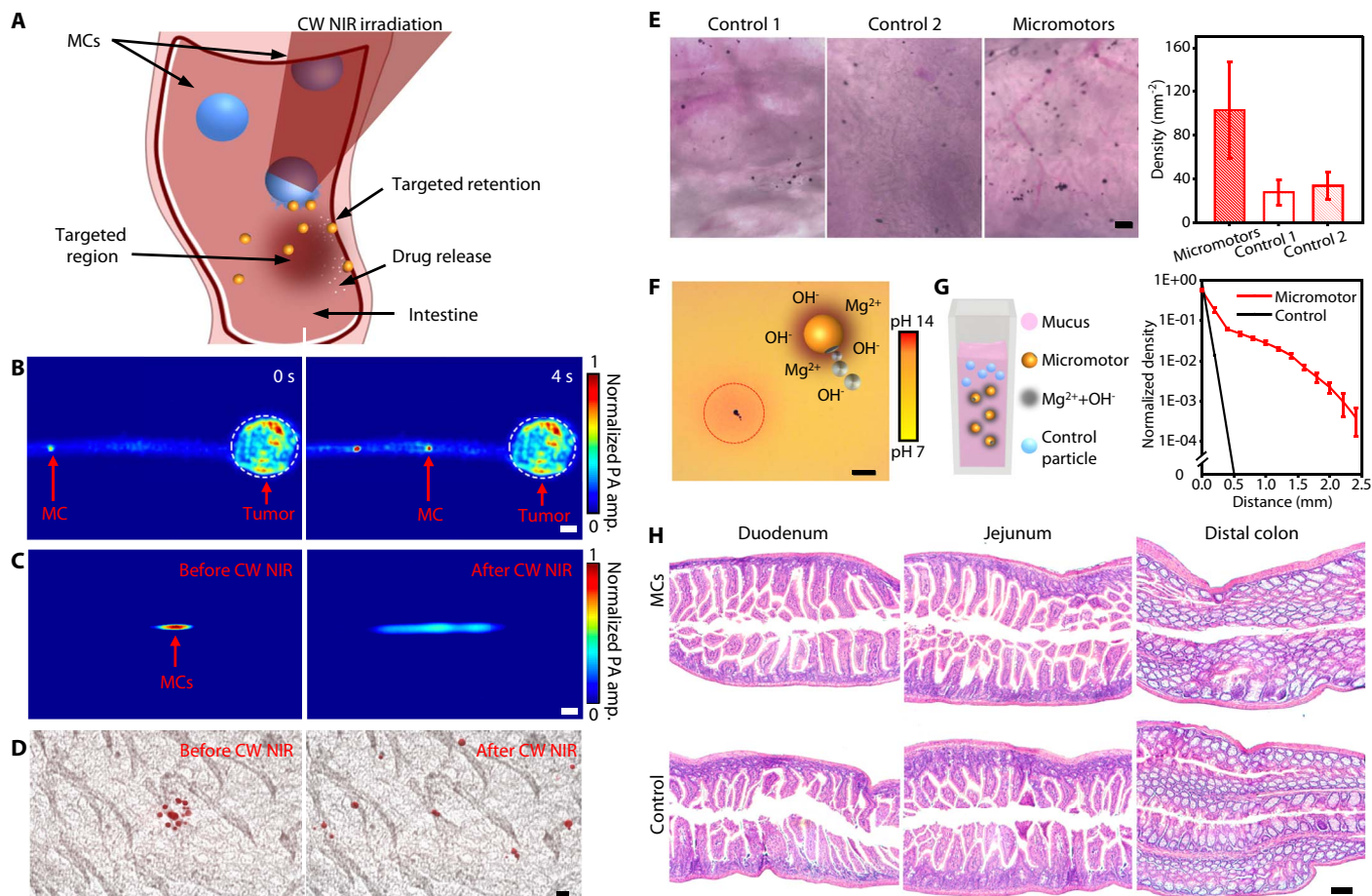
intestines (see Materials and Methods). These results indicate that PACT could precisely monitor and track the locations of the MCs in deep tissues in vivo.

