Properties and applications of nanoparticle/microparticle conveyors with adjuvant characteristics suitable for oral vaccination

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Abstract: Vaccination is one of the most effective approaches in the prevention and control of disease worldwide. Oral vaccination could have wide applications if effective protection cannot be achieved through traditional (eg, parenteral) routes of vaccination. However, oral administration is hampered by the difficulties in transferring vaccines in vivo. This has led to the development of materials such as carriers with potential adjuvant effects. Considering the requirements for selecting adjuvants for oral vaccines as well as the advantages of nanoparticle/microparticle materials as immune effectors and antigen conveyors, synthetic materials could improve the efficiency of oral vaccination. In this review, nanoparticles and microparticles with adjuvant characteristics are described with regard to their potential importance for oral immunization, and some promising and successful modification strategies are summarized.

Keywords: oral vaccine, NP/MP conveyors, intestine, antibody, immune response

Basic features of oral vaccination

Vaccination started in 1796 when Edward Jenner experimented with the cowpox virus.1,2 Vaccination protects against various infectious and noninfectious diseases and has saved countless lives during the past two centuries.3–5 The classical antigens for vaccination are based on various attenuated live pathogens (viruses, bacteria). “Optimized vaccines” (such as those based on the subunits [proteins, nucleotides] of viruses and bacteria) carry, in general, fewer risks of virulence and reversion.6

Usually, vaccines are administered by injection. This has the problem of relatively weak immune responses due to insufficient stimulation of mucous membranes.2,7 Considering the robust immune system in the intestines and increased convenience of oral immunization, the latter has garnered a great deal of attention recently. Compared with the traditional vaccination route, oral vaccination reduces the pain of vaccination and lowers the risk of wound infection/inflammation. Also, based on a rationally designed oral vaccination system, the cost is considerably reduced because robust and integral immune responses can be elicited.

A series of immune-associated tissues and cells has been demonstrated within small- and large-intestinal tracts. These include, for example, the Peyer’s patches in the small intestine and lymphoid follicles within the large intestine, which are important for antigen presentation and appropriate functioning of the immune system to verify and fight infection. Compared with vaccination by subcutaneous routes, the large-intestinal mucosa provides an ideal site for stimulation of effective immune responses...
because of the robust immune systems and greatly reduced number of enzymes within the large-intestinal tract.

Oral vaccines are desirable, but the harsh conditions within the gastrointestinal tract (GIT) mean that they can be degraded by various enzymes, restricted from antigen presentation by the mucosal barrier, and have to compete with microorganisms. Therefore, only a few products are available for oral vaccination for prophylaxis against some infectious diseases. Most licensed oral vaccines or those undergoing manufacture are based on attenuated viruses or bacteria that can survive the GIT environment and produce antigens continuously, which can boost immune responses. For example, Rotateq®, Rotashield®, Tetramune®, Vivovax®, Dukoral®, Kolera, and CVD 103-HgR (known as Orochol® or Mutachol®) are being developed against rotavirus, typhus, or cholera.

Attenuated viruses can revert back to virulent pathogens. Progress in vaccination technology means that adjuvants can be designed to protect nontoxic peptide- or nucleotide-based oral vaccines in the GIT.

**Nanoparticle/microparticle conveyors could be suitable for oral vaccination**

Adjuvants are molecules, compounds, or complexes that promote the potency and longevity of a specific immune response to antigens, but bring about minimal toxicity or long-lasting immune effects on their own. Bacterial factors, plant components, synthetic materials, and other immunity-related factors have been used as efficient adjuvants. With the development of nanoparticle/microparticle (NP/MP) technology, adjuvants can also refer to NPs/MPs that function as antigen-protective vehicles, which function to endow the duration release and presentation of antigens, as well as to act on the immune system directly. In this way, vaccines obtain protection and enhancement from adjuvants. With the integration of vehicles and immunostimulants, rationally designed synthetic NP/MP adjuvants possess the functions of both and may be feasible for oral vaccination due to their targeted release, the capacity for loading and protecting immune effectors, and protection of organisms against large amounts of antigens that might be cytotoxic.

As a common requirement for oral vaccination, adjuvants should be biocompatible, have stable and controlled release, protect antigens from degradation in the GIT tract, have targeted delivery, and be capable of controlled release of antigens to the target immune cells for enhanced presentation of the antigens. These adjuvants would be more efficient if phagocytosis of antigens could be stimulated. As representatives, several successful NPs/MPs with adjuvant functions that are suitable for oral vaccination have been described and their properties and applications summarized.

**Organic polymer adjuvants**

Synthetic organic materials are major choices for adjuvants because of their nonsubstitutable biocompatibility and degradability, ability to protect antigens, and/or ability to directly influence immune responses. Poly(lactic acids) (PLAs), poly(D,L-lactide-co-glycolide) (PLG), poly(lactic-co-glycolic acid) (PLGA), poly(alkyl cyanoacrylates), poly(methyl methacrylates), liposomes, and lipopolysaccharides have demonstrated efficiency in oral vaccination and provided safety, efficacy, and outstanding performance. Recently, novel NPs/MPs based on organic adjuvants with diverse components and structures have been explored with additional properties, such as enhanced stability and targeted delivery of antigens (Figure 1).

