


 Cite this: *RSC Adv.*, 2024, 14, 33952

A doxorubicin loaded chitosan–poloxamer *in situ* implant for the treatment of breast cancer†

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Breast cancer is a serious concern for many women worldwide. Drug-loaded implants have shown several benefits over systemic administrations. To provide anti-cancer drugs with controlled release and reduced systemic toxicity, biodegradable *in situ* implants have attracted a lot of attention. In the present study, we aimed to design and optimize a doxorubicin-loaded chitosan–poloxamer *in situ* implant for breast cancer treatment. Utilizing Box–Behnken Design and a Quality-by-Design (QbD) methodology, the *in situ* implant was prepared with chitosan (X1), poloxamer 407 concentration (X2), and stirring time (X3) as the independent variables. It was characterized for its *in vitro* gelation time, pH, rheology, and morphology, and evaluated based on drug release profile, *in vitro* cytotoxicity activities, *in vitro* anti-inflammatory potential, *in vitro* cellular uptake, and *in vivo* anti-inflammatory and pharmacokinetics to ensure their therapeutic outcomes. The results revealed that the prepared formulation showed a gelation time of 26 ± 0.2 s with a viscosity of 8312.6 ± 114.2 cPs at 37 °C. The developed formulation showed better cytotoxic activity in MCF-7 cell lines compared to the free drug solution. It demonstrated reduced levels of pro-inflammatory cytokines in RAW 264.7 macrophages. Further, the prepared *in situ* implant increases the intracellular accumulation of DOX in the MCF-7 cells. The *in vivo* pharmacokinetic investigations depicted an increase in $t_{1/2}$ and a decrease in AUC of the developed formulation resulting in prolonged drug release and there could be a lower drug concentration in the bloodstream than for the free drug. Therefore, the developed *in situ* implant may offer a viable option for breast cancer treatment.

 Received 29th August 2024
 Accepted 14th October 2024

DOI: 10.1039/d4ra06253a

rsc.li/rsc-advances

1. Introduction

One of the most common and dangerous health issues that women face globally is breast cancer. Every year, over a million new cases are diagnosed. According to the World Health Organization, 670 000 individuals worldwide lost their lives to breast cancer in 2022, while 2.3 million women received a diagnosis.¹ As a result, one of the greatest hazards to humanity is cancer, a global health issue with no known bounds in terms of death and incidence rates. The primary cause of death for patients with breast cancer is metastasis.² The prognosis for metastatic patients remains unsatisfactory and dismal despite advancements in breast cancer treatment.

Systemic treatments like chemotherapy involve intravenous delivery of anti-cancer drugs at maximum tolerable doses that cause serious toxicities in healthy tissues, such as cardiomyopathy and neutropenia^{3,4} and the drug reaches the tumor site in low concentrations, which decreases the tumor treatment's effectiveness.^{5,6} Further, surgery is a non-aggressive procedure that is regarded as the primary strategy in the early stages of breast cancer.⁷ Even with the advancements in surgical methods, there is still a chance that residual tumor cells will stay in the surgical margins or the circulation following the surgical interventions, increasing the risk of metastasis and cancer recurrence.^{8,9} As a result, alternative approaches are required to reduce the chance of distant metastasis, stop the tumor from growing back, and increase the effectiveness of tumor-targeted drugs.¹⁰ The use of targeted drug delivery has the potential to achieve therapeutic concentrations within the tumor site, which would reduce systemic levels and, in turn, reduce the likelihood of unwanted side effects.¹¹

Implantable drug delivery devices are the better approaches for local drug administration to the tumor site.¹² These implants use biopolymer components that are both biodegradable and non-biodegradable. The non-biodegradable

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† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d4ra06253a>



implants require surgical procedures twice, as they need to be inserted and removed. Using biodegradable polymers in implantable devices could be a better approach as the body can naturally remove it after use, eliminating the need for surgical extraction and increasing the patient's acceptance.¹³ In recent decades, many biomaterials have been widely used to develop implantable drug-delivery devices against cancer.¹⁴ Particularly biopolymers have attracted more attention because of their beneficial qualities, which include biocompatibility, biodegradability, and ease of processing.¹⁵ In the past decade, drug delivery systems that are implanted have advanced significantly and have shown to be an extremely effective way to deliver drugs to specified areas.¹⁶ They also reduce the total amount of drug in blood circulation by providing accurate spatial control, recurrence probability can be reduced, enhancing the rate of overall survival, and shielding healthy cells from harm.¹⁷ By maintaining therapeutically relevant drug levels within the tumor for extended durations, they can overcome the limitations of short half-lives, dose-dependent systemic toxicity, and rapid clearance associated with conventional chemotherapy agents.¹⁸ Implantable drug delivery systems include hydrogel, nanofibers, *in situ* gel, and osmotic pumps used for local chemotherapeutics delivery.¹⁹ As an injectable drug delivery technique, *in situ* implants (ISI) have garnered a lot of interest as these systems are injectable into the body with a syringe, and after injection, they solidify to form a semisolid depot.²⁰ When a stimulus (such as ions, temperature, or pH) is applied, the liquid combination precipitates, cross-links, or polymerizes to solidify into an "implant".²¹ Among these, temperature-responsive *in situ* gel has been widely used over the past few years.

Temperature changes cause temperature-sensitive *in situ* gel to shift from a solution to a gel.

Poloxamer 407, a commercially accessible and standardized excipient approved for parenteral use by the FDA, is a well-known thermosensitive component focused by many researchers.²² Poloxamer (PO), being a biocompatible polymer composed of polyoxy ethylene (PEO), polyoxy propylene (PPO), and polyoxy ethylene chains (PEO_n-PPO_n-PEO_n) to form nonionic triblock polymers.²³ It provides flexibility and control in drug delivery system design, allowing for forming formulations with customized release profiles and improved therapeutic efficacy.²⁴ Additionally, the literature suggests that chitosan (CH) has good biodegradability, low toxicity, and suitable biocompatibility.²⁵ Furthermore, chitosan itself has significant effects on several tumor types that are anti-metastatic anti-proliferative, anti-angiogenic, and pro-apoptotic.^{26,27} A thermosensitive chitosan-based injectable hydrogel was developed by Ahsan *et al.* showing an inhibiting effect on tumor development and used as an effective anti-cancer drug carrier.²⁸ Hence, we aimed to develop *in situ* gel as an implant comprising chitosan and poloxamer 407 for treating breast cancer.

Here, we have used doxorubicin (DOX) as a model drug. It is a powerful cytotoxic anthracycline antibiotic that damages DNA and induces cancer cells to undergo apoptosis. This study highlights the potential role of the *in situ* implant incorporating

chitosan and poloxamer 407 using the QbD approach in the treatment of breast cancer by assuring improved cytotoxicity activity, anti-inflammatory activity, and improvised therapeutic potential which would reduce the systemic toxicity of chemotherapy and cancer recurrence after surgical procedure.

2. Materials and methods

2.1. Materials

Chitosan (molecular weight 190–310 kDa, deacetylation degree 75–85%) was obtained from Loba Chemie Pvt Ltd, poloxamer 407 was purchased from Hi Media Pvt Ltd. Doxorubicin was procured as a gift sample from Sisco Research Laboratory Pvt. Ltd (Mumbai). Glacial acetic acid, 3-(4,5-dimethylthiazol2-yl)-2,5-diphenyl tetrazolium bromide (MTT, Sigma), and Dulbecco's modified Eagle's medium (DMEM), dimethyl sulfoxide (DMSO) were obtained from Hi Media Pvt Ltd MCF-7 cells and RAW264.7 cells were bought from National Centre for Cell Science, Pune. All the remaining substances were of analytical grade.

2.2. Optimization

Using Design Expert® Software (version 13, Stat Ease Inc., Minneapolis, USA), the Box-Behnken design was chosen to optimize the *in situ* gel formulation using 3³ factorial design with 3 factors, 3 levels, and 17 runs. In this study, chitosan concentration (X1), poloxamer 407 concentration (X2), and stirring time (X3) were selected as independent variables. The three factorial levels for these variables were designated as follows: -1, 0, and +1 for low, medium, and high levels, respectively as shown in Table 1. The viscosity at 25 °C (Y1), viscosity at 37 °C (Y2), and gelation time (Y3) were chosen as the dependent variables. Using response surface analysis, it was possible to understand the relationship and interaction terms between the parameters as well as the predicted points of the independent variables that would comprise the optimum formulation. A one-way ANOVA was used for the statistical study to ascertain how formulation parameters affected the response variables. Following the software's recommendation, we created one optimal formulation in triplicate. After identifying the range that the ideal conditions might fall into using overlay plots and the desirability function, we measured the dependent variables. To verify the accuracy of the model, we conducted a comparison between the predicted viscosity at 25 °C, viscosity at 37 °C, and gelation time with the observed viscosity at 25 °C, viscosity at 37 °C, and the gelation time.

Table 1 Independent variables and their corresponding levels for DOX-CH-PO *in situ* implant

Independent variables	Symbol	Coded levels		
		-1	0	1
Chitosan concentration (%)	X1	0.3	0.5	0.7
Poloxamer 407 concentration (%)	X2	10	20	30
Stirring time (second)	X3	60	105	150



2.3. Validation of data analysis and response surface technique

Design Expert software from Stat-Ease was modified to utilize the gathered results. By employing Analysis of Variance (ANOVA), the proposed models were verified. The program generated primary and interaction effects of the variables in the form of “3D response surface plots, 2D perturbation curves, and contour plots.” We selected the combination that predicted the highest degree of desirability, and we compared the projected values with the experiment results.

