

Systemic and Mucosal Immune Responses to Sublingual or Intramuscular Human Papilloma Virus Antigens in Healthy Female Volunteers

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Abstract

The sublingual route has been proposed as a needle-free option to induce systemic and mucosal immune protection against viral infections. In a translational study of systemic and mucosal humoral immune responses to sublingual or systemically administered viral antigens, eighteen healthy female volunteers aged 19–31 years received three immunizations with a quadravalent Human Papilloma Virus vaccine at 0, 4 and 16 weeks as sublingual drops (SL, n = 12) or intramuscular injection (IM, n = 6). IM antigen delivery induced or boosted HPV-specific serum IgG and pseudovirus-neutralizing antibodies, HPV-specific cervical and vaginal IgG, and elicited circulating IgG and IgA antibody secreting cells. SL antigens induced ~38-fold lower serum and ~2-fold lower cervical/vaginal IgG than IM delivery, and induced or boosted serum virus neutralizing antibody in only 3/12 subjects. Neither route reproducibly induced HPV-specific mucosal IgA. Alternative delivery systems and adjuvants will be required to enhance and evaluate immune responses following sublingual immunization in humans.

Trial Registration: ClinicalTrials.gov NCT00949572

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Introduction

The mucosal surface is the most common route of infection for a wide range of viral diseases and therefore inducing both mucosal and systemic immunity is a key objective of modern vaccines. The rich infiltration into the sublingual mucosa of antigen-presenting dendritic cells makes it an attractive route of immunization that avoids needles and targets the mucosal immune system [1]. Virus-Like Particles (VLP) comprising the Human Papilloma Virus (HPV) L1 major capsid protein, as well as antigens from other viruses, delivered via the sublingual route have been shown in mice to be highly immunogenic and protective against subsequent viral challenge [2,3,4,5,6]. These observations also support the idea of a "Common Mucosal Immune System" and a link between the genital tract and the systemic immune system [2,3,5,6]. However, while these studies have employed antigen administration as simple sublingual liquid drops, there are characteristics of murine models which need to be considered: the murine sublingual surface is extremely rich in readily accessible dendritic cells [1]; mice are routinely anaesthetized for sublingual immunization, with possible anticholinergic effect on reducing saliva flow and antigen clearance; cholera toxin and related mucosal adjuvants have been employed to enhance responses, which may not be suitable for use in humans [7]. Sublingual immunization with non-toxic cholera toxin B subunit also induces and modulates local and disseminated responses, but this antigen is almost unique in its mucosal immunostimulating

and adjuvant properties [8]. Sublingual delivery has been used for many decades in humans in desensitizing regimes involving prolonged, frequent delivery of high doses of allergens [9]. However, it is only recently that this route has been considered for delivery of prophylactic vaccine antigens, which will require far fewer doses at lower dose levels [1,10]. We report here a preliminary human translational study to determine the character, dissemination and magnitude of systemic and mucosal immune responses to more representative antigens from a vaccine already in widespread use when administered sublingually or intramuscularly to healthy female volunteers. These results are contrasted with data from broadly similar murine studies in which HPV VLPs have been delivered sublingually as simple drops and found to be highly effective in eliciting immune response and protecting against genital HPV infection [3].

Methods

The protocol for this trial and supporting CONSORT checklist are available as supporting information; see Checklist S1 and Protocol S1.

Ethics Statement

Ethical Approval was obtained from the UK National Research Ethics Service, Wandsworth Research Ethics Committee reference

09/80803/77. Written informed consent was obtained from all participants after the nature and possible consequences of the study was explained. Clarification of the legal status of the study was obtained by submitting the protocol to the UK Medicines and Healthcare products Regulation Agency (MHRA) which confirmed it as a “Characterization Study” and not a Clinical Trial of an Investigational Medicinal Product (non-CTIMP/NIMP). Although not a clinical trial, we registered this study protocol on ClinicalTrials.gov (NCT00949572) prior to subject recruitment.

Objectives

We sought to characterize, and contrast, the nature and dissemination of the immune response to sublingual or intramuscular deposition of meaningful viral vaccine antigens in humans, and to compare this with published murine studies [2,3,4,5,6]. The protocol defined no primary or secondary endpoints as this was not a clinical trial. The goal was to describe the immune response following immunization, and the study exploratory endpoint was immune response measured as several immunologic factors and assessed as change in each of these factors from pre to post immunization. The following variables were assessed before and after immunizations: (i) frequency of PBMCs secreting IgG or IgA antibodies to HPV16 L1 VLPs and whole vaccine; (ii) concentration in the serum, cervical secretions and vaginal secretions of IgG to HPV16, HPV6 and HPV18 L1 VLPs; (iii) concentration in the cervical secretions and vaginal secretions of IgA to HPV16, HPV6 and HPV18 L1 VLPs; (iv) titer in the serum, cervical secretions and vaginal secretions of neutralizing antibody to HPV16 or Bovine Papillomavirus control.

Participants

The target recruitment defined by the protocol was 18 healthy female volunteers (in two groups: SL n = 12, IM, n = 6) aged 25–35. Subjects were all recruited at one site, St George’s - University of London, London. As this was a hypothesis-generating study no formal power calculation for sample size was performed. Inclusion criteria included: provide written informed consent; in good health determined by medical history, physical examination, hematology; available for the duration of the study; if of childbearing potential, must have a negative pregnancy test before each immunization; have not donated blood in previous 3 months; eligible for free medical treatment in the UK. Exclusion criteria included: already received HPV vaccine; recent or concurrent participation in another clinical research study; recent or planned use of any investigational or non-registered product; pregnant or breast-feeding; known or suspected ongoing cervico-vaginal disease, malignancy or abnormality; positive results for Human Immunodeficiency Virus or Hepatitis B/C infection; abnormality in hematology; acute or chronic pulmonary, cardiovascular, hepatic, hematologic, renal, blood or neurological disorders, immune dysfunction, autoimmune diseases, diabetes or malignancy; recent immunosuppressive therapy; medications via vaginal route; tongue or frenulum piercings or oral jewelry; recent receipt of blood products or immunoglobulin.

Description of Procedures or Investigations undertaken

Immunization. We purchased the licensed quadravalent Human Papilloma Virus (HPV) vaccine Gardasil® (Sanofi Pasteur), which contains L1-based virus-like particles (VLPs) representing four HPV types: 20 µg each of HPV types 6, 18; 40 µg each of HPV types 16 and 11 per 0.5 mL dose. VLPs are produced in yeast cells (*Saccharomyces cerevisiae* CANADE 3C-5 Strain 1895) by recombinant DNA technology and adsorbed on amorphous aluminum hydroxyphosphate sulphate adjuvant (225

micrograms aluminum per dose). All subjects received three immunizations with 0.5 mL (one standard dose) of vaccine at weeks 0, 4 and 16 (table 1) which is a recommended schedule within the flexibility of the usual 0, 1, 6 months dosing schedule for parenteral immunization. IM immunizations were given into the deltoid muscle. For SL immunization, subjects fasted (except water) for 1 hour prior to challenge, then sat in an upright position, rinsed the mouth with water and expectorated. Absorbent pads (Molnlycke ‘Dry Tips’ small) were applied over parotid duct openings bilaterally to absorb parotid saliva flow. The tongue was raised and the sublingual area gently dried by brief application of a cotton swab without inducing saliva flow from submandibular and sublingual glands. The 0.5 mL contents of a Gardasil® syringe were dispensed drop-wise to the area behind the sublingual fold bilaterally. The tongue was held in gentle opposition to the floor of the mouth for 15 minutes without swallowing, then the cotton pads removed and the subject allowed to swallow. Subjects were fasted completely for 30 minutes under observation and then requested to fast (including fluids) for a further 60 minutes after leaving the clinical site.

Sample collection. Table 1 shows the schedule of immunizations and sample collection. A blood sample was taken before the first immunization, at the time of the first immunization, and then on weeks 1, 4, 5, 8, 16, 17 and 20 after first immunization. Cervical and vaginal wick samples were collected at the time of the first immunization, and then on weeks 4, 16 and 20. Schedules were initiated to accommodate the subjects’ menstrual cycles and a ± 2 day window period was acceptable for all visits, except 1, 5 and 17 which had a ± 1 day window. A protocol amendment was approved during the study to collect cervical and vaginal samples at week 8 for 3/6 subjects in IM group and 6/12 in SL group. To collect mucosal secretions a Weck-Cel surgical spear was placed either in the cervical *os* or against the vaginal wall for 2 minutes, then secretions eluted as described previously [11]. Briefly, spearheads were snipped into the top chamber of a Spin-X tube (Corning) containing 300 µL sterile filtered extraction buffer (250 mM NaCl, 16 protease inhibitor cocktail set 1 (Calbiochem) in phosphate buffered saline (PBS)) and centrifuged at 4°C for 15 minutes at 13,000 g. A repeat extraction was performed by adding additional extraction buffer to the top chamber, and then 8 µL heat inactivated fetal calf serum added to pooled secretions from each sample site, prior to separation into 200 µL aliquots and freezing at -80°C before batch analysis by ELISA as described below.

HPV L1 antigens used in ELISA and ELISPOT assays. HPV11 L1 VLPs were not available and no responses to HPV11 were measured. HPV 6 and 18 L1 VLPs were a kind gift of Shantha Biotechnics Ltd, India. HPV16 L1 VLPs were generated using the Bac-to-Bac® Baculovirus Expression System (Invitrogen) wherein the recombinant bacmid DNA contained an HPV16 L1 gene with a 100% amino acid sequence identity to

Table 1. Schedule of immunizations and sample collection.

	Week							
	0	1	4	5	8	16	17	20
Immunization	x		x			x		
PBMCs sample	x	x	x	x	x	x	x	x
Serum sample	x	x	x	x	x	x	x	x
Cervical & vaginal secretions sample	x		x		x	x		x

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GenBank accession numbers DQ469930 and EU118173. Recombinant HPV16-expressing baculovirus stocks were used to infect Sf21 insect cells (Invitrogen) for 72 hours at 27°C before lysis (IGEPAL® CA-630; Sigma-Aldrich) at room temperature in the presence of protease inhibitors (Complete; Roche). The cell lysate was then subjected to iodixanol gradient fractionation and gradient fractions were collected by bottom puncture and stored at -80°C. The L1 concentration and purity were visualized by SDS-PAGE stained with Colloidal Blue (Invitrogen) and analyzed using ImageJ software (U. S. National Institutes of Health, <http://imagej.nih.gov/ij>). VLP formation was confirmed by electron microscopic analysis of negatively staining particles (Phosphotungstic Acid; Sigma-Aldrich) adsorbed on copper grids coated with formvar (Sigma-Aldrich) and carbon.

Frequency of circulating L1-specific IgG and IgA spontaneously antibody secreting cells (ASCs). The frequency of L1-specific spontaneously antibody-secreting plasmablasts was enumerated in PBMCs separated from heparinized whole blood by Ficol gradient centrifugation in an ELISPOT assay as described previously [12]. PVDF-backed 96 well plates (MAHA S45, Millipore) were coated in advance, and divided into three parts: coating buffer only in wells without any antigen as a background and nonspecific reaction control; L1 HPV16 VLPs; or Gardasil® (as other L1 antigens were not available at this time). The cell density of each sample was adjusted to 5×10^5 , 2.5×10^5 and 1.25×10^5 /well using AIMV medium (Invitrogen, UK) containing penicillin-streptomycin. Each cell concentration was added as duplicate wells on the three antigens or uncoated parts of the plate after blocking. A 2 mg/mL PHA positive control was also added on each plate. After overnight incubation, the specific antibody-secreting cells were recognized by the goat anti-human IgG or IgA conjugated with alkaline phosphatase, and counted and analyzed by the AID EliSpot Reader System. The final results were standardized as positive cell number/ 1×10^5 PBMCs plated.

L1-specific IgG & IgA in serum, cervical & vaginal secretion quantified by ELISA. The concentration of serum, cervical or vaginal L1-specific IgG and IgA was measured by indirect ELISA as described previously [11] using purified L1 HPV16, 6 and 18 VLPs in carbonate-bicarbonate buffer (0.05M pH 9.6) coated individually on MaxiSorp plates (Nunc). One standard curve made by a positive serum with known specific antibody concentration, and positive and negative controls were set-up on each plate, with two blank wells on each plate to monitor background. Serum, cervical and vaginal samples were diluted 1/200 or 1/4 with 0.05% PBS-T20, respectively, and any sample with OD value above the upper limit of the standard curve was further diluted and re-assayed. The specific IgG or IgA was recognized by goat anti-human IgG or IgA conjugated with peroxidase. The concentrations of specific antibodies were measured by an Emax MAXLine Microplate reader at 650 nm after the TMB liquid substrate developed. The raw OD data was analyzed using SoftMax® software.

Functional antibody in serum and cervical and vaginal secretions measured by in vitro pseudovirus neutralization assay. The HPV16 pseudovirus neutralization assay [13] was carried out as previously described [14] and included Bovine Papillomavirus (BPV) as a control for non-specific antibody effects. As a control, the WHO International Standard for HPV16 antibodies, IS16 (code: 05/134; 10 IU/mL; National Institute for Biological Standards and Control, UK; [15]) demonstrated type-specific neutralization of HPV16 at levels consistent with natural infection (median titer 138 [inter-quartile range 115–148]; $n = 3$).