**Classical linear adjuvants**

Poly(D,L-lactic acid) polymers based on lactic acids with meritorious biocompatibility are synthetic biomaterials approved for use in the GIT tract. Except for their degradability, PLA NPs can be phagocytized by immune-associated cells. Oral vaccines based on PLA NP/MP-entraped rotavirus can stimulate long-lasting responses from immunoglobulin (Ig)G and IgA antibodies. Insufficient stability of PLA and weak protection of antigens in the GIT are major issues with regard to NPs/MPs based on PLA adjuvants.

Besides PLA, PLGA and polyethylene glycol (PEG) are the most popular organic materials with biocompatibility and biodegradability. PLGA can also be formulated to NPs or MPs for clinical use. Various antigens can be loaded within PLGA- or PLGA-based conveyors, and their antigenicity maintained fully. In addition, PLGA NPs can be internalized by professional antigen-presenting cells. PLGA NPs are practical pharmaceutical excipients approved by the US Food and Drug Administration. Oral administration of PLGA/bovine serum albumin (BSA) particles has been reported to elicit higher responses by IgG antibodies in serum. By use of targeted ligands, the targeting ability of PLGA vehicles can be realized. PEG has substantial solubility and stability in water and can reduce the degradation of vectors or increase their dispersion under aqueous conditions, so it is frequently used to modify NPs/MPs. The surface antigen of the hepatitis B virus (HBsAg) delivered by PEG–PLA NPs shows enhanced stability in simulated gastric and intestinal
Humoral immunity along with mucosal (secretory IgA) and cellular immune responses have been stimulated through oral vaccination.

**Novel NPs/MPs designed with adjuvant characteristics for oral vaccination**

**Organic adjuvant NPs/MPs**

With the aim to enhance protection of antigens and strengthen stimulation of immune responses, antigens can be encapsulated and released from polymers or complexes with more complicated structures, such as in cross-linked porous materials or assembled NPs or MPs such as micelles, lipid vectors, and complexes based on electrolytes.

The cross-linked materials provide porous structures with large surface areas for efficient loading and controlled release of antigens. Based on these materials, adjuvants for oral vaccination are designed for stability and antigen loading in the stomach, but are exposed to a certain amount of instability at the target position where durative antigen release is allowed. Microspheres based on cross-linked polyacryl starch (PAS) and those based on polymer-grafted starch have been developed as adjuvants for oral vaccines. Human serum albumin (HSA) covalently linked to PAS MPs or HSA entrapped by starch MPs grafted onto 3-(triethoxysilyl)-propyl-terminated polydimethylsiloxane can stimulate effective immune responses. Oral administration of the former adjuvant can elicit strong cellular as well as IgM/IgG-associated humoral and mucosal responses, whereas the latter can stimulate mucosal immune responses.

Other cross-linked adjuvants explored successfully include poly[di(carboxylatophenoxy) phosphazene] (PCPP), poly(di(sodium carboxylatoethylphenoxy) phosphazene) (PCEP), and hydrogels. PCPP is biocompatible and sensitive to pH (Figure 2). It is stable in neutral-to-acidic aqueous conditions, and this cross-linking process is reversed in basic conditions. PCPP is a potent immunoadjuvant for influenza vaccines, and some sodium-salt PCPPs are undergoing clinical trials presently. Polyphosphazene derivatives can enhance the immune responses of vaccines against multiple antigens, such as recombinant HBsAg. For example, PCEP adjuvants are water soluble and can form microparticulates for oral uptake. Co-delivery of PCEP with the influenza X:31 antigen can enhance antigen-specific Ig production and stimulate IgA, IgG1, and IgG2a titers, and enhanced titers of the latter via the oral route have been reported.

Hydrogels with networks can absorb and hold different water contents depending on their components and structures. Polymeric hydrogels can be designed as adjuvants with versatile features, such as a suitable degradation rate, antigen-loading capacity, and environmental sensitivity. For example, the response to pH and temperature of poly(methacrylic acid) and mannann-loaded biodegradable thermosensitive poly (ε-caprolactone)-poly(ethylene glycol)-poly(ε-caprolactone) (PCL–PEG–PCL, PCEC), respectively, have been explored.
for the controlled release of antigens. Both have shown great potential for oral applications in the form of hydrogel or hydrogel microspheres.56–58

Unlike cross-linked materials, branched polymers have complicated structures that can confer versatile properties on polymers for antigen delivery and immunity enhancement. These properties may have suitable amphility, biodegradability, molecular weights/sizes, and target ligands. Sulfobutylated poly(vinyl alcohol)-graft-poly(lactide-co-glycolide) has been developed as an NP adjuvant for the delivery of tetanus toxoid and can stimulate IgG and IgA serum titers through oral administration.59 Among branched polymers, “treelike” dendrimers can be synthesized in a specific order from the core or edge. They have now shown great potential for application in oral vaccination.60–65 For example, tetragalloyl-D-lysine dendrimer conjugated with rhesus C–C chemokine receptor type 5-derived cyclopeptide can target microfold (M) cells, followed by an IgA response in stools and neutralization activity against infection by the simian immunodeficiency virus in rhesus macaque.66

