

Detection of a fluorescent-labeled avidin-nucleic acid nanoassembly by confocal laser endomicroscopy in the microvasculature of chronically inflamed intestinal mucosa

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Abstract: Inflammatory bowel diseases are chronic gastrointestinal pathologies causing great discomfort in both children and adults. The pathogenesis of inflammatory bowel diseases is not yet fully understood and their diagnosis and treatment are often challenging. Nanoparticle-based strategies have been tested in local drug delivery to the inflamed colon. Here, we have investigated the use of the novel avidin-nucleic acid nanoassembly (ANANAS) platform as a potential diagnostic carrier in an experimental model of inflammatory bowel diseases. Fluorescentlabeled ANANAS nanoparticles were administered to mice with chemically induced chronic inflammation of the large intestine. Localization of mucosal nanoparticles was assessed in vivo by dual-band confocal laser endomicroscopy. This technique enables characterization of the mucosal microvasculature and crypt architecture at subcellular resolution. Intravascular nanoparticle distribution was observed in the inflamed mucosa but not in healthy controls, demonstrating the utility of the combination of ANANAS and confocal laser endomicroscopy for highlighting intestinal inflammatory conditions. The specific localization of ANANAS in inflamed tissues supports the potential of this platform as a targeted carrier for bioactive moieties in the treatment of inflammatory bowel disease.

Keywords: confocal laser endomicroscopy, inflammatory bowel disease, diagnostics, dextran sodium sulfate, avidin-nucleic acid nanoassembly, fluorescent nanoparticles, ulcerative colitis

Introduction

Crohn's disease and ulcerative colitis are chronic inflammatory bowel diseases (IBD) with a severe impact on social life, reducing quality of life and the ability to work as a consequence of increasing disability. IBD are chronic diseases with symptoms of bloody diarrhea and abdominal cramp. They generally begin in young adulthood and last throughout life, with nearly 80% of patients with Crohn's disease and 30% of those with ulcerative colitis requiring intestinal surgery. There is a strong need for novel strategies to improve the diagnosis and treatment of IBD. Indeed, the pathogenesis of these diseases is not yet fully understood¹⁻³ and their diagnosis and treatment are often challenging. Standard treatments to induce disease remission include aminosalicylates, corticosteroids, and immunosuppressive agents such as thiopurines and anti-tumor necrosis factor-α. Immunomodulator and combination therapies constitute a significant advance in the management of IBD.4-7 However, the onset of action is delayed with some of these drugs, and/or important adverse systemic effects often

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occur, including anaphylactic/anaphylactoid^{8,9} and lupus symptoms, and bacterial, viral, or fungal infections.^{10–12}

A possible way to reduce side effects without losing efficacy is to use targeting strategies to increase drug tropism to the inflamed sites. This strategy could improve drug therapeutic indices and the development of more effective treatment schedules. In recent years, several nanoparticle and colloidal systems have shown enhanced accumulation at inflamed sites due to hypervascularization¹³ or in tumors because of ineffective lymphatic drainage. These systems have the advantage of being able to carry a high drug payload/circulating unit, and in principle they could address the need for targeted delivery of active agents to IBD sites. Further, the mucosal epithelium is characterized by leaky structure in IBD¹⁵ and this particular feature may offer an opportunity for selective drug targeting through use of colloidal carriers.

In this work, we evaluated the behavior of a recently described nanoparticle platform, ie, an avidin-nucleic acid nanoassembly (ANANAS), when administered parenterally in a mouse model of IBD. ANANAS are fully biodegradable and biocompatible particles consisting of a compositionally defined poly-avidin core capable of accommodating more than 1,000 biotin-linked moieties of different kinds with unique stoichiometric and compositional control through a one-pot synthetic process. 16 Recently, the potential of the ANANAS formulation as a theranostic carrier for intravenous administration was demonstrated. The nanoparticles are capable of circulating for at least 2 hours before being eliminated within the next 2 days through classic scavenging systems with no apparent toxicity and low immunogenicity. 17 These findings paved the way for further exploitation of this nanoparticle tool for in vivo diagnostics and delivery applications.¹⁸ The ANANAS platform is especially interesting because it also meets the current need for convenient and reproducible preparation methods for multifunctional nanocarriers. 19 Multifunctionality in nanoparticles, while offering great potential for targeted treatments, significantly increases the complexity of preparation methods. As a consequence, the availability of standardized and reproducible synthetic procedures to control the multifunctional carrier composition is mandatory for translating research findings into clinical practice.

