Unique insights into the intestinal absorption, transit, and subsequent biodistribution of polymer-derived microspheres

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Edited by Robert Langer, Massachusetts Institute of Technology, Cambridge, MA, and approved July 16, 2013 (received for review April 3, 2013)

Polymeric microspheres (MSs) have received attention for their potential to improve the delivery of drugs with poor oral bioavailability. Although MSs can be absorbed into the absorptive epithelium of the small intestine, little is known about the physiologic mechanisms that are responsible for their cellular trafficking. In these experiments, nonbiodegradable polystyrene MSs (diameter range: 500 nm to 5 μm) were delivered locally to the jejunum or ileum by oral administration to young male rats. Following administration, MSs were taken up rapidly (≤5 min) by the small intestine and were detected by transmission electron microscopy and confocal laser scanning microscopy. Gel permeation chromatography confirmed that polymer was present in all tissue samples, including the brain. These results confirm that MSs (diameter range: 500 nm to 5 μm) were absorbed by the small intestine and distributed throughout the rat. After delivering MSs to the jejunum or ileum, high concentrations of polystyrene were detected in the liver, kidneys, and lungs. The pharmacologic inhibitors chlorpromazine, phorbol 12-myristate 13-acetate, and cytochalasin D caused a reduction in the total number of MSs absorbed in the jejunum and ileum, demonstrating that nonphagocytic processes (including endocytosis) direct the uptake of MSs in the small intestine. These results challenge the convention that phagocytic cells such as the microfold cells solely facilitate MS absorption in the small intestine.

oral delivery | uptake mechanism

Beginning in the 1960s, several groups (1–7) demonstrated that the small intestine could absorb microparticles with a diameter >1 μm, challenging dogma that the small intestine could only absorb small macromolecules. After this discovery, researchers began engineering microspheres (MSs) to deliver drugs with poor water solubility (8–10), poor gastrointestinal permeability (11), or poor oral bioavailability (8, 9, 12–15) to the small intestine. MS-based oral drug delivery systems (ODDSs) can be made from biodegradable polymers (12, 16–20), nonbiodegradable polystyrene (19–21), and polysaccharides (22, 23) and can be engineered to carry polypeptides (18, 24, 25) and other molecules (26, 27). The motivation for using microparticulate-based ODDSs is to improve the oral bioavailability of drugs by enhancing their delivery and transit through the absorptive epithelium of the small intestine. This work may ultimately enable patients to take medications orally instead of relying on daily or weekly injections or other less convenient routes of administration (28).

More than five decades after Sanders and Ashworth discovered that the small intestine can engulf large particles (1), little is known today about the cellular mechanisms or physiologic pathways that facilitate the absorption, transit, and biodistribution of microparticles that are delivered to the small intestine. Nonetheless, the diverse physiology and multitude of cell types in the small intestine likely play a central role in these processes. The absorptive epithelium is comprised primarily of enterocytes, cells that are critical for nutrient absorption (29–33). The distal jejunum and ileum contain a greater density of Peyer’s patches and follicle-associated epithelium (FAE), regions that contain microfold (M) cells, which can facilitate the uptake of antigens (29, 34, 35), viruses (36, 37), and microorganisms (38, 39).

Given the immunological importance and phagocytic activity of M cells, researchers have focused on delivering microparticulate-based ODDSs to the Peyer’s patches and the FAE (40, 41), both of which were once thought to be the regions chiefly responsible for microparticle uptake in the small intestine (42–47). However, non-FAE tissue can also play a role in the absorption and transit of MSs (48, 49). Despite the promise of M cell–targeted delivery, M cells comprise <1% of the total surface area of the absorptive epithelium; their scarcity makes targeting these cells difficult. Many groups have used qualitative or semiquantitative methods to estimate the absorption and biodistribution of MSs in the small intestine, yet very few groups (50) have used quantitative methods to measure the actual percentage of MSs that are absorbed in the small intestine relative to the administered dose. Furthermore, very little research has been done in vivo to understand the specific cellular mechanisms that facilitate MS uptake, the relative importance of each mechanism, or the biodistribution of MSs following oral administration. Understanding the biodistribution and ultimate fate of MSs following ingestion is important for drug targeting, pharmacokinetics, and toxicology.

The objectives of our experiments were threefold: use electron and confocal microscopy to confirm the uptake of polystyrene MSs in the small intestine; approximate the number of polystyrene MSs that are absorbed in the small intestine; and determine the biodistribution of polystyrene MSs following oral or local delivery to the small intestine. In addition, we compared total uptake in the presence and absence of cellular inhibitors to determine which pathways facilitate MS uptake in the small intestine.

