Analysis of Mutagenic Potential of Therapeutic Vaccine Based on BPV-1 E6 Recombinant Protein Combined with Different Adjuvants

Rodrigo Pinheiro Araldi¹,², Diego Grando Módolo¹, Jacqueline Mazzuchelli de Souza¹,², Rodrigo Franco de Carvalho⁰, Lucinéia dos Santos³, Isabel Cristina Cherici Camargo³, Edislane Barreiros de Souza⁴ and Rita de Cassia Stocco⁵

¹Genetics Laboratory, Butantan Institute, São Paulo-SP, Brazil
²Biotechnology Interunit Post-Graduation Program, Biomedical Science Institute, University of São Paulo (USP), São Paulo-SP, Brazil
³Genetics Laboratory, Faculty of Sciences and Languages, São Paulo State University (UNESP), Assis-SP, Brazil

Abstract

Bovine papillomavirus is a worldwide distributed virus that affects at least 60% of Brazilian cattle herd, causing significant economic losses. Considering the great number of BPV-infected bovines, therapeutic vaccines, such those based on E6 protein, are mandatory to control the BPV-related diseases. However, the choice of the adjuvant remains challenging the vaccinology field, since adjuvants are associated with local and systemic reactions. Thus, this study analyzed the cytotoxic and mutagenic potential of BPV-1 E6 protein-based vaccine formulations using different adjuvants: aluminum hydroxide, complete and incomplete Freund’s adjuvants and two anti-oxidant saponin-rich extracts obtained from Agave sisalana - alcoholic and acid hydrolysis extract. Results of these analyzes suggest that acid hydrolysis extract from A. sisalana is an alternative and useful candidate for therapeutic vaccines, being able to reduce the mutagenic potential of antigen.

Keywords: Papillomavirus; Vaccine; E6 protein; Adjuvants; Saponin

Introduction

Vaccines are considered one of the most important medical advances of the last 200 years, reducing the infectious agent-associated deaths [1,2]. Currently, veterinary vaccines are responsible for 23% of all vaccine market [3]. In this sense, with the biotechnological advances, the interest in veterinary vaccines has grown, especially because animal vaccines have a less stringent regulatory and preclinical trials requirement [3]. However, this fact does not dispense the need to evaluate the biosafety of these products through in vitro and in vivo methods.

Bovine papillomavirus (BPV) is the etiological agent of bovine papillomatosis (BP), infectious and neoplastic disease, characterized by the presence of multiple papillomas that can regress spontaneously or persists, leading to malignancies, including urinary bladder and upper digestive carcinomas [4]. BPV is a worldwide distributed virus that causes significant economic losses, especially for Brazil that has the second largest cattle herd in the world, with 215 million of bovines [5]. According to epidemiological data, at least 60% of Brazilian cattle are infected by the BPV [6]. However, this number can be higher, since the viral infection can be asymptomatic [7,8].

Currently, there are few treatment methods against BPV available. Among them are: the papilloma surgical excision [9], self-haemotherapy [10] and control of ectoparasites [11]. However, these methods are few effective, especially for large bovine herds, as verified in Brazil. In this sense, both prophylactic and therapeutic vaccines against BPV are mandatory to control the BPV-related diseases.

The idea to develop a vaccine against BPV had begun in 1940 decade [12]. Since then, different vaccine models were proposed in literature [13-17], however, none of them became a commercial product [5]. Currently, our group demonstrated the efficacy and biosafety of BPV-1 L1 capsomers and virus-like particles (VLPs) produced in Escherichia coli as a candidate for prophylactic vaccine [18]. In contrast to prophylactic vaccines, which stimulates the neutralizing antibodies production, therapeutic vaccines are used to stimulate the cellular immune response by the activation of antigen presenting cells (APCs), CD4+ helper and CD8+ cytotoxic T-lymphocytes (CTLs) [19-21]. For this reason, therapeutic vaccines can eliminate the infection and prevent the cancer progression [22]. In this sense, studies based on HPV have been demonstrated that E6 protein is a useful candidate as antigen for therapeutic vaccines (Table 1).

Based on this data, our group cloned, expressed and purified the BPV-1 E6 protein as candidate to therapeutic vaccine [14]. However, we verified that the BPV-1 E6 protein alone is able to induce DNA damages, neosis [23] and oxidative stress in BPV-free epithelial cells [24]. These data put in check the biosafety of the BPV-1 E6 protein as antigen. Despite that, the capability to eradicate the viral infection, as well as the antigenicity of the protein makes it an important therapeutic vaccine candidate. For this reason, to explore vaccine formulations, evaluating the use of different adjuvants, can be an alternative to improve the biosafety of final products.

Different adjuvants were developed, including emulsions, mineral salt, surfactants, bacterial-derived adjuvants, liposomes, polymeric microspheres and cytokines [25]. Despite this, few adjuvants are used in vaccine formulations due to their toxicity, that can lead to unwanted reactions [26-28]. These reactions can be occur at local level, resulting in pain, inflammation, necrosis, swelling, granuloma, ulcer and sterile abscess formation or, at systemic levels, leading to nausea, fever, uveitis, eosinophilia, allergy, anaphylaxis, adjuvant arthritis and organ-specific toxicity [25].