3. Preparation and physicochemical characterization of DOX–CH–PO *in situ* implant

Poloxamer 407 solutions were prepared by the cold method process. The optimized amount of polymer was dispersed in distilled, deionized water and stirred continuously for one hour at room temperature. The poloxamer solutions that had partially dissolved were kept in the refrigerator at 4 °C until the polymer had entirely dissolved, which took around 24 hours. The chitosan solution was made by adding the optimized amount in a 2% w/v acetic acid solution and stirring continuously until the chitosan was completely dissolved. Aqueous solution of doxorubicin was added to the chitosan solution with continuous stirring. Finally, this drug solution was added to the poloxamer solution at room temperature and evaluated for different parameters.²⁹

3.1. Fourier transform infrared spectroscopy (FTIR)

The drug and polymers in the prepared formulation and solid mixture were examined for possible interactions using infrared spectroscopy (FT/IR-4600, Jasco, Japan). The metal structure of the device was loaded with roughly 10 mg of powdered samples, and FT-IR readings were taken. The spectra of poloxamer 407, pure doxorubicin, chitosan, and the prepared formulation were compared at 400 to 4000 cm^{-1} . A reference spectrum was compared to identify distinctive peaks representing any drug–polymer interaction.

3.2. Differential scanning calorimetry (DSC)

Using reference material as a sample, this approach estimates the temperature-dependent difference in heat required to raise the sample's temperature. Using a differential scanning calorimeter (DSC-60, Shimadzu, Kyoto, Japan), thermograms of poloxamer 407, pure doxorubicin, chitosan, and formulation were recorded. The sample weight ranged from 2 to 3 mg, and the DSC measurements were carried out in a nitrogen-purging atmosphere with a scanning rate of 10 °C min^{-1} . The samples underwent a heating cycle of 10 °C min^{-1} , ranging from 0 to 350 °C.

3.3. X-ray diffraction (XRD)

XRD is an essential analytical tool for determining the crystal-line or amorphous solid state in which drugs may occur. X-ray diffraction investigation of the pure drug, polymers, and

formulation was carried out using a Rigaku X-ray diffractometer. The glass slide containing the powdered materials under evaluation was put on the X-ray diffractometer. Within a 2θ range of 10 to 90°, the scanning rate was 10 minutes.

3.4. Pharmaceutical characterization of DOX–CH–PO *in situ* implant

3.4.1. *In vitro* gelation time. The behavior of the sol–gel transition was determined using the test tube inversion technique.

A 5 mL stoppered test tube with an inner diameter of 10 mm was filled with 1 mL of each prepared solution. Following that, the test tubes were submerged in a water bath with a thermostat set to 37 °C. By flipping the tube once every second until no flow was seen, the sol–gel transition was tracked.³⁰

3.4.2. Investigation of the rheological properties. The measurement of the *in situ* gel's viscosity, which was quantified in centipoises (cPs), was the most significant rheological investigation. A thermo-responsive *in situ* gel transforms from a liquid at room temperature to a gel at physiological temperature. The formulation should have an optimum viscosity, which would allow easy instillation at the tumor site as a solution, but also facilitate the formulation to undergo rapid sol-to-gel transition. Viscosity at 25 ± 2 °C and 37 ± 2 °C temperatures of the developed formulation was measured with a viscometer (DV2T Brookfield Viscometer) equipped with spindle number 63 at 50 rpm.

3.4.3. pH. The ideal pH for an *in situ* gel formulation will allow the formulation to remain stable while also preventing patient irritation upon injection. The pH buffers 4 and 7 were used to calibrate the digital pH meter. A glass electrode was thoroughly dipped into the formulation samples, and twenty milliliters of the optimal formulation were placed in a beaker. Then, the pH of the solution was determined in triplicate.³¹

3.4.4. Morphological study of the prepared formulation. Scanning electron microscopy (ZEISS, Evo 18, Thornwood, NY, United States) was used to assess the morphological characteristics of the *in situ* gel. After the gel formation, the optimized formulation was frozen at -20 °C for 24 h and then lyophilized for 48 h. The SEM was used to investigate the cross-section of the lyophilized sample after it was shattered in liquid nitrogen and created a photomicrograph image by scanning an object with an electron beam in a raster scan pattern at 1000 and 2000 magnification. The sample that had been gold-sputter-coated was held in a tiny area using the SEM sample holder. With an accelerating voltage of 15 kV, the SEM images were taken.

3.4.5. *In vitro* biodegradation analysis. The degradation was measured as a percentage of weight loss (WL) by taking fresh *in situ* implants weighing 0.1 g in 1 milliliter of PBS at two pH values (5.5 and 7.4) and 37 °C. It was then followed with the addition of 13 mg L^{-1} lysozyme solutions (same as those found in human serum). The mixtures were agitated at 60 rpm for various durations (1, 3, 5, 7, 9, and 11 days) at 37 °C. Every day, a new lysozyme solution was maintained *via* replacement. Following the conclusion of every degradation phase, the samples were removed from the medium, cleaned with distilled



water, and dried in a vacuum oven at 50 °C until their weight remained constant. The following equation was used to compute the weight loss (%) after the experiment was run in triplicate.³²

$$\text{WL}(\%) = \frac{W_i - W_f}{W_i} \times 100$$

where W_i denotes the sample's starting weight and W_f its final weight following degradation.

3.4.6. *In vitro* drug release studies. The paddle method was used to monitor drug release with some modest adjustments. Pre-soaked in PBS 7.4 for 12 hours, the dialysis bag has a molecular weight cut-off of 12 000 g mol⁻¹ (6 cm long). Before being put in a dissolution tester, each dialysis bag was fastened with two clamps on either end. Before sealing and placing the dialysis bags in a dissolution tester, DOX-CH-PO solution and free DOX solution were added. Using 500 mL of PBS as a release medium, the test condition involved 50 rpm stirring speed at 37 ± 0.5 °C. Fresh medium was added to the release medium at predetermined intervals. Similarly, it was performed at 0.1 M acetate buffer pH 5.5. The amount of drug released was measured using HPLC at a maximum wavelength of 254 nm.

3.5. Biological characterization of DOX-CH-PO *in situ* implant

3.5.1. Cytotoxicity assay. Using the 3-[4,5-dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide (MTT) technique, the *in vitro* cytotoxicity of free DOX, and DOX-CH-PO ISI was assessed against MCF-7 cells. In a 96-well culture plate (Tarsons, India), cells in the logarithmic phase of culture were plated at a density of 5 × 10⁴ cells per well and allowed to adhere for 24 hours. The entire media was changed to DMEM (Dulbecco's Modified Eagle media) after a day. Then the control cells were treated with free DOX, and DOX-CH-PO ISI (10 to 50 µg mL⁻¹) and incubated for 24 h. After adding the MTT solution (0.5 mg mL⁻¹) and letting it sit for three hours, 100 µL of DMSO was added to dissolve the formazan crystals. A microplate reader was then used to measure the absorbance at 540 nm. The values were presented as mean ± SD and the tests were run in triplicate.³³

3.5.2. *In vitro* anti-inflammatory study. The anti-inflammatory activity of the proposed DOX-CH-PO ISI was determined in RAW 264.7 macrophage cell lines. These cells were cultured in RPMI 1640 medium supplemented with 10% fetal bovine serum, 1% Pen-strep (a combination of streptomycin and penicillin) from a 100 U mL⁻¹ Pen-strep stock solution, and 5% supplied CO₂ at 37 °C. The RPMI 1640 medium containing 100 g mL⁻¹ of the DOX-CH-PO sample completely dissolved. The cells were divided into three groups, *i.e.*, the negative control (NC) group, a disease control group (LPS-treated), and the DOX-CH-PO group. In the next step, 50 µL of LPS (1 g mL⁻¹) was administered to the DC group and DOX-CH-PO group, while 50 µL of RPMI 1640 medium was given to the negative control group. TNF-α, IL-1β, and IL-6 expression levels were measured using ELISA kits, in accordance with the manufacturer's recommendations. The supernatants were subjected to antigen-antibody reactions following

the user's handbook that came with the ELISA kits, and the degree of cytokine expression was assessed. After adding 50 µL of stop buffer to each well, the cytokine level was measured using a microplate reader, and absorbances were recorded at 450 nm.³⁴

3.5.3. Cellular uptake. Fluorescent microscopic imaging was used to assess the cellular absorption of DOX-CH-PO ISI. To be more specific, a coverslip was placed in each well of six-well plates. In 6-well plates, 3 × 10⁵ cells per well were planted on the coverslips with 2 milliliters of culture media. Cells were treated with DOX-CH-PO ISI at equivalent doses of 200 µg mL⁻¹ DOX after a 24 hour incubation period. Following incubation for 4 hours, the drug-containing medium was disposed of. The cells underwent three PBS washes, were fixed for thirty minutes in 75% ice-cold ethanol, and were examined using a fluorescent microscope (Cilika, BT-P-FL). The fluorescence intensity was obtained by using the software ImageJ 1.53.³⁵

3.5.4. *In vivo* anti-inflammatory study. An *in vivo* assessment of the DOX-CH-PO ISI's anti-inflammatory action was conducted to appraise its potential. For this, following the 20th day of treatment, breast tissue samples were taken from recently sacrificed animals. We separated the breast tissues and homogenized them to perform the biochemical study, yielding a 10% homogenate in phosphate buffer. The homogenates underwent a 15 minute centrifugation at 10 000 rpm and 4 °C in an Eppendorf 5425 R G ultracentrifuge. The ELISA kits' instructions for performing antigen-antibody reactions on supernatants were followed to measure the levels of pro-inflammatory cytokines (TNF-α, IL-6, and IL-1β).³⁶ Two groups were assessed for total protein (pg mL⁻¹): group II, which received treatment, and group I, which did not get treatment (disease control).