Results

Transmission Electron Microscopy. Ultrastructural analysis of the small intestine revealed that it was not damaged during the isolated loop surgical procedure. Five minutes after isolating the intestinal loop, MSs (200 nm) were detected in the intestinal lumen adjacent to the microvilli (brush border) of the absorptive epithelium (Fig. L4). MSs (200 nm) were also observed within...
enterocytes (Fig. 1B and C). After 5 min, larger MSs (500 nm) were observed within enterocytes near the lateral membrane of the apical cell boundary (Fig. 1D) and within cytosolic vesicles (Fig. 1E).

**Confocal Microscopy.** Fig. 2 shows representative confocal laser scanning micrographs (CLSMs) of rat jejunum (villus) after completion of a 1-h isolated loop procedure. In Fig. 2A and B, the jejunum was observed after staining the tissue with DAPI and FM 1-43 stain. In Fig. 2C, fluorescent polystyrene MSs (500 nm) were observed within the jejunum. An overlay of Fig. 2A–C revealed that MS traversed from the intestinal lumen through the apical membrane of enterocytes into the villus core (Fig. 2D). 3D fluorescence image analysis confirmed that MSs were colocalized within the tissue and were not separated from the intestine during specimen preparation.

**Gel Permeation Chromatography. Effect of microsphere diameter on absorption in the small intestine.** The uptake of polystyrene MSs following oral administration (fed) or local delivery to the jejunum or ileum is shown in Fig. 3. Total uptake was generally inversely related to MS diameter, independent of delivery method (local vs. oral), yet a greater percentage of 5-μm MSs was absorbed in the jejunum and ileum compared with 2-μm MSs. In the jejunum and ileum, 1- and 2-μm MSs were less likely to be absorbed than 500-nm MSs. A greater number of 500-nm MSs were absorbed in the jejunum compared with the ileum (45.8 ± 8.6% vs. 34.9 ± 9.3%, respectively). In the ileum, total uptake of 500-nm and 1-μm MSs was similar (34.9 ± 9.3% vs. 32.2 ± 11.5%, respectively). Following oral administration, uptake of 500-nm and 1-μm MSs was smaller compared with the uptake observed after delivery to the jejunum and ileum.

**Microsphere biodistribution.** Fig. 4A shows MS uptake following local administration to the jejunum (500 nm) and ileum (1 μm). One hour following the administration of 500-nm MSs, small concentrations of polymer were detected, but a greater concentration of polymer following the administration of 1-μm MSs was detected. After 3 h, greater quantities of 500-nm and 1-μm MSs were detected. After 5 h, a greater concentration of polymer was detected in animals that received 500-nm MSs compared with animals who received 1-μm MSs.

The concentration of polymer (expressed as a percentage of total uptake) following local delivery of 500-nm MSs to the jejunum is shown in Fig. 4B. After 1 h, the concentration of polymer detected in all assayed regions was low. After 3 h, a greater concentration of polymer was detected within the liver than all other regions combined. After 5 h, >30% of the total cumulative administered dose was detected within the liver, and a small but increasing concentration of polymer was detected within the kidneys.

The concentration of polymer (expressed as a percentage of total uptake) following local delivery of 1-μm MSs to the ileum is shown in Fig. 4C. One hour after local administration, more polymer was detected in the lungs than any other region; very
little polymer was detected in the liver. After 3 h, smaller concentrations of polymer were found within the lungs, and more polymer was detected in the liver and kidneys. After 5 h, a higher concentration of polymer was detected in the liver, kidneys, and lungs compared with other assayed regions.

Fig. 4D shows the concentration of polymer following the oral administration of 500-nm and 1-μm MSs to rats. After 5 h, a greater number of 500-nm MSs were detected in the stomach relative to larger MSs, yet a greater number of 1-μm MSs were found within the jejunum, ileum, colon, and cecum (Fig. 4D). After 5 h, very few 500-nm and 1-μm MSs were found within the duodenum.

The biodistribution of MS following oral administration or local delivery to the jejunum or ileum is shown in Table S1. Five hours after administering 500-nm MSs, more polymer was detected in the liver than any other region. Larger MSs (1, 2, and 5 μm) were also detected in the liver, albeit in smaller quantities. Following local MS administration to the jejunum and ileum, a greater concentration of polymer was detected in the liver, kidneys, and lungs compared with other assayed regions. Polymer was detected in almost every tissue sample, including the brain.