Here, in a further attempt to evaluate the clinical potential of this nanoparticle technology in nanomedicine, we generated a near red-emitting fluorophore-labeled ANANAS formulation (ANANAS-red⁶⁸¹), and performed a preliminary investigation of its distribution after intravenous administration in the colonic mucosa of an experimental model of

IBD. Confocal laser endomicroscopy (CLE) is a noninvasive fluorescence-based diagnostic tool for assessment of in vivo histology of the mucosa at subcellular resolution during ongoing endoscopy.²⁰ Probe-based CLE is a system designed for the in vivo characterization of the gastrointestinal mucosa and is currently used in the preclinical^{21,22} and clinical diagnostics^{23–25} of IBD. Intravenous administration of the green fluorescent dye fluorescein as a contrast agent allows visualization of the morphology of mucosal crypts and microvasculature. Dual-band CLE is a research-dedicated system that generates two-color laser excitation (488 nm/660 nm) and permits real-time dual-band fluorescence detection (500-630 nm/680-800 nm). This tool allows detection of two different fluorophore-related probes within the mucosal environment of interest and thus simultaneous visualization of different structures and/or targets.

In this pilot study, ANANAS-red⁶⁸¹ was administered intravenously together with low molecular weight fluorescein solution in order to determine the distribution of nanoparticles within the colonic mucosa, the architecture of which is highlighted by fluorescein. The red⁶⁸¹ fluorophore was used to enable nanoparticle visualization in the colonic microvasculature by CLE. In principle, the fluorophore can be substituted with bioactive compounds to develop a therapeutic application.

Materials and methods

Animal study approval

Procedures involving animals and their care were conducted in compliance with Research Project 27/2013 (protocol number 28893 May 13, 2013). The project proposal was approved by the Institutional Animal Care and Use Committee of the University of Padova according to Italian and European Commission regulations on Laboratory Animal Welfare and Protection (legislative decree number 116/92), which acknowledges the European Directive 86/609 EU.

Induction of dextran sodium sulfate colitis

Chronic colitis was induced in Balb/c mice^{26,27} (female, aged 8–10 weeks) with dextran sodium sulfate (DSS, molecular mass 40,000 Da), according to a protocol specifically optimized and described in detail in the Supplementary material. During treatment, animal body weight, rectal bleeding, stool consistency, and the presence of blood in stools were assessed daily. Assessment of chronic inflammation was confirmed by histological analysis.²⁸ Mice given untreated drinking water were used as controls.

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Synthesis of biotin derivatives used for nanoparticle assembly

The synthetic steps are summarized in Figure 1. Compound 1 and biotin-C₆-NH₂ (compound 2) were obtained as described elsewhere.²⁹ Compound 3 was obtained by reacting compound 2 with fluorescent red⁶⁸¹ succinimidyl ester (red⁶⁸¹-NHS, Sigma Aldrich, St Louis, MO, USA, catalog number 53433) in dry dimethyl sulfoxide at a molar ratio of 1:1.1 in the presence of triethylamine.

In more detail, 45.9 μ L of a biotin- C_6 -NH₂·HCl solution in dry dimethyl sulfoxide (9.3 mg/mL) was added to 49 μ L of red⁶⁸¹-NHS in dry dimethyl sulfoxide (20 mg/mL), followed twice by 7.7 μ L (2×7.7) of triethylamine 1:50 in dry dimethyl sulfoxide. The product (biotin-red⁶⁸¹) was characterized for its ability to bind avidin through gel permeation chromatography and for its fluorescence intensity before and after avidin binding (Supplementary material).

