Anticancer efficacy of perillyl alcohol-bearing PLGA microparticles

Mohammad Farazuddin¹, Bhawna Sharma², Aijaz Ahmed Khan³, Beenu Joshi², Mohammad Owais¹

¹Interdisciplinary Biotechnology Unit, Aligarh Muslim University, Aligarh, Uttar Pradesh, India; ²Immunology Division, NJIL and other Mycobacterial diseases, Agra-282001, Uttar Pradesh, India; ³Department of Anatomy, JN Medical College, Aligarh Muslim University, Aligarh-202002, Uttar Pradesh, India

Abstract: In the present study, a novel poly-lactic glycolic acid (PLGA)-based microparticle formulation of perillyl alcohol (POH) was prepared and characterized. Further, its efficacy was evaluated against di-methyl benzanthracene-induced skin papilloma in Swiss albino mice. The characterization studies showed that POH-bearing PLGA microparticles were of the size 768 ± 215 nm with a ζ-potential value of −7.56 ± 0.88 mV. The entrapment efficiency of the active drug in particles was 42.4% ± 3.5%. POH-bearing PLGA microparticles were stable and released entrapped drug gradually over an extended time period. The in vitro efficacy of POH-bearing PLGA microparticles was evaluated by examining their differential cytotoxicity and assessing their ability to inhibit epidermoid carcinoma cell line (A253). The POH-based microparticles when administered to tumor-bearing animals caused greater tumor regression and increased survival rate (∼80%) as compared with the group receiving free form of POH (survival rate 40%). The superiority of POH-PLGA microparticles over free form of POH was further evident from their ability to modulate apoptosis-regulating factors.

Keywords: poly-lactic glycolic acid, epidermoid cancer cells, skin papilloma, anticancer efficacy

Introduction

Cytotoxicity and other related side effects are the most serious problems associated with the currently available anticancer drugs. Other limitations include widespread systemic distribution and rapid elimination of the administered anticancer drugs from the host body. A worldwide search therefore continues for anticancer drugs that are more potent, less toxic, and manifest minimum untoward effects to the host. Several plant derived compounds have been reported to possess strong anticancer properties and have been shown to delay, inhibit, or reverse cancerous growth in an effective manner. For example, perillyl alcohol (POH), a plant-based compound, has been reported to possess strong anti-cytotoxic properties against several types of cancer including those of breast, pancreatic, and liver.¹⁻³ POH is a monoterpene and constituent of essential oils from a number of plants; namely, perilla (Perilla frutescens), lavendin, peppermint, ginger grass, savin, caraway, and celery seeds.⁴

Before translating the suitability of a novel compound like POH as a potential anticancer agent in the clinical setting, it is desirable to address some of the associated issues like that of its solubility, palatability, and sustained/controlled release in systemic circulation. This requires designing of a suitable drug-delivery system that can release the drug gradually over a long period of time and, in turn, facilitate its uptake by cancer cells and thereby helps in increasing the efficacy of the entrapped drug. Polymeric
microparticles offer a promising technology in this regard. Encapsulation of drugs in the core of microparticles have been shown not only to protect them from the external environment, enroute, but also help in increasing their plasma half-lives in systemic circulation thereby facilitating the attainment of optimum drug availability at the desired target.5–7

To develop promising and effective formulations of anticancer compounds, various polymeric matrices have been investigated and their efficacies worked out.5–10 Drug delivery systems, such as nanoparticle-,5 microemulsion-,6 nanoemulsion-,7 and liposome-based delivery systems,8–10 have been shown to enhance the efficacy of various compounds on systemic as well as topical applications. Among various drug-carrier systems, poly-lactic glycolic acid (PLGA) matrices have been reported to be nontoxic, biodegradable, and shown to release the entrapped drug gradually over a long duration.11

In the present study, POH-bearing PLGA microparticles were prepared and their efficacy against the skin epidermoid cancer cell line (A253) was evaluated. In addition, potential of POH-bearing PLGA microparticles was evaluated in treatment of di-methyl benzo anthracene (DMBA)-induced tumors in Swiss albino mice.