3.5.5. *In vivo* pharmacokinetics study. To examine the absolute bioavailability and pharmacokinetic parameters of three groups of female Wistar rats, an equal dosage of free DOX and DOX-CH-PO ISI (5 mg kg⁻¹ DOX) was intravenously and intramuscularly injected into each group respectively. Rat retro-orbital plexus blood samples (200 µL) were drawn at predetermined intervals (6 h, 12 h, 24 h, 48 h, 72 h), and the plasma was separated by centrifuging the samples for 10 minutes at 5000 rpm. Before analysis, all plasma samples were kept at -20 °C. 100 µL of plasma samples were vortexed for three minutes after adding 1 mL of acetonitrile and water mixture solution. After being collected, the supernatant was evaporated in nitrogen-filled conditions. Following centrifugation and a vortex, the residue was resolved using 100 µL of mobile phase, and the supernatant was subjected to HPLC analysis. A C18-HL reversed-phase column (150 4.6 mm, 3.5 m; Zorbax Ltd) was used to evaluate the samples. At ambient temperature (25–28 °C), a binary gradient solvent system was used for the elution, with a flow rate of 1.00 mL min⁻¹, with acetonitrile (ACN): sodium dodecyl sulphate and orthophosphoric acid as the mobile phase. Plasma from untreated rats was used to prepare standard calibration curves of concentrations 0.2, 0.4, 0.6, 0.8, and 1 µg mL⁻¹. The PK Solver program computed the pharmacokinetic parameters using the linear trapezoidal approach and non-compartmental studies.³⁷



3.6. Statistical analysis

A statistical analysis was performed on the data using version 9 of GraphPad Prism. One-way analysis of variance (ANOVA) was used to compare group variances, with a significance level of $P = 0.05$. The sample sizes range from $n = 3$ to $n = 6$, and the data is shown as mean \pm standard deviation (SD).

4. Results and discussion

4.1. Statistical optimization of DOX-CH-PO *in situ* implant

Based on the Box–Behnken factorial design, the design expert projected a total of 17 runs for three components, which were modified at three distinct levels: chitosan concentration (X1), poloxamer 407 concentration (X2), and stirring time (X3). These runs were varied at three distinct levels (coded as -1 , 0 , and $+1$). The viscosity at $25\text{ }^{\circ}\text{C}$ (Y1), viscosity at $37\text{ }^{\circ}\text{C}$ (Y2), and gelation time (Y3) were studied as responses shown in Table 2.

4.1.1. Effect of independent variables on viscosity at $25\text{ }^{\circ}\text{C}$ (Y1). The polynomial equation (eqn (1)) for response Y1, on viscosity at $25\text{ }^{\circ}\text{C}$, was generated by the model. The primary and secondary effects of each response might be ascertained using factor coefficients. Predicting the reaction for levels of each element is feasible by using an equation stated in terms of coded factors. By default, factors with high levels are coded as $+1$, while those with low levels are coded as -1 . By comparing the factor coefficients in the coded equation, we can ascertain the elements' respective relative importance.

$$Y1 = 151.5 + 2.2375X1 + 19.55 X2 - 3.9625 X3 + 3.275 X1X2 - 0.1 X1X3 + 0.575 X2X3 - 2.725 X1^2 + 9.85 X2^2 - 1.375 X3^2 \quad (1)$$

The equation showed that chitosan (X1) and poloxamer 407 (X2) have a favorable impact on the *in situ* implant while stirring time (X3) has a negative impact on it. The fact that X3's

magnitude was most readily apparent led to the conclusion that its detrimental effects were most severe. The combined interaction effect of X1X2 and X2X3 was also found to be synergistic, with X1X3 having the largest detrimental effect on the gel. As a result, it can be inferred that increasing the stirring time (X3), there is a decrease of viscosity at $25\text{ }^{\circ}\text{C}$ and as the concentration of poloxamer 407 (X2) increases, there is an increase in the viscosity at $25\text{ }^{\circ}\text{C}$ of the *in situ* implant.

With a p -value of less than 0.05 (<0.0001), the ANOVA for the quadratic response surface model of viscosity at $25\text{ }^{\circ}\text{C}$ was determined to be significant. With a p -value of 0.0020 , factor A (chitosan concentration) was found to have a significant impact on viscosity at $25\text{ }^{\circ}\text{C}$. On the other hand, factor C and B's p -values of less than 0.0001 indicate that they significantly affect viscosity at $25\text{ }^{\circ}\text{C}$. There was a significant correlation between the actual and expected results, as evidenced by the variations of less than 0.2 between the adjusted R^2 (0.9925) and predicted R^2 (0.9898) for the viscosity at $25\text{ }^{\circ}\text{C}$ quadratic model. It is recommended to utilize this model to navigate the design space. The contour plot of the reaction and response surface for Y1 are shown in Fig. 1.

4.1.2. Effect of independent variables on viscosity at $37\text{ }^{\circ}\text{C}$ (Y2). The 2D and 3D graphs illustrate the likely impacts of changing independent variables on the degree of viscosity at $37\text{ }^{\circ}\text{C}$ deviation. The viscosity at $37\text{ }^{\circ}\text{C}$ deviation is shown in the actual equation (eqn (2)).

$$Y2 = 8302.8 + 154.675 X1 + 1411.31 X2 - 346.112 X3 + 341.1 X1X2 - 53.65 X1X3 + 126.325 X2X3 - 197.288 X1^2 + 479.437 X2^2 - 121.362 X3^2 \quad (2)$$

The equation showed that the chitosan (X1) and poloxamer 407 (X2) have a positive impact on the viscosity at $37\text{ }^{\circ}\text{C}$, whereas the stirring time (X3) has a negative impact. Similarly, it was found that the interaction between X1X2 and X2X3 was synergistic, while X1X3 had a detrimental effect on viscosity at $37\text{ }^{\circ}\text{C}$.

Table 2 Response values of experimental runs

Formulation code	X1 Chitosan (%)	X2 Poloxamer 407 (%)	X3 Stirring time (s)	Y1 Viscosity at $25\text{ }^{\circ}\text{C}$ (cPs)	Y2 Viscosity at $37\text{ }^{\circ}\text{C}$ (cPs)	Y3 Gelation time (s)
F1	0.3	20	150	141.5	7563.2	23
F2	0.5	20	105	151.8	8245.3	30
F3	0.7	20	150	145.3	7845.6	25
F4	0.7	10	105	138.6	6945.6	35
F5	0.3	20	60	149.3	8015.4	23
F6	0.5	20	105	153.6	8612.3	27
F7	0.5	30	150	176.5	9784.6	10
F8	0.3	30	105	172.1	9542.1	15
F9	0.7	30	105	183.6	10 453.3	12
F10	0.3	10	105	140.2	7398.8	37
F11	0.7	20	60	153.5	8512.4	27
F12	0.5	10	150	135.6	6712.2	40
F13	0.5	20	105	149.2	8098.2	28
F14	0.5	20	105	152.2	8412.3	26
F15	0.5	20	105	150.7	8145.9	28
F16	0.5	30	60	183.2	10 356.9	11
F17	0.5	10	60	144.6	7789.8	35



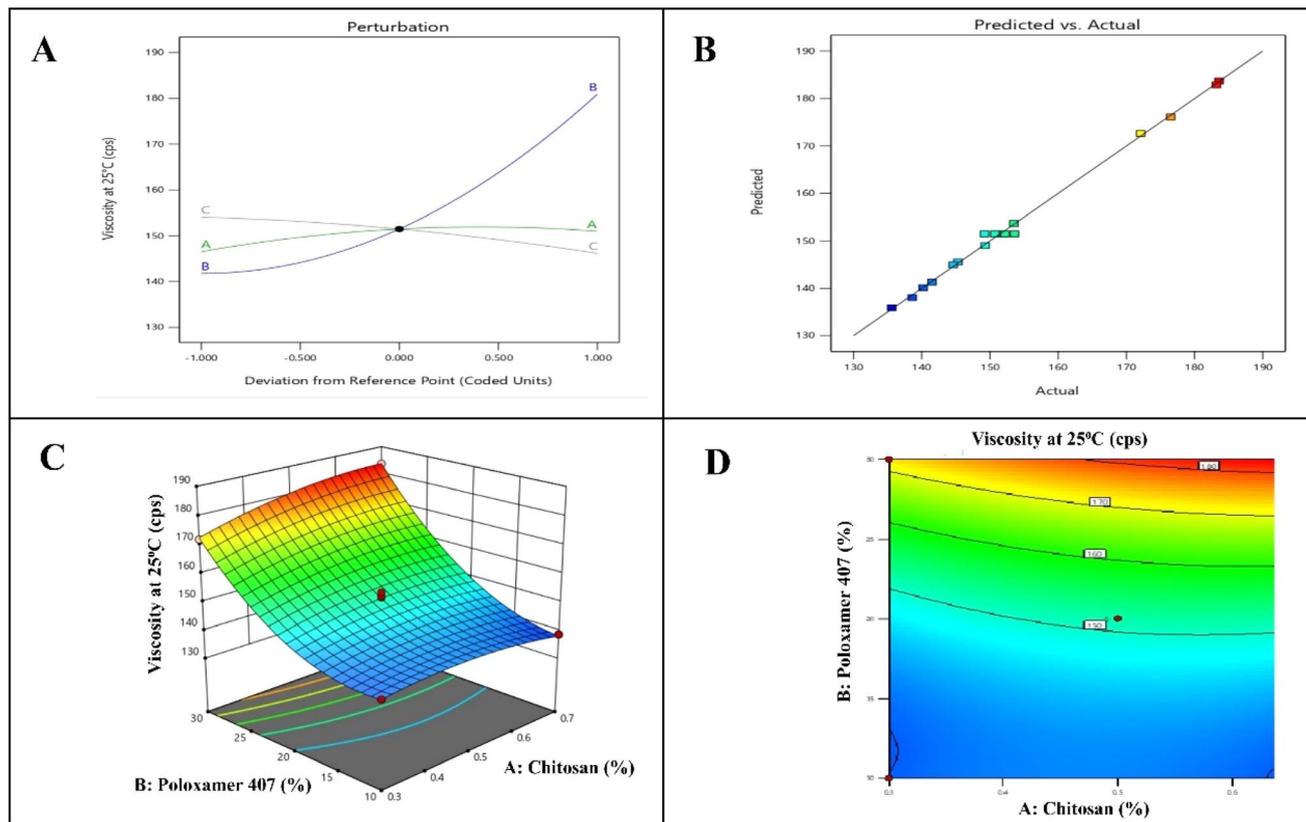


Fig. 1 QBD model graph for viscosity at 25 °C. (A) Perturbation curve, (B) linear correlation plot, (C) response surface plot, (D) contour plot.