*Corresponding author: Rita de Cassia Stocco, Genetics Laboratory, Butantan Institute, 1500, Vital Brazil Avenue, Butantã, São Paulo-SP, 05590-000 Brazil, Fax: +55 11 2627-9701; Tel: +55 11 2627-9701; E-mail: rita.stocco@butantan.gov.br

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Ideally, adjuvants must be safety, stable, biodegradable, immunologically inert and able to produce an appropriated immune response [26,28]. On the one hand the choice of adjuvant can be a challenge for the vaccine success [29], on the other hand, it reduces the antigen quantity and can improve the biosafety of final formulation. For this reason, the discovery of novel adjuvants has aroused a great research interest in vaccinology.

In this context, the saponins (Quill A and QS-21) and semi-synthetic triterpene glycosides from naturally occurring saponins (GPI-0100) are promising adjuvant candidates, since they can elicit a Type I helper T cell response for those diseases in which a cytotoxic T lymphocyte (CTL) response is desired [30-34]. Beside this, the antioxidant property [35-37] make the saponins useful candidate as adjuvant, especially for BPV-1 E6 protein-based vaccines, which antigen (E6 protein) has an oxidant action [24,38,39].

Saponins can be obtained from many different plant species, including Agave sisalana Perrine, popularly known as sisal [40,41]. Currently, Brazil is the world’s largest producer of A. sisalana for the supply of the sisal fiber [41]. However, only four percent of sisal leaves are used for fiber production [42]. The excessive waste of material (mucilage and sisal juice), which comprises 95% of plant byproduct, are generally discarded in the soil [42-44], making the sisal juice an unexplored source of biomolecules with pharmacological interest, including saponins.

However, natural products are not necessary safe [44-47]. Therefore, the safety and efficacy should be evaluated to ensure the best conditions of use [44]. Among the tests recommended to determine the biosafety of drugs candidate to be licensed are the micronucleus test and comet assay [45]. Although commonly used due its high statistical potential, the micronucleus test is less sensitivity than comet assay [45,48,49]. The obtained product was hydrolyzed with HCl and 500 mM NaCl, pH 8.0) for eight hours at 4°C under constant agitation. The BPV-1 E6 recombinant oncoprotein identity was verified by Western blot and mass spectrometry. Results of these analyses confirmed the protein identity, as reported by Araldi et al. [23].

### In silico prediction of BPV-1 E6 protein antigenicity based on bovine MHC-1

The BPV-1 E6 recombinant oncoprotein antigenicity was predicted by in silico analysis. For this, the epitopes for bovine major histocompatibility complex I (MHCI-1) alleles of BPV-1 E6 protein sequence (MDLKPFLARTNPSGIDLCWCREPLSEVDACMVKDFHVI-REGCRYGACTTCLENECLATERLWQGVPVTGREAELHLGTKL-DRILCRCYCGKLTNEHRHVFNFPEEKTANIRGCGYCD-CRHGSRKSY), available in Mazzuchelli-de-Souza et al. [14], were identified using the software NetMHC 4.0 Server (http://www.cbs.dtu.dk/services/NetMHC/). In this analysis, it was identified peptides sequences with 8-11 mers for bovine leucocyte antigen (BoLA) complex (BoLA-D18.4, BoLA-HD6, BoLA-JSP.1, BoLA-T2a, BoLA-T2b and BoLA-T2c). The threshold Rank for strong and weak binders was 0.5 and 2, respectively. The GraphPad Prism 5 (GraphPad Software Inc., USA) software was used for graphical visualization of the results.

### Obtaining the sisaal extracts used as adjuvant

The two saponin-rich extracts used as adjuvant were obtained from Agave sisalana Perrine. Fresh leaves were collected on a sisal farm located in Valente, in the state of Bahia (Brazil). Leaves were dried at 50°C using electric drier and crushed with the aid of a mechanical grinder to the powder form. For this step, 10 mL of purified protein were dialyzed against two liters of dialysis buffer (20 mM Tris-Cl and 500 mM NaCl, pH 8.0) for eight hours at 4°C under constant agitation. The obtained product was hydrolyzed with 2N HCl for four hours, under agitation. The precipitated was separated from acid solution by filtration at room temperature. Both EEAS and AHEAS were kept at 4ºC until use.

### Material and Methods

#### BPV-1 E6 recombinant oncoprotein expression and purification

BPV-1 E6 recombinant oncoprotein was expressed in Escherichia coli BL21, according to Mazzuchelli-de-Souza et al. [14]. The oncoprotein was subjected to dialysis to remove urea and imidazol, used during the BPV-1 E6 recombinant oncoprotein production. Dialysis was performed using Slide-A-Lyzer Dialysis Cassette (3K-12 ml) (Thermo Scientific, Carlsbad, USA). For this step, 10 mL of purified protein were dialyzed against two liters of dialysis buffer (20 mM Tris-Cl and 500 mM NaCl, pH 8.0) for eight hours at 4°C under constant agitation. The BPV-1 E6 recombinant oncoprotein identity was verified by Western blot and mass spectrometry. Results of these analyses confirmed the protein identity, as reported by Araldi et al. [23].