Materials and methods
All reagents used in the study were of highest purity available. PLGA, poly vinyl alcohol (PVA), and POH were purchased from Sigma Chemical Company (St Louis, MO). Dichloromethane (DCM) was of analytical grade of purity and procured locally. Anti-p53 mutant, anti-p53 wild-type (wt), anti-p21/waf1, anti-tubulin and anti-β-actin antibodies were purchased from BD Biosciences (San Diego, CA).

Preparation of POH-loaded PLGA microparticles
Microparticles used in the present study were prepared by oil-in-water-based emulsion solvent evaporation technique using the published protocol as standardized in our laboratory.12,13 Briefly, a known quantity of POH (30 mg) was dissolved in minimum volume of methanol, mixed with PLGA solution (190 mg PLGA dissolved in 1.0 mL DCM), and sonicated in a bath-type sonicator to form the primary emulsion. The primary emulsion was mixed with 100 mL of 10% PVA (w/v) and homogenized using a Silverson L4RT homogenizer (Silverson Machines, East Longmeadow, MA). The resulting oil-in-water emulsion was stirred at 25°C for 18 hours to allow solvent evaporation and formation of POH-entrapped microparticles. The microparticles were centrifuged and thoroughly washed with phosphate buffered saline (PBS) (0.15 M NaCl containing 20 mM sodium phosphate, pH 7.4) to remove surface adsorbed drug. The microformulation was lyophilized and finally stored at 4°C until further use.

Entrapment efficiency of POH in microparticles
Entrapment of POH in microparticles was assessed by dissolving an aliquot of the microparticles in 0.1 N NaOH followed by analysis of POH content by high-performance liquid chromatography (HPLC) following the published procedure.14 Briefly, 10 mg freeze-dried microparticles were dissolved in 1.0 mL of 0.1 N NaOH. The solution was vortexed for 10 minutes followed by centrifugation for 5 minutes at 9168 × g at 25°C. An aliquot (100 µL) of supernatant was mixed with 900 µL methanol. The suitable aliquots of the resultant homogenate solution, were analyzed by reversed phase HPLC using a Symmetry® C-18 column (3.9 mm × 150 mm). The solvent system used was isocratic methanol-water (72:28, v/v). Entrapment of POH was calculated with the help of calibration curve using the pure drug plotted at 220 nm. The percentage entrapment efficiency (% EE) was calculated with the following formula.

% EE = (amount of POH entrapped) /
(total amount of POH used in the beginning) × 100.

Scanning electron microscopy of the microparticles
Scanning electron microscopy (SEM) was performed to characterize the size and surface morphology of POH-loaded microparticles using scanning electron microscope (Zeiss EVO 40; Carl Zeiss SMT AG, Oberkochen, Germany). The lyophilized preparation of POH-loaded microparticles was suspended in 20 mM PBS pH 7.4, and a drop of the formulation was mounted on clear glass stub, air dried, and coated with gold-palladium alloy using a sputter coater. An accelerating voltage of 20.00 kV was used for imaging.

Determination of ζ-potential
ζ-potential of the PLGA microparticles was determined using DTS software (Malvern Instruments Ltd, Worcestershire, UK) based on M3-PALS technology. The formulation was lyophilized in a 2.0 mL microfuge tube, and the samples were reconstituted in 20 mM phosphate buffer, pH 7.4. This dispersion was then rapidly dispersed to a electrophoresis cell to measure the electrophoretic mobility, and ζ-potential...
values were calculated. The experiment was repeated three times, and the average ζ-potential with standard deviation was calculated.

In vitro release kinetics of active POH from PLGA microparticles
To assess the release kinetics of POH from PLGA microparticles, multiple weighed aliquots of the microparticles were dispensed in separate microvials. To each vial, 1.0 mL of 20 mM sterile PBS was added, followed by incubation at 37°C. Aliquots (100 µL) of supernatant were removed after centrifugation at 9168 × g for 10 minutes and analyzed for the POH content.