Effect of Endocytosis Inhibitors on Microsphere Uptake in the Jejunum and Ileum. Fig. 5 shows the total uptake of MS in the jejunum (Fig. 5A) and ileum (Fig. 5B) following local delivery of endocytosis blockers to these regions. After 5 h, the endocytosis blockers chlorpromazine (CPZ), phorbol 12-myristate 13-acetate (PMA), and cytochalasin D (CytD) caused a reduction in the total number of MSs absorbed by the jejunum relative to untreated animals (3.6 ± 2.1%, 10.3 ± 2.3%, and 7.5 ± 2.0% for CPZ, PMA, and CytD, respectively, vs. 45.8 ± 8.6% for untreated animals). Five hours after administering the pharmacologic inhibitors to the ileum, MS uptake decreased relative to untreated animals (12.1 ± 3.6%, 14.4 ± 3.6%, and 4.5 ± 1.6% for CPZ, colchicine, and CytB, respectively, vs. 32.2 ± 11.5% in the absence of inhibitor). In addition to observing a decrease in the quantity of MSs detected within each animal, the biodistribution of MSs was slightly changed relative to nontreated animals following administration of the endocytosis blockers (Fig. S1).

Fig. 2. CLSM images of a single villus 1 h after delivering fluorescent PC-red 500-nm polystyrene MSs to the jejunum is shown with DAPI stain (A), FM 1-43 stain (B), and with illumination of fluorescent PC-red MSs (C). (D) Overlay of all three channels (A–C) is shown. In D, red arrows show where MSs are located within goblet cells; white arrows highlight where MS have penetrated beyond the epithelial cell layer.

Fig. 3. Effect of polystyrene MS diameter on total uptake (expressed as a percentage of total administered dose) 5 h after local delivery to the jejunum or ileum or 5 h after oral administration. n = 4 animals for each cohort.

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Discussion

The transport across the mucus layer is an important aspect in particle uptake. We have recently published a paper where the interaction of adhesive particles with the mucus is discussed in more detail (Fig. 1) (51). There are many factors involved in mucus transport that cannot be fully discussed here. For example, these particles are hydrophobic in nature, so there may be a hydrophobic driving force “pushing” particles toward the gut walls and away from the aqueous lumen. Regardless of the specific mechanisms involved in mucus transport, we feel that it occurs and the fact that orally fed particles (in absence of effects from the artificial isolated loop administration) also experience high uptake is strong evidence of this transport (51).

These experiments (Fig. 1) provide qualitative and quantitative confirmation that polystyrene MSs are taken up by the absorptive epithelium and deposited throughout the rat. Historically, there have been several proposed mechanisms for MS absorption, including endocytosis, which was believed to be limited to particles with a diameter <50 nm. In addition, paracellular transport of MSs between the tight junctions of enterocytes and phagocytosis by M cells were believed to facilitate MS uptake. Although a number of studies have provided evidence that enterocytes can engulf MSs in vitro and in vivo, non-Peyer’s patch uptake by intestinal enterocytes is thought to be minimal. Furthermore, only one literature review in the last decade has recognized the importance of enterocytes and the absorptive epithelium in MS uptake. Delivery to enterocytes would be highly advantageous, especially given that M cells comprise a relatively small proportion of the small intestine’s total surface area.

Data from these studies challenge current dogma in the area of oral drug delivery. Using an in vivo isolated loop technique for mechanistic studies has the advantage of isolating a specific intestinal region and being able to deliver specific pharmacologic uptake inhibitors locally in a controlled manner. In contrast, in vitro cell culture models are used to study the transepithelial transport of small molecule drugs. Several groups have used in vitro cell culture models to study the absorption of starch, polystyrene, poly(acrylic acid), and poly(lactic-coglycolic acid). Many of these studies have used Caco-2 cells derived from a human colon adenocarcinoma cell line that differentiate into cells mimicking enterocytes. HT29 cells mimic goblet cells and incorporate a mucosal layer into the cell monolayers. Cocultures of Caco-2 and HT29 adenocarcinoma cells have been used to approximate the physiology of the small intestine, yet very few studies have looked at specific cellular mechanisms responsible for particle uptake. Furthermore, the predominant cell models (including Caco-2 cell monolayers) vary greatly in morphology, which may undergo even greater change in the presence of MS. Additionally, the relevance of an in vitro cell culture line to the dynamic in vivo environment (i.e., gastrointestinal mobility, fluid flow, mucus secretion, cell turnover) may not be suitable for studying the absorption, translocation, and biodistribution of MSs.