### Table 1: Vaccines based on HPV E6 protein. Previous studies employing the HPV E6 protein as antigen for therapeutic vaccine.

<table>
<thead>
<tr>
<th>Vaccines models</th>
<th>Adjuvant</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>- recombinant vaccinia virus expressing E6 and E7 proteins of HPV-16 and 18 (TA-HPV)</td>
<td>-</td>
<td>Borysiwacz et al. [68]</td>
</tr>
<tr>
<td>- recombinant vaccinia virus expressing either HPV-16 E6 and E7 proteins</td>
<td>-</td>
<td>Meneguzzi et al. [69]</td>
</tr>
<tr>
<td>- long peptide of HPV-16 E6 and E7 proteins</td>
<td>Montanide ISA-51</td>
<td>Welters et al. [71]</td>
</tr>
<tr>
<td>- long peptide of HPV-16 E6 alone and combined with E7</td>
<td>Saponin GPI-0100</td>
<td>Peng et al. [76]</td>
</tr>
<tr>
<td>- vaccination with 9-amino acid of HPV-16 E7 protein</td>
<td>IFA</td>
<td>Mederspach et al. [73]</td>
</tr>
<tr>
<td>- E6 recombinant protein</td>
<td>-</td>
<td>Lin et al. [74]</td>
</tr>
<tr>
<td>- HPV L2, E6 and E7 single fusion protein (TA-CIN)</td>
<td>-</td>
<td>Jons et al. [75]</td>
</tr>
<tr>
<td>- HPV-16 E6, E7 and L2 fusion protein (TA-CIN)</td>
<td>Saponin GPI-0100</td>
<td>Peng et al. [76]</td>
</tr>
<tr>
<td>- DNA vaccine encoding calreticulin linked to E6 protein (CRT/E6)</td>
<td>Peng et al. [76]</td>
<td>Trimmel et al. [77]</td>
</tr>
<tr>
<td>- DNA vaccine targeting HPV-16 and 18 E6 and E7 proteins (VGX-3100)</td>
<td>-</td>
<td>Morrow et al. [78]</td>
</tr>
<tr>
<td>- DNA vaccine encoding HPV-16 E6/E7 fusion antigen</td>
<td>Yan et al. [79]</td>
<td></td>
</tr>
<tr>
<td>- DNA vaccine encoding a single chain trimer of MHC-I linked to HPV-16 E6</td>
<td>Huang et al. [80]</td>
<td></td>
</tr>
<tr>
<td>- HPV-16 uploaded in engineered exomes</td>
<td>ISCOMATRIX</td>
<td>Manfredi et al. [59]</td>
</tr>
</tbody>
</table>

For this reason, this study analyzed the mutagenic potential of BPV-1 E6 recombinant oncoprotein using different adjuvants. The micronucleus test is less sensitivity than comet assay [45,48,49]. Although commonly used due its high statistical potential, the micronucleus test is less sensitivity than comet assay [45,48,49]. For this reason, this study analyzed the mutagenic potential of BPV-1 E6 protein-based therapeutic vaccines using different adjuvants (aluminum hydroxide, complete and incomplete Freund’s adjuvant, ethanolic and acid hydrolysis extracts obtained from A. sisalana) by the micronucleus test, comet and histone γ-H2AX.
Determination of *A. sisalana* dried extract saponin concentration

The saponin concentration was determined by spectrophotometric analysis, according to the literature [50,51]. The *A. sisalana* in powder form was dissolved in distilled water at three concentrations: 0.25, 0.35 and 0.50 mg/mL. An aliquot of 1.0 mL of each solution was incubated for 20 minutes with 1.0 mL of 0.2% cobalt chloride chromogenic reagent and 1.0 mL concentrated sulphuric acid. These solutions were analysed in UV-Vis spectrophotometer at 284 nm. A solution of commercial saponin (Merck, Germany) at 0.2 mg/mL was used as control. The saponin concentration was expressed in: 1) mg/mL for saponins in extract solution and 2) mg/g for saponins in dried extract. These concentrations were obtained from the linear regression curve with the control (saponin, Merck).

Adjuvants and vaccine formulations

The aluminum hydroxide (Alum), complete and incomplete Freund’s adjuvants (CFA and IFA, respectively) were obtained from Sigma (Germany). The *A. sisalana* extracts EEAS and AHEAS were diluted in PBS with 2% Tween 80 (Sigma, Germany) at a final concentration of 50 µg/mL. The cytotoxic and mutagenic potential of these adjuvants were analyzed individually, as well as in association with the antigen (dialyzed BPV-1 E6 recombinant oncoprotein) at a final concentration of 1.0 µg/mL, as proposed by Araldi et al. [23]. The different formulations tested are shown in Table 2.