### The evaluation of the PAMR toward targeted retention and delivery

Of particular biomedical significance is the retention of cargo carriers in the targeted region in intestines. Although most of the previous studies focused on improving the interactions between particles and the mucoadhesives by engineering surface coatings on the passive particles, the biofluid-driven propulsion of the active micromotors can markedly prolong their retention in intestine walls. When the MCs approached the targeted areas in intestines, we could trigger the collapse of the capsules and activate the propulsion of micromotors on demand (see Fig. 5A and Materials and Methods). To investigate the use of the PAMR for targeted delivery, we grew melanoma cells in mouse intestines and coated the intestines with tis-

ues as a model *ex vivo* colon tumor. Owing to the high optical absorption of melanoma cells in the NIR wavelength region, colon tumors could be resolved by PACT. After injection into the intestines, the MCs migrated toward the targeted colon tumor, as illustrated by the time-lapse PACT images in Fig. 5B and movie S6. Once the MCs approached the targeted location, they were irradiated with CW NIR light to trigger a responsive release of the micromotors. The PA signals from the MCs in the intestines were prolonged upon the CW NIR irradiation, suggesting the release of the micromotors (Fig. 5C). The overlaid microscopic images in Fig. 5D show the NIR-triggered release of the micromotors from an MC in the intestines. The DOX-loaded micromotors, observed with red fluorescence, rapidly diffused into the surrounding area after the CW NIR irradiation (see Fig. 5D and Materials and Methods).

To evaluate retention of the micromotors in vivo, we orally administered the enteric polymer-coated micromotors and the paraffin-coated passive Mg and Mg/Au particles (as the Control 1 and Control 2)



**Fig. 5. Evaluation of the PAMR for targeted retention and delivery.** (A) Schematic of the use of the PAMR for targeted delivery in intestines. (B) Time-lapse PACT images of the migration of an MC toward a model colon tumor. Scale bars, 500  $\mu\text{m}$ . (C and D) PACT images (C) and overlaid time-lapse bright-field and fluorescence microscopic images (D) showing the retention of the micromotors in intestines via the NIR-activated propulsion of the micromotors. Scale bars, 200  $\mu\text{m}$  (C) and 20  $\mu\text{m}$  (D). (E) Microscopic images showing the in vivo retention of the control microparticles and the micromotors in intestines (left) and the quantitative analysis of the particle retention in intestines (right). Control 1 and Control 2 represent paraffin-coated passive Mg and passive Mg/Au microparticles, respectively. Scale bar, 100  $\mu\text{m}$ . Error bars represent the SDs from five independent measurements. (F) Microscopic image displaying the change of pH of the surrounding environment upon the micromotors in PBS. (G) Schematic (left) and the experimental (right) diffusion profiles of control silica particles and ingestible micromotors in mucus after 1 hour. Error bars represent the SDs from five independent measurements. (H) Histology analysis for the duodenum, jejunum, and distal colon of the mice treated with the MCs or DI water as the control for 12 hours. Scale bar, 100  $\mu\text{m}$ .

into three mouse groups that underwent a fasting treatment for 8 hours. The mice were euthanized 12 hours after the administration, and their GI tracts were collected to evaluate the retention of the micromotors (see Materials and Methods). The intestines from the mice treated with micromotors retained a much higher number of micromotors than those with passive particles (Fig. 5E, left). The quantitative analysis displays a three- to fourfold increase in the density of the micromotors in the treated intestine segments (Fig. 5E, right). Note that nearly all Mg had already degraded in the retained micromotors 12 hours after administration, as illustrated by the hollow structures of the micromotors in the intestine before and after acid treatment (fig. S12). These results confirm the capability of PAMR for prolonged retention in targeted areas in intestines. Besides the active propulsion, the enhanced retention *in vivo* may also be attributed to the elevated pH and  $Mg^{2+}$  concentration in the surrounding environment caused by Mg-water reactions (see Fig. 5F and Materials and Methods) (33, 34). It has been recently reported that high pH (~8.2 to 12.0) could trigger a phase transition of the mucus and facilitate tissue penetration of the micro/nanoparticles (33–36). To investigate the influence of the micromotors on the pH of the surrounding environment, we dispersed the micromotors in water with phenolphthalein as a pH indicator. The microscopic image in Fig. 5F shows red/orange color in the vicinity of a micromotor, indicating increased pH in the surrounding medium. In addition, an increased concentration of the divalent cation  $Mg^{2+}$  can cause collapse of the mucus gel (37). The enhanced diffusion of the micromotors in the mucus was further validated using a previously reported method (38), as shown in Fig. 5G (see Materials and Methods). Compared with the negligible diffusion of the control silica particles in the mucus, diffusion of the micromotors in the mucus showed a significantly enhanced profile within 40 min. To investigate the cargo release kinetics of the micromotors, we encapsulated a fluorescent anticancer drug, DOX, into the alginate layer of the micromotors (see Materials and Methods). The release of DOX from the micromotors was characterized using an ultraviolet (UV)–visible spectrophotometer. The cross-linking treatment of the hydrogel significantly improved the efficiency of DOX loading (fig. S13A). By increasing the DOX loading amount from 0.5 to 4 mg, the dose of DOX per micromotor could be controlled from ~1 to 20 ng, whereas the encapsulation efficiency (EE) could be improved up to 75.9% (fig. S13B). A higher release rate was observed in the DOX-loaded micromotors in comparison with the DOX-loaded MCs (fig. S14), indicating the promise of using the micromotors for *in vivo* targeted therapy of GI diseases such as colon cancer.