It indicates that as the concentration of poloxamer 407 (X2) and chitosan (X1) increases, there is an increase in the viscosity at 37 °C. Micelle production and micellar aggregation occur in response to an increase in poloxamer 407 concentration. The concentration must be higher than the micellar concentration for the gel phase to occur. Hydrophobic parts of the poloxamer 407 are kept apart in cold water by hydrogen bonding between POP chains and water. With an increase in temperature, the hydrogen bonding is broken and hydrophobic interactions are forming the gel.

The p -value for the model, which is less than 0.05, is less than 0.0001, indicating that the ANOVA for the quadratic response surface model for viscosity at 37 °C is significant. With a p -value of 0.0450, X1 was found to have a significant impact on viscosity at 37 °C. X2 and X3 had p -values of less than 0.0001 and 0.0010, respectively, indicating a substantial impact on viscosity at 37 °C. The adjusted R^2 (0.9728) and projected R^2 (0.9448), which displayed discrepancies of less than 0.2, demonstrated a strong correlation between the expected and actual findings for the viscosity at 37 °C quadratic model. Fig. 2 displays the response surface plot and response contour plot for Y2.

4.1.3. Effect of independent variables on gelation time (Y3). Fig. 3's representation of a 2D and 3D response surface plot showed that gelation time has a positive association with X1 and X3. The inverse relationship between poloxamer 407 and gelation time was also demonstrated by a negative value for the linear coefficient of X2. With increase in the concentration of poloxamer 407 (X2), gelation time decreases.

$$Y_3 = 25.4118 + 0.125 X_1 - 12.375 X_2 + 0.25 X_3 \quad (3)$$

Since the model's p -value is less than 0.05 (<0.0001), the ANOVA for the quadratic response surface model of gelation time was determined to be significant. X1 has p -value of 0.8954 showing that factor A had significant effect on the gelation time. While p -values for X2 and X3 were <0.0001 and 0.7928 showing that it has significant effect on the gelation time. The projected R^2 (0.8836) and adjusted R^2 (0.9155) for the gelation time linear model both showed discrepancies smaller than 0.2, showing a significant correlation between the results of the experiment and those expected.

4.1.4. Optimization and validation of the model. Based on the desired range of response values, independent factors were obtained using statistical and graphical analysis. An optimal *in situ* implant formulation was chosen using the Design Expert Software point prediction based on the desirability factor's proximity to 1. It is determined that the optimized batch with the highest desirability of 1 was chosen, with X1 = 0.5% (chitosan concentration), X2 = 20% (poloxamer 407), and X3 = 105 s (stirring time). The model was validated by observing the responses of an optimal formulation that was developed. The projected and obtained results (viscosity at 25 °C-152.6 cPs, viscosity at 37 °C-8312.6 cPs, and gelation time-26 s) are in good agreement, demonstrating the optimized *in situ* implant formulation's rationality.



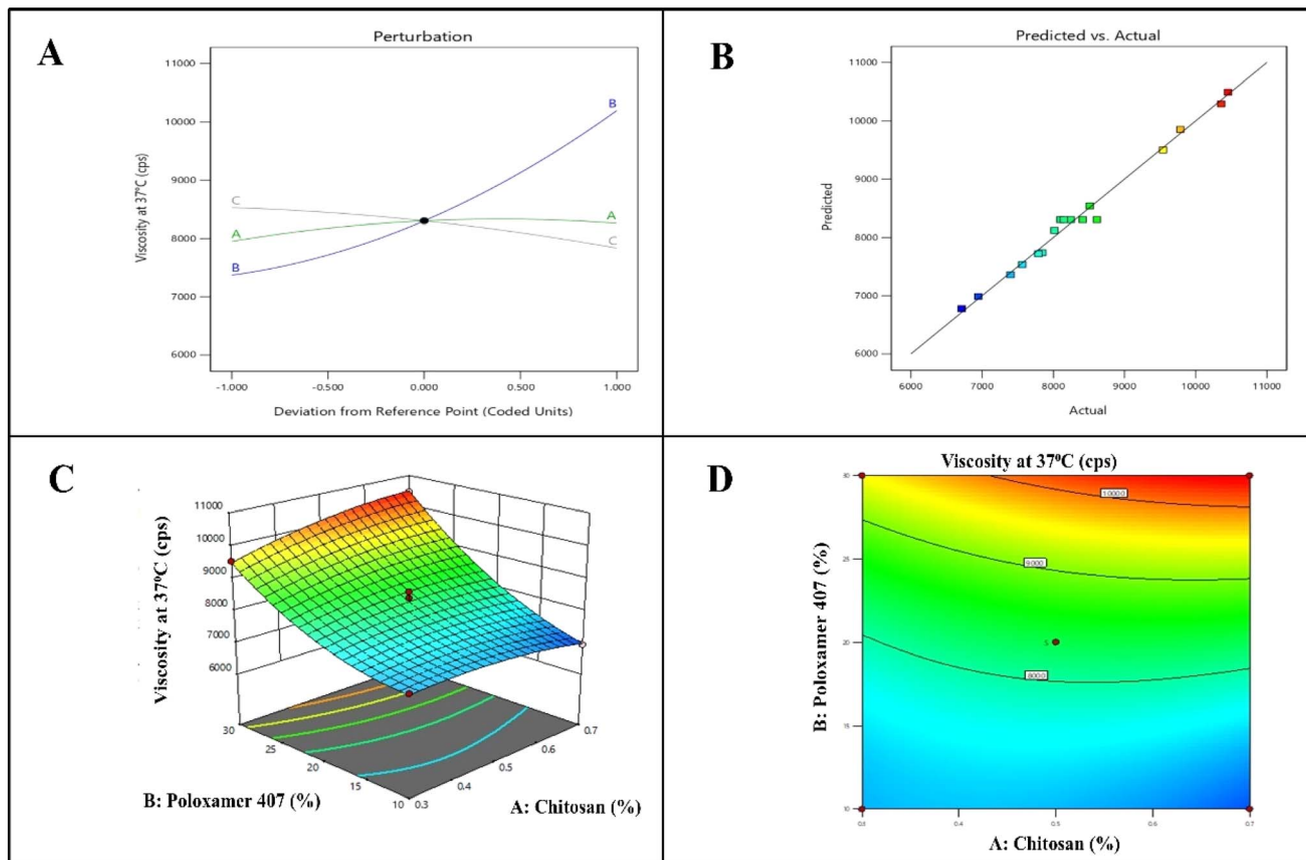


Fig. 2 QBD model graph for viscosity at 37 °C. (A) Perturbation curve, (B) linear correlation plot, (C) response surface plot, (D) contour plot.

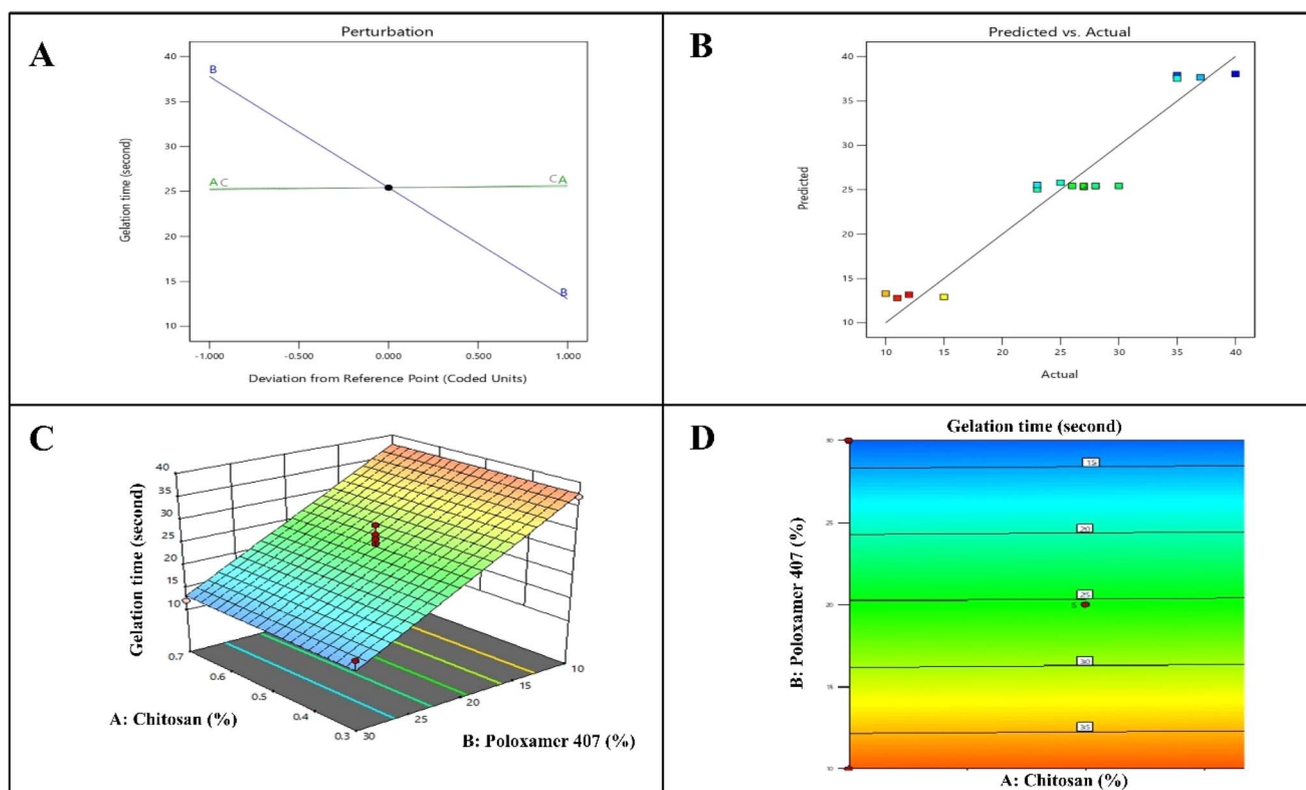


Fig. 3 QBD model graph for gelation time (A) Perturbation curve, (B) linear correlation plot, (C) response surface plot, (D) contour plot.