Self-assembled materials can assemble to NPs or MPs of different shapes automatically. Compared with single molecules, self-assembled vectors can be designed with complicated functions based on relatively simple units that show convenience in terms of polymerization and antigen encapsulation. These self-assembled materials have been explored as novel adjuvants for oral vaccination. Micelles can aggregate in a liquid colloid. Usually, their polar regions are in contact with the surrounding solvent, and the hydrophobic single-tail regions are sequestered in the center. Micelles can load water-soluble and poorly water-soluble antigens, respectively, in hydrophilic blocks and hydrophobic blocks. Through oral administration, a “reverse micelle structure” of phosphatidylcholine encapsulating a baculovirus displaying influenza hemagglutinin-enhanced specific serum IgG and mucosal IgA antibodies in mice has been noted. Meanwhile, 100% protection was conferred on vaccinated mice against 5MLD50 (mouse lethal dose 50%) of highly pathogenic avian influenza heterologous H5N1 strain.67

Different from a micelle, a liposome has a lipid bilayer and an aqueous solution core surrounded by a membrane composed of hydrophobic groups. The sizes of liposomes range from nanometers to micrometers. Water-soluble antigens can be encapsulated within its hydrophilic core, and hydrophobic antigens can be associated with its hydrophobic chemicals. Both forms are stable for the protection and controlled release of antigens. A liposome can stimulate the internalization of the antigen and can target M cells. Therefore, liposomes have been used widely for the oral delivery of drugs and modification of oral vaccines.68 A liposome encapsulating antigens/vaccines,

Figure 2  Synthesis of PCEP.
Notes: Adapted with permission from Andrianov et al. Synthesis, properties, and biological activity of poly[di(sodium carboxylatoethylphenoxy)phosphazene]. Biomacromolecules. 2006;7:394–399. Copyright 2006 American Chemical Society.
Abbreviation: PCEP, poly(di(sodium carboxylatoethylphenoxy) phosphazene).
for example, LiposoMAs (liposomal-antigens), liposomes functionalized with distearoyl phosphatidylcholine and cholesterol, and 1,2-dioleoyl-3-trimethylammonium propane-based liposomes, can stimulate specific mucosal responses through oral immunization and are accompanied frequently with a mixed T-helper (Th1/Th2 immune responses and cellular/humoral immunity. Other synthetic oral vaccines based on liposomes explored include those coated with a bilayer of *Tremella* and acid-induced alginate, or covalently bound with *Ulex europaeus* 1 (UEA1) lectin.

Bilosomes are adjuvants with strengthened stability and protection of peptide antigens as compared with liposomes (Figures 3 and 4). They are stabilized by sodium and potassium bile salts of bile acids conjugated with taurine/glycine and are more resistant to the harsh conditions in the GIT. Bilosomes composed of bile salts and lipid vesicles have been developed as NP or microsphere adjuvants for oral vaccination. Modified bilosomes (eg, glucomannosylated bilosomes) are suitable for oral vaccination, and some bilosomes conjugated with other molecules, such as the B subunit of the cholera toxin, can target M cells.

Charged polymers can affiliate to charged cell surfaces or maintain the dispersion capacity of particles in aqueous solutions. They have been introduced to vaccine vehicles (usually as coating materials) to improve their mucoadhesive properties or the immune responses of organisms. Cationic polymers such as polyethyleneimine (PEI), chitosan (CTS), and polydimethylallyl ammonium (PDDA) have been used as adjuvants because they can delay the appearance of peak IgG titers and improve IgA titers. With strong mucoadhesion and sustained release, calcium–alginate and calcium–yam–alginate MPs coated with N-(4-Н,N-dimethylaminocinnamyl)
CTS stimulate increased IgG and IgA titers. Recently, we developed a “PMMMA” nanolayer with pH sensitivity. This nanolayer was covalently synthesized through free-radical polymerization at the surface of PLGA/antigen NPs using poly(methacrylic acid) harboring carboxyl groups, methyl methacrylate (MMA), and methyl acrylate (MA) as precursors. Ionization of the –COOH group in the MAA blocks in PMMMA occurred under weak alkaline conditions. This leads to repulsive negative surface charges surrounding the NPs and stops the PLGA/antigen NPs from adhering to mucin/epithelial cells in the small intestine. Increasingly, alkaline conditions in the large intestine activate the pH-responsive phase transition of PMMMA and incur antigen release. Through oral vaccination, PMMMA-PLGA/TRX-SIP (surface immunogenic protein) NPs were shown to stimulate durable immunoprotection in *Tilapia* against group-B Streptococcus (GBS) disease, with a high level of enhancement of specific IgM antibodies.