Transmission electron microscopy (TEM) in these studies revealed that 200- and 500-nm MSs were found within cytosolic vesicles. No exocytotic events were observed, which may be a result of the short duration of these experiments. We observed greater uptake of 500-nm polystyrene MSs in the absorptive epithelium of the jejunum than in the Peyer’s patches of the ileum, indicating that an alternative mechanism to M cell phagocytosis facilitates MS uptake. Intestinal enterocytes are not capable of phagocytosis, which suggests that endocytosis facilitates MS uptake in the jejunum. Endocytosis of particles >1 μm has not been described in the literature, and only a single report by Rejman et al. noted that 500-nm MSs could be engulfed by endocytosis. Although the total uptake of 500-nm and 1-μm polystyrene MSs was lower relative to the total uptake observed following delivery of MSs locally to the jejunum and ileum, the uptake of MSs following oral delivery still comprised a relatively high proportion of the total administered dose.

In the jejunum and ileum, MS uptake steadily increased following MS administration, yet the manner in which they increased was slightly different. In the jejunum, MS uptake increased in a linear fashion, whereas in the ileum, MS uptake plateaued early in the study. One possible explanation for the lag time observed in the jejunum could be that more interaction time or intercellular signaling responses are required. In contrast, MS uptake in the ileum appeared to become saturated, leading to a plateau effect. Given these findings—and despite being the least studied

Fig. 4. (A) Effect of time on the uptake of polystyrene microspheres in the jejunum and ileum is shown. The biodistribution of 500-nm MSs 1, 3, and 5 h after local delivery to the jejunum (B) and 1-μm MSs 1, 3, and 5 h after local delivery to the ileum (C) are also shown. D shows quantitative resident time analysis of 500-nm and 1-μm MSs 5 h after oral administration.
mechanism—it is possible that a nonphagocytic mechanism in the jejunum is most responsible for MS uptake.

There are two known types of receptor-mediated endocytosis: clathrin dependent and clathrin independent. Our data suggest that the primary mechanism of MS uptake in the rat small intestine is via clathrin-mediated endocytosis in combination with caveolar-mediated endocytosis and phagocytosis. Clathrin-dependent endocytosis requires adaptor protein (AP) and a GTP-binding protein (dynamin) in addition to clathrin. AP2 facilitates membrane curvature and links the membrane proteins to clathrin, which is a three-limbed shaped protein that polymerizes to form cage-like structures. After transporting particles, clathrin must be recycled to form pits de novo; the binding site of AP2 must be inactivated to release clathrin (52, 53). CPZ activates the recycling AP2 binding sites that are normally inactive and causes the binding of AP2/clathrin. The inhibition of clathrin recycling prevents further endocytosis (53). Okamoto et al. showed that 25 μg/mL of CPZ inhibits clathrin-dependent endocytosis in Chinese hamster ovary cells (54).

Another well-known endocytic pathway involves the formation of caveolae (cholesterol-rich membrane regions) that can easily invaginate due to a localized increase in membrane flexibility. Caveolae can bud off from the membrane, resulting in endocytosis of extracellular content. PMA exerts its effects through a protein kinase C–mediated mechanism, specifically through the phosphorylation of caveolin, a necessary protein in the formation of caveolae, to inhibit caveolin-mediated endocytosis without affecting the return of vesicles to the cell surface (55). Smart et al. showed an absence of folate uptake in M104 cells treated with PMA, specifically through the inhibition of a caveolin-mediated uptake mechanism (56). Similar results with micromolar concentrations of PMA have also been seen inhibiting virus entry into cells (57).

Although MSs were distributed throughout the rat, the majority of particles were found in the liver. Following MS delivery to the ileum, numerous MSs were spread among many organs, including the heart and spleen. This shift in organ distribution supports the theory that there may be different particle uptake mechanisms and distribution pathways. Peyer’s patches are often referred to as gut-associated lymphoid tissue, or GALT, due to the fact that lymph nodes occupy the mucosal layer immediately beneath Peyer’s patches. Increased MS distribution to the heart and spleen may be the result of MS traveling in the lymph, which passes through the lymphatic ducts and returns into the systemic circulation through the thoracic duct. Lymph does not undergo first pass metabolism, which may have allowed some of the MS to bypass the liver via lymphatic circulation. When MSs are delivered to the jejunum, distribution to the spleen and heart diminish due to an absence of FAE and Peyer’s patches; these particles enter the hepatic circulation and are directed to the liver, which may lead to greater particle accumulation in the liver.

Conclusions

The uptake kinetics and biodistribution of polystyrene MSs following oral or local delivery to rat intestine was dependent on MS size, location, and route of administration. This study demonstrated that the nonlymphoid tissue of the absorptive epithelium can absorb MSs and facilitate their biodistribution throughout the rat. The relatively high degree of MS uptake in the absorptive and nonabsorptive epithelium indicates that endocytotic and phagocytic mechanisms are responsible for MS uptake in the small intestine. These findings may potentially guide research aimed at delivering specific molecules to target organs. In addition, the methods and in vivo model used herein may also be useful to toxicologists who are interested in determining the fate or tissue distribution of microparticles, including whether such particles can cross the blood–brain barrier.