4.2. Preparation and physicochemical characterization of DOX-CH-PO *in situ* implant

The *in situ* gel was prepared by optimizing the parameters. The *in situ* gel was formulated using poloxamer 407 (20% w/v) dispersed in distilled water which was kept in the refrigerator at 4 °C for 24 hours for completely dissolving. Chitosan (0.5% w/v) was dissolved in a 2% w/v acetic acid solution. Aqueous solution of doxorubicin was added to the chitosan solution with continuous stirring. Finally, at room temperature, this drug solution was added to the poloxamer solution, it provides ease of injection and *in situ* gelation at the injection site.

4.2.1. FTIR. FT-IR analysis was performed to ascertain whether there is an interaction between poloxamer 407, chitosan, and doxorubicin in the *in situ* gel or not. The FT-IR spectra of poloxamer 407, chitosan, doxorubicin, and formulation are displayed in Fig. 4A. The characteristic peaks of doxorubicin of FTIR spectra at 3747.98 cm^{-1} (O-H stretch), 3313.11 cm^{-1} (N-H stretch), 2894.63 cm^{-1} (C-H stretch), 1729.83 cm^{-1} (C=O stretch), and 1412.62 cm^{-1} (C=C ring stretch). The peak pattern in the formulation spectrum was precisely the same as that of the pure drug. Considerable surface modification is depicted by a small alteration in the fingerprint region's peak intensity. The formulation had several distinct peaks at approximately 1100 cm^{-1} , which corresponded to the typical peaks of poloxamer 407. The spectra of the *in situ* implant

formulation show all of the doxorubicin's distinctive peaks, indicating that there has been no chemical alteration.

4.2.2. DSC. The DSC thermogram of pure doxorubicin showed an endothermic peak at 225.48 °C. Due to the amorphous nature of chitosan, it does not show any peak (Fig. 4B). So, the thermogram of the formulation has not shifted, indicating no change in the physical state between the drug and polymers. It revealed that the drug is in crystalline form in the formulation.

4.2.3. XRD. Fig. 4C display XRD data of poloxamer 407, chitosan, doxorubicin, and formulation. XRD spectrum of doxorubicin indicates peak position (2θ) at 16.49°, 21.79°, 22.43°, and 24.92°, which is crystalline whereas chitosan and poloxamer 407 showed amorphous nature. The XRD results for the formulation showed a peak intensity that was noticeably close to the plain drug. It indicates that the drug retains its property in the formulation and has no change in the physical state.

4.3. Pharmaceutical characterization of DOX-CH-PO *in situ* implant

4.3.1. *In vitro* gelation time. For *in situ* gel systems, gelation time is an essential feature. In general, a shorter gelation period would be preferable to reduce the amount of time needed for the implanted dosage to solidify into a thick gel. Using the test tube inversion method, the behavior of the sol-gel transition

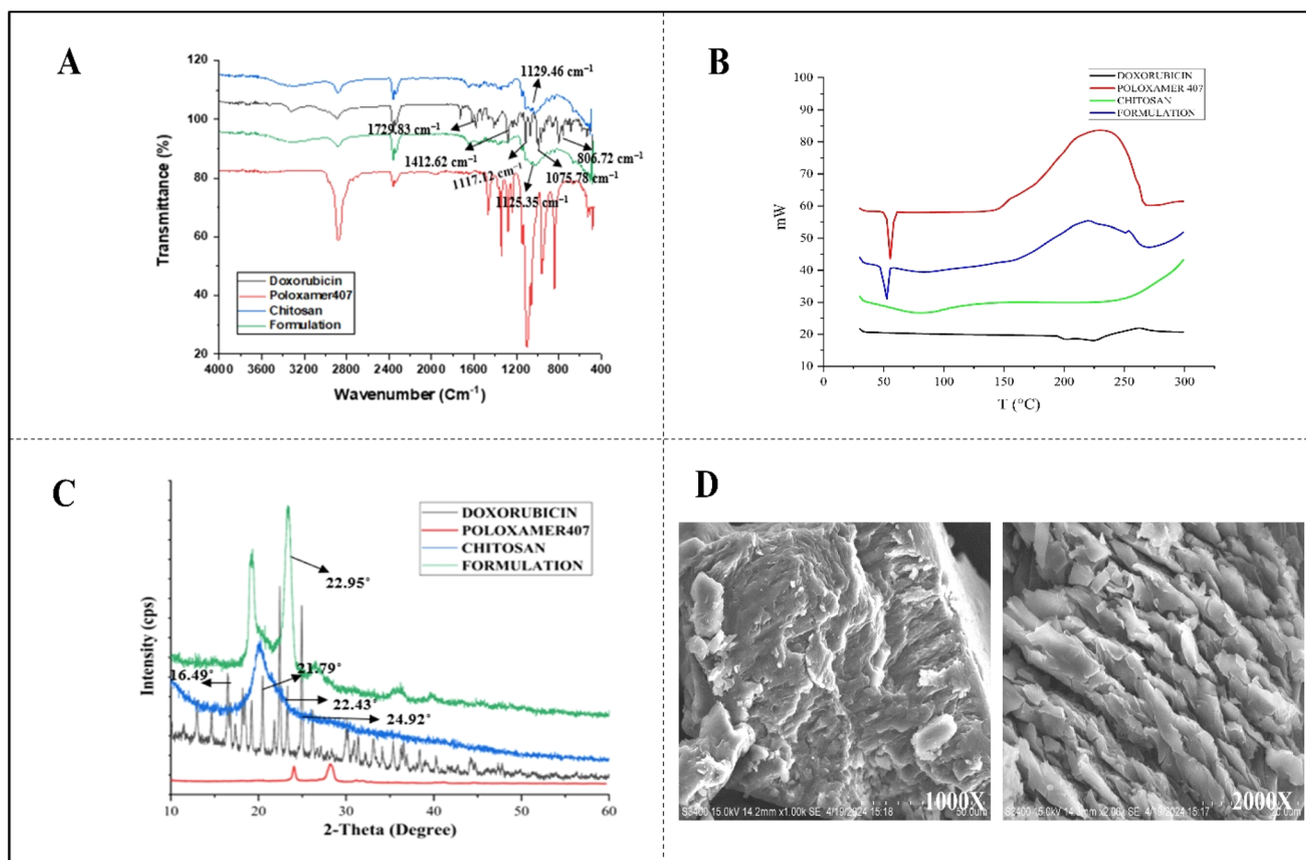


Fig. 4 (A) FT-IR spectrum, (B) DSC thermogram, and (C) XRD spectrum depicting peaks of doxorubicin, poloxamer 407, chitosan, and formulation (D) SEM images of optimized DOX-CH-PO ISI at different magnification.



was found. At 37 °C, the optimum preparation in the vial took $<26 \pm 0.2$ s to stop flowing, indicating rapid gelation which would help to develop a depot drug delivery system. The thermoresponsive behavior of poloxamer 407 and the interaction with chitosan leads to a depot formation at physiological temperatures.³⁸ Z. Fathalla *et al.* reported that a combination of poloxamer 407 with chitosan showed a gelation time of 17.9 ± 3.5 s for ophthalmic delivery systems.³⁹ The optimized formulation was further characterized for various physicochemical parameters.

4.3.2. pH. A pH meter calibrated using standard buffers of pH 4 and pH 7 under the protocol was used to check the optimized batch's pH. The developed formulation was measured using a digital pH meter and was found to be 6.81. This pH value is compatible with the site and there would be no irritation as the pH of solid tumors is typically between 6.5 to 7.2, indicating a slightly acidic environment.⁴⁰

4.3.3. Investigation of rheological properties. Viscosity has a great influence on thermosensitive *in situ* gels. As indicated, a formulation should ideally have a low viscosity at application and a high viscosity afterward to enable it to remain at the application site. The prepared formulations F1–F17 of the temperature-triggered *in situ* implant were used to measure the viscosity of the preparation by using Brookfield Viscometer (DV-II + Pro). After raising the temperature from 25 to 37 °C, the viscosity of the prepared formulation was measured in the range of 6712.26 to 10 453.3 cPs. The optimized batch showed a viscosity of around 8312.6 ± 114.2 cPs at 37 °C. When chitosan interacts with poloxamer 407 through hydrogen bonding and electrostatic interactions, it enhances the formulation viscosity.⁴¹ Upon increasing the temperature, poloxamer 407 forms micelles due to the hydrophobic nature of poly(propylene oxide) blocks, and these micelles aggregate into a gel network at body temperature.⁴²

4.3.4. Morphological study of the prepared *in situ* implant. The morphology study of the optimized batch was carried out in SEM (Hitachi S300N) at two different magnifications. At 1000× magnification, the gel matrix surface gave the impression of

being rough and corrugated as displayed in Fig. 4D. At 2000× magnification, it showed cylindrical crystal surface morphology but a structure with a multilayer. Phase separation or polymer separation was not evident.