Preparation of ANANAS-red⁶⁸¹

The term ANANAS refers to a series of poly-avidin nanoparticle formulations originating from the high affinity interaction of avidin with nucleic acids.³⁰ These buffersoluble nanoparticles are formed by a spontaneous double self-assembly in which a nucleic acid molecule acts as a central element around which several avidins (one for each 14 base pairs) nucleate, leading to toroidal assemblies (diameter approximately 120 nm). In a second self-assembly process, a defined amount of biotinylated poly(ethylene glycol) (PEG) is added to permit buffer solubility and increase stability.²⁹ Finally, the ANANAS assemblies are polymerized forms of avidin. In this study, red⁶⁸¹-labeled ANANAS nanoparticles were synthesized in collaboration with ANANAS Nanotech (Padova, Italy) according to a procedure described elsewhere.¹⁷ In detail, a highly PEGylated ANANAS nanoparticle formulation was prepared and stabilized by mixing plasmid DNA and a complex of avidin and compound 1 at 1:0.25 molar ratio. 17,29 After purification and ultravioletvisible characterization of the nanoassembly, biotin-C_z-red⁶⁸¹ (compound 3) was added to at 1:3 avidin:biotin molar ratio. Finally, the mixture was purified from low molecular weight derivatives by gel permeation chromatography using a Sepharose 6-FF column on a fast protein liquid chromatography system (AKTA purifier, GE Healthcare, Little Chalfont, UK) with 10 mM phosphate, 150 mM NaCl pH 7.4 (phosphatebuffered saline) as the eluent. Four hundred and seventy five fluorophores/nanoparticles were found to be stably attached, as determined from the assembly ultraviolet-visible spectrum on the basis of the 280 nm (avidin, DNA, and red⁶⁸¹) and 681 nm (red⁶⁸¹) absorption values. The assembly size

Figure I Synthesis of biotin derivatives used in assembly of ANANAS-red^{sel}. **Abbreviations:** ANANAS, avidin-nucleic acid nanoassembly; DMSO, dimethyl sulfoxide; TEA, triethylamine; RT, room temperature; NHS, N-hydroxysuccinimide; PEG: poly(ethylene glycol).

was approximately 120 nm as measured by dynamic light scattering (Supplementary material).

Confocal laser endomicroscopy

We used the Cellvizio Dual Band system integrated with Proflex Ultra Mini O/Z probes (Mauna Kea Technologies, Paris, France) and a flexible endoscope (MAS-GM, Olympus, Italia S.r.l., Milano, Italy). Post-processing imaging analysis was performed using IC-viewer software. CLE was carried out on anesthetized healthy and diseased mice after endoscopic confirmation of mucosal damage. Fluorescein in phosphate-buffered saline solution, in the presence or absence of ANAN-AS-red⁶⁸¹ (100 $\mu g/mouse$), was administered intravenously (100 μL) in healthy and DSS-treated animals (three animals per condition) and CLE images were acquired 15–30 minutes after injection. Longer observation times could not be used due to the rapid fluorescein clearance.

Results and discussion

Optimization of fluorescein dosage for dual-band CLE investigation

Preliminary experiments were carried out to optimize experimental conditions to provide the most valuable biological information using the dual-band probe. When using dual-band fluorescence imaging, if the signal generated by one fluorophore is too high, signal cross-over into the second channel may occur, resulting in false-positive readings. In our experiments, the concentration of red⁶⁸¹ dye in the injected nanoparticle formulation was several orders of magnitude lower (0.00167% w/v) than that of the fluorescein dye normally used for single-band CLE-assisted visualization of the microvasculature (between 10% and 0.5% w/v).31 Since this was the first application of dual-band CLE for preclinical research in IBD, we tested a range of fluorescein concentrations to eliminate the risk of a falsepositive signal from fluorescein interfering with specific red⁶⁸¹-related signals.

In this preliminary investigation, healthy and inflamed animals were injected with different fluorescein solutions (0.1%, 1.0%, and 10% w/v in phosphate-buffered saline) at $100\,\mu\text{L/dose}$ to identify the fluorophore concentration allowing the morphology of the vasculature (in the green channel) to be reliably distinguished without generating cross-over of fluorescent emission in the red channel. Only the intermediate fluorescein concentration tested was found to be suitable for our purposes (Figure 2).

Administration of the 10% dye solution generated a strong and diffuse signal in the green channel, together with

a significant bleed-through of fluorescent emission in the red channel. The diffuse signal, likely due to diffusion of the dye from the vessels to the surrounding stroma, impaired visualization of crypt and microvasculature architecture. In addition, the strong spectral cross-over effect prevented reliable identification of a positive signal due to the red-emitting dye in the red channel. Using 0.1% fluorescein, no bleed-through was observed in either healthy or inflamed mucosae, but the microvasculature pattern was poorly defined. Finally, 1% fluorescein was found to be suitable for our purposes, because it allowed the crypt and microvascular architecture to be visualized in a manner similar to that possible with use of 10% fluorescein in human clinical practice, 23 with negligible spectral cross-over in either healthy or chronically inflamed mucosae.