Biocompatibility and biodegradability of the PAMR are important for biomedical applications. The materials of the MCs—such as Mg, Au, gelatin, alginate, and enteric polymer—are known to be biocompatible. To evaluate the toxicity profile of the PAMR *in vivo*, we orally administered MCs or deionized (DI) water to healthy mice once a day for two consecutive days. Throughout the treatment, no signs of distress—such as squinting of eyes, hunched posture, or lethargy—were observed in either group. Initially, the toxicity profile of the MCs in mice was evaluated through changes in body weight. During the experimental period, the body weights of the mice administered with MCs had no significant difference from those of the control group (see fig. S15 and Materials and Methods). The histology analysis was performed to evaluate further the toxicity of the PAMR *in vivo*. No lymphocytic infiltration into the mucosa or submucosa was observed, indicating no inflammation (see Fig. 5H and Materials and Methods).

## DISCUSSION

Two key challenges should be addressed for applying synthetic micromotors to practical biomedical applications: (i) advanced imaging techniques to locate micromotors in deep tissue at high spatiotemporal resolution with high contrast and (ii) precise on-demand control of micromotors *in vivo*. With high molecular sensitivity at depths, PACT allows real-time monitoring of micromotors in intestines at high spatial resolution for subsequent control. Here, micromotors with partial coating of functional multilayers were designed as both the imaging contrast agents and the controllable drug carriers. An Au layer was used to significantly increase the optical absorption for PA imaging and the reaction rate for efficient propulsion simultaneously. A gelatin hydrogel layer was used to enlarge the loading capacity of different functional components, such as therapeutic drugs and imaging agents. A parylene layer was applied to maintain the geometry of the micromotors during propulsion. Our current platform, integrating real-time imaging and control of micromotors in intestines *in vivo*, may lead to the next generation of intelligent microrobotic systems and provide opportunities for precise microsurgery and targeted drug delivery.

Although the current platform has been demonstrated in small animals, human clinical translations may require tens of centimeters of tissue penetration. PACT can provide up to 7-cm tissue penetration, which is limited by photon dissipation. By using a more penetrating excitation source—microwave and acoustic detection—thermoacoustic tomography (TAT) promises tissue penetration for clinical translations (39, 40). Moreover, incorporation of a gold layer in the micromotor design provided an excellent microwave absorption contrast for TAT owing to the high electrical conductivity and thus enhanced the deep tissue imaging capability of the microrobots for clinical applications. Focused ultrasound heating may increase the depths of thermally triggered microrobot release to the whole-body level of humans.

Currently, passive diffusion-based delivery suffers from complex designs, particle size constraints, low precision, and poor specificity. Our platform allows micromotors to reach any targeted regions in mice intestines with high precision. It can be tailored to particles of any sizes and can be applied to any biological media without additional design efforts. Our platform can also be easily modified to carry various cargos, including therapeutic agents and diagnostic sensors, with real-time feedback during delivery.

Biocompatibility and biodegradability of the micromotors are essential for practical biomedical applications. The components of our micromotors, widely used as therapeutic agents and in implantable devices, were found to be safe for *in vivo* applications (41–43). The micromotors have been eventually cleared by the digestive system via excrement, without any adverse effects.