**Porous inorganic particles**
Adjuvants based on organic materials are degraded frequently as they travel through the GIT. Traditional calcium- or aluminum-based adjuvants given parenterally are also unstable. Porous inorganic adjuvants based on silica and carbon have

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**Figure 4** Bilosome-based adjuvants for oral vaccination.
**Notes:** Assembly of the bilosome-based oral vaccines comprising liposome, bile salts, and the hydrophilic antigens. Protection of bilosome on the antigens enhance their resistance against the harsh environment in GIT.
**Abbreviation:** GIT, gastrointestinal tract.
been developed recently for oral vaccination because they have improved consolidation, low toxicity, large inner spaces for antigen loading, and well-designed antigen release. The particle sizes, degree of porosity, pore sizes, and decorations for improving the dispersity or other properties of the adjuvants can be controlled through modulation of synthetic approaches.\textsuperscript{86–89} With relatively lower toxicity, some nanometer-to-micrometer-sized carbon- and silica-originated porous particles have demonstrated safety and efficacy for oral administration.\textsuperscript{82}

Porous carbon NPs can be “tailored” with attractive properties for oral vaccination. Their porous structures allow them to load and release antigens at suitable velocities. Due to their insolubility in aqueous media and their cytotoxicity, modification of carbon NPs can improve their properties for application in oral immunization.\textsuperscript{90,91} A porous hydrophobic carbon NP termed “C1” (470 nm) can protect antigens in the GIT and promote their uptake by M cells. Oral vaccination of C1 encapsulating a BSA system was shown to stimulate IgG titers efficiently; mucosal IgA responses and Th1/Th2-mediated responses were also enhanced.\textsuperscript{92}

CaCO\textsubscript{3} MPs with porous structures have been explored as inorganic adjuvants for oral administration (Figure 5). Dissolving chicken egg ovalbumin peptide linked to self-assembling peptide KFE8 (Ac-FKFEFKFE-Am) (OVA-KFE8) nanofibers in CaCl\textsubscript{2} and Na\textsubscript{2}CO\textsubscript{3} solutions can achieve production of OVA-KFE8/CaCO\textsubscript{3} composite MPs, with OVA-KFE8 encapsulated stably within the core. Mucosal antibody responses were elicited through oral vaccination due to efficient penetration of the mucus barrier.\textsuperscript{93}

Porous silica particles can adsorb antigens and shield them from exposure to harsh GIT environments such as low pH in the stomach and enzymes in the intestines. As representatives, Santa Barbara Amorphous SBA-15 nanostructured silica, and S1 and S2 silica NPs can modulate the immune responsiveness of mice via the oral route.\textsuperscript{84} By use of organic structure-directing agents such as poly(ethylene oxide)-poly(propylene oxide)-poly(ethylene oxide) (PEO–PPO–PEO), mesoporous SBA-15 NPs can be developed as cylinders (length: 2 µm; mean diameter: 10 nm).\textsuperscript{85} Recovering organic polymers at high temperature can generate thermally stable products. BSA-15 can evoke immunologic memory and provide sustained release of antigens.\textsuperscript{94,96} Application of S1 and S2 for oral vaccines has been reported. Oral immunization of spherical S1 (430 nm) and S2 (130 nm) encapsulating BSA can induce higher levels of IgG and IgA titers, as well as Th1- or Th2-mediated responses. These silica particles are usually stable and biocompatible.\textsuperscript{96} Obtaining advanced features by modification of silica particles with biocompatible organic polymers is promising. For example, Aerosil R972 microspheres composed of CTS and nanometer-sized hydrophobic silica have been endowed with enhanced acid stability, improved moisture absorption, improved thermal

Figure 5 SEM scanning analysis.
Notes: (A) SEM micrographs of control CaCO\textsubscript{3} microparticles depicting spherical morphology and a highly porous interior. (B) OVA-KFE8 nanofiber-loaded composite microparticles with a denser surface and core. Adapted from Snook JD, Chesson CB, Peniche AG, et al. Peptide nanofiber-CaCO\textsubscript{3} composite microparticles as adjuvant-free oral vaccine delivery vehicles. J Mater Chem B. 2016;4:1640–1649.\textsuperscript{93} http://pubs.rsc.org/en/content/articlelanding/2016/tb/c5tb01623a#!divAbstract, with permission of the Royal Society of Chemistry.

Abbreviation: OVA-KFE8, chicken egg ovalbumin peptide linked to self-assembling peptide KFE8 (Ac-FKFEFKFE-Am).
In this respect, most single-component adjuvants, such as those antigens can be released to stimulate immune responses. Conversely, they should be bioutilizable or biodegradable so as to protect antigens and transport them to immune cells by enhancing transporter-mediated cell-active absorption. NP/MP adjuvants are expected to be used as adjuvants for oral vaccination based on molecules or polymers that can stabilize vaccines through antigen encapsulation.

To conquer the harsh conditions in the GIT, most adjuvants are based on molecules or polymers that can stabilize vaccines. PLA microspheres are known to enhance the stability of vaccines through antigen encapsulation.26 PEG can shield particles from enzyme digestion and is used frequently to modify oral vaccines. Combined use of PEG and PLA can achieve stimulation of mucosal uptake and immune responses through oral administration. Hence, this is dependent on the characterization of the functional components and specific structures of the NPs/MPs.

Based on their chemical constituents and structures, the NP/MPs used for oral vaccination can act as vaccine stabilizers, antigen modulators, immunity enhancers, target tracers, environmental sensors, and enhancers of cell uptake. Adjuvants can be designed with high biocompatibility, appropriate stability, antigen-loading and maintenance ability, targeting properties, and environmentally responsive release. Through rational design, synthetic adjuvants are expected to have integrated plasticity and functionality.28 Some representative synthetic adjuvants with specific roles in oral vaccination are discussed below.