Materials and Methods

Polystyrene MSs with mean diameters of 500 nm and 1, 2, and 5 μm were purchased from Polysciences and administered both locally using an isolated loop procedure (with and without the addition of pharmacologic inhibitors CPZ, PMA, CybT and Colchicine) and oral gavage. Total MS uptake following oral administration was quantified after 5 h. All animals were killed and had the following samples collected: portal vein blood (∼1 mL), celiac arterial blood (∼1 mL), lungs, heart, spleen, kidneys, liver, stomach, stomach rinse, duodenum, duodenum rinse, jejunum, jejunum rinse, ileum, ileum rinse, cecum, cecum rinse, colon, colon rinse, and brain. Gel permeation chromatography, TEM, and confocal microscopy were used to analyze the uptake of MS from collected tissues. For more details, see SI Materials and Methods.


Supporting Information

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SI Materials and Methods

Materials. Polystyrene microspheres (MSs) with mean diameters of 500 nm and 1, 2, and 5 μm were purchased from Polysciences and were stored at 4 °C until further use. Suspending polymers were used as received from the manufacturer (25 mg MS/mL). A dosage volume of 1 mL (total microsphere weight ~ 25 mg) was used in all studies. All of the particles used in the study are part of the carboxylated Polybead product line from Polysciences. They are a series of monodisperse polystyrene MSs that are provided as aqueous suspensions with minimal surfactant in the final preparation. The particles have a slight anionic charge from sulfate ester and have been synthesized to achieve precise size distributions. These particles were selected so that all of the particles would be identical with the exception of particle size for the different study groups.

The uptake inhibitors chlorpromazine (CPZ), phorbol 12-myristate 13-acetate (PMA), cytochalasin B (CytB), and CytD were purchased from Sigma-Aldrich and stored according to the manufacturer’s instructions. Dosing concentrations and intervals were selected for each inhibitor by using material safety data sheets and in vitro cell culture protocols published in the literature (1–8). CPZ and colchicine were dissolved into phosphate buffer solution (concentration = 50 μg/mL and 0.1 mM, respectively) immediately before administration. CytB, CytD, and PMA were dissolved into dimethyl sulfoxide and stored at −80 °C until use. Immediately before administration, dimethyl sulfoxide and inhibitor solutions were thawed and diluted in PBS to concentrations of 60, 4, and 2 μM for CytB, CytD, and PMA, respectively.

Isolated Loop Procedure. Male Sprague–Dawley rats (200–250 g) were anesthetized with 3% (vol/vol) isoflurane. After sedation, a 3-cm midline incision was made, and the desired section of the small intestine was identified. The proximal jejunum was located by identifying the ligament of Trietz; the distal jejunum and the beginning of the proximal ileum were located by visualizing an increase in the density of GALT. The distal ileum was identified by locating the cecum. Once the desired sections of the small intestine were identified, a 6-cm segment was isolated with two ligatures that were created using monofilament silk 0-0 suture. Care was taken not to disrupt normal blood flow from the mesenteric vasculature.

After isolating the intestinal segments, polystyrene MSs with a diameter of ~500 nm and 1, 2, and 5 μm were administered locally by direct injection into the jejunum and ileum. The isolated loop was returned to the abdomen, which was closed during the procedure to maintain normal physiologic conditions. After various time points (5 min to 5 h), samples were collected in the following order: portal vein blood (~1 mL), celiac arterial blood (~1 mL), lungs, heart, spleen, kidneys, liver, stomach, stomach rinse, duodenum, duodenum rinse, jejunum, jejunum rinse, ileum, ileum rinse, cecum, cecum rinse, colon, colon rinse, and brain. The intestinal rinses were combined with their respective sections and were treated as one sample. Additionally, all urine was aspirated from the bladder, and fecal matter was removed from the rectum. Urine and feces were added to urine and fecal samples that were collected throughout the experiment. All samples were stored at −18 °C until further processing. Each experimental condition was repeated four times.

Tissue Processing and Polymer Detection. After killing the animal, gel permeation chromatography (GPC) was used to detect the concentration of polymer within each sample. Excised tissue was minced into small pieces, added to ~5 mL PBS, and homogenized using a Cole-Palmer Ultrasonic Homogenizer CV26 with a high gain Q horn and extender (40% amplitude for 30 s). Homogenized tissue samples were lyophilized for 48 h, leaving behind a powdered tissue digest. Following tissue sample dehydration, polymer was extracted from the tissue by adding chloroform (10 mL), mixing the slurry on an end-over-end mixer for 9 h, and filtering the tissue samples using a 0.1-μm polytetrafluoroethylene (PTFE) syringe filter to remove nonsoluble biological debris. Filtered samples were lyophilized for an additional 24 h and stored at −18 °C. We have validated this method [that was also used by others (9)] by doping tissues with known amounts of polystyrene beads and achieving greater than 90% recovery.