Statistical methods

As this was not a clinical trial, no randomization was performed and no safety data (adverse events) were solicited, no primary or secondary endpoints were specified. Subjects were allocated to one of two sequential cohorts: “IM” who received all immunizations via the intramuscular route ($n = 6$); and “SL” who received all immunizations as sublingual drops ($n = 12$). Subjects were not randomized as we wished to develop and evaluate B cell assays carried out on fresh blood samples by recruiting the first subjects into the intramuscular delivery group (and for whom measurable antibody secreting cells (ASCs) were likely to be seen). As the immunization routes could not be blinded the study was not blinded. There were no protocol deviations. As this hypothesis-generating study was not powered to detect significant differences between groups or between time points, no statistical testing was performed and descriptive statistics only are presented.

Results

Subjects enrolled

Eighteen female subjects aged 19–31 years (IM group mean 24.2, median 25; SL group mean 26.3, median 27.5) were enrolled and completed the protocol. There were no protocol deviations (figure 1).

Circulating Anti-L1 HPV16 B cell responses measured by ELISPOT

HPV16 L1 VLPs and whole vaccine (VLP HPV 6, 11, 16 and 18) were used as a coating antigens. From previous studies using oral, nasal or intramuscular immunization [12,16], a transient increase in the frequency of cells spontaneously secreting anti-L1 IgG and IgA was expected, peaking around 7 days after each immunization (on weeks 0, 4 and 16) and then falling back to baseline, reflecting the generation and maintenance kinetics of plasmablasts. An increase in IgG and IgA antibody secreting cell (ASC) frequencies was seen after IM immunizations (figure 2), but not after SL immunization.

Anti-L1 HPV6, HPV16 and HPV18 serum & cervico-vaginal IgG & IgA responses measured by ELISA

The kinetics of the anti-HPV L1 VLP antibody response in serum and in cervico-vaginal secretions was measured before and at various time points after each immunization using an antigen-specific antibody binding ELISA, with purified L1 HPV16, 6 and 18 VLPs as coating antigens. As expected, IM immunization induced an increase in serum anti-HPV6, HPV18 and HPV16 L1 VLP IgG from baseline (figure 3). IM immunization also induced an increase in cervical anti-HPV6 and HPV16 L1 VLP IgG, and to a lesser extent in anti-HPV18 L1 VLP IgG. An increase in vaginal anti-HPV6 and HPV16 L1 VLP IgG was seen, but not anti-HPV18 L1 VLP IgG. SL immunization induced an increase in anti-HPV16 L1 VLP IgG in serum, cervical secretions and vaginal secretions, and a slight increase in cervical anti-HPV18 L1 VLP IgG. However, the serum anti-HPV16 L1 VLP IgG at week 20 after IM immunization was $219 \mu\text{g/mL}$ (SEM ± 57.3 , a 38.9-fold rise from week 0), compared with $5.73 \mu\text{g/mL}$ (± 2.9 , 3.4-fold rise) after SL immunization. In contrast, relative levels of specific IgG in mucosal secretions were not as dissimilar as in serum: mean 45.5 ng/mL (± 10.8 , 2.2-fold rise) in cervical secretions after SL immunization, 76.7 ng/mL (± 19.6 , 9.8-fold rise) after IM immunization. Similarly, in vaginal secretions the values were 56.1 ng/mL (± 13.8 , 3.2-fold rise) and 115.5 ng/mL (± 34.9 , 10.9-

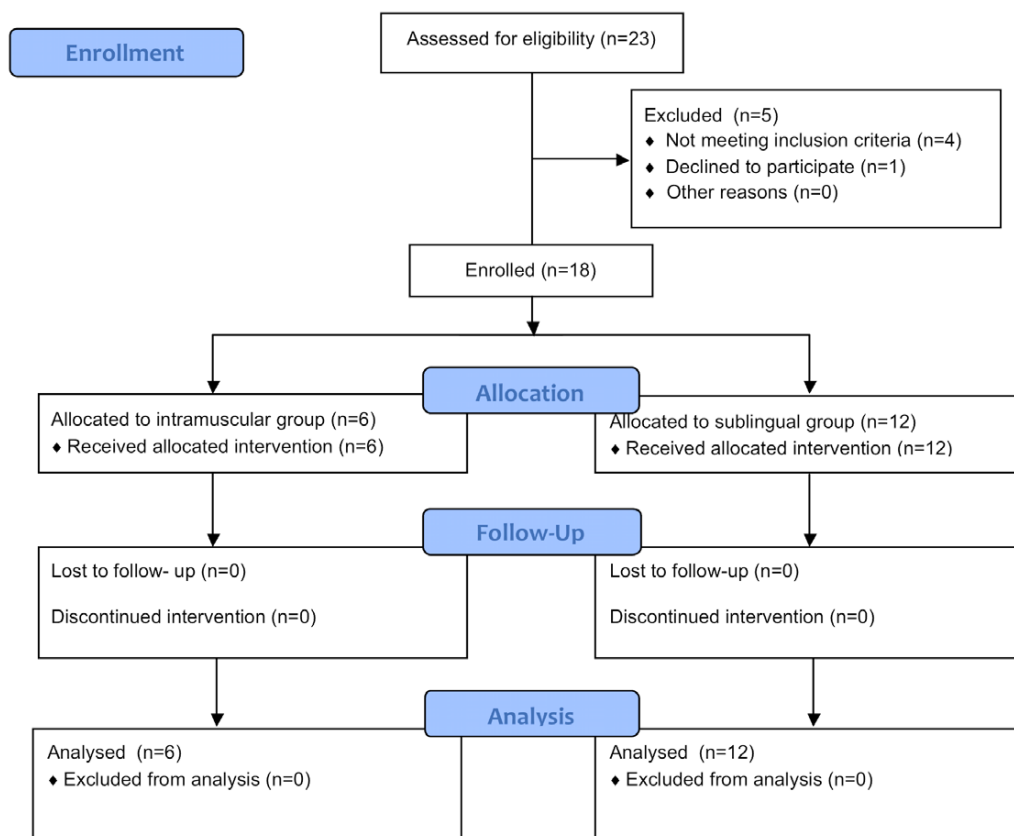


Figure 1. CONSORT diagram.

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fold rise) for SL and IM immunizations, respectively. IgA responses in secretions were variable at all time points (figure 4).

Neutralization of HPV16 by serum and cervical and vaginal secretions

The detection of antibodies capable of neutralizing HPV16 pseudoviruses was examined using serum and cervical and vaginal samples taken at week 0 and 20 (4 weeks post last immunization) for all subjects (table 1). Some intermediate time points were also evaluated for selected subjects (data not shown). The control Bovine Papillomavirus (BPV) was used as a control for non-specific antibody reactivity. All subjects had undetectable virus neutralization titers in mucosal secretions at week 0 (table 2), but 2/6 and 1/12 subjects in the IM and SL groups, respectively, had detectable serum virus neutralizing activity at week 0, which is in line with the ~12% prevalence estimates for previous HPV16 infection expected in this population [17,18]. IM immunization induced or boosted serum neutralizing antibodies in all subjects at week 20, and very low level neutralizing titers also appeared in mucosal secretions at week 20 after IM immunization in the 3/6 subjects with the highest serum neutralizing titers. Sublingual immunization did not induce any neutralizing activity in mucosal secretions. However, sublingual immunization did boost pre-existing serum neutralizing activity in one subject (013) to a level similar to that seen in subjects without pre-existing neutralizing activity who received IM immunization and induced weak serum neutralizing titers in two others (015, 023). This suggests that, unlike IM immunization, while SL immunization may not be very

effective at priming the immune response it may be able to boost pre-existing immunity.

Discussion

Parenteral immunization with vaccines containing HPV L1 VLPs and formulated using alum and/or TLR agonist adjuvants is highly effective at inducing both serum and mucosal antibodies, and conferring long-lasting protection against HPV infection by the homologous or related HPV genotypes [19,20,21,22]. Several murine models have shown that simple drops placed under the tongue can induce functional antibody and T cell responses to viruses such as Herpes simplex virus (HSV), influenza, Human Immunodeficiency Virus (HIV) and HPV, and protection against genital challenge with HSV and HPV [2,3,4,5,6]. The possibility to develop a needle-free sublingual human vaccine, specifically targeting the induction of mucosal immunity and applicable to a wide range of genital viral infections is compelling [1]. However, these murine models often incorporate features that are not compatible with real-world human vaccine strategies, such as the use of anti-cholinergic anesthetics that may block saliva flow, and mucosal adjuvants based on cholera toxin-related proteins that are unsafe in humans when given nasally [7]. Sublingual desensitization regimes use frequent, prolonged high doses of allergens [9]. Non-toxic cholera toxin B subunit antigens induce disseminated antibody responses after sublingual immunization, but this molecule has intrinsic mucosal immunogenicity and adjuvanticity not seen in the majority of protein antigens [8], and like allergens may therefore not be representative of real world sublingual human

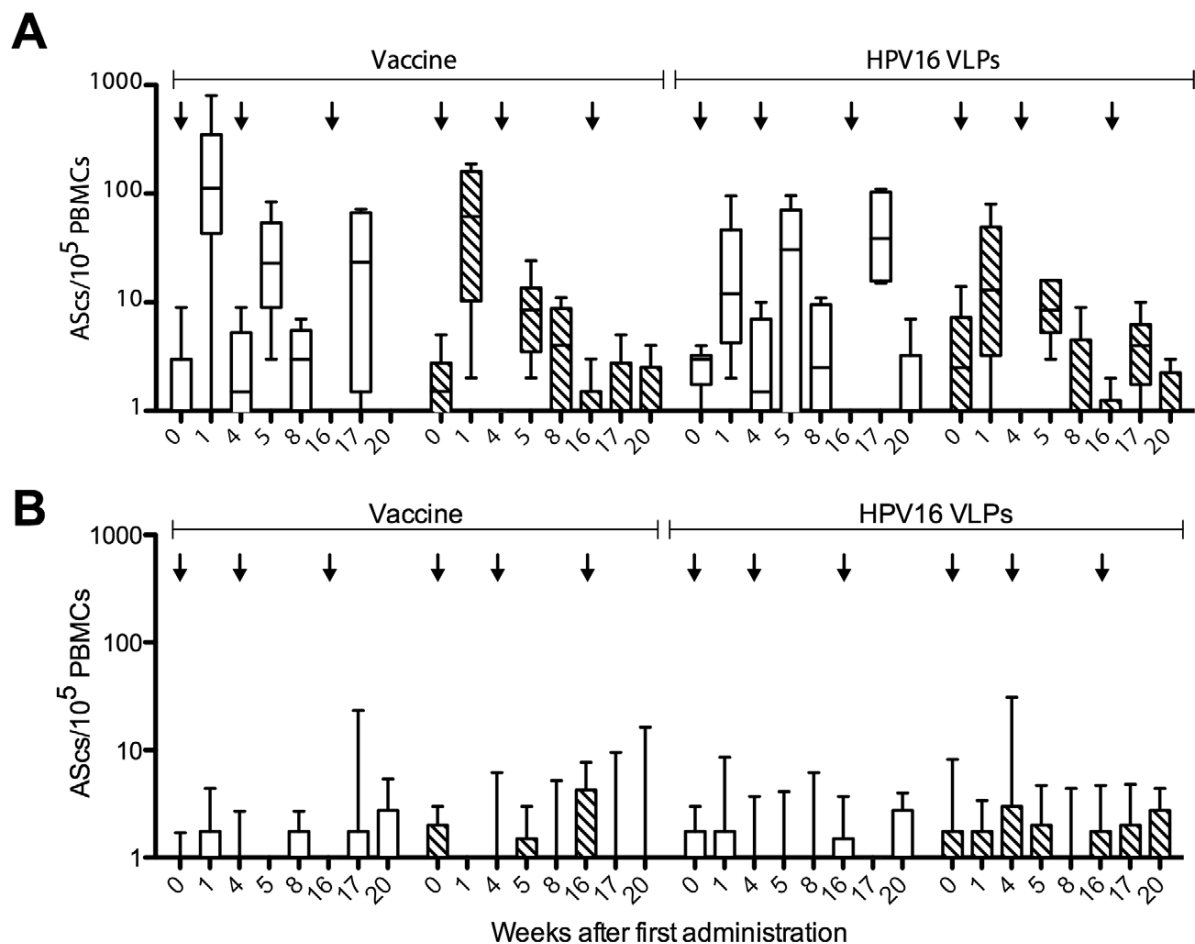


Figure 2. Circulating antibody secreting cell responses. The Y axis values indicate the group median frequency of antibody secreting cells (ASCs) per 10⁵ PBMCs plated, secreting IgG (white bars) or IgA (hatched bars) against Gardasil vaccine or L1 HPV16. Panel A: subjects immunized intramuscularly. Panel B: subjects immunized sublingually. Arrows indicate immunizations. Box: 25th to 75th percentiles, whiskers: 10 to 90 percentiles.

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vaccine antigens. We carried out a preliminary translational study to characterize immune responses to the sublingual or parenteral administration of a viral antigen that is already in widespread use as a human vaccine [23], and therefore representative of the human application of sublingual immunization. As this was the first use of this vaccine sublingually in humans we followed a 0, 1, 4 month prime-boost schedule which is an acceptable schedule for IM immunization using Gardasil®.

We observed that sublingual immunization generally induced a similar pattern of immune responses to intramuscular, but at much lower magnitudes. An increase in serum anti-HPV16 L1 VLP IgG was detected after both IM and SL immunization (figure 3). However, the serum anti-HPV16 L1 VLP IgG at week 20 after IM immunization was ~38 times higher than after SL immunization. Similarly, while IM immunization was able to both prime or boost serum virus neutralizing activity in all subjects, sublingual immunization could only boost serum neutralizing activity in a subject with pre-existing activity at week 0, and induce low levels of serum neutralizing antibody in two other subjects with undetectable neutralizing activity at day 0. This suggests that with optimization, the sublingual route may have a role in boosting pre-existing immunity induced by another route, and is capable of inducing functional antibody.