Cell culture

Both cytotoxic and mutagenic tests were performed in *vitro*, using the CRIB cell line, which was already used to determine the mutagenic potential of BPV-1 E6 recombinant oncoprotein [23]. CRIB cells were extended in culture flasks of 25 cm², containing 5.0 mL of complete medium (Eagle's minimal essential medium - MEM, supplemented with 10% of fetal bovine serum) (Cultilab, Brazil) as proposed by Flores and Donis [52]. All analysis was performed without antibiotic supplementation, since the antibiotics can lead to DNA damages, resulting in false-positive results [53].

Annexin V-PI assay

The annexin V-PI assay was used to analyze the cytotoxic potential of antigen, adjuvants and vaccine formulations. Cells were expanded in culture flasks of 25 cm² with 5.0 mL of complete medium until a confluence of 70%. Cells were subjected to different treatments (Table 2) for 48 hours at 37°C, time necessary for two duplication rounds. After, the medium was transferred for Falcon tube of 15.0 mL and the cells and incubated with 2.0 mL of trypsin/EDTA solution (Cultilab, Brazil) for five minutes at 37°C to promote the monolayer disaggregation. Cell suspension was transferred to Falcon tube containing the medium removed and centrifuged at 200 x g for five minutes, discarding the supernatant. The pellet was homogenized with 1.0 mL of cold sterile PBS and transferred for polypropylene tube of 1.5 mL. Cells were centrifuged at 200 x g for five minutes, discarding the supernatant. Cells were homogenized with 100 µL of binding buffer and incubated with 5.0 µL of Annexin V-FITC and 5.0 µL of propidium iodide (PI) (Quatro G, Brazil) for 15 minutes at room temperature. Cells were homogenized with 100 µL of binding buffer and centrifuged at 200 x g for five minutes, discarding the supernatant. Cells were homogenized with 100 µL of binding buffer and centrifuged at 200 x g for five minutes, discarding the supernatant. Cells were homogenized with 100 µL of binding buffer and incubated with 5.0 µL of Annexin V-FITC and 5.0 µL of propidium iodide (PI) (Quatro G, Brazil) for 15 minutes at room temperature. Cells were homogenized with 100 µL of binding buffer and centrifuged at 200 x g for five minutes, discarding the supernatant. Cells were homogenized with 100 µL of binding buffer and incubated with 5.0 µL of Annexin V-FITC and 5.0 µL of propidium iodide (PI) (Quatro G, Brazil) for 15 minutes at room temperature. Cells were homogenized with 100 µL of binding buffer and centrifuged at 200 x g for five minutes, discarding the supernatant. Cells were homogenized with 100 µL of binding buffer and incubated with 5.0 µL of Annexin V-FITC and 5.0 µL of propidium iodide (PI) (Quatro G, Brazil) for 15 minutes at room temperature. Cells were homogenized with 100 µL of binding buffer and centrifuged at 200 x g for five minutes, discarding the supernatant. Cells were homogenized with 100 µL of binding buffer and incubated with 5.0 µL of Annexin V-FITC and 5.0 µL of propidium iodide (PI) (Quatro G, Brazil) for 15 minutes at room temperature.

**Table 2: Drugs tested, Concentration, Abbreviation**

<table>
<thead>
<tr>
<th>Drugs tested</th>
<th>Concentration</th>
<th>Abbreviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Negative control</td>
<td>-</td>
<td>Control</td>
</tr>
<tr>
<td>A. Sisalana extracts diluent (PBS + 2% Tween 80)</td>
<td>5 µL/mL</td>
<td>T80</td>
</tr>
<tr>
<td>Positive control (cyclophosphamide)</td>
<td>50 µg/mL</td>
<td>C+</td>
</tr>
<tr>
<td>Positive control (H₂O₂)</td>
<td>100 µM</td>
<td>H₂O₂</td>
</tr>
<tr>
<td>BPV-1 E6 recombinant protein</td>
<td>1 µg/mL</td>
<td>E6</td>
</tr>
<tr>
<td>Aluminium hydroxide</td>
<td>5 µL/mL</td>
<td>Alum</td>
</tr>
<tr>
<td>Complete Freund's adjuvant</td>
<td>5 µL/mL</td>
<td>CFA</td>
</tr>
<tr>
<td>Incomplete Freund's adjuvant</td>
<td>50 µg/mL</td>
<td>IFA</td>
</tr>
<tr>
<td>Ethanolic extract of <em>A. sisalana</em></td>
<td>50 µg/mL</td>
<td>EEAS</td>
</tr>
<tr>
<td>Acid hydrolysis extract of <em>A. sisalana</em></td>
<td>50 µg/mL</td>
<td>AHEAS</td>
</tr>
<tr>
<td>BPV-1 E6 recombinant protein + aluminium hydroxide</td>
<td>1 µg/mL + 50 µg/mL</td>
<td>E6 + Alum</td>
</tr>
<tr>
<td>BPV-1 E6 recombinant protein + complete Freund's adjuvant</td>
<td>1 µg/mL + 50 µg/mL</td>
<td>E6 + CFA</td>
</tr>
<tr>
<td>BPV-1 E6 recombinant protein + incomplete Freund's adjuvant</td>
<td>1 µg/mL + 50 µg/mL</td>
<td>E6 + IFA</td>
</tr>
<tr>
<td>BPV-1 E6 recombinant protein + ethanolic extract of <em>A. sisalana</em></td>
<td>1 µg/mL + 50 µg/mL</td>
<td>E6 + EEAS</td>
</tr>
<tr>
<td>BPV-1 E6 recombinant protein + acid hydrolysis extract of <em>A. sisalana</em></td>
<td>1 µg/mL + 50 µg/mL</td>
<td>E6 + EHEAS</td>
</tr>
</tbody>
</table>

*Positive control employed in mutagenic analyses (micronucleus test and histone γH2AX)*

*Positive control employed in ROS production analysis using the DCF-DA assay diluted in PBS + 2% Tween 80.