ANANAS-red⁶⁸¹ reaches mucosal microvasculature in inflamed but not healthy bowel

ANANAS-red⁶⁸¹ were coinjected with 1.0% fluorescein. Selective detection of nanoparticles was observed exclusively in the inflamed tissue (Figure 3). The red signal appears in the inflamed vessels (highlighted by the green fluorescein) only, and is clearly related to the presence of ANANAS-red⁶⁸¹, as confirmed by the merged images. The speckled red signal is likely due to the low fluorophore concentration (0.00167% red⁶⁸¹ versus 1% fluorescein w/v), the discrete concentration of nanoparticles, and the lower relative brightness of red⁶⁸¹ as compared with fluorescein (not shown). The results therefore confirm that, like other colloidal drug carriers, ANANAS nanoparticles are capable of selectively reaching inflamed tissues. Of note, when CLE probing was performed on healthy regions of the patchily inflamed colon, no red⁶⁸¹ signal was observed (not shown). This indicates that the system can discriminate between healthy and inflamed areas in the same organ.

Conclusion

In these preliminary studies, dual-band CLE was used for the first time to characterize the architecture of the microvasculature and visualize fluorescent-labeled nanoparticles simultaneously in an in vivo model. A distinctive pattern of distribution of intravascular nanoparticles was observed in chronic inflamed mucosa compared with healthy controls. The results confirm that ANANAS nanoparticle-based delivery strategies have the ability to reach the microvasculature of inflamed bowel mucosa together with their cargo. This result further supports the potential of this novel soft biodegradable

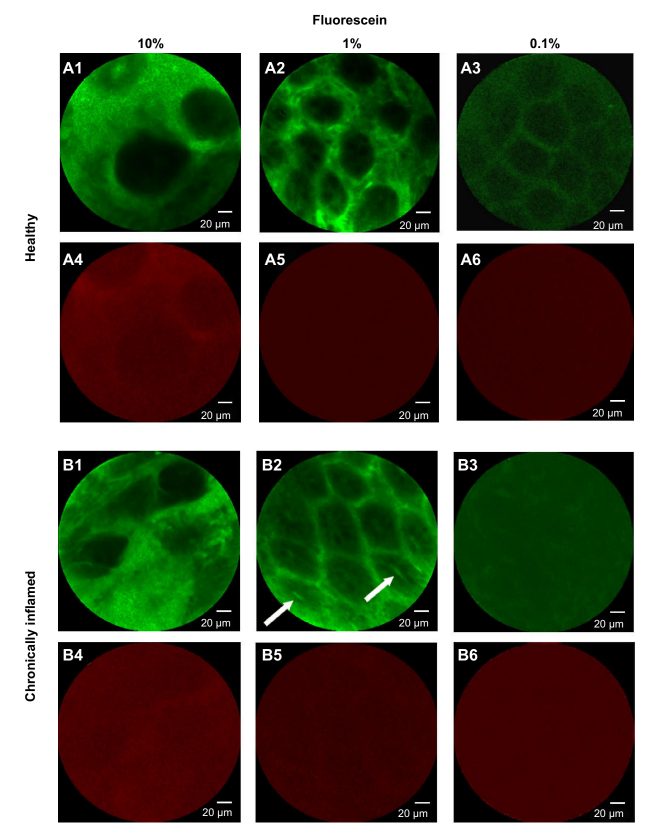


Figure 2 Confocal laser endomicroscopic images of healthy and chronically inflamed mucosa in the green and red channels.

Notes: The images were obtained after intravenous administration of fluorescein at different concentrations (10%, 1.0%, and 0.1% w/v). Each image represents one example of three per group of animals per condition. The images obtained in the green channel (A1–A3 and B1–B3) show the hexagonal honeycomb appearance of the mucosa. Nonideal fluorescein concentrations yield either overexposed images (10%, A1 and B1) or loss of details (0.1%, A3 and B3). Normal mucosa shows a network of capillaries surrounding the crypt openings with a honeycomb appearance. In inflamed mucosa, the crypt lumen is large and variably shaped, with fluorescein leakage into the crypt openings (arrows in B2). In the red channel (A4–A6 and B4–B6), any visible structure (A4 and B4) is due to fluorescein-related signal cross-over.