The GIT has a vast immune system. The small intestine possesses gut-associated lymphoid tissue (GALT) comprising Peyer’s patches.99 The large intestine and rectum possess lymphoid follicles. Lymphoid tissues and other types of immune cells are also found in the lamina propria of the gut, the esophagus, and the appendix. The small and large intestines provide efficacious target sites for oral vaccination. Peyer’s patches can present antigens from the digestive tract to the immune system. M cells are found in GALT and mucosa-associated lymphoid tissue of other parts of the GIT. Mucosal immune responses can be initiated on the apical membrane of M cells. With the aid of M cells, antigens in the gut lumen can be transported across the epithelial cell layer to the lamina propria to interact with immune cells.

Targeting vaccines to the small intestine is a popular strategy. However, oral vaccines that can target small intestines filled with enzymes (proteases, esterase, trypsin) have rarely achieved success in practice.8 Vectors and antigens can be inactivated severely. NP/MP adjuvants are expected to protect antigens and transport them to immune cells by enhancing transporter-mediated cell-active absorption.100 Conversely, they should be bioutilizable or biodegradable so that antigens can be released to stimulate immune responses. In this respect, most single-component adjuvants, such as PLGA, lose their protective function or ability to release antigens in the GIT before cell uptake.

The large intestine contains fewer enzymes than the small intestine and is considered more suitable for oral vaccination.101 Immune responses are elicited in regional lymph nodes in the mucosa of the large intestine.102 The conditions are quite different in the stomach, small intestine, and large intestine, so tailoring of adjuvants becomes more complex. The adjuvants’ sensitivity to pH and their ability to escape from the small intestine should be considered.

Characteristics and mechanisms of functional NPs/MPs used as adjuvants for oral vaccination

The rational design and tailoring of adjuvants can realize versatile functions that facilitate their use for oral vaccination. Some NPs/MPs suitable for oral vaccination can enhance the immune responses of organisms and help them to develop cellular, mucosal, and systemic immunostimulatory responses through oral administration. However, this is dependent on the characterization of the functional components and specific structures of the NPs/MPs.

Antigen protection

To conquer the harsh conditions in the GIT, most adjuvants are based on molecules or polymers that can stabilize vaccines. PLA microspheres are known to enhance the stability of vaccines through antigen encapsulation.26 PEG can shield particles from enzyme digestion and is used frequently to modify oral vaccines. Combined use of PEG and PLA can achieve stimulation of mucosal uptake and immune responses (including humoral immunity) as well as secretory IgA mucosal and Th1 immune responses.48

Controlled release of antigens

Antigens must be presented to immune cells for the development of immune responses. The stability and release of
antigens are critical in this process. The particle size, dispersity in water, degradation velocity, and hydrophilicity/hydrophobicity are associated with antigen release in this context. PEG of different lengths can modulate water dispersion and shield antigens from enzymes. Solid lipids of suitable degradation rate can release antigens at a relatively lower speed. Resins can downregulate the degradation rate and have been studied for oral vaccination. Using the methacrylate-based polymer Eudragit® FS30D encapsulating PLGA MPs (>10 μm), particles can escape absorption and degradation in the stomach and small intestine, but release the inner vaccines in the large intestine, where the pH is increased (Figure 6).8

Promotion of cell uptake
Successful presentation of antigens requires an efficient cell uptake of antigens by immune cells. The more intact the epitope, the more efficient and specific are the immune responses. Bioutilizable polymers can promote transporter-mediated transmembrane movement of encapsulated antigens.100 Polyesters, polylipids, polysaccharides, and some biomolecule-based polymers are ideal candidates.103–106 Particle size is a critical factor for cell uptake. In general, particles of size <300 nm are preferred for cell internalization. Through modification of some positively charged polymers, such as liposomes and PEI, cell uptake can be stimulated and toxicity lowered.107 With the aid of adjuvants such as PLGA NPs and porous silica particles, the concentration of the internalized antigens can be increased considerably.

Integration of targeted constituents
Targeting strategies are expected to resolve the problems associated with the low concentration and continuous degradation of antigens in the GIT. The 9-amino M-cell-homing peptide CKSTHPPLSC (CKS9) can affiliate to M cells. For oral vaccines, the M-cell targeting can be realized by conjugation to NPs/MPs (such as CTS NPs) or antigens.108–109 Based on M-cell-homing peptide-fused antigens, oral administration of thiolated Eudragit microparticulate antigens results in strong secretory IgA, IgG, and cluster of differentiation (CD)4-positive and CD8-positive T-cell responses. Cholera toxin and the recombinant nontoxic pentameric B subunit of CT (rCTB) can target GM1 (monosialotetrahexosylganglioside).110 Multiple-mutated CT adjuvants can enhance immune responses,111 CTS-glucosaminan nanoassemblies with increased numbers of mannose molecules on the surface can target antigen-presenting machinery precisely.112 Other target ligands, such as UEA1 lectin, tetanus toxoid, or botulinum toxoid targeting M cells, cholera toxin and rCTB targeting GM1, and granulocyte-macrophage colony-stimulating factor ± interleukin-12 DNA, have been used in oral vaccination and promoted IgG and IgA responses.110,113–115

Most ligands for oral vaccination target small-intestinal immune cells. The large intestine-targeting properties of these vaccines are based mainly on the ability to bypass the small intestine.8,84 There are considerably fewer enzymes in the large intestine, and thus, studies on ligands that can target the large intestine and development of oral vaccines with prolonged duration in the large intestine are needed.