ROS detection by DCF-DA assay

Considering that previous study showed that BPV-1 E6 promotes the oxidative stress [24], the reactive oxygen species (ROS) generation was determined using the dichlorofluorescein diacetate (DCF-DA) (Sigma, Germany), according to the de-Sá-Júnior et al. [46]. CRIB cells were expanded in culture flasks of 25 cm² with 5.0 mL of complete medium until a confluence of 80%. The medium was discarded by inversion and the cells were washed with 2.0 mL of PBS sterile at 37°C to remove death cells. A volume of 5.0 mL of complete medium was transferred to the culture flasks with 100 mM of DCF-DA diluted in dimethylsulfoxide (DMSO) (Merck, Germany). Cells were incubated at 37°C at 5% CO₂ for 30 minutes. Cells were washed with 2.0 mL of PBS sterile at 37°C. A volume of 5.0 mL of complete medium was transferred to the culture flasks and subjected to different treatments (Table 2) for 24 minutes at 37°C at 5% CO₂. The monolayer was disaggregated using 2.0 mL of trypsin/EDTA solution (Cultilab, Brazil). The cell pellet was homogenized with 1.0 mL of cold sterile PBS and transferred for
polypropylene tube of 1.5 mL. Cells were centrifuged at 200 x g for five minutes and homogenized in 400 μL of PBS. As positive control, it was employed 100 mM of hydrogen peroxide. Cells were analyzed in BD Accuri C6 flow cytometry (BD Biosciences, USA), using the FL1 channel in a total of 10,000 events. This analysis was performed in triplicate.

**Micronucleus test**

To verify the genotoxic potential of antigen, adjuvants and vaccine formulations (Table 2), it was performed the micronucleus test. A total of 1 x 10⁵ CRIB cell was transferred to six-well plate, containing a sterile coverslip of 24 x 24 mm with 2 mL of MEM medium, supplemented with 10% of fetal bovine serum. After one hour, cultures were subjected to different treatments (Table 2), remaining incubated for 48 hours, time necessary to two replication cycles. After this time, medium was removed, and the cells were washed three times with PBS for five minutes and fixed with methanol for 30 minutes. Cells were stained with solution 1:4 Giemsa-PBS for 3 minutes and, after, washed twice with PBS. Coverslips containing the biological material were mounted on slides using Entellan (Merck, Germany). Slides were analyzed in a fluorescence microscope (Carl Zeiss, Germany) to observe the frequency of micronucleated cells in a total of 1,000 cells, according to Araldi et al. [53]. As negative control, it was employed cell without any drug and, as positive control, cells were treated with 50 μg/mL of saponin. As positive control, cells were treated with 50 μg/mL of to Araldi et al. [53]. As negative control, it was employed cell without any drug and, as positive control, cells were treated with 50 μg/mL of saponin. As positive control, cells were treated with 50 μg/mL of saponin. As positive control, cells were treated with 50 μg/mL of saponin. As positive control, cells were treated with 50 μg/mL of saponin. As positive control, cells were treated with 50 μg/mL of saponin. As positive control, cells were treated with 50 μg/mL of saponin.

**Histone γ-H2AX assay**

The histone H2AX variant is unique in eukaryotes due to its carboxyl tail, that include a high conserved sequence, comprised by one serine residue at position 139, which is phosphorylated in the presence of DNA double strand-breaks (DSBs) [54]. The histone H2AX (p Ser139) is also known as γ-H2AX [54]. Thus, the immunodetection of this histone is recognized as a hallmark of DSBs and, therefore, clastogenesis [55-57]. Currently studies show that the histone γ-H2AX assay is 100 times more sensitivity than comet assay [54,58]. For this reason, this method was additionally performed to evaluate the clastogenic potential of therapeutic vaccines.