One of the potential translational advantages proposed for mucosal immunization is that it appears to specifically induce mucosal immunity [10]. However, while IM immunization was capable of inducing measurable virus neutralizing activity in cervical and/or vaginal secretions in 3/6 subjects (concomitant with high serum neutralizing titers suggesting transudation of serum IgG), no mucosal virus neutralizing activity was induced by SL immunization in any subject. Similarly, while IM immunization induced increases in mucosal anti-HPV6 and HPV16 L1 VLP IgG, sublingual immunization only induced an increase in mucosal anti-HPV16 L1 VLP IgG. However, it is intriguing that relative levels of specific IgG in mucosal secretions were not as dissimilar as in serum with only ~1.7-fold higher levels in cervical secretions after IM immunization compared with SL immunization. Similarly, in vaginal secretions IM immunization gave ~2-fold higher levels. This can be interpreted as SL immunization preferentially favors mucosal over systemic responses, or that neither route is very efficient at inducing mucosal immunity. Mucosal IgA responses were infrequent, low level, transient and extremely variable after either SL or IM immunization, despite HPV-specific IgA ASC responses after IM immunization. The kinetics of the ASC response to sublingual immunization is not well defined in humans and it is possible that we missed the

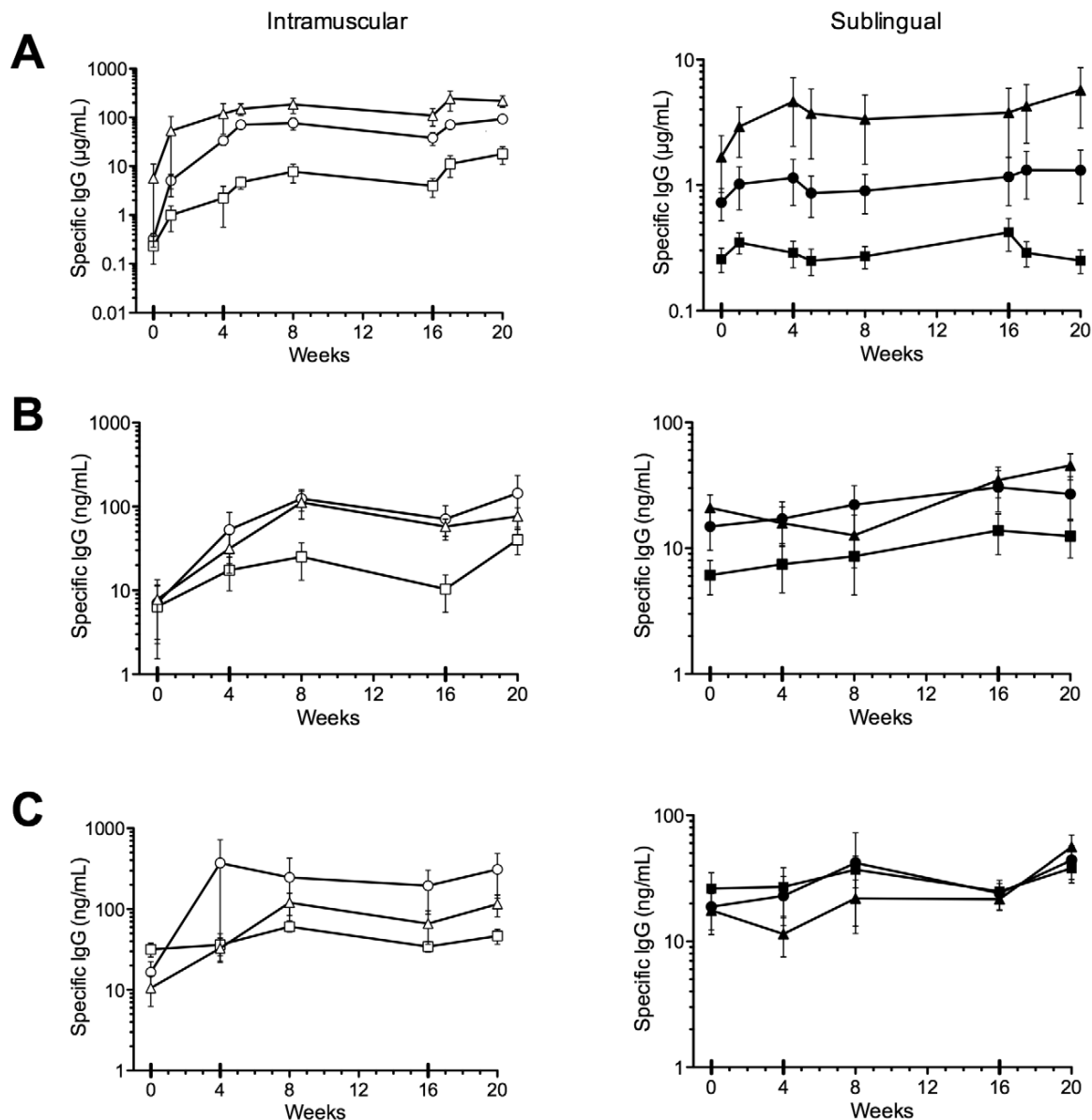


Figure 3. Serum, cervical and vaginal IgG responses. The Y axis values (note different scales) indicate group mean anti-L1 HPV6 (circles), HPV16 (triangles) and HPV18 (squares) IgG concentration in serum (panel A), cervical secretions (panel B) or vaginal secretions (panel C), for subjects immunized intramuscularly (left, open symbols), or sublingually (right, closed symbols). Error bars SEM.
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response to SL immunization. Additionally, although ASC responses to the first IM immunization appeared higher than subsequent immunizations, previous studies [16] have shown a shift to a slightly earlier timing of the peak response to booster immunizations (around day 5), which may explain the apparent fall in frequency measured 7 days after the booster immunizations.

Due to the considerable volume of saliva produced despite covering the parotid ducts, it is highly likely that some sublingually administered VLPs would have been removed from the mucosal surface within minutes of application. To be effective, extensive optimization of sublingual delivery will be required to improve penetration of the sublingual mucosa of humans, perhaps by making use of mucoadhesives or other delivery systems designed to resist salivary degradation [24]. In addition, the size of VLPs and

VLP-alum aggregates may have restricted access across the mucosal barrier. Manipulation of the particle size may therefore optimize responses. However, despite these caveats, some HPV-specific immunity was generated *de novo* in 2/12 subjects whose day 0 serum neutralizing antibody titer was below the cut-off in our assay, and boosted the day 0 titer in another subject. These observations are encouraging for potential future vaccine strategies based upon sublingual delivery. It is also possible that the bivalent vaccine, Cervarix[®], which makes use of a TLR-4 agonist in the vaccine formulation and has been shown to induce higher levels of HPV-specific antibodies when administered parenterally, may have induced significantly higher antibody levels [25], especially if combined with mucosal delivery systems that enhance sublingual contact times [24]. We allowed subjects to swallow retained saliva,

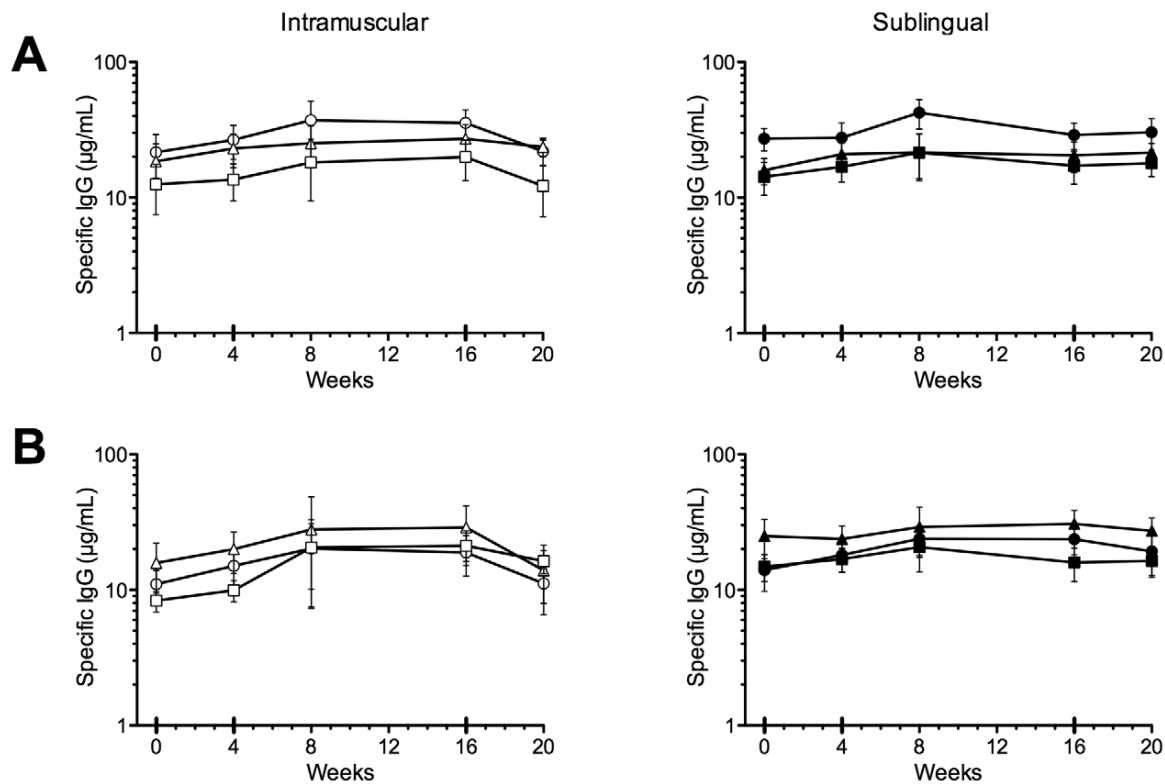


Figure 4. Cervical and vaginal IgA responses. The Y axis values indicate group mean anti-L1 HPV6 (circles), HPV16 (triangles) and HPV18 (squares) IgG concentration in cervical secretions (panel A), or vaginal secretions (panel B), for subjects immunized intramuscularly (left, open symbols), or sublingually (right, closed symbols). Error bars SEM.
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Table 2. HPV16 pseudovirus neutralization by serum and genital antibodies.

		HPV16 Neutralization Titer using in indicated sample					
		Cervix		Vagina		Serum	
Immunization Route	Subject ID	Week 0	Week 20	Week 0	Week 20	Week 0	Week 20
Intramuscular	001	- ^a	-	-	-	-	4,367
	004	-	-	-	26	-	34,255
	005	-	-	-	-	-	4,055
	017	-	-	-	-	-	3,856
	018	-	51	-	39	189	61,743
	020	-	42	-	53	5,086	44,494
Sublingual	006	-	-	-	-	-	-
	007	-	-	-	-	-	-
	008	-	-	-	-	-	-
	009	-	-	-	-	-	-
	010	-	-	-	-	-	-
	011	-	-	-	-	-	-
	013	-	-	-	-	213	2,249
	014	-	-	-	-	-	-
	015	-	-	-	-	-	632
	019	-	-	-	-	-	-
	021	-	-	-	-	-	-
	023	-	-	-	-	-	92

^a indicates reciprocal neutralization titers <20. All samples tested negative for neutralizing antibodies against the control Bovine Papillomavirus, BPV.
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and theoretically this may have allowed some immune responses to be induced in the small intestine. However, it is unlikely that 20–40 µg of VLPs would survive gastric acid, and indeed even the uniquely immunogenic mucosal antigen and adjuvant cholera toxin or its B subunit given at doses of 1–5 mg requires buffering with bicarbonate solution to retain immunogenicity via the oral route in humans [26,27].

In conclusion, this preliminary translational human study indicates that, in marked contrast to murine studies, SL delivery of a representative virus vaccine antigen formulated with alum is only modestly immunogenic in humans. This route can, however, induce low level serum and mucosal antibodies, and functional serum neutralizing antibody. The observation that SL immunization could boost pre-existing serum neutralizing activity also points to the possible use of IM prime/SL boost schedules. For this approach to be advanced, the next steps require significant optimization of the SL delivery system for human use, and the investigation of optimal SL-IM prime-boost schedules. Once this is achieved the benefits of sublingual delivery on the character, magnitude and dissemination of responses should be compared in clinical trials.

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Supporting Information

Protocol S1 Trial Protocol.
(PDF)

Checklist S1 CONSORT Checklist.
(DOC)

Acknowledgments

We are grateful for the kind gift of HPV 6, 11 and 18 VLPs from Shantha Biotech, Hyderabad, India. We acknowledge the critical review of the protocol and advice from Raphaelle El-Habib, Sanofi-Pasteur. We are indebted to John T. Schiller and Christopher B. Buck (National Cancer Institute, Bethesda, MD) for access to the pseudo virus clones used in this study.

Author Contributions

Conceived and designed the experiments: SB DJML. Performed the experiments: ZH SLB RG CO. Analyzed the data: ZH SLB SB DJML. Contributed reagents/materials/analysis tools: SLB SB. Wrote the paper: ZH SLB RG SB CO DJML.