In summary, we report an ingestible microrobotic platform with high optical absorption for imaging-assisted control in intestines. The encapsulated micromotors survived the erosion of the stomach fluid and permitted propulsion in intestines. PACT noninvasively monitored the migration of the micromotors and visualized their arrival at targeted areas *in vivo*. CW NIR irradiation toward targeted areas induced a phase transition of the capsules and triggered the propulsion of the micromotors. The mechanical propulsion provided a driving force for the micromotors to bind to the intestine walls, resulting in extended retention. The proposed platform lays a foundation for targeted delivery in tissues and opens a new horizon for precision medicine.

**MATERIALS AND METHODS****Materials**

Commercially available magnesium microparticles with a diameter of  $20 \pm 5 \mu\text{m}$  were purchased from TangShan WeiHao Magnesium Powder. Agarose, FITC-albumin, alginate, gelatin, and DOX hydrochloride were purchased from Sigma-Aldrich. Paraffin liquid, paraffin wax, hexane, glutaraldehyde, phenolphthalein, hydrochloric acid, glass coverslip, and gene frame were purchased from Thermo Fisher Scientific. Acrylic polymers (Eudragit L 100-55) were purchased from Evonik Industries. Silicone rubber tubes (inner diameter, 0.5 mm) were purchased from Dow Silicones.

**Fabrication of the micromotors**

The Mg-based Janus micromotors were constructed with an embedding method (fig. S1). The Au, alginate, and parylene were deposited in a layer-by-layer manner. Mg particles were first washed with acetone three times and dried at room temperature before use. Mg particles were dispersed in acetone with a particle concentration of  $\sim 0.1 \text{ g ml}^{-1}$  and then spread on the glass slides at room temperature. After the acetone evaporated, Mg particles were attached onto the surface of the glass slides through physical adsorption, exposing the majority of the surface areas of the particles to air. Subsequently, the glass slides coated with Mg particles were deposited with a Au layer ( $\sim 100 \text{ nm}$  in thickness) using an electron beam evaporator (Mark 40, CHA Industries). After the deposition, a mixture containing alginate (2%, w/v) and DOX was dropped on the glass slides and then dried with  $\text{N}_2$  gas. Aqueous  $\text{CaCl}_2$  (0.2 ml of 5%, w/v) was then dropped onto the glass slides to cross-link alginate. After 30 min, the glass slides were washed with pure water and dried with  $\text{N}_2$  gas. The glass slides were coated with a parylene C layer (750 nm in thickness) using a parylene coater (LabTop 3000, Curtiss-Wright). The resulted micromotors were collected by scratching from the glass slides.

**Preparation of the MCs**

MCs were fabricated on the basis of a controlled emulsion technique according to the previous reports (fig. S2) (44, 45). A mixture containing gelatin (5%, w/v) and micromotors (5%, w/v) at  $40^\circ$  to  $60^\circ\text{C}$  was extruded from a 30-gauge needle into 50 ml of liquid paraffin at  $\sim 60^\circ\text{C}$ . Pure water was used as the solvent here, in which micromotors remained stable due to the formation of a compact hydroxide passivation layer on the Mg surfaces. Subsequently, an enteric polymer solution consisting of 100 mg of Eudragit L-100 in 2 ml of organic solvent mixture (acetone:methanol = 1:1, v/v), as previously reported (46), was extruded into the liquid paraffin. The extruded solution was kept at  $60^\circ\text{C}$  for 4 hours to evaporate the acetone and methanol, and then the temperature was lowered to  $0^\circ\text{C}$  with an ice bath. To harvest the MCs from the liquid paraffin, we added cold water ( $\sim 4^\circ\text{C}$ ) to the liquid paraffin with magnetic stirring for more than 20 min, and most MCs were separated from the liquid paraffin into the water. The water containing MCs was extracted and then washed three times with hexane. The size of the MCs could be controlled by varying the rotational speed of magnetic stirring between 100 and 1000 rpm (fig. S5). The collected MCs were rinsed with an aqueous hydrochloric acid solution (pH 2) and then washed with pure water to remove the hydrochloric acid. Subsequently, the MCs were cross-linked through incubation with glutaraldehyde for 1 hour followed by water rinse.