Toxicity tests for POH-bearing PLGA microformulation
A new formulation of a given drug molecule has to be tested for any inherent toxicity before being examined for its efficacy. To settle this issue, toxicity of the in-house prepared formulation was tested both in vitro and in vivo. Preliminary acute drug toxicity was based on in vitro erythrocyte lysis test, wherein hemoglobin, released as a result of membrane leakage or disruption caused by exposure to low doses of the drug, is measured.15 Briefly, fresh blood was obtained from a healthy rabbit and collected in anticoagulant solution (ethylenediaminetetraacetic acid), followed by centrifugation at 1000 × g for 10 minutes at 4°C. Buffy coat as well as plasma was discarded. The washed erythrocytes were diluted with isotonic buffer (20 mM PBS), and 50% hematocrit was prepared. To study the extent of hemolysis, the suspension of red blood cells (RBCs) was incubated with 1.0 mL of free form as well as POH encapsulated in PLGA microparticles (10 mg/mL) at 37°C for 1 hour. Free POH was dissolved in 50 µL of dimethyl sulfoxide (DMSO), and finally volume was made up to 1.0 mL with PBS (final 5% DMSO). After 1 hour, the reaction mixture was centrifuged at 1500 × g, and supernatant was collected and analyzed by ultraviolet-visible spectroscopy (λmax = 576 nm) for released hemoglobin. The percentage hemolysis was determined by the following equation:

\[
\left(\frac{(Abs_t - Abs_c)}{Abs_{100\%} - Abs_c}\right) \times 100
\]

where Absc is the absorbance of the supernatant from samples incubated with the drugs, Absc is the absorbance of the supernatant from controls (PBS), and Abs100% is the absorbance of the supernatant of controls incubated in the presence of 1% Triton® X-100, which causes complete lysis of RBCs (total hemolysis).

Hepatic and renal toxicities were monitored by applying multidose regimen (total seven doses, at alternate days) to determine biochemical profiles of serum creatinine and alkaline phosphatase (ALP). The blood was collected by retro-orbital puncture from the mice of different groups after the last administered dosage. The blood was allowed to clot at room temperature, and serum was separated for investigation of creatinine and ALP as per respective guide provided by the manufacturer. MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay
Epidermoid carcinoma cell line A253 (ATCC HTB-41™) was purchased from ATCC (Manassas, VA) and maintained in Roswell Park Memorial Institute (RPMI) 1640 culture medium supplemented with 10% heat-inactivated fetal calf serum. To perform MTT assay, 5 × 10³ cells were transferred to each well of a 96-well plate. The plate was incubated for 24 hours, and then increasing concentrations (0–60 µg/mL) of POH were added to each well, three wells received medium only with no POH and served as control. The plate was incubated for 72 hours, and cell proliferation was measured by adding 20 µL MTT dye (5 mg/mL in PBS) per well. After further incubation for another 4 hours at 37°C in a humidified chamber with 5% CO₂, the formazan crystals formed due to reduction of dye by viable cells in each well were dissolved in 150 µL DMSO, and optical density (OD) was read at 620 nm in a SpectraMax M2 plate reader (Molecular Devices, Sunnyvale, CA).

Effect of POH-PLGA-microparticles in induction of apoptosis in the human epidermoid cancer cell line
To examine the in vitro inhibitory effect of POH-bearing PLGA microparticles, the microformulation as well as free form drug was incubated with A253 cells (1 × 10⁶ cells) for 12 and 24 hours. After incubation, the cells were scraped, centrifuged, and washed with RPMI medium and lysed with TNN lysis buffer. The lysate was subjected to 12% sodium dodecyl sulfate polyacrylamide gel electrophoresis, and gels were electroblotted on polyvinylidene fluoride (PVDF) membrane following published protocol.16

Animals
Female Swiss albino mice of weight 20 ± 2 g were obtained from the institute’s animal house facility. The animals were housed in poly propylene cages on wood powder bedding in an air-conditioned ambience. Animals were quarantined
on equal light/dark cycles (12/12 hour) and were kept on a pellet diet (Ashirwad, Chandigarh, India) and water ad libitum. Animals were examined for their mortality and morbidity prior to commencement of the study, and only healthy animals were included in the experiments. The techniques used for administration of various formulations as well as sacrifice of the animals were strictly performed following mandates approved by the animal ethics committee (Committee for the Purpose of Control and Supervision of Experiments on Animals, Government of India).