Before sample analysis, the lyophilized powders were reconstituted in 1 mL of chloroform by mixing samples on an end-over-end mixer for 1 h. Each solution was filtered again using a 0.1-μm PTFE syringe filter. Filtered samples were run through a GPC (Shimadzu Corporation) equipped with Styragel HR5E and HR4E columns (Waters Corporation) and a Shimadzu RID-10A refractive index detector. A peak for polystyrene was identified, the area under the curve for various polymer concentrations was calculated, and a linear standard curve was constructed (R² = 0.9993).

Using the polystyrene calibration curve, the concentration of polystyrene in each lyophilized sample was calculated. Percent uptake was calculated by taking the sum of all quantities of polystyrene detected in each tissue (excluding isolated loop and
loop rinse samples), divided by the total dose administered and multiplying these values by 100. We excluded the rinse sample from this calculation as it represents the portion of the formulation that is not absorbed. Similarly, the isolated loop is also excluded despite the fact that there is absorption [as evidenced by transmission electron microscopy (TEM) and confocal] because the method cannot distinguish absorbed from adsorbed particles. Therefore, this calculation is the most conservative approach to quantify uptake. Inclusion of the isolated loop sample would increase the stated uptake toward a possible overestimate. The biodistribution was compared for each sample across study groups. Additionally, mass balance was measured by dividing the concentration of polymer in each sample by the total administered dose. Positive controls verified that 99.8 ± 1.3% of polystyrene MSs were recovered from tissue samples. No MSs were detected in the negative control samples.

**TEM.** After completion of the isolated loop experiments, gross anatomical sections of intestine were harvested, washed in warm PBS (37 °C), and fixed in a solution of 20% (vol/vol) paraformaldehyde/20% (vol/vol) glutaraldehyde in sodium cacodylate buffer for 2 h at 37 °C. After fixation, samples were rinsed with sodium cacodylate buffer (0.2 M) and postfixed with 1% osmium tetroxide dissolved in 0.2 M sodium cacodylate buffer for 3 h at room temperature. Immediately following fixation in osmium tetroxide, samples were stained in a solution of 1% uranyl acetate overnight at room temperature. Following fixation and tissue staining, samples were dehydrated using progressively increasing concentrations of ethanol (30–100% [vol/vol]). Dehydrated samples were embedded and polymerized in hard-grade LR White embedding medium (Electron Microscopy Sciences). Tissue blocks were sectioned to a thickness of 75–85 nm using a Reichert Ultra microtome and diamond knife. Ultrathin sections were placed on copper grids and viewed on a Phillips 410 transmission electron microscope equipped with an Advantage HR CCD camera. Images were acquired with Advanced Microscopy Techniques imaging software.

**Confocal Microscopy.** Following a 1-h isolated loop experiment, sections of rat jejunum were harvested, washed in PBS, fixed in 75% (vol/vol) formaldehyde for 1 h, and stored at 4 °C until further processing. Before imaging, intestinal sections were stained with FM-143 fluorescent membrane stain (excitation wavelength: 488 nm; emission wavelength: 600 nm) and DAPI fluorescent nuclei stain (excitation wavelength: 358 nm; emission wavelength: 461 nm). The polystyrene MSs were stained with the fluorescent red dye PC-red (excitation wavelength: 377 nm; emission wavelength: 479 nm).

In preparation for imaging, representative intestinal tissue samples were embedded into optimal cutting temperature (OCT) media for sectioning and cut into 20-μm sections with a Leica cryostat. Intestinal tissue samples were imaged using a Leica TCS SP2 AOBS spectral confocal laser scanning microscope using UV excitation (fluorescence emission of 450 nm). Multiple optical sections of each sample were taken. Images were captured, and the resulting images were analyzed using OpenLab software to measure and compare the fluorescence intensity of the tissue samples.

**Statistical Analyses.** For all experiments, the mean ± SEM is reported. A one-way ANOVA was performed using Microcal Origin Graphical Software. Statistical significance was set at the 0.05 level.