**Characterization of the structures of the micromotors and the MCs**

Scanning electron microscopy (SEM) images of the Mg-based micromotors were acquired with a field emission scanning electron microscope (FEI Sirion) at an operating voltage of 10 keV (Fig. 2A). The samples were coated with a 5-nm carbon layer to improve the conductivity (Leica EM ACE600 Carbon Evaporator). The bright-field and fluorescence microscopic images of the micromotors and the MCs were taken with a Zeiss AXIO optical microscope (Fig. 2B, and figs. S3 and S4). To observe the structure of the DOX-loaded micromotors and MCs using fluorescence imaging, we stained the micromotors and the MCs with FITC-albumin. Labeling of FITC-albumin onto the micromotors was carried out by dip-coating the micromotors-loaded glass slides in 0.2 ml of FITC-albumin solution ( $0.2 \text{ mg ml}^{-1}$ ), followed by dip coating in an alginate solution (2%, w/v). Labeling of the FITC-albumin onto the MCs was conducted by adding FITC-albumin into the gelation solution.

**Characterization of the PA performances of the MCs**

Characterization of the PA performances of the MCs was conducted using a PACT system (19). The MCs, bare Mg microparticles, and blood were separately injected into three silicone tubes. Both ends of the tubes were sealed with agarose gel (2%, w/v). The PACT system used a 512-element full-ring ultrasonic transducer array (ring radius, 50 mm; central frequency, 5.5 MHz; more than 90% one-way bandwidth; Imasonic SAS) for two-dimensional panoramic acoustic detection. Each element has a cylindrical focus (numerical aperture, 0.2; element elevation size, 20 mm; pitch, 0.61 mm; inter-element spacing, 0.1 mm). A laboratory-made 512-channel preamplifier (26-dB gain) was directly connected to the ultrasonic transducer array housing, minimizing cable noise. The preamplified PA signals were digitized using a 512-channel data acquisition system (four SonixDAQs, 128 channels each, 40-MHz sampling rate, and 12-bit dynamic range; Ultrasonix Medical ULC) with programmable gain up to 51 dB. The digitized radio frequency data were first stored in the onboard buffer, then transferred to a computer, and reconstructed using the dual-speed-of-sound half-time universal back-projection algorithm (Fig. 2, C to G, and figs. S6 and S7) (19).

**PACT of the migration of the MCs**

PACT of the migration of the MCs in model intestines was carried out by injecting the MCs into a silicone tube, which was covered by chicken breast tissues. Migration of the MCs in the tube was driven by microfluidic pumping and was captured by PACT (Fig. 3B and movie S2).

For in vivo experiments, all experimental procedures were conducted under a laboratory animal protocol approved by the Office of Laboratory Animal Resources at California Institute of Technology. Three- to 4-week-old nude mice (20- to 25-g body weight; Hsd: Athymic Nude-FoxlNU, Harlan Co.) were used for in vivo imaging. Before the imaging experiments, the mice were fasted for  $\sim 8$  hours, followed by the oral administration with the MCs. The mouse was then fixed to a laboratory-made imaging platform by taping the fore and hind legs on the top and bottom parts of the holder in the PACT system. During imaging, the mice were under anesthesia with 1.5% vaporized isoflurane. The administered mice were imaged continuously for  $\sim 8$  hours to monitor the MC migration process (Fig. 4 and movie S5).

To study the migration of the MCs toward the targeted diseased areas, melanoma cells as the model tumor were cultivated, and the

cells were injected into intestines. A silicone tube filled with the MCs was connected to the intestine and was sealed with agarose. A syringe pump (Fisher Scientific 78-01001) was also connected with the tube to drive the MCs into intestines. The migration process was captured by PACT (Fig. 5B and movie S6).

### Quantify speeds of MC migration

The acquired frames containing MC migration were first averaged to project the trajectories of the MCs. The migration paths of MCs were manually identified from the averaged image. Time traces at points along the migration paths were then extracted, forming images in which one dimension was the distance along the migration paths ( $x$ ) and the other dimension was the elapsed time ( $t$ ). Median filter (3 pixels by 3 pixels) was then used to smooth the  $x$ - $t$  images. Applying a threshold (one-third of the maximum) segmented out the pixels containing MCs. The center positions of MCs along the path were estimated by calculating the geometric centers of the segmented pixels for given times. The center positions at the elapsed time points were fitted linearly to compute the migration speeds.