**Treatments**

Animals in the resting phase of hair cycle were used in the study. The interscapular region (over an area of 2 cm²) of the experimental animals were shaven using non lubricated electric clippers. The mouse skin tumors were induced by using DMBA as carcinogen following the procedure standardized in our laboratory. The skin of the shaven dorsal portion of the mice was exposed to DMBA (52 µg in 200 µL acetone) that was applied topically three times a week for 12 weeks. Animals were routinely examined for gross morphological changes on skin and development of tumors. It has been reported that POH at 10 mM concentration does not cause toxicity in a mice model. This fact was kept in mind when selecting specific dosage regimens of various cream-based POH formulations. The formulations of POH were applied daily with cream base for a period of 7 days after induction of papilloma, with a dose of 10 mM POH in 500 µL cream base per animal. Animals were observed for 1 month to examine the effect of various formulations. The animals were divided into the following groups; each consisting of 15 animals: Group I, untreated control (DMBA applied, followed by treatment with cream base only); Group II, sham PLGA microparticles; Group III, control (healthy animals); Group IV, POH (free form); Group V, POH-bearing PLGA microparticles.

**Tumor measurement**

Tumor measurement was carried out after 1 month of last dosage application of various POH-based formulations. The diameters of the tumors were measured using a Vernier Caliper, and the tumor volume was determined using the following formula:

\[ V = D \times d^2 \times \frac{\pi}{6}, \]

where \( V \) = tumor volume, \( D \) = biggest dimension, and \( d \) = smallest dimension.

**Histopathological studies**

Animals were sacrificed and their excised papillomas were immersion fixed in 10% formalin. Next, the tissue blocks of 3 × 6 × 5 mm³ dimensions were processed for paraffin embedding. Thick sections (10 micrometer) were cut with rotary microtome and stained with H and E stain. Observations were made under light microscope (Olympus-BX 40-Japan), representative photomicrographs with final magnification of X100 were used for comparative study.

**Preparation of nuclear fraction**

The skin/tumor tissues were removed from experimental mice using sharp scalpel blades. The tissue samples were placed on ice, and fat was scraped off before further processing. The samples were homogenized in the presence of protease inhibitor cocktail, and the nuclear fraction was prepared according to the method published elsewhere.

**Western blotting**

The tissue homogenate was analyzed for the presence of various apoptotic factors by Western blotting. Briefly, the protein content of the homogenate was determined by Lowry et al method using bovine serum albumin as a standard. The homogenate (30 µg/well) was subjected to PAGE under nondenaturing conditions. The gels were electroblotted onto PVDF membranes, blocked overnight with 5% non-fat dry milk and probed with appropriate antibodies at the dilutions recommended by the suppliers. To quantify equal loading, membranes were re-probed with α-tubulin antibody to determine housekeeping protein tubulin.

**Statistical analysis**

One-way analysis of variance was used for comparing the mean values of tumor volume between various groups after ascertaining the homogeneity of variance between treatments. Post-hoc analysis for comparing the two groups was done using the least statistical difference technique. The Kaplan-Meier analysis was used to determine survival of tumor-free animals, and differences among various experimental groups were analyzed by log-rank test.

**Results**

**Entrapment efficiency, size, and ζ-potential of PLGA microparticles**

Various PLGA-based microparticle formulations have been widely used for the delivery of antibiotics as well as anticancer drugs to accomplish sustained release of the entrapped
The in-house developed PLGA microparticles had entrapment efficiency of 42.4 ± 3.5, and size was 768 ± 215 nm, as revealed by SEM and Nanophox (Sympatec GmbH, Clausthal-Zellerfeld, Germany) size analyzer (Figure 1A and B). As ζ-potential acquired by a small sized particle regulates its half-life in vitro as well as in vivo, it was considered important to determine ζ-potential of the in-house prepared microparticles. The POH-PLGA microparticles used in the present study had a ζ-potential of −7.6 ± 0.8 mV.