4.3.5. *In vitro* degradation analysis. Fig. 5 shows the biodegradability of the optimized formulation in the presence of a lysozyme enzyme. The findings showed that the formulation underwent biodegradation due to the reduced sugar released from the samples. Maximum weight loss values of $74.24 \pm 2.2\%$ (pH = 5.5) and $67.58 \pm 1.7\%$ (pH = 7.4) were observed when the degradation rate was progressively raised for up to 11 days due to the presence of chitosan and poloxamer 407. The existence of all the hydrophilic water-soluble components and their great biodegradability supports the current finding. This gel's ability to adsorb lysosomes is explained by several functional hydrophilic NH₂, OH⁻ and COOH groups in these substances. At pH 5.5, chitosan is in a more protonated and soluble form, leading to a faster degradation process than at pH 7.4.^{43,44}

4.3.6. *In vitro* drug release studies. Fig. 6 displays the cumulative percent of DOX release *versus* time profiles for pure DOX, and DOX–CH–PO ISI. At pH 7.4, it was observed that about $85.32 \pm 1.2\%$ of DOX was released from pure DOX over 72 hours whereas $29.01 \pm 2.1\%$ of DOX was released from the gel within 16 h and $58.44 \pm 2.5\%$ after 72 hours displaying a slow-release profile. In contrast, there was $76.52 \pm 1.3\%$ of DOX release over 72 hours from the gel at pH 5.5. A controlled release of doxorubicin was seen in both pH conditions. The drug doxorubicin exhibits a notable increase in release at pH 5.5, owing to its improved solubility under slightly acidic pH circumstances. In pure DOX, the release pattern depends upon the intrinsic physicochemical nature of the drug. Additionally, the developed formulation formed a rigid matrix, slowing the drug's diffusion out of the gel. The combination of the thermosensitivity of poloxamer 407 and the pH sensitivity of chitosan affects the delayed drug release pattern. The gelation helps to localize the drug at the tumor site, creating a depot that slowly releases the drug over time, ensuring a sustained therapeutic effect.⁴⁵ This

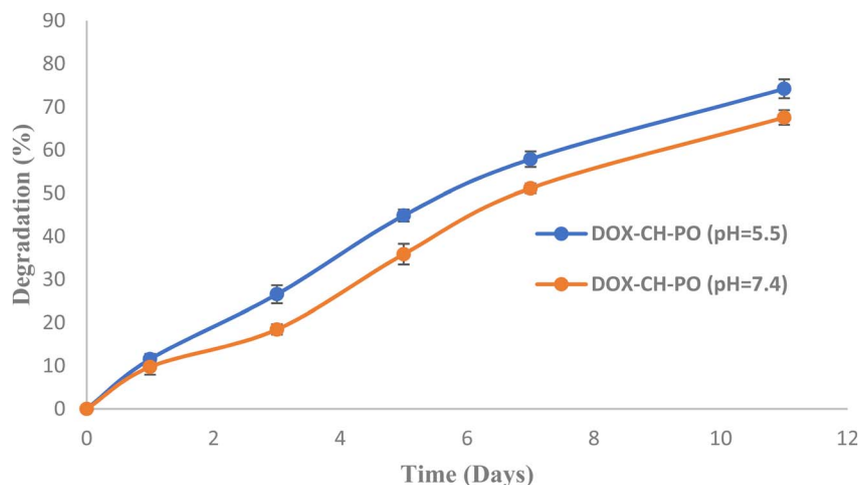


Fig. 5 Degradation percentage of DOX–CH–PO ISI at pH = 5.5 and pH = 7.4. Values represent mean \pm SD ($n = 3$).



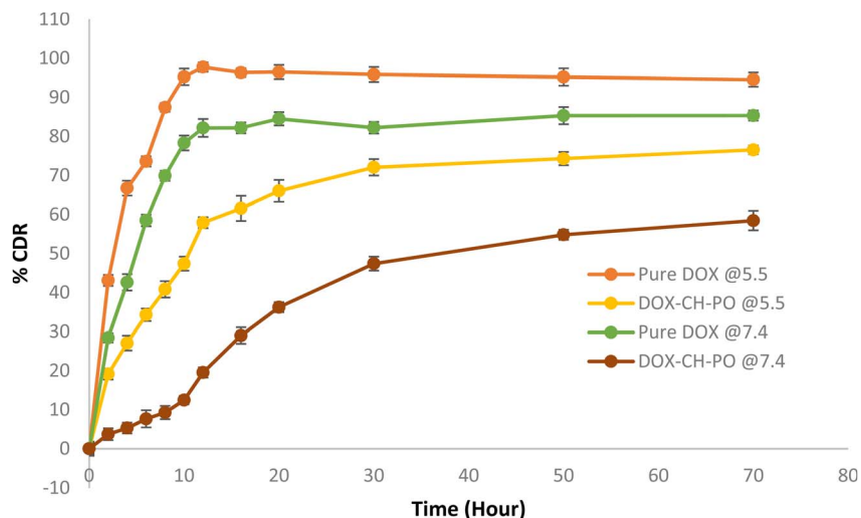


Fig. 6 *In vitro* drug release of pure DOX and DOX-CH-PO ISI. Results are presented as the mean \pm SD, $n = 3$.

delayed release helps maintain the effective concentration for a longer period, reduce dose dependency, and increase drug efficacy. J. Varshosaz *et al.* reported similar results in the combination of chitosan with poloxamer 407 showing sustained release of the drug for the ocular delivery of ciprofloxacin.⁴⁶

4.4. Biological characterization of DOX-CH-PO *in situ* implant

4.4.1. Cytotoxicity assay. *In vitro* cell viability or cytotoxicity of prepared DOX-CH-PO ISI and free DOX in MCF-7 cells were investigated. The cell growth inhibition activity of free DOX and

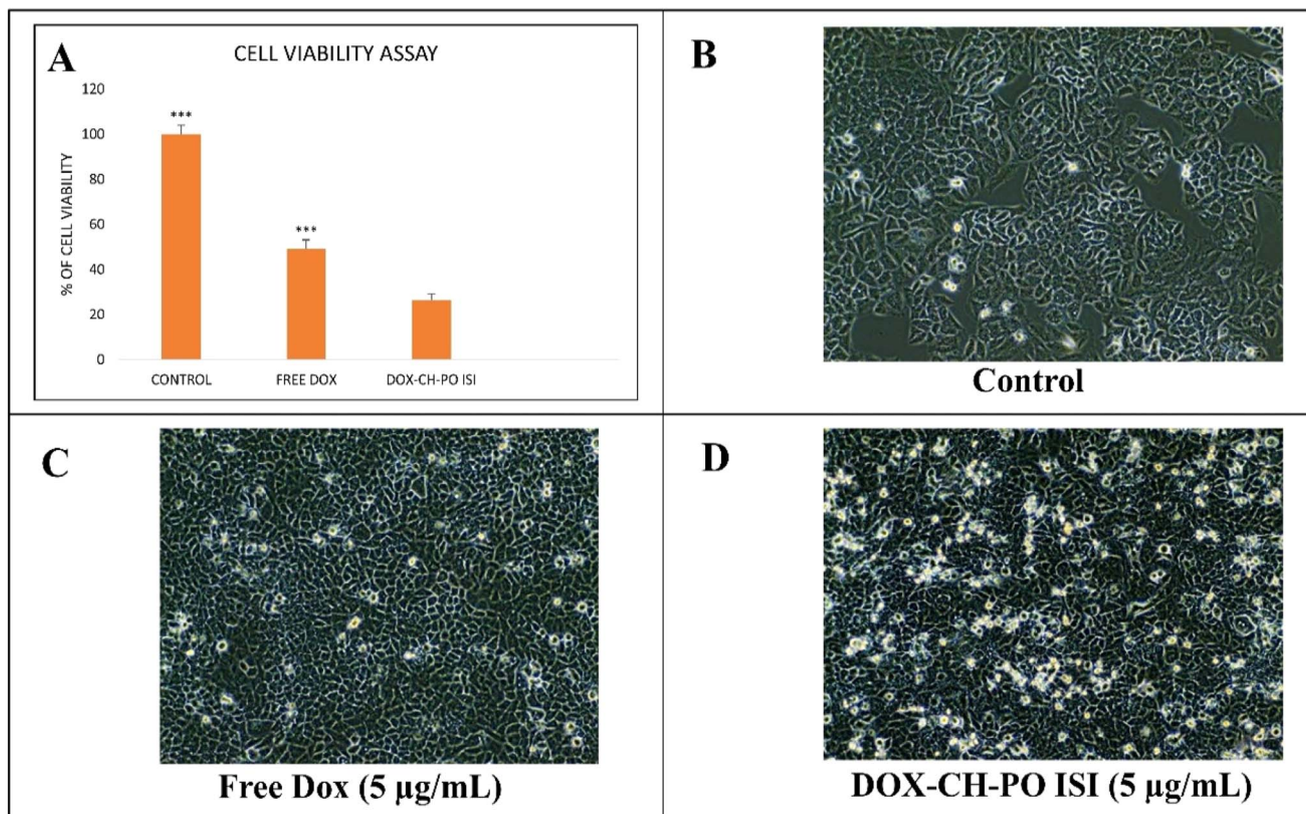


Fig. 7 (A) Effect of free DOX and DOX-CH-PO ISI on the viability of the MCF-7 cells (B) microscopical images of cells of control (C) microscopical images of cells of free DOX $5 \mu\text{g mL}^{-1}$ (D) microscopical images of cells of DOX-CH-PO ISI $5 \mu\text{g mL}^{-1}$; $200\times$ magnification data represent mean \pm SD, $n = 3$ (***) $p < 0.001$.



DOX-CH-PO against breast cancer is shown in Fig. 7A. The free DOX and DOX-CH-PO ISI killed $49.09 \pm 3.9\%$, and $26.47 \pm 2.8\%$ of the MCF-7 cells respectively. When chitosan interacts with the negatively charged cell membranes of cancer cells, it

may be more likely to accumulate in tumor cells rather than in healthy tissues, reducing off-target effects. Poloxamer enhances the penetration of the drug into the cancer cells due to its ability to disrupt cell membranes, facilitating drug uptake in cells.