CONSORT 2010 checklist of information to include when reporting a randomised trial*

Section/Topic	Item No	Checklist item	Reported in section
Title and abstract			
	1a	Identification as a randomised trial in the title	NA
	1b	Structured summary of trial design, methods, results, and conclusions (for specific guidance see CONSORT for abstracts)	NA
Introduction			
Background and objectives	2a	Scientific background and explanation of rationale	Introduction
	2b	Specific objectives or hypotheses	Introduction
Methods			
Trial design	3a	Description of trial design (such as parallel, factorial) including allocation ratio	Methods
	3b	Important changes to methods after trial commencement (such as eligibility criteria), with reasons	NA
Participants	4a	Eligibility criteria for participants	Methods
	4b	Settings and locations where the data were collected	Methods
Interventions	5	The interventions for each group with sufficient details to allow replication, including how and when they were actually administered	Methods
Outcomes	6a	Completely defined pre-specified primary and secondary outcome measures, including how and when they were assessed	Methods
	6b	Any changes to trial outcomes after the trial commenced, with reasons	NA
Sample size	7a	How sample size was determined	Methods
	7b	When applicable, explanation of any interim analyses and stopping guidelines	NA
Randomisation:			
Sequence generation	8a	Method used to generate the random allocation sequence	NA
	8b	Type of randomisation; details of any restriction (such as blocking and block size)	NA
Allocation concealment mechanism	9	Mechanism used to implement the random allocation sequence (such as sequentially numbered containers), describing any steps taken to conceal the sequence until interventions were assigned	NA
Implementation	10	Who generated the random allocation sequence, who enrolled participants, and who assigned participants to interventions	NA
Blinding	11a	If done, who was blinded after assignment to interventions (for example, participants, care providers, those	NA

		assessing outcomes) and how	
	11b	If relevant, description of the similarity of interventions	NA
Statistical methods	12a	Statistical methods used to compare groups for primary and secondary outcomes	NA
	12b	Methods for additional analyses, such as subgroup analyses and adjusted analyses	NA
Results			
Participant flow (a diagram is strongly recommended)	13a	For each group, the numbers of participants who were randomly assigned, received intended treatment, and were analysed for the primary outcome	NA
	13b	For each group, losses and exclusions after randomisation, together with reasons	NA
Recruitment	14a	Dates defining the periods of recruitment and follow-up	NA
	14b	Why the trial ended or was stopped	NA
Baseline data	15	A table showing baseline demographic and clinical characteristics for each group	NA
Numbers analysed	16	For each group, number of participants (denominator) included in each analysis and whether the analysis was by original assigned groups	Methods
Outcomes and estimation	17a	For each primary and secondary outcome, results for each group, and the estimated effect size and its precision (such as 95% confidence interval)	NA
	17b	For binary outcomes, presentation of both absolute and relative effect sizes is recommended	NA
Ancillary analyses	18	Results of any other analyses performed, including subgroup analyses and adjusted analyses, distinguishing pre-specified from exploratory	Methods
Harms	19	All important harms or unintended effects in each group (for specific guidance see CONSORT for harms)	NA
Discussion			
Limitations	20	Trial limitations, addressing sources of potential bias, imprecision, and, if relevant, multiplicity of analyses	NA
Generalisability	21	Generalisability (external validity, applicability) of the trial findings	NA
Interpretation	22	Interpretation consistent with results, balancing benefits and harms, and considering other relevant evidence	NA
Other information			
Registration	23	Registration number and name of trial registry	Methods
Protocol	24	Where the full trial protocol can be accessed, if available	NA
Funding	25	Sources of funding and other support (such as supply of drugs), role of funders	Financial Disclosure

*We strongly recommend reading this statement in conjunction with the CONSORT 2010 Explanation and Elaboration for important clarifications on all the items. If relevant, we also recommend reading CONSORT extensions for cluster randomised trials, non-inferiority and equivalence trials, non-pharmacological treatments, herbal interventions, and pragmatic trials. Additional extensions are forthcoming: for those and for up to date references relevant to this checklist, see www.consort-statement.org.



CLINICAL STUDY PROTOCOL

Characterisation of human disseminated cellular and humoral immune responses following sublingual or intramuscular deposition of antigens

Short title: Measuring responses to sublingual antigens

Study Identification: SG09-EN01

REC number: 09/80803/77

Version number: 1.04.2c

Date: 21 APRIL 2009 / 06 August 2009

Amendment 2. 27 Jan 2010

Amendment 3 23rd Feb 2010

This study will be conducted in accordance with ICH GCP Guidelines (Directive CPMP/ICH/135/95) and the Declaration of Helsinki (1964) and subsequent amendments.

SYNOPSIS

Title	Characterisation of human disseminated cellular and humoral immune responses following sublingual or intramuscular presentation of antigens
Short Title	Immune response to sublingual antigens
Sponsor	St. George's University of London
Principal Investigator	Prof. David J. M. Lewis
Planned Study Dates	October 2009 - September 2010
Study type	Physiology study with challenge agents
Objectives	<p>To explore in humans the:</p> <ol style="list-style-type: none"> 1. Concentration and isotype profile of antigen-specific antibody in serum and cervico-vaginal secretions 2. Frequency and isotype profile of antigen-specific antibody secreting cells in blood 3. Frequency and expression profile of mucosa-associated homing, memory and regulatory markers on antigen-specific T cells in blood in response to <i>in vitro</i> antigen stimulation 4. Profile of cytokine secretion by peripheral blood mononuclear cells in response to <i>in vitro</i> antigen stimulation measured by ELISA <p>following presentation of an immunological challenge agent via sublingual or axillary lymph nodes</p>
Challenge Agents	<p>Purified, recombinant L1 protein antigens from Human Papilloma Virus serotypes 6, 11, 16 & 18 in the following amounts:</p> <p>Human Papillomavirus Type 6 L1 protein 20 micrograms Human Papillomavirus Type 11 L1 protein 40 micrograms Human Papillomavirus Type 16 L1 protein 40 micrograms Human Papillomavirus Type 18 L1 protein 20 micrograms</p> <p>adsorbed on amorphous aluminium hydroxyphosphate sulphate adjuvant (225 micrograms Al)</p>
Study design	<ul style="list-style-type: none"> • Physiology study using challenge agents, non-CTIMP • Open label, non-randomised, hypothesis generating study
Sample size	<p>EIGHTEEN subjects in all, in two groups:</p> <p>(1) SIX receiving challenge agents by intramuscular injection of deltoid muscle</p> <p>(2) TWELVE receiving challenge agents sublingually</p>
Study population	Healthy female adult volunteers aged 18-35

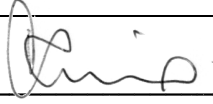
Route of Administration	<ol style="list-style-type: none"> 1. Intramuscular (deltoid injection): 6 subjects 2. Sublingual (as drops under the tongue): 12 subjects <p>Given on three occasions at 0, 1 and 4 months</p>
Duration of Treatment	<p>The entire duration of the study will be 20 weeks (up to 24 weeks including screening period).</p> <p>During this period subjects will make a total of 9 outpatient visits to the clinical site</p>
Primary & Secondary endpoints	<p>No primary and secondary endpoints are defined in this hypothesis-generating study.</p> <p>A number of exploratory variables will be measured.</p>
Exploratory variables	<ol style="list-style-type: none"> 1. Concentration, neutralising activity and isotype profile of antigen-specific antibody in serum and cervico-vaginal secretions measured by ELISA and/or LUMINEX assay and virus neutralisation assay. 2. Frequency and isotype profile of antigen-specific antibody secreting cells in blood measured by ELISPOT assay 3. Frequency and expression profile of mucosa-associated homing, memory and regulatory markers on antigen-specific T cells in blood in response to <i>in vitro</i> antigen stimulation measured by Flow Cytometry (FACS) and CFSE proliferation assay 4. Profile of cytokine secretion by peripheral blood mononuclear cells in response to <i>in vitro</i> antigen stimulation measured by ELISA
Safety evaluations	None, not a CTIMP
Efficacy evaluation	None, not a CTIMP

Signatures

PROTOCOL APPROVED BY

Prof. David JM Lewis, St George's University of London

Signature



Date

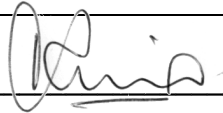
22 FEB 2010

**I AGREE TO CONDUCT THE STUDY IN ACCORDANCE WITH THE INFORMATION
CONTAINED IN THIS STUDY PROTOCOL:**

Principal Investigator, St George's University of London

Prof. David JM Lewis

Signature



Date

22 FEB 2010

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1 General Information

Protocol title	Characterisation of human disseminated cellular and humoral immune responses following sublingual or intramuscular presentation of antigens
Protocol title acronym	SG09_EN01
Sponsor	St George's University of London
Host site	St George's University of London
Name(s), address(es) and telephone number(s) of clinical laboratory (ies) & other medical and/or technical department(s) and /or institutions involved in the study	<p>Abraham Roodt, Clinical Trials and Data Manager The Doctors Laboratory 60 Whitfield Street London W1T 4EU Tel: +44 (0)20 7307 7373</p> <p>Zhiming Huo MD PhD St. George's Vaccine Institute Centre for Infection St. George's - University of London Cranmer Terrace London SW17 0RE Tel: 0 20 87 25 32 11</p> <p>Simon Beddows Ph.D. Principal Scientist HPV R&D Section Head Virus Reference Department Centre for Infections Health Protection Agency 61 Colindale Avenue London NW9 5HT Tel: +44 (0) 20 8327 6169 Fax: +44 (0) 20 8200 1569</p> <p>R. Karl Malcolm BSc (Hons) PhD CChem MRSC PGCHET Senior Lecturer in Pharmaceutics School of Pharmacy Medical Biology Centre 97 Lisburn Road Belfast BT9 7BL Northern Ireland T: +44 (0)28 9097 2319 E: k.malcolm@qub.ac.uk</p>

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2 Study Background

2.1 Research Background

Please provide any relevant background information to support the research area/disease

This study builds on and extends a series of studies we have undertaken using licensed vaccines as safe human challenge agents to induce an immune response that can then be characterised. Using this approach we have characterised the profile of B and T cell responses following oral and nasal administration and intramuscular injection of model antigens¹⁻⁴.

It is well recognised that uptake and presentation of antigens either locally or in lymph nodes draining mucosal surfaces such as gut, respiratory, or genital tracts, often induces an immune response with a different profile that when the same antigens are presented in systemic lymph nodes following intramuscular injection⁵. In our previous studies we found for example that while oral delivery of antigen induces mainly an IgA response amongst circulating B cells, and surface markers indicating a gut homing tendency, nasal administration of antigens induces a mixed IgG and IgA response, and both systemic and mucosal homing markers¹⁻⁴. This, and others' work, has contributed the concept of a "common mucosal immune system" (CMIS) distinct from the systemic immune system. Injected (intramuscular or subcutaneous) antigens tend to interact with the systemic immune system, induce IgG, and lymphocytes with surface markers (such as L-selectin) that home to systemic but not mucosal tissue. In contrast, application of antigen to a mucosal surface spreads responses throughout the mucosal immune system by the expression of mucosal-homing markers such as ($\alpha 4\beta 7$ integrin for gut and $\alpha 4\beta 1$ integrin and CCR10 to genital tract⁶). There are limitations to dissemination within the CMIS: for example nasal immunisation preferentially induces immunity in the respiratory and genital tracts, whereas rectal immunisation preferentially induces immunity in the large bowel⁵. A better understanding of how the common mucosal immune system is "wired-up", and an improved knowledge of the cell surface markers that guide lymphocytes to mucosal or systemic sites, will be of value in better understanding inflammatory and immune-mediated diseases such as post reactive arthritis, and designing novel vaccines and vaccine delivery systems.

Whereas 'oral' delivery of antigens has tended to involve the ingestion, swallowing and presentation of antigen to small bowel immune induction sites, interest has focussed recently on sublingual delivery of antigen. The sublingual mucosa is rich in dendritic cells that can take up antigens and present them to the mucosal immune system⁷. It has been known for many years that repeated sublingual administration (usually daily application for many weeks) of high doses of *allergens* (such as grass pollen) is safe, and induces a non-allergic immune response, which is as clinically effective in reducing allergic responses to the same allergens as desensitisation by repeated subcutaneous injections⁷. Different patterns of cell surface markers associated with regulatory function, and cytokine secretion in response to stimulation have also been observed after nasal and sublingual routes of allergen delivery⁸, which is an area we wish to explore using the pathogen-derived antigen challenge agents in this study.

While the conversion of an allergic-type of immunity to a non-allergic type of immunity appears to require frequent sublingual application of high doses of allergens, it has recently been shown that only a few sublingual exposures of pathogen-derived *antigens* can induce protective immunity. For example, sublingual administration of inactivated influenza virus to mice on two occasions induced both systemic and mucosal antibody responses and conferred protection against a lethal intranasal challenge with influenza virus⁹. These responses could be enhanced when cholera and E. coli toxin-derived adjuvants were co-administered⁹. Cholera toxin subunit B itself induces systemic and genital tract immune responses in mice¹⁰ and humans (J Holmgren personal communication) after sublingual

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immunisation. However, molecules such as cholera toxin are somewhat unusual in that they are potent immunogens and mucosal adjuvants in their own right, with powerful effects on immune cells that polarise and alter immune response profiles⁵, and so are not very representative of protein antigens in general.

It is therefore of interest to observe the profile of more representative protein *antigens* following sublingual delivery, and to compare this with our and others' experience with nasal, oral and injected delivery of antigens. We have selected the L1 protein of Human Papilloma Virus (HPV) as a representative protein antigen to use as a challenge agent to induce immune responses disseminated within the CMIS, and the systemic immune system, which can then be studied and characterised. Each strain of HPV has a unique L1 protein antigen: strains 16 and 18 are associated with cervical cancer in women, and strains 6 and 11 with anogenital warts in both sexes.