In vitro release kinetics of POH-loaded PLGA microparticles

In vitro release kinetics of POH-loaded PLGA microparticles was studied at 37°C in PBS, pH 7.4. POH-PLGA microparticles showed sustained release, with only 12% of the entrapped drug leaking out in the initial 18 hours. In the next 12 hours, around 17% of the total drug was released. PLGA microparticles showed an initial burst release pattern followed by sustained release kinetics for extended time period. In the present study, PLGA microparticles were found releasing 22% in the initial 72 hours with an overall total 30% release in 168 hours (Figure 1C).

In vitro and in vivo toxicity of POH-based PLGA microparticle formulation

Before determining the efficacy of any formulation, it is desirable to assess its intrinsic toxicity. With this aim, POH-bearing PLGA microparticle (in-house prepared) toxicity was evaluated, and it was found that POH-bearing PLGA microparticles had induced relatively less percent RBC lysis than free form of the drug (Figure 2). Sham PLGA also induced very negligible lysis.

In another set of experiments, animals treated with multiple dosages of POH-bearing PLGA microparticles were
analyzed for liver as well as renal function test parameters to evaluate in vivo toxicity. As shown in Table 1, POH-bearing PLGA microparticles had comparatively low levels of ALP and creatinine than the free form of the drug. The results established that POH-bearing PLGA microparticles did not have any in vitro and in vivo toxicity and are safe to use.

Cytotoxic effect of POH-bearing PLGA microparticles

The cytotoxic effect of POH formulations against epidermoid cancer cells was studied using MTT assay as shown in Figure 3. The data shows that the cytotoxic effect of POH was significant on epidermoid cancer cells at a concentration of 25 µg/mL and resulted in killing of about 50% of the cell population.

Nanoparticles are endocytosed by cells faster, enabling high payload of drug molecules, and exhibit more cytotoxic effect. With this hypothesis, the differential cytotoxicity of free and PLGA microparticle-encapsulated POH at its half-maximal inhibitory concentration (IC50) was investigated. As shown in Figure 4, free POH caused 50% cell death, whereas it increased to ~65% in the POH-PLGA microparticle-treated group in 48 hours incubation (POH-PLGA-microparticle versus free POH; P < 0.001). In addition, the time-dependent efficacy of POH-bearing microparticles on epidermoid cancer cell system was examined. As shown in Figure 5A, POH-PLGA microparticles enhanced the expression of p21/waf1 and bax at 12 hours post-incubation, whereas POH in free form was not very effective. Similarly, 24 hours post-incubation, the expression of p21/waf1 in cells treated with POH-PLGA microparticles increased further, while the free form drug was not found to be effective (Figure 5B). The results clearly suggest that microparticle encapsulated POH is delivered efficiently to the cancer cells and can easily modulate various apoptotic factors and eventually results in apoptosis of the cancer cells.

Effect of POH-PLGA microparticles on regression of tumors and survival of animals

After establishing anticancer efficacy of inhouse-prepared microparticles against cancer cells in vitro, the present study was extended in vivo using a mouse model of skin carcinoma. For this purpose, regression in the volume of DMBA-induced tumors was measured after treatment with various POH formulations. As shown in Figure 6, the percent regression was much higher in the POH-PLGA microparticle-treated group as compared with those receiving the free form of the drug.

Table 1 Concentrations of creatinine and ALP in plasma of animals treated with POH-bearing PLGA microparticle formulation

| Groups                      | Creatinine (mg %) | ALP (IU/L) |
|-----------------------------|-------------------|------------|
| Control                     | 0.32 ± 0.052      | 32.46 ± 2.12 |
| Sham PLGA microparticles    | 0.28 ± 0.034      | 36.89 ± 3.41 |
| POH (free form)             | 0.35 ± 0.038      | 44.28 ± 2.32 |
| POH-PLGA microparticles     | 0.31 ± 0.011      | 34.25 ± 2.14 |

Abbreviations: ALP, alkaline phosphatase; PLGA, poly-lactic glycolic acid; POH, perillyl alcohol.