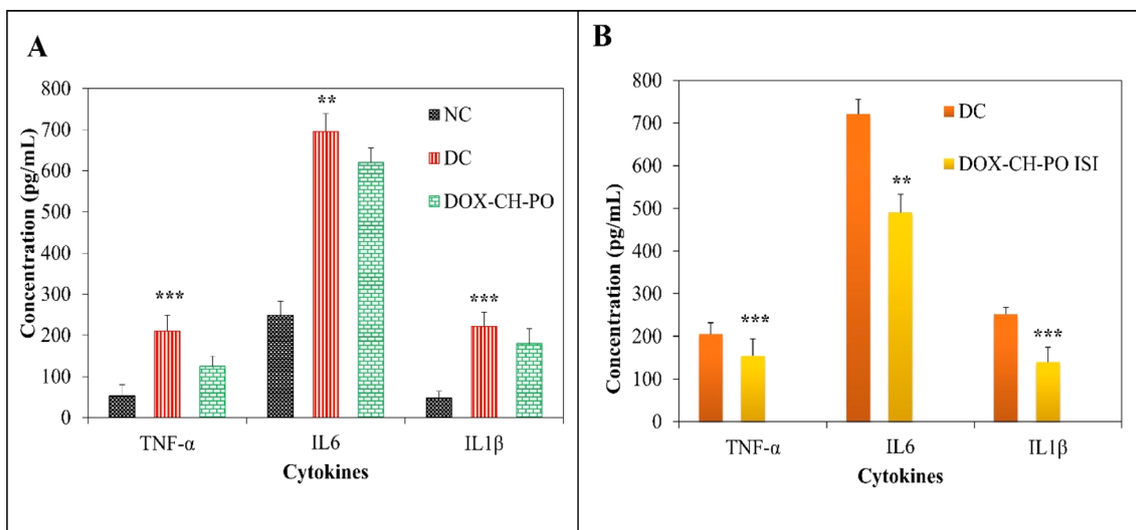


Fig. 8 (A) *In vitro* anti-inflammatory effect of normal cells, disease control cells, and DOX-CH-PO ISI. (B) *In vivo* anti-inflammatory study of DOX-CH-PO ISI on different cytokines levels. Data represent mean \pm SD, $n = 3$ (** $p < 0.01$, *** $p < 0.001$).

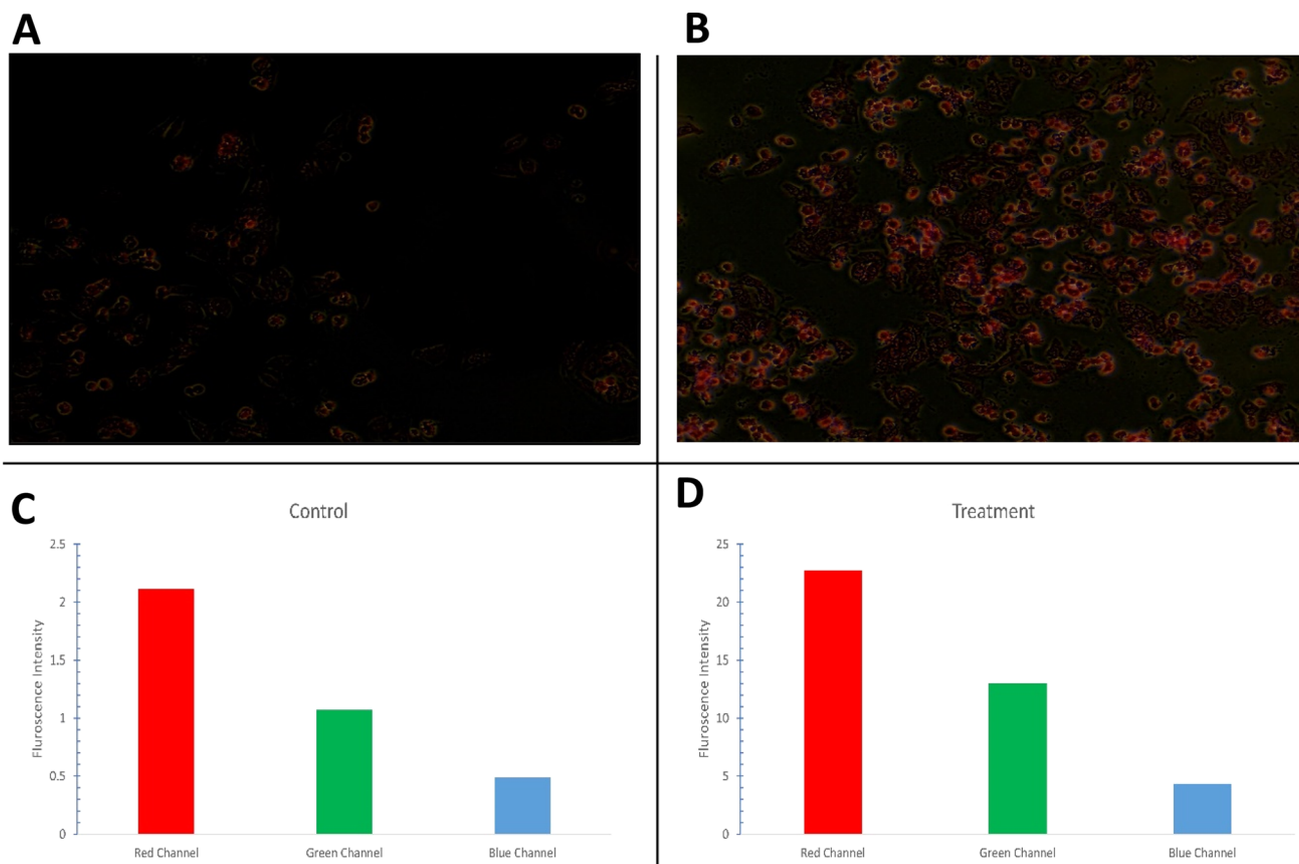


Fig. 9 *In vitro* cellular uptake of (A) negative control (B) DOX-CH-PO ISI in MCF-7 cells; 200 \times magnification (C) fluorescence intensity of control group (D) fluorescence intensity of formulation group.



This data demonstrates that DOX-CH-PO ISI exhibited better anticancer cytotoxicity as combining the drug with poloxamer and chitosan has a synergistic effect of killing the MCF-7 cells compared to free DOX. According to microscopic photographs, live cells are shaped differently from dead cells, which are bright and refractile. Typical images of living and dying cells taken under a microscope (Fig. 7C and D). Y. I. Cho *et al.* reported that the thermosensitive hydrogels made of chitosan-doxorubicin conjugates showed that chitosan-doxorubicin also had significant cytotoxicity comparable to free doxorubicin.⁴⁷ Previous reports also indicated that a combination of P407 and doxorubicin showed the synergistic killing of MC-38 tumor cells and reduced tumor cell proliferation.⁴⁸

4.4.2. *In vitro* anti-inflammatory study. The *in vitro* anti-inflammatory study of the DOX-CH-PO ISI was investigated using RAW264.7 cells. The present investigation involved pre-treating RAW264.7 cells' LPS-induced inflammatory response with DOX-CH-PO ISI. Fig. 8A indicates that the DOX-CH-PO ISI

has anti-inflammatory properties by downregulating pro-inflammatory cytokines in RAW264.7 cell lines including TNF- α , IL-6, and IL-1 β . Chitosan has excellent anti-inflammatory properties inhibiting pro-inflammatory cytokines, such as IL-1 β , IL-6, and TNF- α . Previously, it has been demonstrated chitosan attenuates inflammation through multiple mechanisms, such as altering cytokine expression, scavenging free radicals, and regulating immunological responses.⁴⁹ It has been earlier reported on the impact of chitosan and its oligosaccharides on LPS-stimulated RAW 264.7 cells, the researchers found that exposure to chitosan and its oligosaccharides resulted in a reduction of LPS-induced TNF- α and IL-6 release in the culture medium.⁵⁰ The results indicated that DOX-CH-PO ISI pre-treated LPS-induced RAW264.7 cells may serve anti-inflammatory properties and lower inflammatory factor levels.

4.4.3. Cellular uptake. The MCF-7 cell lines were used to study the cellular uptake of DOX-CH-PO ISI. As displayed in Fig. 9B, the fluorescence microscopy image demonstrated that the