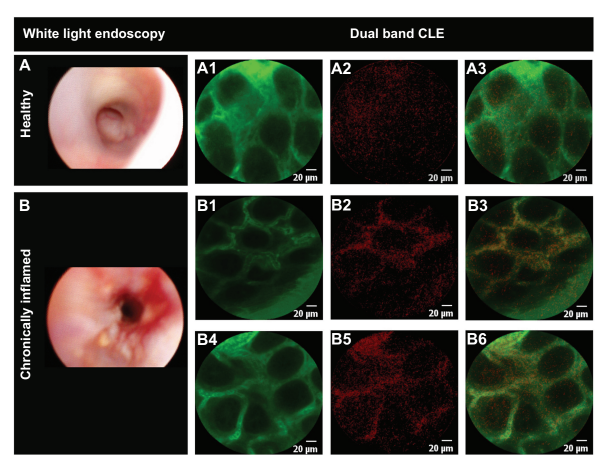


Figure 3 Differential nanoparticle distribution in inflamed and healthy mucosa.

Notes: Selected images of healthy (A) (n=3) and chronically inflamed (B) (n=3) mouse mucosa, upon intravenous administration of ANANAS-red⁶⁸¹ in phosphate-buffered saline containing 1% fluorescein. CLE images were acquired in the green (A1, B1, B4) and red (A2, B2, B5) channels. Merged red and green images are shown in panels A3, B3, and B6. The yellow color corresponds to the merged signal from green and red channels.

Abbreviations: ANANAS, avidin-nucleic acid nanoassembly; CLE, confocal laser endomicroscopy.

nanoparticle platform as a tool for targeted delivery of diagnostic agents. It should be noted that due to the high compositional versatility of the platform and its ease of formulation and tuning, part or all of the diagnostic elements could be substituted with therapeutic agents, making the nanosystem a potential carrier for targeted drug delivery to inflamed mucosa. The results are encouraging for future development of the platform, although additional studies are necessary before translation into clinical practice. On the diagnostic side, future experiments aimed at determining the duration of distribution and optimization of nanoparticle concentration are warranted. In the context of targeted drug delivery, research should focus on the development of linker technologies to enable efficient local release of the biotin-linked bioactive agent as soon as the nanoparticles reach their target.

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Disclosure

MM is among the inventors of the ANANAS technology, and is a shareholder in ANANAS Nanotech s.r.l., a spinoff company that holds the intellectual property related to the ANANAS formulation. The other authors report no conflicts of interest in this work.

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Supplementary material

Figure S1 shows the gel permeation chromatography analysis of biotin-C₆-red⁶⁸¹ (compound 3) before and after addition of avidin. Figure S2 shows the fluorescence intensity calibration curves of the same compound in phosphate-buffered saline, alone and in the presence of avidin. Figure S3 shows the results of dynamic light scattering analysis of the avidin-nucleic acid nanoassembly (ANANAS)-red⁶⁸¹ nanoparticles. Figure S4 shows histological images of colonic mucosa in dextran sodium sulfate (DSS)-treated and healthy mice.

Characterization of biotin-C₆-red⁶⁸¹

For analysis, a small aliquot of the coupling reaction solution (0.5 μ L) was diluted to 218 μ L with phosphate-buffered saline alone or phosphate-buffered saline containing avidin 1 mg/mL (Belovo Chemicals, Bastogne, Belgium), and analyzed by fast protein liquid chromatography (Superdex peptide column with phosphate-buffered saline as the eluent, 0.5 mL/min, ultraviolet absorption 681 nm). Figure S1 shows the overlapped chromatograms of the starting red⁶⁸¹-NHS, the reaction product as before and after the addition of avidin, with peak assignments. The hydrolyzed red⁶⁸¹-NHS (Red⁶⁸¹-COOH) and its biotin derivative were tested for fluorescence intensity. The biotin derivative was also tested upon addition of avidin. Fluorescence intensity calibration curves

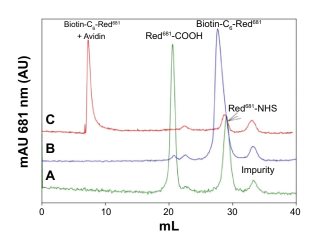


Figure S1 Chromatographic analysis of red⁶⁸¹ derivatives. (**A**) Starting NHS reagent, (**B**) biotin- C_e -red⁶⁸¹ derivative (compound 3), and (**C**) compound 3 in the presence of avidin.

Notes: Analyses were carried out using fast protein liquid chromatography apparatus (Akta Purifier, GE Healthcare, Little Chalfont, UK) equipped with a Superdex peptide column and eluted with 10 mM phosphate, 150 mM NaCl, pH 7.4 (phosphate-buffered saline) at 0.5 mL/min. Chromatograms were registered at 681 nm

Abbreviations: ANANAS, avidin-nucleic acid nanoassembly; NHS, N-hydroxysuccinimide.