CRIB cells were expanded in culture flasks of 25 cm² with 5 mL of complete medium until a confluence of 60-70%. Cells were incubated with different tested drugs (Table 2) for 24 hours at 37°C. Cells were subjected to monolayer disaggregation with 2.0 mL of trypsin/EDTA solution (Cultilab, Brazil). Cells were transferred to 1.5 mL polypropylene tubes and fixed in 1.0 mL of 1.0% formalin solution at 4°C for 2 hours. Cell suspension was centrifuged and washed twice with 1.0 mL of PBS at 4°C to remove the formalin residues. Cells were incubated with 1.0% BSA at 4°C for 20 minutes, washed with 200 μL of PBS, and incubated overnight at 4°C with the polyclonal anti histone γ-H2AX (p Ser139) antibody produced in rabbit (Novusbio, USA) at a dilution of 1:200 in PBS with 0.01% Triton X-100 (Sigma, Germany). Cells were centrifuged under described conditions, washed twice with PBS and incubated at 4°C for two hours with anti-rabbit conjugated with Alexa Fluor 488 secondary antibody (Invitrogen, USA) at 1:2000 dilution in 1% BSA. Next, cells were washed with PBS, centrifuged and homogenized in 100 μL of PBS. Cells were analyzed in BD Accuri C6 cytometer (BD Bioscience, USA), employing the FL1 channel. A total of 10,000 events were analyzed. This analysis was performed in triplicate. Results were analyzed in FlowJo software (TreeStar, USA), using the mean of percentage of immune-labeled cells. CRIB cells incubated exclusively with secondary antibody and non-incubated with any primary, neither secondary antibodies were used as controls.

**Statistical analyses**

Statistical analyses of annexin V-PI, DCF-DA and histone γ-H2AX assays were performed based on ANOVA followed by Tukey’s multiple comparison tests. Statistical analyses of micronucleus test were performed by non-parametric methods, using the Friedman test followed by the Dunn post-hoc test. These analyses were performed using the GraphPad Prism version 5 software (GraphPad Software Inc., USA), with 5% of significance level.

**Results**

**Saponin concentration of Agave sisalana dry extract**

Based on the linear regression curve (R²=0.9963) of absorbance values of spectrophotometric analysis of commercial saponin (Merck), it was obtained the following equation: A=1.984[s] + 0.0995 (A - absorbance and [s] - saponin concentration in mg/mL), which was used to determine the saponin concentration in dry extract of A. sisalana. Results of this analysis are showed a concentration of 908.4 mg saponins/g of dry extract, indicating that saponins represent 90.84% of weight of the dry extract (Table 3).

**Cytotoxic potential analysis of adjuvants and vaccine formulations**

Results of annexin V-PI assay (Figure 1A-C and Table 4), employed to analyze the cytotoxicity, showed significant statistical differences among the different vaccine formulations (F=1446, R²=0.9986 and p<0.0001). Based on this result, it was performed the Tukey’s multiple comparison test, which showed that both the A. sisalana extracts (EEAS and AHEAS), as well as the vaccine formulations using these extracts as adjuvants (E6+EEAS and E6+AHEAS) are cytotoxic in relation to control (Figure 1 and Table 4). A similar cytotoxicity was verified in cell treated with the A. sisalana extracts diluent (T80) (Figure 1B and Table 4). These data were reinforced by the statistical analysis based on the live cell percentage, that pointed out that the A. sisalana extracts (EEAS and AHEAS), the vaccines formulations based on these saponin-rich extracts (E6+EEAS and E6+AHEAS) and the diluent (T80) showed a cytotoxic potential (Figure 1C and Table 4). By the contrast, the Alum, CFA, IFA and the vaccine formulations based on these adjuvants did not show significant statistical differences when compared with the control (Figure 1C and Table 4), indicating that these adjuvants are safe.

**ROS production**

Based on the ROS-positive cell percentage (Figure 2 and Table 4) it was performed an ANOVA analysis, that pointed out significant statistical differences among the formulations tests (F=13930, R²=0.9986 and p<0.0001). For this reason, it was performed the Tukey’s multiple comparison test. Result of this test showed that the BPV-1 E6 recombinant protein, the adjuvants CFA and IFA, as well as the diluent of A. sisalana extracts T80 increased the percentage of ROS-producing cells, suggesting an oxidative potential (Figure 2A and Table 4). Similar results were also verified in vaccine formulations E6+CFA and E6+IFA (Figure 2B and Table 4). By the contrast, the Alum, EEAS

<table>
<thead>
<tr>
<th>Concentration (mg/mL)</th>
<th>Absorbance</th>
<th>Saponin concentration (mg/mL)</th>
<th>Saponin mg/g</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25</td>
<td>0.537</td>
<td>0.22</td>
<td>882.68</td>
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<tr>
<td>0.35</td>
<td>0.737</td>
<td>0.32</td>
<td>918.01</td>
</tr>
<tr>
<td>0.5</td>
<td>1.017</td>
<td>0.46</td>
<td>924.52</td>
</tr>
</tbody>
</table>

*Results expressed in "mg" of saponins per "mL" of dry extract solutions of A. sisalana,* "results expressed in "mg" of saponins per "g" of dry extract of A. sisalana.

**Table 3**: Saponin concentration. Analysis of saponin content in A. sisalana.
Figure 1: Results of annexin V-PI assay: A) dot plots showing the live (Q2-LL), early (Q2-LR), late apoptotic (Q2-UR) and necrotic cell percentage (Q2-UL); B) histograms showing the mean of live (black), early and late (gray) and necrotic cell percentage (red) verified in triplicates, C) Statistical analysis based on the live cell percentage, showing the absence of cytotoxic potential of BPV-1 E6 recombinant oncoprotein, Alum, CFA, IFA, as well as the vaccine formulations based on these adjuvants. Results show that AHEAS and E6+AHEAS present a cytotoxic potential similar to the diluent (T80). The EEAS and E6+EEAS show the highest cytotoxic potential. Total of 10,000 events analyzed.

