I-Ascorbic Acid and Thymoquinone Dual-Loaded Palmitoyl-Chitosan Nanoparticles: Improved Preparation Method, Encapsulation and Release Efficiency

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Keywords: thymoquinone, I-ascorbic acid, hydrophobic modification, co-loaded nanoparticles, chitosan

Abstract:

Encapsulation of dual compounds of different characters (hydrophilic and hydrophobic) in single nanoparticles carrier could reach the site of action more accurately with the synergistic effect but it is less investigated. In our previous findings, combined-compounds encapsulation and delivery from chitosan nanoparticles were impaired by the hydrophilicity of chitosan. Therefore, hydrophobic modification on chitosan with palmitic acid was conducted in this study to provide an amphiphilic environment for better encapsulation of antioxidants; hydrophobic thymoquinone (TQ) and hydrophilic I-ascorbic acid (LAA). Palmitoyl chitosan nanoparticles (PCNPs) coloaded with TQ and LAA (PCNP-TQ-LAA) were synthesized via the ionic gelation method. Few characterizations were conducted involving nanosizer, Fourier-transform infrared spectroscopy (FTIR), field-emission scanning electron microscopy (FESEM) and high-resolution transmission electron microscopy (HRTEM). UV?VIS spectrophotometry was used to analyze the encapsulation and release efficiency of the compounds in PCNPs. Successfully modified PCNP-TQ-LAA had an average particle size of 247.7 \pm 24.0 nm, polydispersity index (PDI) of 0.348 \pm 0.043 and zeta potential of 19.60 \pm 1.27 mV. Encapsulation efficiency of TQ and LAA in PCNP-TQ-LAA increased to 64.9 \pm 5.3% and 90.0 \pm 0%, respectively. TQ and LAA in PCNP-TQ-LAA system showed zero-order release kinetics, with a release percentage of 97.5% and 36.1%, respectively. Improved preparation method, encapsulation and release efficiency in this study are anticipated to be beneficial for polymeric nanocarrier development.

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Article

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Abstract: Encapsulation of dual compounds of different characters (hydrophilic and hydrophobic) in single nanoparticles carrier could reach the site of action more accurately with the synergistic effect but it is less investigated. In our previous findings, combined-compounds encapsulation and delivery from chitosan nanoparticles were impaired by the hydrophilicity of chitosan. Therefore, hydrophobic modification on chitosan with palmitic acid was conducted in this study to provide an amphiphilic environment for better encapsulation of antioxidants; hydrophobic thymoquinone (TQ) and hydrophilic L-ascorbic acid (LAA). Palmitoyl chitosan nanoparticles (PCNPs) co-loaded with TQ and LAA (PCNP-TQ-LAA) were synthesized via the ionic gelation method. Few characterizations were conducted involving nanosizer, Fourier-transform infrared spectroscopy (FTIR), field-emission scanning electron microscopy (FESEM) and high-resolution transmission electron microscopy (HRTEM). UV–VIS spectrophotometry was used to analyze the encapsulation and release efficiency of the compounds in PCNPs. Successfully modified PCNP-TQ-LAA had an average particle size of 247.7 \pm 24.0 nm, polydispersity index (PDI) of 0.348 \pm 0.043 and zeta potential of 19.60 \pm 1.27 mV. Encapsulation efficiency of TQ and LAA in PCNP-TQ-LAA increased to $64.9 \pm 5.3\%$ and $90.0 \pm 0\%$, respectively. TQ and LAA in PCNP-TQ-LAA system showed zero-order release kinetics, with a release percentage of 97.5% and 36.1%, respectively. Improved preparation method, encapsulation and release efficiency in this study are anticipated to be beneficial for polymeric nanocarrier development.

Keywords: chitosan; co-loaded nanoparticles; hydrophobic modification; L-ascorbic acid; thymoquinone

1. Introduction

Tremendous illnesse treatments like those of cancers [1–3] and tuberculosis (TB) [4–6] require a combination of drugs to achieve positive progress, whether to cure, control or palliate the pain.



Although the standard therapies have been successful to some extent, multi-drug transportation could be eased by formulating them together rather than taking them individually, without reducing their effectiveness. Researchers have started to look at multi-drug delivery from single system formulation for various treatments [7–15]. Aside from knowing drug compatibility, a system definitely needs an outstanding carrier to content and deliver them excellently.