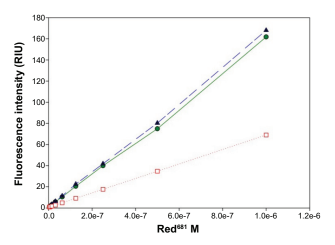


Figure S2 Fluorescence intensity versus concentration of red⁶⁸¹-COOH (\bullet) and biotin-C_c-red⁶⁸¹ as before (\triangle), and in the presence of avidin (\square). **Abbreviation:** RIU, relative intensity units.

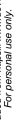
were generated on serially diluted phosphate-buffered saline solutions, starting from 1 μ M red⁶⁸¹, with $\lambda_{\rm exc}$ 681 nm and $\lambda_{\rm em}$ 708 nm. The slope of the fluorescence intensity versus concentration curve was used to calculate the relative fluorescence intensity with respect to the starting red⁶⁸¹-COOH. The fluorescence intensity of avidin-linked biotin-C₆-red⁶⁸¹ was 43% that of the same reagent when analyzed free in solution. The same result was obtained using ANANAS instead of avidin (not shown).

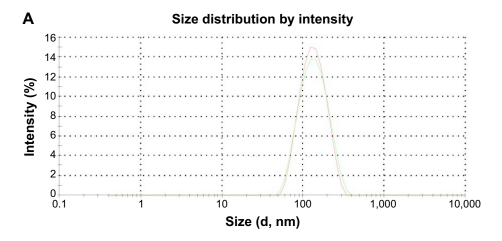
Size distribution of ANANAS nanoformulation

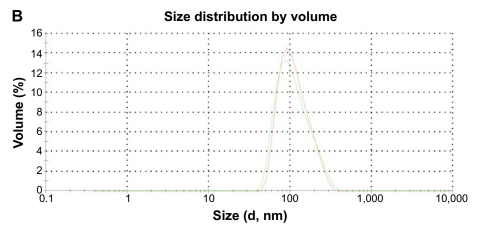
Dynamic light scattering measurements were performed using a Malvern Zetasizer Nano ZS (Malvern Instruments Ltd, Malvern, UK) and the results are shown in Figure S3.

Induction of DSS chronic colitis

Chronic colitis was induced in Balb/c female mice^{1,2} (aged 8–10 weeks) by using multiple cycles of dextran sodium sulfate (DSS, 40 kDa, Sigma-Aldrich, St Louis, MO, USA). We performed preliminary optimization experiments to identify the optimal DSS dose and administration schedule yielding chronic inflammation. During treatment, animal body weight, rectal bleeding, stool consistency, and the presence of blood in stools were assessed daily. Assessment of chronic inflammation was confirmed by histological analysis. The final induction protocol identified is reported below: initially, we administered two cycles comprising 7 days of 3% DSS in drinking water and 3 days of untreated drinking water. Next, 5% DSS was administered for 3–4 days until a 10% weight







ANANAS formulation size distribution (nm)		
By intensity	By volume	Z-average
143±55	118±57	123.0

Figure S3 (A) Intensity and (B) volume weighted size distribution of ANANAS assembly prepared for this work, as determined by dynamic light scattering. Abbreviation: ANANAS, avidin-nucleic acid nanoassembly

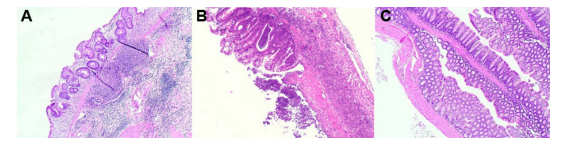


Figure S4 Histology images of dextran sodium sulfate-treated (A, B) and healthy (C) mouse colonic mucosa. Notes: Colonic segments, fixed in 4% formaldehyde, were stained with hematoxylin and eosin and reviewed by a pathologist in a blinded fashion. Original magnification for histology, 40×.

loss was achieved, then substituted with untreated drinking water for 3 days or until full body weight recovery. The 5% DSS/untreated drinking water cycle was repeated four times. Inflammatory changes³ (crypt architectural disarray, transmural inflammation with lymphoid follicles) were confirmed histologically in colonic sections from DSS-treated mice compared with control mice (Figure S4).

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