Figure 2: Results of DCF-DA assay: A) histograms showing the cell percentage labelled with the DCF-DA probe (indicating the ROS production); B) histograms showing the percentage of cells producing ROS; C) histograms of mean of fluorescence intensity (MFI) of cells producing ROS. Results show that BPV-1 E6 recombinant oncoprotein has a high oxidative action, being able to increase both the percentage and MFI of cells producing ROS. Both Freund's adjuvant (CFA and IFA) increased the percentage of cell producing ROS, but only CFA increased the MFI. Although the results show a MFI increase in cells treated with EEAS and AHEAS, it is verified a reduced percentage of ROS-producing cells. Moreover, the vaccine formulations with these extracts of A. sisalana were able to reduce the E6-induced oxidative stress, demonstrating the antioxidant action of these adjuvants. Total of 10,000 events analyzed.
and AHEAS reduced significantly the percentage of ROS-producing cells, suggesting an antioxidant potential (Figure 2B and Table 4). Similar results were also verified in vaccine formulations E6+Alum, E6+EEAS and E6+AHEAS (Figure 2B and Table 4).

**Micronucleus test**

Based on the number of micro-nucleated cells (Figure 3 and Table 5), it was performed the Friedman test, which pointed out significant statistical differences among the adjuvants, neither among the vaccine formulations using E6 alone (CFA, IFA and EEAS) as well as the vaccine formulations based on these adjuvants and formulations (Figure 2B and Table 4). However, the results did not show statistical differences among the adjuvants, neither among the vaccine formulations using the BPV-1 E6 recombinant protein in relation to negative control (Figure 3 and Table 5). This data suggest that the tested adjuvants can reduce the antigen genotoxicity.

**Expression levels of histone γH2AX**

Based on the mean of histone γH2AX-expressing cell percentage (Figure 4A, B and Table 5), it was performed the ANOVA analysis, which results pointed out significant statistical differences among the different formulations tested (Friedman statistic=38.28, p<0.0001). Considering this result, it was performed the Turkey's multiple comparison test, which results showed that BPV-1 E6 recombinant protein has a similar genotoxic potential to cyclophosphamide, employed as positive control (Figure 4 and Table 5). Moreover, the AHEAS reduced the clastogenic action of these adjuvants and formulations (Figure 4B and Table 5). These data suggest that the tested adjuvants can reduce the antigen genotoxicity.

**Discussion**

There are several techniques for the therapeutic vaccines production, but the vaccines based on E6 and E7 protein are the most promising [20]. In this sense, the E6 protein stands out for its stability and relative obtaining facility [14,59,60]. Moreover, the protein contains diverse epitopes for human leukocyte antigens (HLA - HPV E6) [61], as well as for bovine leukocyte antigens (BoLA - BPV E6) identified by *in silico* analysis [14]. For these reasons, since 2013 we are studying the BPV-1 recombinant protein as antigen candidate to therapeutic vaccine [14,23]. In order to confirm the antigenicity of BPV-1 E6 protein, we analyzed the protein binding-sites with bovine MHC-I alleles using *in silico* tools. Results of this analysis showed the antigenic potential of BPV-1 E6 recombinant oncoprotein (Figure 5), reinforcing its use as antigen for therapeutic vaccines.

However, previous analysis showed that BPV-1 E6 recombinant oncoprotein alone is able to induce DNA damages in CRIB cells and bovine lymphocytes [23,62]. We also verified that BPV-1 E6 recombinant oncoprotein promotes metabolic deregulations, resulting in ROS production [24] through a supposedly mechanism homologous to HPV-16 E6 oncoprotein [63].

Nevertheless, vaccines based on recombinant protein require the use of adjuvants to increase the immune response, reducing the antigen quantity in final vaccine formulation [26-28]. The addition of adjuvants can promote a better presentation for MHC class I, increasing the endogenous processing of the vaccine, and consequently the uptake by MHC class I, driving more efficiently to dendritic cells, which increase the presentation of MHC class I and the activation of CD8+ cells. In this context, previous study showed that saponin-based adjuvants can stimulate de CD8+ cells activation [64,65]. For this reason, the saponins emerge as additional candidate to adjuvants for E6-based protein vaccines.

Based on these data, we analyzed the cytotoxic and mutagenic potential of the most employed adjuvants (Alum, CFA and IFA) alone and combined with the BPV-1 E6 recombinant protein. Moreover, considering that the mutagenic potential BPV-1 E6 protein is related to their oxidant activity [24], we included two antioxidant saponin-rich extracts obtained from *A. sisalana*: EEAS and AHEAS.
The cytotoxic potential of antigen, adjuvants and vaccines was evaluated through the annexin V-PI assay. Results of this analysis pointed out that EEAS shows the highest cytotoxic potential, which was verified by the elevate number of necrotic cells (Figure 1B and Table 4). The cytotoxicity of EEAS can be attributed to the surfactant presence, that can bind to cholesterol present in cell membrane, leading to pore formation and hemolysis [27]. The hemolytic action of EEAS also explains the high percentage of necrotic cells verified in the vaccine formulation using this extract as adjuvant (E6+EEAS) (Figure 1B and Table 4). On the one side, the necrotic action of EEAS represents an unwanted effect, by the other side it suggests an unexplored antineoplastic action of ethanolic extract of A. sisalana.