The HPV L1 antigens have been selected from a number of potential immunological challenge agents, as they possess several advantageous characteristics for a human immunological challenge agent:

1. They are antigens from the capsid of a human pathogenic virus that infects the genital tract, and therefore representative of a wide number of viral protein antigens.
2. They are available as a pure preparation, suitable for human use, in the licensed HPV vaccine "Gardasil".
3. Their safety via injection has been demonstrated in numerous clinical trials of HPV vaccines, and the UK Dept of Health is engaged in a mass immunisation campaign employing HPV L1 antigen vaccines in healthy young adult women.
4. The optimum safe and immunogenic human dose of L1 antigens has been determined after injection, which provides a guide to the dose suitable for sublingual application.
5. They spontaneously form "Virus Like Particles" in the "Gardasil" preparation. These larger structures are expected to be more readily taken up by sublingual dendritic cells than small proteins or peptides, and so increase the probability of a detectable immune response to the challenge.
6. They are adsorbed onto alum, which has been used for decades in many licensed vaccines. This will aggregate and is expected to further enhance dendritic cell uptake.
7. They are known to reliably induce both serum and genital tract antibody responses in women after injection, and assays to measure responses in genital tract secretions are described¹¹. This ensures we will have a reliable positive response group, which we will recruit first to enable assay development for subsequent application to the sublingual challenge group in which immune responses are expected to be lower and less frequent.
8. The "Gardasil" preparation has the advantage of containing 4 different L1 antigens (from strains 6, 11, 16 and 18). Seroepidemiology of women aged 23 (the median age of subjects generally recruited into our studies) in England¹² shows that around 45% already have immunity to at least one of the 4 antigens, but only 15% have immunity to two or more. This means that in our sample we will expect half the subjects not to be immune to any of the antigens, in whom we can study priming and boosting responses to a neoantigen, which is the most interesting aspect of this study - as previous genital tract infection may bias the subsequent response to the sublingual immune challenge.

However, we will also have around half of the subjects in whom we can simultaneously study responses to recall antigens that they have seen before (thereby increasing our prospects of seeing any response at all to sublingual challenge) and neoantigens. While this hypothesis generating study is not powered to analyse subgroup responses to the 4 antigens in detail, by determining subject's serostatus at entry it will be interesting to see if there is any suggestion that prior genital tract infection modulates the response to

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cross-reacting antigens. Prior genital infection is essentially a mucosal immunological prime, which may be boosted by the mucosal (sublingual) or parenteral (intramuscular) boost in different ways. It has recently been shown that around 1% of young women have oral infection with HPV¹³, which might theoretically modulate the immune response. However the low prevalence is unlikely to significantly affect the number of subjects being studied. Any observed differences in immune responses between subjects seropositive or seronegative at entry could be further investigated in larger studies.

2.2 Challenge Agents

Please provide details of challenge agents(s) to be used in the study (including name, manufacturer, country of manufacture, form)

The challenge agents will consist of purified, recombinant L1 protein antigens from Human Papilloma Virus serotypes 6, 11, 16 & 18.

A single, 0.5 mL pre-filled syringe of "Gardasil" Human Papillomavirus Vaccine [Types 6, 11, 16, 18] (Recombinant, adsorbed) will be used for each challenge, containing:

Human Papillomavirus Type 6 L1 protein 20 micrograms

Human Papillomavirus Type 11 L1 protein 40 micrograms

Human Papillomavirus Type 16 L1 protein 40 micrograms

Human Papillomavirus Type 18 L1 protein 20 micrograms

L1 proteins are in the form of virus-like particles produced in yeast cells (*Saccharomyces cerevisiae* CANADE 3C-5 (Strain 1895)) by recombinant DNA technology, adsorbed on amorphous aluminium hydroxyphosphate sulphate adjuvant (225 micrograms Al), together with the following excipients: Sodium chloride, L-histidine, polysorbate 80, Sodium borate, Water for injections.

Each dose is presented as a suspension in a pre-filled syringe.

The Marketing Authorisation Holder is Sanofi Pasteur MSD SNC, 8 rue Jonas Salk, F-69007 Lyon, France. Vaccines will be purchased through an approved supplier in the UK.

Syringes will be stored in the Pharmacy Area of the Vaccine Institute in a refrigerator (2°C - 8°C).

Vaccine Institute SOPs for receipt, storage, accountability and reconciliation will be followed.

2.3 Objectives and Purpose

Please Insert a detailed description of the objectives and the purpose of the study

The purpose of this hypothesis-generating study is to explore in healthy humans the profile of the mucosa-associated and systemic immune systems' response to pathogen-derived protein antigens.

We will induce an immune response in healthy volunteers by the application of immunological challenge agents to the mucosal immune system (locally or via the sublingual lymph nodes), and the systemic immune system (probably via axillary lymph nodes). This immune response will then be characterised by a number of immune assays, and any differences and similarities in the pattern of responses induced by the two systems explored.

We will apply techniques that we have developed and published in previous studies of oral,

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nasal and intramuscular challenge, as well as developing new techniques to characterise T cell responses and regulatory, memory and genital tract-associated surface markers.

Initially we will recruit and challenge subjects via the intramuscular route, as this is known to reliably induce systemic and vaginal secretion antibody responses. This will enable us to develop exploratory cellular assays. Once we are confident that the novel assays are performing with acceptable accuracy and reliability, we will recruit subjects to receive sublingual challenge, as this route of challenge is expected to give lower level and less frequent responses due to reduced antigen access across the sublingual mucosa.

Blood will be taken at several time-points for serum antigen-specific antibody levels and enumeration and characterisation of antigen-specific T and B cells using well-established techniques. We will collect cervical and vaginal secretions using well-established techniques within the Vaccine Institute in which secretions are collected onto clinical ophthalmic sponges by gentle application against the wall of the vagina and cervix under direct speculum examination. Secretions are extracted by centrifugation into a preservative buffer that prevents protease degradation, prior to analysis for antigen-specific antibody in ELISA or Luminex assay and neutralisation assay.

As this is a hypothesis generating study it is not expected that we will be able to characterise all the parameters, in all the assays, in all the subjects. The core assays will consist of measurement of antigen-specific antibody in serum and cervico-vaginal secretions as these techniques are well established in the literature. The other assays will be exploratory and methodological development.

Specifically we will characterise:

1. Concentration, neutralising activity and isotype profile of antigen-specific antibody in serum and cervico-vaginal secretions

Using published ELISA and Luminex^{11, 12} assays with L1 proteins as coating antigens, and virus neutralisation assay, we will measure the levels of IgG and IgA L1-specific antibody in serum and mucosal secretions. A 'mucosal' response will be characterised by IgA>IgG, and a systemic/nasal response by the reverse.

2. Frequency and isotype profile of antigen-specific antibody secreting cells in blood

We will adapt published ELISpot assays^{3, 14-16} (to detect single cells secreting antigen specific antibody *in vitro*) we have developed to detect responses to nasal or oral administration of antigen, to employ L1 proteins as coating antigens. By enumerating IgG and IgA L1-specific antibody secreting cells trafficking in the blood a 'mucosal' response will be characterised by IgA>IgG, and a systemic/nasal response by the reverse. Expression of mucosa-associated homing markers on antigen-specific B cells in blood can be measured by bead extraction, but this is not planned in this study.

3. Frequency and profile of regulatory, memory and mucosa-associated homing marker expression on antigen-specific T cells in blood measured by CFSE proliferation assay

We will update our published T cell assays¹ to detect antigen specific T cells trafficking in the blood following challenge responding *in vitro* to antigen stimulation, using the flow cytometry based CFSE proliferation assay running on a Guava flow cytometer. In addition to enumerating numbers of T cells we will characterise intracellular cytokine expression and cell surface markers of responding cells including $\beta 7$ integrin, CCR10, L-selectin and other markers associated with mucosal or systemic homing⁶, and markers of memory (CD45RO/RA) and regulatory function (Foxp3).

4. Profile of cytokine secretion by peripheral blood mononuclear cells in response to *in vitro*

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antigen stimulation measured by ELISA. By measuring secretion of cytokines into the supernatant of cultures of PBMCs stimulated with L1 antigens we will be able to determine mucosal patterns of response (IL17) and the Th1: Th2 bias.

2.4 Preclinical and clinical data

Please insert a summary of findings from non-clinical studies that potentially have clinical significance and from clinical studies relevant to the study (please reference all published findings)

This study builds on and extends a series of published studies we have undertaken using licensed vaccines as safe human challenge agents to induce an immune response that can then be characterised. Using this approach we have characterised the profile of B and T cell responses following oral and nasal administration and intramuscular injection of model antigens¹⁻⁴.

Extensive preclinical and clinical data in support of the suitability of the antigens in the Gardasil preparation as challenge agents, and their safety evaluation, are presented in detail in section 5 of the Gardasil SPC (Appendix 3)

Gardasil is an adjuvanted non-infectious recombinant quadrivalent vaccine prepared from the highly purified virus-like particles (VLPs) of the major capsid L1 protein of HPV types 6, 11, 16 and 18. The VLPs contain no viral DNA, they cannot infect cells, reproduce or cause disease. HPV only infects humans, but animal studies with analogous papillomaviruses suggest that the efficacy of LI VLP vaccines is mediated by the development of a humoral immune response.

In clinical studies, 99.9%, 99.8%, 99.8%, and 99.6% of individuals who received Gardasil became anti-HPV 6, anti-HPV 11, anti-HPV 16, and anti-HPV 18-seropositive, respectively, by 1 month Postdose 3 across all age groups tested. Gardasil induced high anti-HPV Geometric Mean Titres (GMTs) 1 month Postdose 3 in all age groups tested.

The efficacy for HPV 16/18 related CIN 2/3 or AIS is based on data from protocols 005 (16-related endpoints only), 007, 013, and 015. The efficacy for all other endpoints is based on protocols 007, 013, and 015. Results of individual studies support the results from the combined analysis. Gardasil was efficacious against HPV disease caused by each of the four vaccine HPV type.

2.5 Population

Please insert a description of the population to be studied (including sample size, participant group, patient group (if applicable)). Please indicate if a vulnerable population is to be studied

- We will recruit a total of 18 healthy female volunteers, aged 18-35.
- All subjects will be healthy, and will provide written informed consent.
- No vulnerable groups will be included.
- From previous experience we expect the median age to be around 23 years old.
- We expect around half the subjects to be naive for the challenge agents¹², while the remainder will have had some exposure to one or more of the antigens.

As group size is small in this hypothesis generating study we have elected to recruit a narrow age range, to avoid possible bias due to age-related (hormonal) changes in immune response, especially in the genital tract.

Group 1:

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We will initially recruit SIX subjects into a positive response group that will receive INTRAMUSCULAR challenge. This will enable us to generate positive reference samples and develop the novel assays, as it is known that intramuscular immunisation with L1 antigens induces systemic and mucosal immune responses.¹¹

Group 2:

Once the novel assays are performing satisfactorily, we will recruit a further TWELVE subjects into the group to receive SUBLINGUAL challenge. It may not be necessary for subjects in group 1 to have completed the protocol before group 2 is recruited.

2.6 Dose Rationale

Please insert a description of, and justification for the, route of administration, dosage, dosage regimen and treatment period(s)

A single 0.5 mL vial of "Gardasil" vaccine has been shown through extensive clinical trials to be the optimum safe and immunogenic dose for intramuscular immunisation, and is licensed for use in healthy adult women the UK.

To enable this scientific study to complete within the time and resources available, a shortened 0, 1, 4 month challenge regimen will be employed, which is based on the recommendation of the Gardasil SPC section 4.2: "*If an alternate vaccination schedule is necessary, the second dose should be administered at least one month after the first dose and the third dose should be administered at least 3 months after the second dose*".

Although sublingual application of antigen is likely to be less efficient at stimulating immune responses due to the physical barrier to passive antigen entry across the sublingual mucosa, the extensive human safety profile of this dose level and dose regimen when injected makes it an appropriate choice for this initial hypothesis generating study.

The tiny (microgram) amounts of L1 protein and alum present no credible risk if swallowed, when compared with the huge (gram) amounts of protein (and other particulates such as toothpaste) swallowed daily. Any swallowed L1 or alum is expected to be immediately degraded by gastric acid and digestive enzymes.

2.7 Risk/Benefits

Please insert a summary of the known and potential risks & benefits, if any, to human subjects

2.7.1 Potential Risks

2.7.1.1 General risks

Gardasil does not contain thiomersal preservative. It does not contain live organisms and cannot cause HPV infection or disease. The VLPs contain no viral DNA, they cannot infect cells, reproduce, or cause disease.

Venepuncture may be associated with bruising, localised discomfort and fainting. Dedicated clinical facilities, including phlebotomy chairs will be used and staff must be trained and experienced in venepuncture. The blood volumes drawn over 20 weeks do not present a significant risk of causing anaemia in a healthy person, and subjects must be screened for anaemia before entry.

Vaginal speculum examination can be uncomfortable, and staff must be experienced in the technique. Collection of secretions onto single-use, sterile, soft ophthalmic sponges (developed to collect tears) is painless and risk-free. A dedicated room complete with colposcopy couch, private changing and showering area, must be used to minimise subject discomfort and ensure privacy.

Clinical examination and screening blood tests may disclose medical conditions such as HIV, sexually transmitted disease, cervical cancer. Staff taking consent must be trained in pre-test discussions and the Principal Investigator is an Infectious Diseases clinician who can inform subjects of the diagnoses, answer questions or concerns, and arrange appropriate follow-up with the subject's consent.