Figure 2 Erythrocyte lysis test: In vitro toxicity was measured by erythrocyte lysis caused by different POH formulations.

Notes: Hemolysis test was performed as described in the Materials and methods section. Data represented here are means of three different experiments ± standard deviations.

Abbreviations: DMSO, dimethyl sulfoxide; PLGA, poly-lactic glycolic acid; POH, perillyl alcohol.
drug (P value <0.001; POH-PLGA-microparticle versus free POH). The treatment with microparticle-based POH formulation resulted in 80% tumor regression, while the free form of drug was able to regress tumor by 65% only. The sham microparticles were devoid of any significant anticancer activity.

Next, the efficacy of POH-PLGA microparticles was assessed in terms of the survival of treated animals. Survival graph shows the augmentation of anticancer efficacy of PLGA-encapsulated POH microparticles as well as free form POH against DMBA-induced tumorogenesis at different time points. POH-encapsulated PLGA microparticles showed 80% survival, whereas the group receiving free form of POH resulted in only 40% survival in 13 weeks (POH-PLGA microparticles versus free POH; P < 0.05). None of the animals survived beyond 12 weeks in the control group that was treated with cream base only (P < 0.01) (Figure 7).

**Histopathological analysis**

For histopathological studies, skin tissue samples were isolated from animals of various groups and analyzed following the protocol described in Materials and methods. Histopathological studies of tumors identified papillomas in various groups. However, the relative frequency of the individual tumor types differed among animals receiving different treatments. In the skin of healthy animals, there were only a few cell layers thick keratinocytes with mild keratin and pilosebaceous units (Figure 8A), whereas mice treated with DMBA followed by no POH treatment (positive control) had obvious profuse papillomatous growth with complex fibrovascular core, prominent acanthosis and keratin perl (Figure 8B). On the other hand, in the free POH treated group, a large amount of keratin and marked acanthosis was visible as depicted in Figure 8C, while hyperkeratosis, only mild acanthosis, and thin but long papillary growth was observed in animals treated with POH-PLGA microparticles (Figure 8D).
significant upregulation of p53wt and p21/waf1 expression in comparison with free form POH (POH-PLGA microparticles versus free POH; \(P < 0.05\)).

**Discussion**

Cell-cycle progression in eukaryotes is regulated by several key factors that determine whether the cell will re-enter the cycle, withdraw from it, or undergo differentiation. A slight deviation from the normal course may result in uncontrolled proliferation of the cells that eventually form the basis of cell transformation. In general, malignant state is preceded by several discernible stages, including initiation of DNA damage followed by tumor promotion both in vivo and cancer cells growing in vitro.

A large number of phytochemicals have been reported to prevent or arrest uncontrolled growth of cells through a variety of operative mechanisms. Current global focus is on natural products as a means to control various types of cancers. Some plant-based products possess strong antioxidant properties and have the potential to control or even reverse the process of carcinogenesis and emerge as potential alternatives to chemotherapeutic anticancer agents.23 However, before developing such potential phytochemicals as prospective therapeutic agents, certain challenges such as palatability, poor solubility, and other related problems have to be addressed. Some of these can be generally circumvented through the use of an appropriate delivery system that has potential to modify the pharmacokinetics of the drug and facilitate its controlled and even preferential release at the desired tumor site.
POH, a dietary constituent and a mono-terpene isolated from cherries, had been used widely as an anticancer agent against various types of cancer in animal models.4,24,25 POH is known to arrest cells in G0/G1 phase and induce apoptosis.26 Although POH is a very effective chemotherapeutic agent, its anticancer properties have not been explored completely. In fact, certain intrinsic properties of POH such as small size, poor solubility in aqueous solution, and bioavailability restrict attainment of its effective concentration at the targeted site and thus limit its application in cancer treatment. Earlier studies suggest that localized delivery at the targeted site could be enhanced significantly by the use of microparticle-based drug delivery and eventually can be successfully employed in tumor therapy.27,28 It has recently been demonstrated that naturally occurring phytochemicals can be encapsulated, covalently attached, and adsorbed onto microparticles in order to surmount drug-solubility problems.10,29,30 The constituents of the microparticle-based delivery system can also facilitate co-solubilization of the compound/drug in question. This approach has significant implication in cancer therapeutics because more than nearly half of the phytochemicals showing anticancer activity have solubility constraints.31 Nanosize range of the microparticles entails a high surface area that not only provides sustained drug release but also provides a useful strategy for their functionalization. The present study focuses on development and characterization of various POH-bearing microparticle formulations and their evaluation in chemotherapy against DMBA-induced skin papilloma in Swiss albino mice.