Nanoparticles (NPs) come to fulfil this request by acting as a promising carrier due to its large surface area that causes it to be more reactive [11,16,17]. Having a size of 10–1000 nm [18–21], NPs are classified as potent drug carrier as it can move more freely in the body. NPs uptake by cells was reported to be 15–250 times higher than 1–10 μ m particles [22]. Furthermore, NPs have been broadly used in the biomedical sector to treat diseases like diabetes [23–25] and cancers [26–29]. As NPs carrier is now drawing promising outcomes for multi-drug therapy, the present study aims at developing smart chitosan-based nanoparticles (CNPs) for combined antioxidants, hydrophobic thymoquinone (TQ) and hydrophilic L-ascorbic acid (LAA).

The use of chitosan (CS) as a polymer for NPs production has been widely discovered [13,30–32]. Abundant sources of CS from chitin of crustacean shells, fungi and insects makes it accessible and cheap [33–35]. It is one of the most auspicious polymers for drug delivery, which is biodegradable and biocompatible [33,36,37]. CS can be degraded in vertebrates by enzymatic reactions, depending on the degree of deacetylation (DD) and molecular weight (MW); low DD and MW could assist in faster CS degradation [34,38]. Moreover, the cationic property of CNPs enriched its efficacy in internalizing negatively charged cell plasma membranes [18,39]. CNPs can be synthesized by several routes, but the ionic gelation method has been more meticulously discovered, involving crosslinking reactions of CS amine groups with various anionic crosslinkers [36,40]. However, ionic gelation implementation in producing CNPs was typically done by using sodium tripolyphosphate (TPP) crosslinker [30,36,41,42].

In addition, the presence of amino and hydroxyl groups in CS makes it highly modifiable for moieties addition [43]. Palmitic acid is one of the potential modifiers for hydrophobic sites addition. Studies by Sharma et al. stated that the transfection efficiency of CS for gene delivery was successfully enhanced once modified with palmitic acid [44]. In another study, palmitic acid was used in modifying CS for an improved controlled release of tamoxifen, a breast cancer drug [45]. Kuen et al. reported on the modification of CS with palmitic acid for more effective encapsulation of hydrophobic Silibinin into NPs. The encapsulation efficiency (EE) of Sibilibin in modified palmiltoyl-chitosan nanoparticles (PCNP) was increased twice as much, compared to the EE of Silibinin in unmodified chitosan nanoparticles (CNP), because the palmitoyl anchor provides a hydrophobic core for Silibinin to stay in [41]. The concept of having both hydrophilic and hydrophobic sites in one NP system could be implemented to contain a wider range of compatible drugs or compounds to elevate synergistic effects. Therefore, this study aims at developing modified-chitosan nanoparticles by using palmitic acid *N*-hydroxysuccinimide (NHS) ester for hydrophobic thymoquinone (TQ) and hydrophilic L-ascorbic acid (LAA) antioxidants, as illustrated in Figure 1 below.

TQ is an active hydrophobic component of *Nigella sativa*, known as black seed, that shows several pharmacological properties and potential therapeutic effects such as anti-inflammatory and antioxidant properties [46,47]. LAA, known as vitamin C, is a universal antioxidant that possesses various benefits in preventing and reducing the common cold [48]. Its potential for alleviating acute viral infections treatments [49,50] was also reported. The combination of LAA and TQ was proven to have an anticonvulsant property by showing synergistic effects in lessening pentylenetetrazole-induced seizures [51]. As TQ and LAA are both antioxidants, this powerful combination can scavenge reactive oxygen species (ROS). ROS are unstable radicals containing oxygen that can easily react with molecules in a cell. Its inhibition is important to maintain good health because an excessive amount of ROS in the body could lead to oxidative stress. Prolonged oxidative stress state could then prime the emergence of complications such as neurological disorders, hypertension and acute respiratory distress syndrome [52]. Hence, by taking sufficient antioxidants in a more effective formulation, along with practicing healthy life style, numerous illnesses could be prevented and a more productive life awaits.

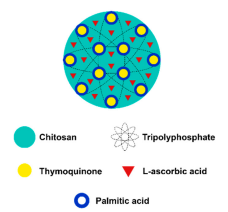


Figure 1. Proposed illustration of hydrophobically modified palmitoyl-chitosan nanoparticles encapsulated with hydrophobic thymoquinone and hydrophilic L-ascorbic acid, palmiltoyl-chitosan nanoparticles (PCNP)-thymoquinone (TQ)-L-ascorbic acid (LAA).