2.7.1.2 Intramuscular injection

The following text is taken from sections 4.8 and 4.9 of the Gardasil SPC (Appendix 3):

The following vaccine-related adverse reactions were observed among recipients of Gardasil at a frequency of at least 1.0% and also at a greater frequency than observed among placebo recipients. They are ranked under headings of frequency using the following convention: [Very Common (1/10); Common (1/100, <1/10); Uncommon (1/1,000, <1/100); Rare (1/10,000, <1/1,000); Very Rare (<1/10,000), including isolated reports]

General disorders and administration site conditions:

Very common: pyrexia.

Very common: at the injection site: erythema, pain, swelling.

Common: at the injection site: bruising, pruritus.

In addition, in clinical trials adverse reactions that were judged to be vaccine- or placebo-related by the study investigator were observed at frequencies lower than 1%:

Respiratory, thoracic and mediastinal disorders:

Very rare: bronchospasm.

Skin and subcutaneous tissue disorder:

Rare: urticaria.

Seven cases (0.06%) of urticaria were reported in the Gardasil group and 17 cases (0.18%) were seen in the adjuvant-containing placebo group.

In the clinical studies, subjects in the Safety Population reported any new medical conditions during the follow-up of up to 4 years. Among 11,778 subjects who received Gardasil and 9,686 subjects who received placebo, there were 26 cases of non-specific arthritis/arthropathy reported, 19 in the Gardasil group and 7 in the placebo group.

Post Marketing Experience

Post Marketing adverse events have been spontaneously reported for Gardasil and are not listed above.

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Because these events were reported voluntarily from a population of uncertain size, it is not possible to reliably estimate their frequency or to establish, for all events, a causal relationship to vaccine exposure.

Blood and lymphatic system disorders: lymphadenopathy.

Immune system disorders: hypersensitivity reactions including anaphylactic/anaphylactoid reactions.

Nervous system disorders: Guillain-Barré syndrome, dizziness, headache, syncope.

Gastrointestinal disorders: nausea, vomiting.

Musculoskeletal and connective tissue disorders: arthralgia, myalgia.

General disorder and administration site conditions: asthenia, fatigue, malaise

Overdose

There have been reports of administration of higher than recommended doses of Gardasil. In general, the adverse event profile reported with overdose was comparable to recommended single doses of Gardasil.

Anaphylaxis

Staff administering challenge agents must be trained in resuscitation and management of anaphylaxis, with regular refreshers. The clinical site will maintain a crash trolley, oxygen, defibrillator and SOPs to manage acute medical emergencies associated with immunisation, including access to emergency services. Adequate numbers of trained staff must be present during challenges.

2.7.1.3 Topical sublingual application

The risk profile associated with topical sublingual application has not been specifically evaluated, but is expected to be considerably less than after injection given the low level of protein expected to passively cross the oral mucosa, compared with the dose delivered intramuscularly. Single-dose and repeated-dose toxicity and local tolerance studies revealed no special hazards to humans (Gardasil SPC section 5.3 Appendix 3). Natural infection with HPV (which can also be oral¹³) is initially asymptomatic and not associated with localised reactions to L1 proteins on the viral surface that contact the mucosal surface (warts develop after 3 weeks due to the effects on cell growth of internal viral proteins and cancer-promoting genes - not present in the Gardasil preparation).

The tiny (microgram) amounts of L1 protein and alum present no credible risk if swallowed, when compared with the huge (gram) amounts of protein (and other particulates such as toothpaste) swallowed daily. Any swallowed L1 or alum is expected to be immediately degraded by gastric acid and digestive enzymes.

As daily sublingual application of allergens over many months has been shown to be required to induce tolerance to allergic responses, no tolerising effect is expected from the three applications of these virus-derived protein antigens at 0, 1 and 4 months proposed here.

2.7.2 Potential Benefits

By better understanding the common mucosal immune system we will learn more about how the body handles pathogen antigens presented via systemic or mucosa-associated lymph nodes. This may advance our understanding of immune responses to viral infections, and inform the design of mucosal vaccine delivery systems. We will also develop novel assays, and a model of mucosal immune challenge with model viral antigens.

Each subject screened will receive a free basic medical health check.

2.7.2.1 Intramuscular injection

Section 5.1 of the Gardasil SPC (Appendix 3) details the level of benefit associated with intramuscular immunisation.

In clinical studies, 99.9%, 99.8%, 99.8%, and 99.6% of individuals who received Gardasil became anti-HPV 6, anti-HPV 11, anti-HPV 16, and anti-HPV 18-seropositive, respectively,

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by 1 month Post-dose 3 across all age groups tested. Gardasil induced high anti-HPV Geometric Mean Titres (GMTs) 1 month Post-dose 3 in all age groups tested. This level of immunity was associated with high levels (100%) of protection against cervical cancer lesions. Subjects over 18 are not eligible for the free NHS mass immunisation programme and so will benefit from a free vaccination (currently approximately £120 per injection).

Subjects already infected with HPV can expect to gain no benefit against the serotype with which they are infected, but will be protected against other serotypes. Although we will use an alternative schedule of immunisation, it is one recommended by the Gardasil SPC and subjects can expect to receive a high level of benefit in protection against HPV infection and associated cervical cancer or genital warts.

2.7.2.2 Topical sublingual administration

No data is available for the level of immunity induced by sublingual immunisation. However, due to physical barriers limiting antigen penetration across the oral mucosa, and destruction of swallowed antigen by gastric acid and digestive enzymes, it is expected that any immune responses induced will be low level and non-protective.

It is theoretically possible that topical sublingual exposure may prime subjects for an enhanced immune response to subsequent intramuscular injection, however this is speculative. Subjects should therefore not expect to gain any benefit from inclusion in this arm of the study.

2.8 Subject population(s) for analysis

Please insert the number of subjects planned to be enrolled. In multi-centre studies, the number of enrolled subjects projected for each study site should be specified. Provide a reason for choice of sample size, including reflection on (or calculations of) the power of the study and clinical justification

All subjects will be recruited at one site.

We will recruit a total of 18 healthy female volunteers, aged 18-35.

From previous experience we expect the median age to be around 23 years old.

We expect around half the subjects to be naive for the challenge agents¹², while the remainder will have had some exposure to one or more of the antigens.

We will initially recruit the first SIX subjects into a group that will receive INTRAMUSCULAR challenge.

We will subsequently recruit a further TWELVE subjects into the group to receive SUBLINGUAL challenge.

The numbers have been selected according to our previous experience with mucosal antigen delivery, and the expected circa 100% response rate seen with intramuscular immunisation with HPV vaccines¹⁷. As this is a hypothesis generating study it is not powered to enable between-group subgroup analysis. A response after challenge in any of the assays will constitute a valid response to challenge. In addition to providing data on antigen handling via systemic or mucosal routes, this study will generate data on the feasibility of sublingual challenge studies, and guide the choice of suitable assays to characterise immune responses in subsequent larger studies.

2.9 Study design/type
Please insert a description of the type/design of the study to be conducted (e.g. double-blind placebo controlled, parallel design) and a schematic diagram of the study design, all procedures and stages.

- **Physiology study using a challenge agent, not a CTIMP**
- **Open label, non-randomised, hypothesis generating study**

2.9.1 Schedule of visits and procedures

Visit no.	1	2	3	4	5	6	7	8	9
Study day no.	-28 to -2	0	7	28	35	56	112	119	140
Study week no.		0	1	4	5	8	16	17	20
Study month no.		0		1		2	4		5
Informed consent	X								
In/exclusion criteria	X	X		X			X		
Demography	X								
Pulse, temperature	X	X		X			X		
Heart (including blood pressure), lungs, abdomen, oral cavity; cervical speculum	X								
Pregnancy test									
Blood	X								
Urine		X		X			X		
Urinalysis (dipstick)	X								
Blood for exclusions (15 mL): FBC, HBV sAg, HIV1&2 Ab, HCV Ab*	X								
Challenge									
Group 1: 0.5 mL Gardasil IM		X		X			X		
Group 2: 0.5 mL Gardasil sublingual		X		X			X		
Blood for B / T cell analysis (20-50 mL)†		BT	B	TB	B	TB	BT	B	TB
Blood for serum antibody (5 mL)		X	X	X	X	X	X	X	X
Cervico-vaginal secretions for antibody		X		X		X	X		X
Volume of blood drawn (Total: 380 mL)	15	55	25	55	25	55	55	25	55

* FBC: Full Blood Count. HBV sAg: Hepatitis B virus surface antigen. HIV1&2 Ab: Antibody for HIV1 & 2 viruses. HCV Ab: Antibody to Hepatitis C virus
 † 50 mL total split as 15 mL to B cell assay / 35 mL to T cell assay on days when both taken. Otherwise 20 mL for B cell assays and 50 mL for T cell assays on days when taken alone

2.9.2 Acceptable Visit Windows

Visits within the following window periods will not constitute a protocol deviation:

Visits 3, 5 and 8: +/- 1 day

Visits 4, 6, 7 and 9: +/- 2 days

Given the exploratory nature of this study, it will be acceptable to collect and analyze samples taken at the *closest possible time* to the scheduled visit, if the subject is unable to make the scheduled visit. No additional visits are allowed.

2.9.3 Details of study visits

The length of visits will vary according to the purpose of each particular visit.

After initial contact expressing interest in the study, subjects will receive verbal information and an Information Sheet which they are encouraged to discuss this with their GP, friends and family. After at least 24 hours subjects will be eligible to sign written consent, having had a pre-test discussion relating to HIV testing. They are then eligible to make Visit 1.

Visit 1: Eligibility check

(Between 28 and 2 days before visit 2)

Basic demographic information will be collected to identify subject, and to check inclusion/exclusion criteria. No information relating to race, height, weight will be collected. Past and current medical and medication history will be recorded to check exclusion criteria. A general medical examination (heart, lungs, abdomen, mouth, vagina-cervix (with speculum) will be performed. Pulse and temperature recorded.

Blood will be taken and tested for Full Blood Count, HIV and Hepatitis B and C infections, pregnancy. Urine dipstick analysis.

Visits 2, 4 and 7: Antigen delivery and immunology response follow-up

(Day 0, 28 and 112)

Pulse and temperature recorded. General state of health, concomitant medications elicited to determine any exclusion/withdrawal criteria. Urine pregnancy test carried out.

Blood taken for immunology assays:

- Serum on all visits
- B cell assays visits 2 & 7
- T cell assays on all visits

Vaginal-cervical secretions collected onto a soft ophthalmic sponge using a speculum, for immunology assays.

The HPV antigens are given sublingually (as drops placed onto the undersurface of the tongue) or by injection (into the upper arm muscle) - according to study group.

Visits 3, 5, 6 and 8: Immunology response follow-up

(Day 7, 35, 56, 119)

Blood taken for immunology assays:

- Serum on all visits
- B cell assays visits 3, 5 & 8
- T cell assays visit 6

Vaginal-cervical secretions collected onto a soft ophthalmic sponge using a speculum at visit 6, for immunology assays

Visit 9: Final immunology response follow-up

(Day 140)

At the final visit blood will be taken for T cell and serum immunology assays. Vaginal-cervical secretions collected onto a soft ophthalmic sponge using a speculum, for immunology assays.

On **one only** of visits 2, 4, 6, 7 **or** 9 subjects will be invited to collect a vaginal sample using the self-inserted and self-removed 'Instead cup' in addition to weck-cel samples. Participation in this procedure is voluntary and not required for protocol compliance. This sample may be used for immunology assays or by collaborators investigating the *in vitro* effects of healthy cervico-vaginal samples on vaccine antigen integrity. Samples will be processed according to laboratory SOPs of the collaborating laboratories.

This concludes a subject's participation.

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2.10 Data Identification

Please insert the identification of any data to be recorded directly on the Case Report Forms (CRFs) (i.e. no prior written or electronic record of data), and to be considered to be source data

Source data will consist of the subject demographics, results of screening exclusion assays, physical examination, administration of challenge agents, details of vials used, allocation to group, and will be entered directly onto the CRFs.

Results of immunological assays will be entered onto paper worksheets, which will be considered source data. Electronic data will be used for subsequent analysis.

2.11 Primary and/or Secondary Study Endpoints

Please insert a specific statement of the primary endpoints and the secondary endpoints, if any, to be measured during the study

No primary and secondary endpoints are defined in this hypothesis-generating study.

A number of exploratory variables will be measured:

1. Concentration, neutralising activity and isotype profile of antigen-specific antibody in serum and cervico-vaginal secretions measured by ELISA and/or LUMINEX assay and virus neutralisation assay
2. Frequency and isotype profile of antigen-specific antibody secreting cells in blood measured by ELISPOT assay
3. Frequency and expression profile of mucosa-associated homing, memory and regulatory markers on antigen-specific T cells in blood in response to *in vitro* antigen stimulation measured by Flow Cytometry (FACS) and CFSE proliferation assay
4. Profile of cytokine secretion by peripheral blood mononuclear cells in response to *in vitro* antigen stimulation measured by ELISA

2.12 Randomisation

Please insert a description of the measures taken to minimize/avoid bias, including randomisation (please include a description of the method) and blinding (if applicable)

There will be no randomisation.

The nature of the challenge makes blinding impossible and no placebo or comparator will be used. Each subject acts as their own control, comparing pre-challenge with post-challenge time points.

The first 6 subjects will be recruited into the intramuscular challenge group, to allow assays to be developed and positive control samples generated, as this route of challenge is known to reliably induce mucosal and systemic immune responses.