The POH-bearing PLGA microparticle formulation was characterized for sizing using electron microscopy, and its surface charge properties were assessed on the basis of \( \zeta \)-potential. The biophysical properties of the novel formulation were also assessed. Further, in vitro anticancer efficacy of PLGA based POH formulation was examined against human epidermoid cancer cell line (A253). Finally, the in vivo therapeutic potential of POH-PLGA microparticle formulations was evaluated on the basis of tumor size regression, survival, histology, and expression profiles of various apoptotic molecules.

To optimize the size and loading efficiency of the active components, a series of PLGA-based microparticles with varying compositions were prepared. The size of the in-house prepared microparticles was standardized by incorporating PLGA of various molecular masses. Further, the entrapment efficiency at various added weight percentage values of POH to PLGA was examined. It was found that 50%–70% weight of POH to that of PLGA leads to entrapment efficiency of around 42.4 \( \pm \) 3.5 with the size of prepared microparticles in the range of 768 \( \pm \) 215 nm (Figure 1A and B). This formulation of POH-PLGA was used in subsequent studies.
Usage of the free form drug does not allow attainment of effective concentration unless a large payload is administered, which ironically results in many untoward effects. In such cases, drug-delivery systems offer help by providing continuous and constant supply of the therapeutics for an extended time period. The hydrophobic POH is likely to be physically dispersed by its encapsulation throughout the matrix of PLGA particle. The release kinetics of the entrapped drug showed an overall slow and sustained release that was likely to be regulated by rate of polymer biodegradation. The release kinetics of entrapped drug was determined by incubating the drug-bearing microparticles in PBS, pH 7.4 at 37°C to simulate physiological conditions. The amount of released POH was estimated by reverse phase HPLC analysis.

A linear release of POH accounted for around 22% of the total POH entrapped in PLGA microparticles. In concurrence with earlier reports, where PLGA with the 50:50 ratio of lactic:glycolic acid composition was found to degrade in 1–2 months, the in-house-prepared PLGA microparticles acquired steady release and showed 30% release of entrapped molecules in a time span of 7 days (Figure 1C).

Both in vitro and in vivo toxicity data revealed that POH-bearing PLGA microparticles had negligible toxicity when compared with free form of the drug and thus assures its safety (Figure 2 and Table 1). The cytotoxicity of free POH was next examined, and it was found that POH had an IC$_{50}$ of 25 µg/mL against A253 human epidermoid cancer cells (Figure 3). This concentration was used to analyze differential cytotoxicity of free and PLGA microformulation-encapsulated POH and their comparative potential in induction of apoptosis in epidermoid cancer cell line. Due to faster uptake and subsequent release of the drug by POH-PLGA microparticles, it exerts more cytotoxic effect on epidermoid cancer cells than its free form (Figure 4). This observation could be explained on the premise that application of POH-PLGA microparticles delivers a high amount of active drug at the tumor site mainly. Further, polymer conjugation of low molecular weight drugs alters their biodistribution; enabling their passive targeting and reducing access to sites of toxicity. Microparticle-based drug formulations entertain a special status in improving the efficacy of drug against many types of cancers. Therefore, the present study was extended to gain insight in POH-PLGA microparticle-mediated modulation of various proteins involved in the apoptosis of A253 cells. Western blot analysis was performed to determine levels of various cell cycle and apoptosis regulating factors. As evident from Figure 5A, incubation of POH-microparticle formulation with epidermoid cancer cells resulted in enhanced expression of p21/waf1 when compared with free form of POH. The expression of another important pro-apoptotic factor Bax was also analyzed in A253 cells upon their incubation with POH-PLGA microparticle formulations. The POH-microparticle formulation up-regulates the expression of Bax, thereby enhancing apoptosis induction in cancer cells (Figure 5A). In the control groups (cells treated with sham PLGA microparticles), the expression profile of these molecules was not significantly affected when compared with POH-PLGA microparticles, suggesting that microparticle-based formulation is effective in upregulating p21/waf1 and Bax gene expression. The effect of POH-PLGA microparticles on expression of pro-apoptotic molecules was more prominent after 24 hours incubation (Figure 5B). Interestingly, the POH-PLGA microparticle was
equally successful in downregulating p53-mutant expression in A253 cancer cells (data not shown). Of note, both p53 and/or p21-dependent and -independent pathways have been previously reported to be involved in POH-induced cell cycle arrest and apoptosis in cancer cells; however, the extent of the cell responses varied and can be attributed to the mechanism operative in a specific cell type involved and the doses of the drug used in the study.34,35