Modulation of the immune response
Some adjuvants for oral vaccines can modulate the immune response directly. Upon modification with PDDA, PEI, or CTS, IgG immune responses can be delayed, and mucosal immune responses induced, as compared with BSA or BSA/Freund’s complete adjuvant (FCA) adjuvants.82 Antigen modified with the cholera toxin can enhance
humoral, cellular, and mucosal immune responses.\textsuperscript{105} PLGA can stimulate durative immunization because it prolongs the time needed for antigen release.\textsuperscript{83,84,100,105} Toll-like receptors (TLRs) can induce the synergistic activation of T cells. CpG oligodeoxynucleotides (CpG-ODNs) can stimulate the functions of several types of immune cells and enhance transepithelial delivery, and can improve mucosal and systemic immunity.\textsuperscript{116} Class A ODN can induce the maturation of plasmacytoid dendritic cells (pDCs) and activation of natural killer cells through indirect cytokine signaling, while Class B ODN can strongly stimulate human B-cell and monocyte maturation, and stimulate the maturation of pDC.

**pH sensitivity of the intestinal environment**

The conditions within the GIT are complicated. In principle, oral vaccines must bypass gastric acidic fluids and enzymes in the small intestine before antigens are released to immune cells in the GIT. pH-sensitive biomaterials can serve as protective vehicles and an environment-stimulated release platform. Multicomponent systems can integrate different pH-sensitive characteristics for effective delivery of antigens to expected sites.

In our study, PMMMA was constructed on the surface of PLGA/antigen NPs (Figure 7). PMMMA was stable under the gastric and small-intestinal conditions of \textit{Tilapia} and could bypass the small intestine and release PLGA/antigens in the large intestine due to phase transition. Strong IgM responses were enhanced in \textit{Tilapia} against the surface immunogenic protein of GBS pathogenic bacteria.\textsuperscript{84}

**Potential of NPs/MPs for DNA-based oral vaccines**

With the knowledge of the genetic information of many pathogens, various DNA-, RNA-, peptide-, or recombinant protein-based vaccines can be constructed and utilized with NPs/MPs. Immunization with such vaccines (eg, plasmid DNA vaccine for the human immunodeficiency virus) instead of traditional attenuated viruses or protein scan leads to greater safety and lower costs.\textsuperscript{117}

Several aluminum-based compounds have been administered with DNA vaccines and PLGA particles via the oral route with their properties demonstrated. Aluminum phosphate can enhance immune responses upon mixing with DNA plasmids. Aluminum hydroxide cannot form complexes with DNA plasmids directly, but they can exert their effects on antigens after expression in situ.\textsuperscript{88} Furthermore, DNA plasmids encoded for the proteins required by foot and mouth disease (FMD) viral capsid assembly can be delivered via the oral route in mice and guinea pigs with calcium phosphate NPs, which induce immune responses and contribute to protection from FMD.\textsuperscript{118}

Figure 7 (Continued)
Polymers have been used in DNA vaccination. PLGA NPs can be carrier adjuvants for DNA and peptide vaccines. For instance, DNA plasmids encoded from the outer capsid proteins of rotavirus can elicit rotavirus-specific effects in mice with PLGA NPs. Also, Spf66 (the peptide from merozoite stage-specific protein fragments of *Plasmodium falciparum*) can be encapsulated in a mixture of two types of PLGA particles for vaccination against malaria.\(^{119-121}\) DNA vaccine for infectious bursal disease caused by the infectious bursal disease virus (IBDV) has been delivered via the oral route in chickens within cationic PLGA MPs.\(^{122}\) The design of the DNA vaccine was based on virus protein VP2 (which can induce neutralizing antibodies in chickens), and IBDV-VP2 DNA vaccine-loaded cationic PLGA MPs improved protection against the disease.\(^{122}\) Other cationic polymers, such as PEI and liposomes, harbor positive charges to absorb negatively charged DNA, generate complexes to protect DNA from rapid degradation by enzymes, and help DNA avoid lysosomes possibly through a “proton sponge” effect. The efficacy of these vaccines may not be satisfactory, but their reliability has garnered increasing attention upon them and their adjuvants.\(^{123,124}\)

**Prospects in oral vaccination**

Detrimental mutation, acute toxicity, and delayed side effects may prevent routine vaccination. For oral vaccination, the barrier is even greater because of the harsh environment in the GIT. Fortunately, adjuvants can help to overcome these difficulties.\(^{125}\)

It is not common for oral vaccines to be substitutes for injection-based vaccination, but achievements have been made utilizing NP/MP carriers with adjuvant activities for oral vaccination. NP/MP conveyors for oral vaccination are biodegradable and biocompatible; they can protect, elicit controlled release of antigens, and enhance immune responses, and their modification can strengthen targeting to specific sites. Considering the lack of natural carriers for oral vaccines, these characteristics can be the principles for selecting potential NP/MP materials, simplify the search strategy, and accelerate the discovery of novel oral vaccines. For example, one can decide if a pH-sensitive polymer-coated silica oral drug carrier is worthy of further investigation for oral vaccination, and then modify it for a “new mission”. The development of NP/MP conveyors with adjuvant properties can promote progress in the oral route of vaccination, which is beneficial to people’s health and the vaccination industry.