The remaining 12 subjects will then be entered into the sublingual group.

2.13 Maintenance of randomisation codes

Please insert a description of the maintenance of the randomisation codes and the procedure for breaking codes

Not applicable.

2.14 Inclusion criteria

Please insert subject inclusion criteria

All subjects must satisfy the following criteria at study entry:

1. A female adult volunteer aged between 18 and 35 years old.
2. Subjects who the investigator believes can and will comply with the requirements of the protocol.
3. Provide written informed consent following a detailed written explanation of participation in the protocol.
4. They are in good health as determined by medical history, physical examination, haematology testing, and clinical judgement before entering into the study.
5. They are available for the whole duration of the study.
6. If of childbearing potential, must have a negative pregnancy test before each immunisation.
7. They have not donated blood during 3 months prior to study entry and agree to not donate for 3 months after the end of their participation in the study.
8. They are eligible for free medical treatment

2.15 Exclusion criteria

Please insert subject exclusion criteria

Subjects will be considered ineligible to enter the study if they present a condition which *could interfere with normal immune responses* to the challenge agent, including if they meet any of the following exclusion criteria:

1. They have already been vaccinated with an HPV vaccine
2. They have participated in a clinical trial in the last 6 months in which they have been exposed to an investigational product (pharmaceutical product or placebo or device) or concurrent participation in another clinical research study at the time of enrolment.
3. Use of any investigational or non-registered product (drug or vaccine) within 30 days preceding the first dose of challenge agent, or planned use during the study period.
4. They are pregnant or breast-feeding.
5. They have a known or suspected ongoing cervico-vaginal disease, malignancy or abnormality discovered at time of screening.
6. They present in the samples obtained at the screening visit: positive results for HIV, HBs Ag, anti-HBc and anti-HCV antibody, a clinically significant abnormality in haematology. Normal ranges will be defined by the pathology laboratory undertaking assays.
7. They have a clinically significant acute or chronic pulmonary, cardiovascular, hepatic or renal functional abnormality, blood or neurological disorders, immune dysfunction, autoimmune diseases, diabetes (excluding history of gestational diabetes), or malignancy at the time of enrolment, as determined by medical history, physical examination or laboratory screening tests.
8. They have received any form of immunosuppressive therapy in the past 6 months.
9. They are receiving any medications via vaginal route (as this may interfere with collection of samples).
10. They have any tongue or frenulum piercings or oral jewellery that may interfere with

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sublingual delivery.

11. They have received blood products or immunoglobulins 120 days prior to enrolment.
12. They have thrombocytopaenia or any coagulation disorder (because bleeding may occur following an intramuscular administration in these individuals).
13. Any other medical, psychiatric or social condition, drug treatment, occupational or other responsibility that, in the judgement of the investigator, would interfere with or serve as a contradiction to adherence to the study protocol or ability to give informed consent.
14. Individuals who cannot read or speak fluent English.

2.16 Subject withdrawal criteria

Please insert subject withdrawal criteria (i.e. terminating study treatment) and procedures specifying:

(a) When and how to withdraw subjects from the study.

(b) The type and timing of the data to be collected for withdrawal of subjects.

(c) Whether and how subjects are to be replaced.

(d) The follow-up for subjects withdrawn from study treatment.

(a) When and how to withdraw subjects from the study. (b) The type and timing of the data to be collected for withdrawal of subjects.

According to the Gardasil SPC (Appendix 3):

(i) "Individuals who develop symptoms indicative of **hypersensitivity** after receiving a dose of Gardasil should not receive further doses of Gardasil." - These subjects will be withdrawn from the study at that point. No further data will be collected.

(ii) "Administration of Gardasil should be postponed in subjects suffering from an **acute severe febrile illness**." - Such subjects will not be withdrawn but challenges will be postponed until the subjects fully recover. This will be recorded as a protocol deviation and subsequent timepoints will be calculated relative to the date of the postponed immunisation.

(iii) In addition to the above, subjects experiencing a **suspected drug reaction**, including those listed on the Gardasil SPC, will be withdrawn if in the opinion of the Principal Investigator the severity of the reaction may interfere with the study objectives, or if there is a possibility that the reaction may re-occur on subsequent challenge. No further data will be collected.

(iv) Any subject that becomes **pregnant**, or who tests positive in a pregnancy test taken during the protocol, will be immediately withdrawn. The subject will not undertake any further investigations, sample collections or challenges. The subject will be followed up and the outcome of the pregnancy recorded in the source notes.

(v) If new information becomes available that unfavourably alters the risk-benefit analysis of Gardasil, the Principal Investigator may halt the study and withdraw all subjects that require further challenges, as this is a hypothesis generating study and subjects will not be exposed to increased risk from premature discontinuation. The Principal Investigator will discuss the situation with subjects in the group receiving intramuscular immunisation, to determine whether they would benefit from continuing any incomplete schedule via their GP or travel clinic. However no further Gardasil challenges will be given by clinical staff. Subjects who have completed all scheduled challenges will not be withdrawn.

(vi) "However, the presence of a **minor infection**, such as a mild upper respiratory tract infection or low-grade fever, is not a contraindication for immunisation." Subjects will be challenged according to the expected study schedule in this situation. A note of the intercurrent illness will be made in the source notes.

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(c) Whether and how subjects are to be replaced.

There will be no replacement of withdrawn subjects, as this hypothesis generating study is not powered to detect any specific frequency of events.

(d) The follow-up for subjects withdrawn from study treatment.

Subjects withdrawn due to an adverse drug reaction will be followed-up according to the ▼Yellow Card scheme.

There will be no follow-up of other withdrawn subjects.

2.17 Study treatment

Please insert a description of the study treatment(s) and the dosage and dosage regimen. Also include a description of the dosage form, packaging, and labelling. Please indicate storage conditions and outline regulatory requirements (where applicable)

Name of challenge agents

Gardasil, suspension for injection in a pre-filled syringe. Human Papillomavirus Vaccine [Types 6, 11, 16, 18] (Recombinant, adsorbed).

Qualitative and quantitative composition

1 dose (0.5 ml) contains approximately:

Human Papillomavirus Type 6 L1 protein 20 micrograms

Human Papillomavirus Type 11 L1 protein 40 micrograms

Human Papillomavirus Type 16 L1 protein 40 micrograms

Human Papillomavirus Type 18 L1 protein 20 micrograms

L1 protein in the form of virus-like particles produced in yeast cells (*Saccharomyces cerevisiae* CANADE 3C-5 (Strain 1895)) by recombinant DNA technology. Adsorbed on amorphous aluminium hydroxyphosphate sulphate adjuvant (225 micrograms Al)

Dosage regimen

Three challenges with 0.5 mL of Gardasil will be given either sublingually or intramuscularly on months 0, 1 and 4 (as recommended in the Gardasil SPC).

Formulation

Suspension for injection in a pre-filled syringe. Prior to agitation, Gardasil may appear as a clear liquid with a white precipitate. After thorough agitation, it is a white, cloudy liquid.

Excipients

Sodium chloride L-histidine Polysorbate 80 Sodium borate Water for injections.

Incompatibilities

In the absence of compatibility studies, Gardasil must not be mixed with other medicinal products.

Shelf life

3 years.

Special precautions for storage

Store in a refrigerator (2°C - 8°C). Do not freeze. Keep the pre-filled syringe in the outer carton in order to protect from light.

Nature and contents of container

0.5 ml suspension in a pre-filled syringe (Type 1 glass) with plunger stopper (siliconized FluroTec-coated bromobutyl elastomer or non-coated chlorobutyl elastomer) and tip cap (bromobutyl) without needle.

Regulatory requirements

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Review of this protocol by MHRA confirmed that it is not a Clinical Trial of an Investigational Medicinal Product (IMP) as defined by the EU Directive 2001/20/EC. There is no requirement to submit a Clinical Trial Authorisation (CTA) to the MHRA.

Gardasil will be purchased from an approved supplier, and although this is not a clinical trial it will be used under the general direction and supervision of the Principal Investigator who is a registered medical practitioner.

Standard reporting of adverse drug reactions to MHRA will be conducted under the ▼Yellow Card Scheme.

2.18 Duration

Please insert the expected duration of subject participation and a description of the sequence and duration of all study periods, including follow-up, if any

Subjects will participate for up to 6 months including: a 1 maximum month screening period, followed by a 5 month follow-up period after the first immunological challenge.

2.19 Treatment of subjects

Please insert the treatment(s) to be administered, including the name(s) of all the product(s), the dose(s), the dosing schedule(s), the route/mode(s) of administration, and the treatment period(s), including follow-up period(s) for subjects for each treatment/study treatment group/arm of the study

Although this is not a clinical trial, the Principal Investigator shall maintain a Delegations Log of study staff authorised to administer challenge agents, who will have the appropriate training in immunisation and resuscitation techniques (in compliance with section 3 "Characteristics of staff" of the Department of Health "Patient Group Direction (PGD) for the supply and administration of Human Papillomavirus Vaccine" [Appendix 2]) which will be recorded on an up to date CV and training record held in the site file.

A single 0.5 mL dose of Gardasil HPV vaccine will be used as an immunological challenge in all subjects in both groups.

All subjects in both groups will receive a challenge on month 0, 1 and 4.

Group 1: Six subjects will receive the challenge via intramuscular injection into the deltoid muscle (on the contralateral side to their dominant hand), following the instructions in section 6.6 of the Gardasil SPC (Appendix 3).

Group 2: Twelve subjects will receive the challenge via drops applied topically to the sublingual surface of the tongue according to a predefined SOP (Appendix 1)

2.20 Discontinuation

Please insert a description of the stopping rules or discontinuation criteria for individual subjects, parts of the study, and entire study. Please include any information on dose modification procedures

Individual subject stopping/discontinuation is detailed in section 2.16

The study will be stopped if a suspected adverse drug reaction that is not listed on the "Gardasil" SPC is reported to the MHRA using the ▼Yellow Card scheme. In this case no further challenges will be administered and subjects already entered will be withdrawn from immunological follow-up at the discretion of the Principal Investigator.

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If new information becomes available that unfavourably alters the risk-benefit analysis of Gardasil, the Principal Investigator may halt the study and withdraw all subjects that require further challenges, as this is a hypothesis generating study and subjects will not be exposed to increased risk from premature discontinuation. The Principal Investigator will discuss the situation with subjects in the group receiving intramuscular immunisation, to determine whether they would benefit from continuing any incomplete schedule via their GP or travel clinic. However no further Gardasil challenges will be given by clinical staff. Subjects who have completed all scheduled challenges will not be withdrawn, and will complete all scheduled follow-up visits. No new subjects will be recruited.

2.21 Challenge Agents Accountability

Please insert accountability procedures including the placebo(s) and comparator(s), if any

Vaccine Institute GCP-compliant SOPs will be used to record receipt, administration, loss, and destruction of unused vials of Gardasil.

2.22 Accountability procedure

Please insert the procedure for accounting for missing, unused and spurious data

The SOPs and laboratory worksheets of the Vaccine Institute will be used to record and monitor data.

2.23 Monitoring subject compliance

Please insert the procedures for monitoring subject compliance

All challenges will be administered directly by trained and delegated study staff.

2.24 Permitted medication

Please insert medication(s)/treatment(s) permitted (including rescue medication) and not permitted before and/or during the study (e.g. concomitant medicines)

Subjects will be withdrawn at the point that they commence any medications or transfusions referred to in the Exclusion Criteria.

All other concomitant medications are permitted.

See also section 4.5 of Appendix 3 Gardasil SPC "Interaction with other medicinal products and other forms of interaction"

2.25 Efficacy parameters

Please insert the specifications of the efficacy parameters

Not a CTIMP: no efficacy parameters will be measured.

2.26 Method and timing for efficacy parameters

Please insert methods and timing for assessing, recording and analyzing efficacy parameters

Not applicable.

2.27 Safety parameters

Please insert specifications for safety parameters

Not a CTIMP: no safety parameters will be measured.

2.28 Adverse event reporting

Please insert details of expected adverse events and the procedures for eliciting reports of and for recording and reporting of these adverse events and intercurrent illnesses.

As the challenge agents are not IMPs there is no specific reporting of adverse events or SUSARS.

Suspected adverse drug reactions will be reported to MHRA using the s ▼Yellow Card Protocol 04-02c

scheme.

2.29 Adverse event follow-up

Please insert the type and duration of the follow-up of subjects after adverse events

Suspected adverse drug reactions will be followed to resolution or stabilisation and reported to MHRA using the ▼Yellow Card scheme.

2.30 Statistical methods

Please insert a description of the statistical methods to be employed including timing of any planned interim analysis(es), level of significance to be used

No primary and secondary endpoints are defined in this hypothesis-generating study.

A number of exploratory variables will be measured. As this is a hypothesis generating study it is not powered to enable between-group subgroup analysis. A response after challenge in any of the assays will constitute a valid response to challenge.

Appropriate statistical tests of significance (parametric or non-parametric) will be used for between-group, or between-time-point analyses of exploratory variables, according to the nature and size of the dataset, and in consultation with a statistician before publication.

2.31 Subject analysis

Please insert the selection of subjects to be included in the analyses (e.g. all randomised subjects, all dosed subjects, all eligible subjects, evaluable subjects)

Data will be analysed for all subjects receiving at least one challenge.

2.32 Method and timing for safety parameters

Please insert methods and timing for assessing, recording and analyzing safety parameters

Not applicable.