DMBA, a polycyclic aromatic hydrocarbon, is metabolized to chemically reactive electrophile and initiate a cascade of reaction that eventually results in carcinogenesis by covalent interaction(s) with DNA.22 Further investigations suggest that it induces skin papilloma in animal models by mutagenesis in Ha-ras oncogene.36 Upon topical application of POH-PLGA microparticles, 80.8 ± 5.2 regression in tumor size was observed after the treatment, whereas free form POH showed only 65.1 ± 7.1 regression (Figure 6). Increased regression in the POH-PLGA microparticle group could be attributed to sustained release of POH. Interestingly, the tumor regression results were found to have great correlation with survival rate of the treated animals. In case of POH-PLGA microparticles, survival rate was highest (80%) over a period of 13 weeks when compared with survival rate of the animals treated with free form POH (40%) (Figure 7). Further, histopathological analysis of the skin tissues isolated from treated animals was performed and the results clearly demonstrated that free form POH had more acanthosis and papillary growth than POH-bearing PLGA microformulations (Figure 8). The pro-apoptotic gene p53 regulates the balance between cell proliferation and cell death.37,38 It has
been observed that the p53 gene gets mutated in most types of the malignancies, including sarcomas. Upon exposure to a potent carcinogen, the toxic insult of the cell is generally nullified by a chain of cell cycle regulatory events programmed to check the uncontrolled proliferation by a suitable repair mechanism. The failure to induce expression of functional p53wt leads to disregulation of cell cycle arrest or cell death and tumor progression. The results of the present study showed decreased p53-mutant expression upon treatment with POH-bearing PLGA microparticles. Also, higher efficacy of the microparticle-based formulation of POH can be correlated with upregulated expression of p53wt and p21/waf1 genes (Figure 9).

The survival data further establish higher supremacy of POH-PLGA microparticles over its free form. The observed higher efficacy of POH-bearing PLGA microparticle formulations can possibly be attributed to the greater bioavailability of POH and its accumulation at the tumor site. Finally, it can be inferred that microparticle-based formulations not only overcome the solubility constraints of the poorly water soluble POH but also facilitate its release in regulated fashion that eventually results in better efficacy of POH in terms of tumor regression and survival of treated animals.

**Conclusion**

Entrapment of POH in PLGA microparticles makes its release more sustained, thereby facilitating accumulation of required concentration at the target site. As revealed by cytotoxic assay, POH-bearing PLGA microparticles showed greater growth inhibition of human epidermoid cancer cell line than free form POH. Further, higher tumor regression, increased survival, histopathological study, and Western blot profiles of various cell cycle regulatory proteins clearly suggest that POH-PLGA microparticles offer better efficacy than free form POH against DMBA-induced skin papillomas in animal models.

**Acknowledgments**

The authors are thankful to their coordinator, Professor M Saleemuddin, for providing the facilities. Mohammad Farazuddin acknowledges financial support in the form of SRF provided from CSIR (India). Financial assistance provided by UGC and DST under the FIST program is greatly acknowledged.

**Disclosure**

The authors claim that there is no conflict of interest in this work.

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