The present work demonstrates the synthesis and optimization of stable and homogenously dispersed palmitoyl-chitosan nanoparticles co-loaded with thymoquinone and L-ascorbic acid (PCNP-TQ-LAA). This study explores how the dual loaded compounds, hydrophobic TQ and hydrophilic LAA in a modified chitosan nanoparticles system is possible. Development and study of such systems are important, as a lot of treatment regimens involve a concoction of drugs/compounds of different physico–kinetic properties and dual administration of their combination are often plagued with many problems. This study attempts to be a proof-of-concept to pave way for finding a solution to this. As suggested in Figure 2, ammonium cations (from the chitosan chains) form ionic interaction with the phosphate anions (from the TPP crosslinker). Such interaction is possible because the positively charge chitosan can form an electrostatic attraction with negatively charge TPP. A similar interaction was suggested in previous research work [53]. Analysis on particle size, dispersion and zeta potential by the nanosizer, investigation on the presence of functional groups by Fourier-transform infrared spectroscopy (FTIR), surface morphologies by field-emission scanning electron microscopy (FESEM) and high-resolution transmission electron microscopy (HRTEM) together with encapsulation and release efficiency studies by UV-Vis spectrophotometry were conducted to characterize the NPs.

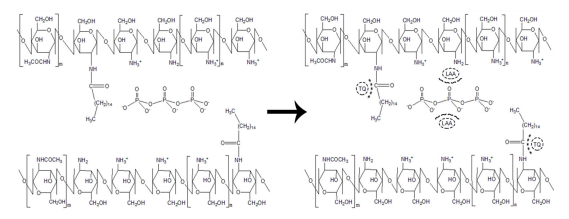


Figure 2. Proposed ionic interaction of pamiltoyl-chitosan nanoparticles, PCNP (**left**) and palmitoyl-chitosan nanoparticles encapsulated with thymoquinone and L-ascorbic acid, PCNP-TQ-LAA (**right**). "m" and "n" denote repetition of acetylated and deacetylated group of chitosan, respectively. (*) denotes hydrophobic–hydrophobic interaction between TQ and palmitic acid, while (~) denotes hydrophilic–hydrophilic interaction between L-ascorbic acid and tripolyphosphate.

2. Materials and Methods

2.1. Materials

CS with MW = 50,000–190,000 Da, \geq 75% (deacetylated) was used as a carrier, and TPP with MW = 367.86 Da was used as the crosslinking agent. Both were purchased from Sigma-Aldrich (St. Louis, MO, USA). Glacial acetic acid, sodium hydroxide pallets, 5% w/v hydrochloric acid, L-ascorbic acid, dimethylsulfoxide (DMSO), sodium bicarbonate, sodium dodecyl sulfate and methanol were purchased from R&M Chemicals (Semenyih, Selangor, Malaysia). Thymoquinone, palmitic acid *N*-hydroxysuccinimide ester and picrylsulfonic acid solution were purchased from Sigma Aldrich (St. Louis, MO, USA). Absolute ethanol was purchased from Systerm. Phosphate Buffered Saline (PBS) pellet was purchased from MP Biomedicals (Solon, OH, USA). All chemicals are of analytical grade and were used without any further purification.

2.2. Synthesis of Palmitic Acid-Modified Chitosan, PCS

For the purpose of modifying chitosan to be more hydrophobic, palmitic acid NHS ester was added in the synthesis process. Chitosan (CS) powder was dissolved in 1.0% acetic acid and distilled water to form a concentration of 1.0 mg/mL CS master solution (CS MS). Separately, palmitic acid NHS ester was dissolved in absolute ethanol to a concentration of 0.36 mg/mL. Then, the CS solution was adjusted to pH 6. Palmitic acid NHS ester solution was added into the CS solution dropwise, with a volume ratio of 1:2 and the mixture was left for 20 h at 50 °C to react. After the incubation completed, the solution was adjusted to pH 9 by using 1 M sodium hydroxide. Then, it was centrifuged at 2200× *g* for 45 min to form a precipitate. Following precipitation, it was washed once with 50:50 acetone: absolute ethanol and twice with distilled water. The supernatant was removed from each centrifugation cycle. The precipitate was dried in an oven for 72 h at 50 °C. This process produced pellet, which is called palmitoyl-chitosan (PCS). The PCS pellet will be used to synthesize PCNP, PCNP-LAA, PCNP-TQ and PCNP-LAA-TQ.