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References
1. Han S. Clinical vaccine development. Clin Exp Vaccine Res. 2015;4:46–53.
2. Silin DS, Lyubomska OV, Jirathitikal V, et al. Oral vaccination: where are we? Expert Opin Drug Del. 2007;4:323–340.
3. Mitragoti S, Lahann J. Physical approaches to biomaterial design. Nat Mater. 2009;8:15–23.
4. Shah PK, Chyu KY, Fredrikson GN, Nilsson J. Immunomodulation of atherosclerosis with a vaccine. Nat Clin Pract Cardiovasc Med. 2005;2:639–646.
5. Singh S, Yang G, Schluns KS, Anthony SM, Sastry KJ. Sublingual vaccines.
6. Hilleman MR. Vaccines in historic evolution and perspective: a narrative of vaccine discoveries. Vaccine. 2000;18:1436–1447.
7. Joseph A, Itskovitz-Coooper N, Samira S, et al. A new intranasal influenza vaccine based on a novel polyacylanionic lipid-carboxyl-spermine (CCS) I. Immunogenicity and efficacy studies in mice. Vaccine. 2006;24:3990–4006.
8. Zhu Q, Talton J, Zhang G, et al. Large intestine-targeted, nanoparticle-releasing oral vaccine to control genitorectal viral infection. J Control Release. 2015;210:48–57.
9. Kreuter J, Berg U, Liehl E, Soliva M, Speiser P. Influence of the particle size on the adjuvant effect of particulate polymeric adjuvants. Vaccine. 1986;4:125–129.
10. Anderson J, Smith BP, Ulrich JT. Vaccination of calves with a modified bacterin or oil-in-water emulsion containing alkali-detoxified Salmonella-typhimurium lipopolysaccharide. Am J Vet Res. 1991;52:596–601.
11. Igartua M, Hernandez RM, Esquisabel A, Gascón AR, Calvo MB, Pedraz JL. Enhanced immune response after subcutaneous and oral immunization with biodegradable PLGA microspheres. J Control Release. 1998;56:63–73.
12. Prieto MB, Delie F, Fattal E, et al. Characterization of V3 BRU peptide-loaded small PLGA microspheres prepared with a (W1/O/W2) emulsion solvent evaporation method. Int J Pharm. 1994;117:137–145.
13. Chaicumpa W, Paraio JR, New RC, et al. Immunogenicity of liposome-associated oral cholera vaccine prepared from combined Vibrio-cholerae antigens. Asian Pac J Allergy. 1990;8:87–94.
14. Nayak B, Panda AK, Ray P, Ray AR. Formulation, characterization and evaluation of rotavirus encapsulated PLA and PLGA particles for oral vaccination. J Microencapsul. 2009;26:154–165.
15. Primard C, Rochereau N, Luciani E, et al. Traffic of poly(lactic acid) microparticles into immune competent cells. Biomaterials. 2010;31:6069–6074.
16. Kim SY, Doh HJ, Ahn JS, et al. Induction of mucosal and systemic immune response by oral immunization with H. pylori lysates encapsulated in poly(D,L-lactide-co-glycolide) microcapsules. Vaccine. 1999;17:607–616.
17. Kim SY, Doh HJ, Jang MH, Ha YJ, Chung SI, Park HJ. Oral immunization with Helicobacter pylori-loaded poly(D, L-lactide-co-glycolide) nanoparticles. Helicobacter. 1999;4:33–39.
41. Kotlof KL, Sztein MB, Wasserman SS, Losonsky GA, DiLorenzo SC, Walker RI. Safety and immunogenicity of orally inactivated whole-cell Helicobacter pylori vaccine with adjuvant among volunteers with or without subclinical infection. *Infect Immun.* 2001;69:3581–3590.

42. Joshi VB, Geary SM, Salem AK. Biodegradable particles as vaccine antigen delivery systems for stimulating cellular immune responses. *Hum Vaccines Immunother.* 2013;9:2584–2590.

43. Gutierrez I, Hernandez RM, Igartua M, Gascon AR, Pedraz JL. Size dependent immune response after subcutaneous, oral and intranasal administration of BSA loaded nanospheres. *Vaccine.* 2002;21:67–77.

44. Samadi N, Abbadesa A, Di Stefano A, et al. The effect of lauryl capping group on protein release and degradation of poly(D,L-lactic-co-glycolic acid) particles. *J Control Release.* 2013;172:436–443.

45. Carcabo AM, Hernandez RM, Igartua M, Rosas JE, Patarroyo ME, Pedraz JL. Potent, long lasting systemic antibody levels and mixed Th1/Th2 immune response after nasal immunization with malaria antigen loaded PLGA microparticles. *Vaccine.* 2004;22:1423–1432.

46. Jiang T, Singh B, Li HS, et al. Targeted oral delivery of BmpB vaccine using porous PLGA microparticles coated with M cell homing peptide-coupled chitosan. *Biomaterials.* 2014;35:2363–2373.

47. Mundargi RC, Babu VR, Rangaswamy V, et al. Nano/micro technologies for delivering macromolecular therapeutics using poly(D,L-lactide-co-glycolide) and its derivatives. *J Control Release.* 2008;125:193–209.