### Highlight MCs using temporal frequency filtering

The frames of interest were first smoothed by a Gaussian filter ( $\delta = 3$  pixels). Then, Fourier transformation with respect to time was applied to all frames. An empirical band-pass filter was used to eliminate signals from either the static background or the respiration motion-affected pixels, and thus the slowly moving pixels containing MCs were highlighted.

### CW NIR-activated propulsion of the micromotors

PBS solution (30  $\mu$ l) mixed with MCs was dropped on a piece of gene frame. A glass coverslip was then carefully placed over the gene frame. A CW NIR laser (808 nm, 2 W) was used to irradiate the micromotors obliquely with the light beam aligned to the focus of the microscope. The focal diameter of the beam was  $\sim 0.8$  cm. Once the position of the laser spot on the glass slide was marked, the NIR laser was turned off, and the MCs were moved to the marked spot. The MCs were irradiated before they completely sank to the bottom of the glass slide. The disintegration of the MCs occurred within a 0.1-s exposure of the CW NIR light. In addition, during each respiration cycle, the resting time (the duration free of respiration motion) is typically longer than 0.3 s (19). Thus, once the real-time PACT detects that MCs have reached the targeted area, the CW NIR light can trigger the release during the resting time, avoiding the influence of respiration motion. The process of the NIR-triggered disintegration of the MCs and the propulsion of the micromotors were captured using a high-speed camera (AxioCam 720 mono) at 100 and 25 frames  $s^{-1}$ , respectively (movie S3). ImageJ with the plugin Manual Tracking was used to track the micromotors (Fig. 3, D and E).

### Characterization of the propulsion of the micromotors

To simulate the gastric and intestinal environments, we prepared 0.01 M HCl (pH 2) as the model gastric fluid and 50 mM potassium phosphate buffer (pH 6.5) as the model intestinal fluid. To characterize the movement of the micromotors, we placed  $\sim 10$   $\mu$ l of model fluid with 1% Triton X-100 on a glass slide. Then, a  $\sim 2$   $\mu$ l aqueous micromotor suspension in water was added to the model solution on the glass slide. The movement of micromotors was captured using a high-speed camera (AxioCam 720 mono) at  $\sim 25$  frames  $s^{-1}$  (figs. S10 and S11 and movie S4). ImageJ with the plugin Manual Tracking was

used to track the micromotors. At least 20 micromotors were tracked to calculate the average speeds and the SDs.

To investigate the pH change of the surrounding environment upon micromotor propulsion, we added 5  $\mu$ l of phenolphthalein (0.5% in alcohol, w/v) into 1 ml of PBS solution (pH 6.5) as an indicator. An optical microscope (Zeiss AXIO) was used to capture the movie in color mode (Fig. 5F).

### Retention of the micromotors in vivo

Mice underwent a fasting treatment for  $\sim 8$  hours before the retention investigation. A 0.1-ml suspension containing the enteric polymer-coated micromotors ( $\sim 10^6$   $ml^{-1}$  in water) was orally administered to the mice. During the experiment, water feeding was maintained. As the controls, paraffin-coated passive particles (Mg particles and Mg/Au particles as Control 1 and Control 2, respectively) were prepared by incubating 0.05 g of particles with 1 g of paraffin wax at 75°C overnight and then sequential washing with chloroform, acetone, and pure water (47). Then, the control sample was orally administered. After 12 hours, both groups of mice were euthanized, and the intestines were collected. Retention of micromotors and passive particles was observed using a Zeiss AXIO optical microscope at  $\times 5$  magnification. The retained micromotors and control particles were counted using ImageJ (Fig. 5E). Dissolution of Mg after administration was characterized by optical imaging before and after acid treatment (fig. S12).

### Diffusion of the micromotors in mucus

Diffusion of micromotors in mucus was investigated following a reported method (38). A cuvette was filled with 3.5 ml of porcine mucus, and then 100  $\mu$ l of micromotors suspension ( $\sim 10^6$   $ml^{-1}$  in water) was pipetted into the mucus. Silica microparticles of the same size were used as control. Optical images were captured every 2 min. During the observation, the cuvettes were treated with sonication for 5 s with an ultrasound bath cleaner to remove bubbles. ImageJ was used to count particles in the mucus (Fig. 5, F and G). The numbers were normalized by the number of particles injected at the start, and the ratios were calculated at distances away from the initial point.