**Cytotoxicity of Uptake Inhibitors.** To assess the cytotoxicity of uptake inhibitors, a blue-LIVE/DEAD fixable dead cell stain kit from Molecular Probes was used. For each inhibitor (CPZ, PMA, CytB, CytD, and colchicine), an isolated loop with 5.5 h of inhibitor incubation was performed as described above without the administration of MSs. Additionally, a solution of PBS and a 75% (vol/vol) solution of ethanol was used for negative and positive controls, respectively. Following the 5.5-h incubation period, 5 μL of the blue fluorescent reactive dye solution was mixed with 195 μL of phosphate buffer solution, injected into the isolated loop, and incubated for an additional 30 min. The intestinal section was washed and was rinsed with fresh phosphate buffer solution and placed in 37% (vol/vol) paraformaldehyde for 1 h. Fixed and stained intestinal samples were embedded in OCT media, cut into 50-μm sections with a Leica cryostat, and prepared for confocal microscopy. Samples were imaged on a Leica TCS SP2 AOBS spectral confocal microscope with UV excitation. Emission was read at 450 nm at multiple optical sections. OpenLab software was used to quantitatively compare the relative intensity of staining in all samples.

**SI Results and Discussion**

As mentioned in the Introduction, no in vivo studies have been published to systematically investigate uptake mechanisms and pathways for the purpose of studying polymeric translocation across the small intestine. Using chemical blockers allows inhibition of specific pathways and, when used in conjunction with quantitative uptake studies, allows investigation of specific pathways. Although the blockers have been chosen for their specificity, each blocker has the potential to be cytotoxic. To ensure that any observed changes in uptake are a result of pathway inhibition, a study was performed to evaluate cytotoxicity. Development of a LIVE/DEAD fluorescent staining assay for in vivo cell viability was used.

The assay described in SI Materials and Methods was successful in yielding quantitative results regarding in vivo cell viability. Results are shown in Fig. S2. Representative samples in each study group are shown in blue, corresponding intensity maps are shown in red, and a bar graph of average intensity for each image is shown in the bottom right of Fig. S2. Negative controls of fresh intestinal tissue had a low level of fluorescence intensity (Fig. S2A). A small fluorescence signal is expected because intestinal epithelium has a high level of turnover (<24 h) and a basal amount of cell death would be expected.

As a positive control, 70% (vol/vol) ethanol was injected into the intestine. Ethanol induced cytotoxicity, which is evident from the high signal intensity observed in Fig. S2C. Additionally, the isolated loop section with ethanol appeared shrunken and had a dark purple to black appearance. This observed change was seen only in tissue that was exposed to ethanol, whereas all other isolated loops included in the study did not undergo any observable changes in appearance. Although the isolated loop technique, and many variations of it, has been implemented for many years, no previous study has investigated cell viability with the technique. In an isolated loop that received a bolus of phosphate buffer solution, a moderate amount of intensity was observed (Fig. S2B). A small fluorescence signal is expected because intestinal epithelium has a high level of turnover (<24 h) and a basal amount of cell death would be expected.

Performing the fluorescent cytotoxicity assay on isolated loops that received pharmacologic blockers demonstrated a low level of toxicity that was comparable to, and never exceeded, the amount of cytotoxicity observed in the isolated loop control (Fig. S2). Therefore, quantitative uptake results gained by the use of these chemical blockers can be considered to be the effect of blocking specific pathways and are not the result of cell toxicity.


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**Fig. S1.** Biodistribution of 1-μm polystyrene MSs is expressed as a percentage of the total administered dose 5 h after local delivery of to the ileum in the presence of CPZ, colchicine, and CytB and in the absence of drug.
Fig. S2. LIVE/DEAD fluorescent stain was used on isolated intestinal loops 5.5 h after the administration of (A) no material (control); (B) PBS; (C) 70% (vol/vol) ethanol; (D) chlorpromazine; (E) colchicine; (F) CytB; (G) CytD; and (H) PMA. Intensity maps are shown in red, and the bar graph in the bottom right figure plots the average relative intensities (20 images analyzed for each group; maximum intensity = 255).
Table S1. Biodistribution of particles in specific organs