2.33 Termination criteria

Please insert the criteria for the termination of the study

The study will be terminated if:

1. Insufficient subjects have been recruited within 12 months of administering the first challenge to the first subject. In this case all subjects entered will complete the protocol.
2. A suspected adverse drug reaction that is not listed on the "Gardasil" SPC is reported to the MHRA using the ▼Yellow Card scheme. In this case no further challenges will be administered and subjects already entered will be withdrawn from immunological follow-up at the discretion of the Principal Investigator.

The study conduct shall comply with all relevant laws of the EU if directly applicable or of direct effect and all relevant laws and statutes of the UK country in which the study site is located including but not limited to, the Human Rights Act 1998, the Data Protection Act 1998, the Medicines Act 1968, and with all relevant guidance relating to medicines and clinical studies from time to time in force including, but not limited to, the ICH GCP, the World Medical Association Declaration of Helsinki entitled 'Ethical Principles for Medical Research Involving Human Subjects' (2008 version).

This study will be conducted in compliance with the protocol approved by the REC and according to GCP standards. No deviation from the protocol will be implemented without the prior review and approval of the sponsor and REC except where it may be necessary to eliminate an immediate hazard to a research subject. In such case, the deviation will be reported to the sponsor and REC as soon as possible.

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2.34 Ethical considerations

Please insert description of ethical considerations relating to the study

The informed consent shall explicitly make reference to the possibility that an abnormality may be incidentally discovered such as a haematological disorder, immunological deficiency, cervico-vaginal disease or malignancy, chronic viral infection, etc.

Subjects must be eligible for free medical treatment as part of the inclusion criteria. Any medical conditions discovered at the time of screening may therefore be managed at no cost to the subject.

Subjects will be fully informed of any abnormal conditions discovered by screening tests, the physical examination, or the immunology assays (e.g. an immunoglobulin or lymphocyte deficiency) and the consequences and necessary management explained by the Principal Investigator who is a Consultant Physician. The Principal Investigator will seek the permission of the subject to communicate the findings to their General Practitioner for further action, and ensure that the communication has been received. Subjects unwilling to have details of any condition reported to their GP will be encouraged to reconsider, but their wishes will be respected.

Subjects will be tested for HIV, hepatitis B and C at screening. This is required as HIV or chronic hepatitis virus infections will considerably affect the immune response to the challenge agents and skew the data. Clinical samples from infected subjects also pose a risk to laboratory staff as several of the exploratory assays cannot be run under Universal Precautions. Subjects will undergo a pre-test discussion by a qualified GUM nurse experienced in HIV and HIV-related issues, as part of the informed consent process. Any positive results will be given to the subject in person by the Principal Investigator who is a Consultant Infectious Diseases/HIV Physician, and who will personally ensure that necessary follow-up and management is arranged if the subject gives their permission. Subjects unwilling to have details reported to their GP will be encouraged to attend a GUM clinic where anonymity and confidentiality may be assured, but their wishes will be respected.

Subjects must consent to allow study staff to inform their General Practitioner of their participation in the study, nature and number of challenge agents administered, and to seek information that may contraindicate their safe exposure to the challenge agents.

Subjects entering the sublingual administration group will need to be fully aware, through written information in the informed consent sheet reinforced by verbal communication from study staff, that although they will have been exposed to the Gardasil vaccine they cannot expect to have protective immunity, and should consider themselves unimmunised in the event of being offered HPV immunisation in the future. Although specific trials have not been conducted, it is current practice that boosting with any of the available HPV vaccines is acceptable.

2.35 QC & QA

Please insert details of procedures for generation, recording and reporting of study data in compliance with the protocol, GCP standards and any other applicable regulatory requirements

Results of screening pathology tests will be generated by TDL on printed sheets which will be initialled and dated by the Principal Investigator and filed in the Source Notes.

Any clinically significant result will be marked "CS", and non-clinically significant results "NCS" by a trial Physician on the original test result printout. All annotations will be initialled

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and dated by a trial Physician.

Results of immunology assays will be recorded and checked by technicians undertaking the tests and signed, and dated by them on paper worksheets.

Study staff will be responsible for data entry and verification for those data they enter into the study documentation.

All data, both electronic and paper, will be held in secure (locked / password protected as appropriate) facilities within the Vaccine Institute under the provisions of the SGUL Data Protection Act registration and local governance requirements.

2.36 Deviation reporting

Please insert the procedures for reporting any deviation(s) from the original statistical plan. Any deviation(s) from the original statistical plan should be described and justified in the protocol and/or in the final report as appropriate

Not applicable.

2.37 Subject analysis

Please insert a statement that the investigator(s) will permit study related monitoring, audits, REC review, and regulatory inspection(s), providing direct access to source data/documents*

The investigators will permit study related monitoring, audits, REC review, and regulatory inspection(s), providing direct access to source data/documents.

* Countersignature of this research protocol by the sponsor (or on behalf of the sponsor) indicates sponsor agreement with the above statement

2.38 Data storage

Please insert details of procedures for data handling, record keeping and archiving arrangements both during and post study, including location of data, accessibility rights and security provisions

All data, both electronic and paper, will be held in secure (locked / password protected as appropriate) facilities within the Vaccine Institute under the provisions of the SGUL Data Protection Act registration and local governance requirements. No archiving of data will be undertaken once the study has completed, and a study report and any publication in the literature completed.

2.39 Finance and insurance arrangements

Please insert details of source of finance for study and provide details of insurance arrangements/provider to cover the study

Finance will be provided by a grant from the European Commission Seventh Framework Programme through the EURONEUT41 consortium of which St George's is a full partner (grant reference REE0024).

Indemnity will be provided by the St George's University of London clinical studies and trials insurance.

2.40 Publication rights

Please insert details of publication policy (e.g. ownership of data). Please note, if Trust/SGUL are sponsoring the research then the ownership rights of intellectual property lie with the institution

Publication of data will be in accordance with the provisions of the Consortium Agreement entered into between St George's and the EURONEUT-41 Consortium funding this project.

2 Supplementary Information

2.41 Literature

Please provide reference to any literature and data that are relevant to the study and that provide background for the study

1. Castelo-Branco LR, Griffin GE, Poulton TA, Dougan G, Lewis DJ. Characterization of the circulating T-cell response after oral immunization of human volunteers with cholera toxin B subunit. *Vaccine* 1994;12:65-72.
2. Cosgrove CA, Castelo-Branco LR, Hussell T, et al. Boosting of cellular immunity against Mycobacterium tuberculosis and modulation of skin cytokine responses in healthy human volunteers by Mycobacterium bovis BCG strain Moreau Rio de Janeiro oral vaccine. *Infect Immun* 2006;74:2449-52.
3. Huo Z, Sinha R, McNeela EA, et al. Induction of protective serum meningococcal bactericidal and diphtheria-neutralizing antibodies and mucosal immunoglobulin A in volunteers by nasal insufflations of the Neisseria meningitidis serogroup C polysaccharide-CRM197 conjugate vaccine mixed with chitosan. *Infect Immun* 2005;73:8256-65.
4. Lewis DJ, Novotny P, Dougan G, Griffin GE. The early cellular and humoral immune response to primary and booster oral immunization with cholera toxin B subunit. *Eur J Immunol* 1991;21:2087-94.
5. Holmgren J, Czerkinsky C. Mucosal immunity and vaccines. *Nat Med* 2005;11:S45-53.
6. Mestecky J, Moldoveanu Z, Russell MW. Immunologic uniqueness of the genital tract: challenge for vaccine development. *Am J Reprod Immunol* 2005;53:208-14.
7. Frati F, Moingeon P, Marcucci F, et al. Mucosal immunization application to allergic disease: sublingual immunotherapy. *Allergy Asthma Proc* 2007;28:35-9.
8. Wu HY. Induction of mucosal tolerance in SLE: a sniff or a sip away from ameliorating lupus? *Clin Immunol* 2009;130:111-22.
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10. Cuburu N, Kweon MN, Song JH, et al. Sublingual immunization induces broad-based systemic and mucosal immune responses in mice. *Vaccine* 2007;25:8598-610.
11. Nardelli-Haeffiger D, Wirthner D, Schiller JT, et al. Specific antibody levels at the cervix during the menstrual cycle of women vaccinated with human papillomavirus 16 virus-like particles. *J Natl Cancer Inst* 2003;95:1128-37.
12. Jit M, Vyse A, Borrow R, Pebody R, Soldan K, Miller E. Prevalence of human papillomavirus antibodies in young female subjects in England. *Br J Cancer* 2007;97:989-91.
13. D'Souza G, Agrawal Y, Halpern J, Bodison S, Gillison ML. Oral Sexual Behaviors Associated with Prevalent Oral Human Papillomavirus Infection. *J Infect Dis* 2009;199:1263-9.
14. Launay O, Sadorge C, Jolly N, et al. Safety and immunogenicity of SC599, an oral live attenuated Shigella dysenteriae type-1 vaccine in healthy volunteers: Results of a Phase 2, randomized, double-blind placebo-controlled trial. *Vaccine* 2009;27:1184-91.
15. Mills KH, Cosgrove C, McNeela EA, et al. Protective levels of diphtheria-neutralizing antibody induced in healthy volunteers by unilateral priming-boosting intranasal immunization associated with restricted ipsilateral mucosal secretory immunoglobulin a. *Infect Immun* 2003;71:726-32.
16. Sadorge C, Ndiaye A, Beveridge N, et al. Phase 1 clinical trial of live attenuated Shigella dysenteriae type-1 DeltaicsA Deltaent Deltaefp DeltastxA:HgR oral vaccine SC599 in healthy human adult volunteers. *Vaccine* 2008;26:978-87.
17. Joura EA, Kjaer SK, Wheeler CM, et al. HPV antibody levels and clinical efficacy following administration of a prophylactic quadrivalent HPV vaccine. *Vaccine* 2008;26:6844-51.

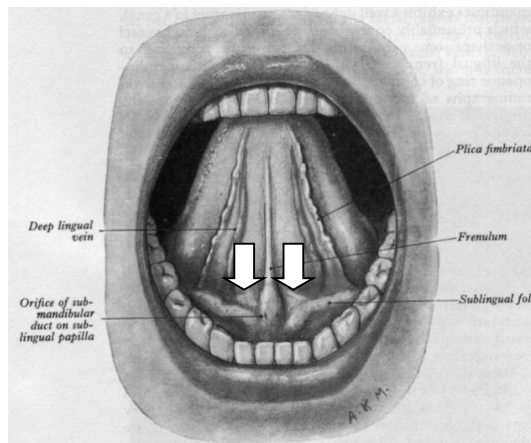
2.42 Supplementary details

Please insert any further relevant information to this research protocol (e.g. common toxicity criteria, tools to be used in study)

- Appendix 1: Standard Operating Procedure for sublingual application of Gardasil challenge agent
- Appendix 2: Characteristics of staff delegated to administer challenge agents
- Appendix 3: Summary of Product Characteristics for Gardasil

Appendix 1: Standard Operating Procedure for sublingual application of Gardasil

1. Subjects must fast for 1 hour prior to challenge, but may take water freely.
2. Operator to wear gloves and plastic apron. Masks and eye protection not required.
3. One 0.5 mL pre-filled syringe of Gardasil is opened, needle not attached, and placed conveniently to hand.
4. Subjects sit in a dental chair in an upright position with the head in a level position and supported by headrest.
5. Subjects rinse the mouth with bottled water and expectorate.
6. Absorbent pads specifically designed for the purpose (e.g. "Molnlycke 'Dry Tips' small) applied over parotid duct openings bilaterally to absorb parotid saliva flow.
7. Tongue raised by the subject to reveal the sublingual area which is gently dried by brief application of a cotton swab without aggravating saliva flow from submandibular and sublingual glands.
8. The complete contents of the Gardasil syringe immediately (without delay to prevent saliva accumulation) dispensed to the under surface of the tongue in the area behind the sublingual fold bilaterally (white arrows on diagram below), as gentle, individual drops (not squirted) with the syringe held just above the surface and not touching the surface of the mouth. Drops to be as carefully as possible contained within the area behind the sublingual fold.



9. Subject lowers tongue and holds in gentle opposition to the floor of the mouth, trapping the contents as much as possible under the tongue. Subject to avoid swallowing. This position to be held for 15 minutes.
10. After 15 minutes has passed, cotton pads over parotid ducts removed.
11. Subject to remain in clinical site for further 30 minutes without taking anything orally.
12. Subject allowed to leave clinical site and asked to fast, including fluids, for 1 hour.

Appendix 2: Characteristics of staff delegated to administer challenge agents

(taken from Department of Health “Patient Group Direction (PGD) for the supply and administration of human papillomavirus vaccine”)

Qualifications required

- Registered healthcare professional/nurse with current Nursing and Midwifery Council registration level 1 or 2.
- Assessed as competent to work with this patient group direction.

Additional requirements

- Will have undertaken training in the role, care and administration of the medicine specified in the PGD.
- Must be competent in the recognition and management of anaphylaxis.
- Must have access to a current copy of the BNF and Immunisation against infectious disease (‘Green book’) and comply with its recommendations (available on DH website – www.dh.gov.uk/greenbook).
- Must have access to all relevant DH advice, including the relevant CMO letters

or

- training and competent in all aspects of immunisation including contraindications and recognition and treatment of anaphylaxis.

Continued training requirements

- Annual attendance at the PCT’s or workplace update on resuscitation skills and the management of anaphylaxis within the community.
- Maintenance of own level of updating with evidence of continued professional development (PREP requirements)

or

- regular updates in immunisation, vaccination, anaphylaxis and cardiopulmonary resuscitation

or

- to reinforce and update knowledge and skills in this area of practice, including basic resuscitation and anaphylaxis training, with particular reference to changes and national directives.