### Encapsulation and release of DOX from the micromotors

The EE and release profile of DOX on the MCs and micromotors were elevated using previous methods (48, 49). To encapsulate DOX into the micromotors, we dropped 1.0 ml of alginate solution (2%, w/v) with different concentrations of DOX onto the glass slides containing Au layer-coated Mg microparticles, and then 1.0 ml of  $CaCl_2$  solution was dropped onto the glass slide to cross-link alginate, followed by coating of a parylene layer and water rinse three times. Micromotors without cross-linking were also prepared. The amount of DOX was measured through a UV-visible spectrophotometer at 485 nm (fig. S13). The EE of DOX on the micromotors was determined using the following equation

$$\begin{aligned} \text{EE of DOX (\%)} &= \frac{\text{Initial amount of DOX used} - \text{amount of DOX in supernatant}}{\text{Initial amount of DOX used}} \\ &\times 100\% \end{aligned} \quad (1)$$

For the drug release study,  $\sim 10$  mg of DOX-loaded micromotors were suspended in 5 ml of PBS with magnetic stirring at 37°C and

8000 rpm. At different time intervals, the supernatant was removed and replaced with fresh PBS. The concentration of DOX was determined by measuring its absorbance using a spectrophotometer at a wavelength of 485 nm (fig. S14).

### Toxicity estimation of the micromotors

To estimate the toxicity of the MCs *in vivo*, we administered 0.1 ml of micromotor suspension to 5- to 6-week-old nude mice (20- to 30-g body weight; Hsd: Athymic Nude-Fox1NU, Harlan Co.) via oral gavage. Healthy mice treated with DI water were used as a negative control. The body weight of mice was measured daily during the experiment (fig. S15). Mice were euthanized, and the intestines were collected for histological characterization 6 days after administration. To prepare the intestine sample for histology investigation, the intestines were treated with 10% (v/v) buffered formalin for 15 hours. The intestines were cut to smaller sections as duodenum, jejunum, and distal colon. The longitudinal tissue sections were washed in tissue cassettes and embedded in paraffin. The tissue sections were sliced into 8- $\mu$ m-thick sections using a freezing microtome (CM1950, Leica) and stained with hematoxylin and eosin assay. The samples were imaged with an optical microscope (Zeiss AXIO) (Fig. 5H).

### SUPPLEMENTARY MATERIALS

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Text S1. Small-animal whole-body imaging modalities and PACT

Fig. S1. The fabrication flow of the ingestible micromotors.

Fig. S2. The preparation of the MCs.

Fig. S3. Bright-field and fluorescence microscopic images of the micromotors confirming the successful drug loading in micromotors.

Fig. S4. Bright-field and fluorescence microscopic images of the MCs confirming the successful drug loading in the MCs.

Fig. S5. Dependence of the size of the MCs on the rotation speed of magnetic stirring.

Fig. S6. Long-term stability of the PA signals of the MCs under the NIR illumination used in the PACT *in vitro* and *in vivo*.

Fig. S7. Fluorescence imaging of the MCs in a silicone tube under tissues with different depths.

Fig. S8. Long-term structure stability of the MCs in the gastric fluid and the intestinal fluid.

Fig. S9. Velocities of Mg-based micromotors in the different media.

Fig. S10. Velocities of bare Mg microparticles in the different media.

Fig. S11. Quantification of MC migration speeds.

Fig. S12. Characterization of Mg dissolution in micromotors 12 hours after administration.

Fig. S13. Effects of cross-linking and DOX loading amount on the EE of the micromotors and dose per micromotor.

Fig. S14. Profile of DOX release from MCs and micromotors as a function of time.

Fig. S15. The weight changes of the mice after the oral administration of the MCs and the control (DI water).

Movie S1. Animated illustration of the PAMR *in vivo*.

Movie S2. PA imaging of the migration of a MC in model intestines.

Movie S3. NIR-triggered destruction of the MC and activated autonomous propulsion of the ingestible micromotors.

Movie S4. Propulsion of the micromotors in biofluids.

Movie S5. PA imaging of the MCs *in vivo* for 7.5 hours.

Movie S6. PA imaging of the migration of an MC toward a model colon tumor in intestines.

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## A microrobotic system guided by photoacoustic computed tomography for targeted navigation in intestines in vivo

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