<table>
<thead>
<tr>
<th>Organ</th>
<th>500 nm</th>
<th>1 μm</th>
<th>2 μm</th>
<th>5 μm</th>
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</thead>
<tbody>
<tr>
<td>Brain</td>
<td></td>
<td></td>
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<tr>
<td>Jejunum</td>
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<td>0 ± 0</td>
<td>0 ± 0</td>
<td>0 ± 0</td>
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<tr>
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<td>0 ± 0</td>
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<td>0 ± 0</td>
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<tr>
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<td>0 ± 0</td>
<td>—</td>
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<tr>
<td>Central blood</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jejunum</td>
<td>0.07 ± 0.05</td>
<td>3.51 ± 3.02</td>
<td>0.09 ± 0.09</td>
<td>0.17 ± 0.09</td>
</tr>
<tr>
<td>Ileum</td>
<td>0.86 ± 0.50</td>
<td>0.04 ± 0.03</td>
<td>1.17 ± 0.71</td>
<td>0.10 ± 0.08</td>
</tr>
<tr>
<td>Fed</td>
<td>0.07 ± 0.07</td>
<td>0.09 ± 0.09</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Heart</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jejunum</td>
<td>0.14 ± 0.05</td>
<td>0.87 ± 0.20</td>
<td>0.87 ± 0.54</td>
<td>0.28 ± 0.26</td>
</tr>
<tr>
<td>Ileum</td>
<td>3.73 ± 3.03</td>
<td>0.41 ± 0.23</td>
<td>0.12 ± 0.08</td>
<td>0.30 ± 0.18</td>
</tr>
<tr>
<td>Fed</td>
<td>0.20 ± 0.12</td>
<td>0.27 ± 0.16</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Kidneys</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>Jejunum</td>
<td>3.62 ± 2.69</td>
<td>2.97 ± 1.61</td>
<td>2.82 ± 1.03</td>
<td>0.19 ± 0.19</td>
</tr>
<tr>
<td>Ileum</td>
<td>0.35 ± 0.35</td>
<td>1.01 ± 1.01</td>
<td>0.11 ± 0.10</td>
<td>3.98 ± 0.96</td>
</tr>
<tr>
<td>Fed</td>
<td>1.80 ± 0.73</td>
<td>1.12 ± 0.65</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Liver</td>
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</tr>
<tr>
<td>Jejunum</td>
<td>36.73 ± 11.04</td>
<td>9.01 ± 3.23</td>
<td>11.97 ± 4.06</td>
<td>15.46 ± 5.59</td>
</tr>
<tr>
<td>Ileum</td>
<td>26.26 ± 14.20</td>
<td>27.78 ± 10.35</td>
<td>2.12 ± 1.23</td>
<td>7.80 ± 4.02</td>
</tr>
<tr>
<td>Fed</td>
<td>22.89 ± 9.81</td>
<td>17.37 ± 7.33</td>
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</tr>
<tr>
<td>Lungs</td>
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</tr>
<tr>
<td>Jejunum</td>
<td>0.48 ± 0.36</td>
<td>8.49 ± 6.03</td>
<td>0.31 ± 0.25</td>
<td>0.81 ± 0.60</td>
</tr>
<tr>
<td>Ileum</td>
<td>0.13 ± 0.08</td>
<td>1.55 ± 0.99</td>
<td>0.51 ± 0.34</td>
<td>0.21 ± 0.21</td>
</tr>
<tr>
<td>Fed</td>
<td>0.77 ± 0.56</td>
<td>0.06 ± 0.06</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Portal blood</td>
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</tr>
<tr>
<td>Jejunum</td>
<td>2.26 ± 1.31</td>
<td>0.47 ± 0.28</td>
<td>0.24 ± 0.20</td>
<td>0.97 ± 0.60</td>
</tr>
<tr>
<td>Ileum</td>
<td>1.63 ± 1.63</td>
<td>0.28 ± 0.28</td>
<td>0.12 ± 0.12</td>
<td>0.11 ± 0.07</td>
</tr>
<tr>
<td>Fed</td>
<td>0 ± 0</td>
<td>2.65 ± 1.71</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Spleen</td>
<td></td>
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</tr>
<tr>
<td>Jejunum</td>
<td>0.63 ± 0.28</td>
<td>1.24 ± 1.06</td>
<td>0.20 ± 0.16</td>
<td>0.43 ± 0.22</td>
</tr>
<tr>
<td>Ileum</td>
<td>3.72 ± 2.62</td>
<td>0.47 ± 0.24</td>
<td>0.16 ± 0.16</td>
<td>0 ± 0</td>
</tr>
<tr>
<td>Fed</td>
<td>0.01 ± 0.01</td>
<td>0.05 ± 0.05</td>
<td>—</td>
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</tr>
</tbody>
</table>

Five hours after administering 500 nm MS, more polymer was detected in the liver than any other region. Larger MS (1, 2, and 5 μm) were also detected in the liver, albeit in smaller quantities. Following local MS administration to the jejunum and ileum, a greater concentration of polymer was detected in the liver, kidneys, and lungs compared to other assayed regions. Polymer was detected in almost every tissue sample, including the brain. MSs with diameters of 2 and 5 μm were delivered locally, but not by oral administration. n = 4 animals for each cohort. Data are shown for regions of the rat that were assayed following local delivery to the rat small intestine (jejunum and ileum) or following oral administration (fed). —, not measured.