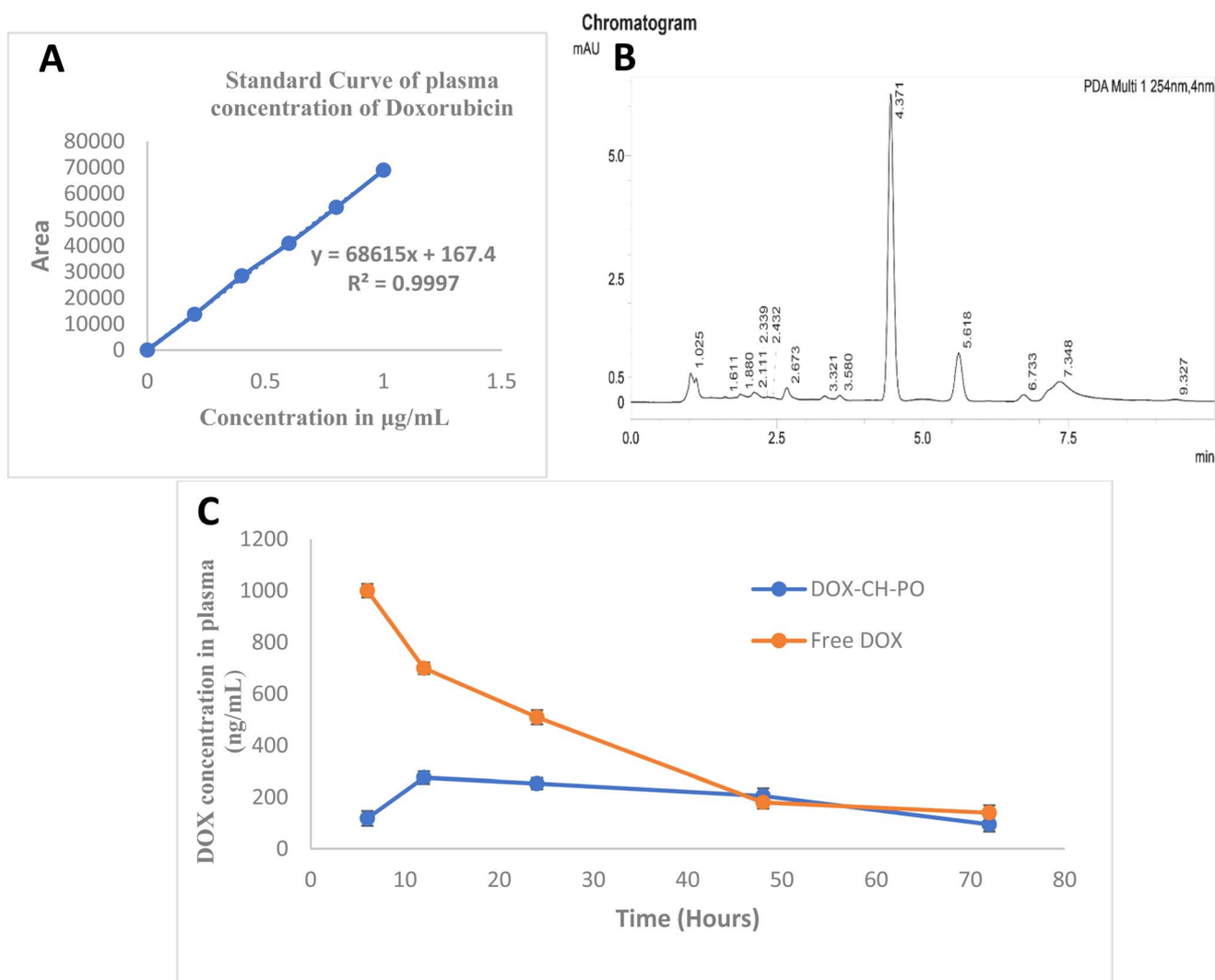


Fig. 10 (A) Standard curve of plasma concentration of doxorubicin (B) chromatogram of plasma sample after *in situ* implant administration of DOX (5 mg kg^{-1}). (C) The DOX mean plasma concentration–time profiles of the free DOX and the DOX-CH-PO ISI at a dose of 5 mg kg^{-1} DOX in rats. Data were presented mean \pm SD ($n = 3$).



Table 3 Plasma pharmacokinetic parameters of DOX in the free DOX and DOX-CH-PO ISI at an equal dose of 5 mg kg⁻¹ DOX in rats (n = 3)

Sr. No.	Pharmacokinetic parameters	Free DOX	DOX-CH-PO ISI
1	C _{max} (ng mL ⁻¹)	1000 ± 24.69	276 ± 21.86
2	T _{max} (h)	6 ± 2.45	12 ± 1.86
3	t _{1/2} (h)	22.64 ± 3.78	40.26 ± 4.92
4	AUC _{0-t} (ng h mL ⁻¹)	31 765.71 ± 245.32	14 160 ± 312.85
5	AUC _{0-∞} (ng h mL ⁻¹)	36 339.70 ± 296.87	19 678.14 ± 321.56
6	MRT _{0-∞} (h)	31.34 ± 4.52	59.93 ± 2.81
7	CL (L ⁻¹ h ⁻¹)	4.12 ± 2.84	7.62 ± 1.32

cell lines incorporated DOX-CH-PO ISI (represented by the red dots). After 4–6 hours of observation, the formulation group showed a higher mean fluorescence intensity of 23.2 (red channel) than the control group (2.1) (Fig. 9C and D). The DOX-CH-PO ISI conjugates exhibited higher fluorescence intensity, suggesting a higher amount of cellular uptake compared with the control MCF-7 cell line. The positively charged chitosan makes it easier for it to interact with the negatively charged cell membranes, which improves cell adhesion and uptake.⁵¹ Additionally, poloxamer interacts with the lipid bilayer of the cell membrane to increase permeability, which amplifies this impact even further. This combination may improve the drug's ability to penetrate cells. Recently, it has been reported that doxorubicin-loaded poly (methacrylamide) based copolymeric nanoparticles functionalized with chitosan showed significant cellular internalization.⁵²

4.4.4. *In vivo* anti-inflammatory study. Published research indicates that chitosan can reduce the production of TNF- α , IL-1 β , and IL-6 in LPS-stimulated RAW264.7 cells. In the current study, DOX-CH-PO ISI significantly reduced the levels of cytokines, including IL-6, TNF- α , and IL-1 β (Fig. 8B). The study suggests that DOX-CH-PO ISI may lessen the generation of pro-inflammatory cytokines when used in anticancer treatment as there was a decrease in the infiltration of inflammatory cells in the treated groups.

4.4.5. *In vivo* pharmacokinetics study. The objective of this study was to effectively accumulate DOX in the tumor while lowering the drug's concentration in the blood and other organs. Fig. 10A and B represent the standard curve of plasma concentration of doxorubicin and the chromatogram of the plasma sample after *in situ* implant administration of DOX (5 mg kg⁻¹) respectively. Fig. 10C shows the mean plasma concentration–time profiles of the free DOX and DOX-CH-PO ISI. Table 3 provides an overview of some key pharmacokinetic parameters. The t_{1/2} values of the free DOX and the DOX-CH-PO ISI were 22.64 ± 0.23 and 40.26 ± 0.37 h, respectively. These results suggested the t_{1/2} of DOX was prolonged from the DOX-CH-PO *in situ* implant resulting in the drug remaining in the body for a longer period, potentially extending its therapeutic effects. With an increase in the T_{max} value, it suggested that DOX from the prepared formulation has a prolonged absorption phase. The C_{max} of DOX-CH-PO ISI was found to be 276 ng mL⁻¹ compared to free DOX 1000 ng mL⁻¹. The drug is released slowly when administered *via* an *in situ* implant. The AUC_{0-t} of the developed formulation and the free DOX were 31 765.71 and 14 160 (ng mL⁻¹ × h) respectively. Therefore, the prepared *in*

situ implant decreases the likelihood of the drug entering the bloodstream which could indicate the drug remains in the tumor for a prolonged time. G. Zibin *et al.* reported similar results that there was an effective accumulation of DOX in the tumor with reduced drug concentrations in the blood and healthy organs when doxorubicin-loaded zein *in situ* gel for interstitial chemotherapy.⁵³

5. Conclusion

In the present study, a biodegradable and sustained-release DOX-CH-PO *in situ* implant was successfully prepared for the treatment of breast cancer. The *in situ* implant's physicochemical, pharmacological, and biological potential for treating breast cancer has been effectively developed and characterized. The QbD method was used to prepare seventeen formulations (F1–F17) from 3³ factorial designs and optimize the amounts of chitosan, poloxamer 407, and stirring time. The results showed that a liquid formulation that gels upon injection at 37 °C in 26 ± 0.2 s was successfully produced using 0.5% w/v chitosan and 20% w/v poloxamer 407 (optimized DOX-CH-PO ISI). The optimized formulation showed good rheological properties and extended drug release. The prepared formulation showed better cytotoxicity due to the synergistic potential of chitosan and poloxamer against the MCF-7 cells compared to the free drug. The DOX-CH-PO ISI exhibits anti-inflammatory characteristics by inhibiting pro-inflammatory cytokines, such as TNF- α , IL-6, and IL-1 β , in RAW264.7 cell lines. It has been observed that DOX-CH-PO *in situ* implant increases the cellular uptake in the MCF-7 cells. The *in vivo* pharmacokinetic the developed formulation results showed an increase in the plasma half-life, a decrease in the peak plasma concentration, and AUC which may indicate maintaining high concentrations of the drug within the tumor and less exposure of the drug to the bloodstream. Therefore, it is possible to enhance the effectiveness and decrease the systemic toxicity of chemotherapy drugs by using the sustained-release DOX-CH-PO *in situ* implant which may be a viable option for the treatment of breast cancer. Further, the intrinsic toxicological and repeated dose kinetics profiling is key to predict the clinical outcomes.

Data availability

The authors declare that the data supporting the findings of this study entitled “Doxorubicin loaded chitosan–poloxamer *in situ*



implant for the treatment of breast cancer” available within the paper. Should any raw data files be needed in another format, they are available from the corresponding author upon reasonable request. Source data are provided in this paper.

Author contributions

Guru Prasanna Sahoo: manuscript writing, data collection and proof reading. Vineet Kumar Rai: data collection, experimental work, compilation, and manuscript writing. Deepak Pradhan: proofreading, software and language editing. Jitu Halder: data analysis, language editing and critical analysis. Tushar Kanti Rajwar: compilation, manuscript writing. Ritu Mahanty: data collection and experimental work. Ivy Saha: data collection, experimentation and statistics. Ajit Mishra: data collection and experimental work. Priyanka Dash: animal experimentation and proof reading. Chandan Dash: software analysis, data collection and proof reading. Jameel Al-Tamimi: data analysis, language editing. Salim Manoharadas: data analysis, language editing and funding support. Biswakanth Kar: animal experiment and data interpretation. Goutam Ghosh: conceptualization of the topics and critical analysis. Goutam Rath: conceptualization of the topic and design of the table of content, supervision, data analysis and paper writing, editing and proof reading.

Conflicts of interest

The authors declare no conflict of interest

Acknowledgements

The DBT Builder project funded the research with order no (BT/INF/22/SP45078/2022). The authors would also like to thank the Researchers Supporting Project Number (RSPD2024R1082), King Saud University, Riyadh, Saudi Arabia. King Saud University, Riyadh, Saudi Arabia for funding this research work. The authors also acknowledge the financial support of the Department of Biotechnology (DBT), Govt. India.

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