Although the live cell percentage of AHEAS had showed a significant statistical difference in relation to control, it was not verified significant statistical differences between AHEAS/E6+AHEAS and the diluent (T80) (Figure 1C and Table 4). Considering that the acid hydrolysis reduces the hemolytic action of A. sisalana extract [40], the results suggest that the cytotoxicity verified in cells treated with AHEAS alone or in association with BPV-1 E6 recombinant protein can be attributed to the Tween 80, used as diluent. By the opposite, the other adjuvants analyzed (Alum, CFA and IFA) did not show cytotoxic potential in relation to control (Figure 1C and Table 4).

Although not cytotoxic, the Freund’s adjuvants (CFA and IFA) alone or combined with BPV-1 E6 recombinant oncoprotein increased the percentage of ROS-producing cells (Figure 2 and Table 4). This result is alarming, once the BPV-1 E6 recombinant protein has a
The genotoxic potential analysis showed a high frequency of micronuclei in CRIB cells treated with BPV-1 E6 recombinant protein (Figure 3), reinforcing the mutagenic potential of this protein, previous described in Araldi et al. [23]. The genotoxic activity can be attributed to the fork replication stress and the oxidant action of the BPV-1 E6 recombinant protein. The adjuvants analyzed did not show genotoxic activity when tested alone or combined with the antigen (Figure 3 and Table 5). These data suggest that the antigen genotoxic potential is directly related with the oxidant activity of BPV-1 E6 recombinant oncoprotein.

Results of flow cytometry showed that the treatment with BPV-1 E6 recombinant protein, CFA, IFA, EEAS, E6+CFA, E6+IFA and E6+EEAS promoted an expressive increase in γ-H2AX expression levels (Figure 4). Results of statistical analyses showed these treatments present a clastogenic potential similar to cyclophosphamide, an alkylating drug used as positive control (Figure 4 and Table 5). These data reinforce the antigen mutagenic potential, also verified by micronucleus test (Figure 3). In this sense, it was not verified significant statistical differences between AHEAS and E6+AHEAS and control (Figure 4 and Table 5). The antioxidant mechanisms involve direct inhibition of the ROS generation or the scavenging of the free radical. According to Dini et al. [66], the antioxidant mechanisms of prominent oxidant capability [24], which was also verified in this study (Figure 2 and Table 4). However, the Alum, EEAS and AHEAS reduced significantly the percentage of ROS-producing cells (Figure 2 and Table 4), indicating the antioxidant property of A. sisalana extracts. Interesting, the diluent of A. sisalana extracts (T80) showed to increase the ROS production in relation to control, but when added to EEAS and AHEAS, it was verified a reduction of cell percentage labelled by the probe, reinforcing the antioxidant capability of these extracts (Figure 2 and Table 4). The antioxidant activity of A. sisalana extracts can be justified by the flavonoids present in these extracts [41].

**Table 5:** Micronucleus test and histone H2AX assay. Results of mutagenic analyses (micronucleus test and histone γH2AX assay).

Results of flow cytometry showed that the treatment with BPV-1 E6 recombinant protein, CFA, IFA, EEAS, E6+CFA, E6+IFA and E6+EEAS promoted an expressive increase in γ-H2AX expression levels (Figure 4). Results of statistical analyses showed these treatments present a clastogenic potential similar to cyclophosphamide, an alkylating drug used as positive control (Figure 4 and Table 5). These data reinforce the antigen mutagenic potential, also verified by micronucleus test (Figure 3). In this sense, it was not verified significant statistical differences between AHEAS and E6+AHEAS and control (Figure 4 and Table 5). The antioxidant mechanisms involve direct inhibition of the ROS generation or the scavenging of the free radical. According to Dini et al. [66], the antioxidant mechanisms of
saponins involves both antiradical and reducing property, which is related to the metal ion chelating activity [67].

In summary, the results suggest that acid hydrolysis extract from A. sisalana (AHEAS) is a useful candidate as adjuvant for BPV E6 recombinant protein-based therapeutic vaccine, being able to increase the vaccine biosafety. Considering the functional homologies among the BPV and HPV E6 protein, and the antigenicity of these proteins, our results indicate that AHEAS could be a useful adjuvant candidate for human vaccines against HPV. In addition, the antiinociceptive and anti-inflammatory action of AHEAS [40], could also reduce the eventual local reactions related to the cytotoxicity. Moreover, considering that sisal juice is rich in saponin, but it is discarded as a residue of sisal fibre industry, to use this sisal juice as source of adjuvants emerges as an ecological and cheaper alternative.

Ethical Statement

This study was approved by Ethical Committee of Butantan Institute (Permission number 1319/14).

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Conflict of Interest

The authors declare that there is no conflict of interest regarding the publication of this paper.

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