48. Jain AK, Goyal AK, Mishra N, Vaidya B, Mangal S, Vyas SP. PEG-PLA-PEG block copolymeric nanoparticles for oral immunization against hepatitis B. *Int J Pharm.* 2010;387:253–262.

49. Artursson P, Edman P, Laakso T, Sjoholm I. Characterization of polyacryl starch microparticles as carriers for proteins and drugs. *J Pharm Sci.* 1984;73:1507–1513.

50. Wikingsson L, Sjoholm I. Polyacryl starch microparticles as adjuvant in oral immunisation, inducing mucosal and systemic immune responses in mice. *Vaccine.* 2002;20:3355–3363.

51. Heritage PL, Underdown BJ, Brook MA, McDermott MR. Oral administration of polymer-grafted starch microparticles activates gut-associated lymphocytes and primes mice for a subsequent systemic antigen challenge. *Vaccine.* 1998;16:2001–2007.

52. Alcock HR, Kwon S. An ionically cross-linkable polyphosphazene-[PCPP] is a potent immunoadjuvant for an influenza vaccine formulated as a cationic liposome. *Biomacromolecules.* 1998;1:2010–2007.

53. Payne LG, Jenkins SA, Woods AL, et al. Poly[d(carboxylatophenoxyl)phosphazene] (PCPP) is a potent immunoadjuvant for an influenza vaccine. *Vaccine.* 1998;16:92–98.

54. Andrianov AK, Marin A, Chen J. Synthesis, properties, and biological activity of poly[d(sodium carboxylatoethylenophenoxyl)phosphazene] and its hydrogels and membranes. *Macromolecules.* 1989;22:75–79.

55. Payne LG, Jenkins SA, Woods AL, et al. Poly[d(carboxylatophenoxyl)phosphazene] and its hydrogels and membranes. *Macromolecules.* 2006;7:394–399.

56. Eng NF, Garlapati S, Gerdt V, Babiuk LA, Mutwiri GK. PCEP enhances IgA mucosal immune responses in mice following different immunization routes with influenza virus antigens. *J Immune Based Ther Vaccines.* 2010;8:4.

57. Hariyadi DM, Ma Y, Wang Y, et al. The potential for production of freeze-dried oral vaccines using alginate hydrogel microspheres as protein carriers. *J Drug Deliv Sci Tec.* 2014;24:178–184.

58. Bowersock TL, Shahaly WSW, Blevins WE, et al. Poly(methacrylic acid) hydrogels for rumen bypass and the delivery of oral vaccines to ruminants. *Acs Sym Ser.* 1994;545:214–220.

59. Wu QI, Gong CY, Shi S, et al. Mannan loaded biodegradable and injectable thermosensitive PCL-PEG-PCL hydrogel for vaccine delivery. *Soft Mater.* 2012;10:472–486.

60. Jung T, Kamn W, Breitenbach A, Hungerer KD, Hundi E, Kissel T. Tetanus toxoid loaded nanoparticles from sulfobutylated poly(vinyl alcohol)-graft-poly(lactide-co-glycolide): evaluation of antibody response after oral and nasal application in mice. *Pharm Res.* 2001;18:352–360.

61. Beg S, Samad A, Nazish I, et al. Colloidal drug delivery systems in vaccine delivery. *Curr Drug Targets.* 2013;14:123–137.

62. Tomalia DA. Dendrimer research. *Science.* 1991;252:1231–1232.
98. Zhang Q, Zhao Q, Zhang Y, et al. Investigation of 3-D ordered mesoporous silica SBA-15: a new effective adjuvant to induce antibody response. Small. 2006;2:254–256.

99. Ulmer JB, DeWitt CM, Chastain M, et al. Enhancement of DNA vaccine potency using conventional aluminum adjuvants. Vaccine. 1999;18:18–28.

100. Zhang Q, Zhao Q, Zhang Y, et al. Investigation of 3-D ordered materials with a high adsorption capacity for BSA and their potential application as an oral vaccine adjuvant. J Colloid Interface Sci. 2014;434:113–121.

101. Ogue S, Takahashi Y, Onishi H, Machida Y. Preparation of double hydrophobic silica in modifying their properties. Int J Pharm. 2014;462:269–274.

102. Liu P, Sun Y, Wang Q, Sun Y, Li H, Duan Y. Intracellular trafficking and cellular uptake mechanism of mPEG-PLGA-PLL and mPEG-PLGA-PLL-Gal nanoparticles for targeted delivery to hepatomas. Biomaterials. 2014;35:760–770.

103. Zhu Q, Thomson CW, Roseenthal KL, McDermott MR, Collins SM, Gauldie J. Immunization with adenovirus at the large intestinal mucosa as an effective vaccination strategy against sexually transmitted viral infection. Mucosal Immunol. 2008;1:78–88.
123. Chu DS, Schelling JG, Shi J, Convertine AJ, Stayton PS, Pun SH. Application of living free radical polymerization for nucleic acid delivery. *Acc Chem Res.* 2012;45:1089–1099.
124. Ulmer JB, Mason PW, Geall A, et al. RNA-based vaccines. *Vaccine.* 2012;30:4414–4418.
125. Zhu J, Wang Y, Huang H, et al. Upregulation of KPNβ1 in gastric cancer cell promotes tumor cell proliferation and predicts poor prognosis. *Tumour Biol.* 2016;37:661–672.