2.3. Synthesis of Palmitic Acid-Modified Chitosan Nanoparticles, PCNP

Firstly, the PCS pellet was dissolved with 1.0% acetic acid and distilled water to a concentration of 1.0 mg/mL. It was diluted to two-fold to get 0.5 mg/mL PCS working solution. Then, the solution was adjusted to pH 5 by adding 1 M aqueous sodium hydroxide solution. The crosslinker, TPP was dissolved in distilled water to make a concentration of 0.7 mg/mL and altered to pH 2.0 by using 1.0 M hydrochloric acid. PCNP was formed by adding 600 μ L of PCS to 250 μ L of TPP. Then, the mixture was centrifuged at 13,000 rpm for 20 min to get purified PCNP (only for unencapsulated PCNP). Previously, it was found that 250 μ L of TPP was an optimum volume to synthesize CNP and therefore, the same volume was used in this study to synthesize modified PCNP. That optimum TPP volume was mainly determined based on the smallest empty carrier produced [42]. This is because the expansion after encapsulation will be most minimal, which will enhance in vivo biological delivery.

2.4. Encapsulation of L-Ascorbic Acid and Thymoquinone into PCNP

For a single encapsulation process, LAA was prepared by first pouring approximately 10.0 mg of LAA into 10.0 mL of 0.7 mg/mL TPP solution, making an LAA stock concentration of 5.7 mM. Then, 250 μ L of diluted LAA was added into 600 μ L of 0.5 mg/mL PCS solution, to produce PCNP-LAA. Final LAA concentration that was used is 160 μ M. Another single-loaded PCNP, PCNP-TQ, was prepared by dissolving approximately 2.0 mg of TQ in 2.0 mL of 99.0% DMSO, making a stock concentration of 6.1 mM. Only 100 μ L of diluted TQ was added into 600 μ L of 0.5 mg/mL of PCS. Lastly, 250 μ L of 0.7 mg/mL of TPP was dropped into the PCS-TQ mixture to produce PCNP-TQ. Final TQ concentration that was used is 150 μ M.

The encapsulation of both compounds, LAA and TQ, used to make PCNP-LAA-TQ is shown in Figure 1. Basically, the LAA and TQ solutions were prepared as mentioned previously. PCNP-TQ-LAA

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was prepared by mixing 100 μ L of diluted TQ solution with 600 μ L of 0.5 mg/mL of PCS for a while. Then, 250 μ L of LAA-diluted solution was added into the mixture of TQ and PCS.

2.5. Quantification of Primary Amine Content in CNP and PCNP

Primary amine content determination was done by using a chemical called picrylsulfonic acid (TNBS reagent) according to modified methods [30,54]. The process started with preparation of the TNBS reagent, NaHCO₃, SDS and HCl at concentration of 0.05% (v/v), 0.1 M, 10.0% (w/v) and 1.0 M, respectively. First, standard solutions of CS were prepared by performing two-fold dilution of 0.5 mg/mL CS solution in 0.1 M NaHCO_3 ; each sample contained 50 μ L of CS solution and 50 μ L of NaHCO₃, serially diluted. This was followed by the addition of 50 μ L of 0.05% (v/v) TNBS. Similar steps were applied to prepare standard solutions for PCS. Second, for sample solutions, 100 µL of CNP of different TPP volume (0 to 300 µL) were added into 100 µL of 0.05% (v/v) TNBS. All standards and sample solutions were incubated at 37 $^{\circ}$ C water bath for 3 h. Next, 100 μ L of all standard and sample solutions were transferred into 96 well plates. This was followed by addition of 100 μ L of 10.0% (w/v) sodium dodecyl sulfate and 75 μ L of 1.0 M HCl into each occupied well and they were mixed well. Samples were then read with on a μ Quant microplate reader (Bio-Tek Instruments, Winooski, VT, USA) at 335 nm. Absorbance values were analyzed by using Equation (2) to find the primary amine percentage available: of CND A 1

$$Percentage of available amines = \frac{\frac{Abs. of CNP}{Abs. of PCNP}}{\frac{Abs. of CS}{Abs. of CS}} \times 100$$
(1)

2.6. Physicochemical Characterizations of PCNP Samples

2.6.1. Detection of Functional Groups in PCNP Samples

Emergence and disappearance of functional groups in nanoparticle samples were detected by FTIR. Prior to analyzing samples by FTIR, they were freeze-dried using a Coolsafe 95-15 PRO freeze drier (ScanVac, Lynge, Denmark) for 48 h to pull out liquid content. FTIR was performed using a Perkin Elmer Spectrum 100 FTIR Spectrometer (Shelton, CT, USA) with a universal attenuated total reflectance (ATR) technique to identify the functional groups in the PCNPs. Samples were measured in the range of 280–4000 cm⁻¹ at 25 °C.

2.6.2. Particle Size Distribution, PSD Study

PSD study was performed by using dynamic light scattering (DLS) technique to analyze particle size, dispersity in the sample and zeta potential. Prepared NPs were diluted with distilled water to produce 40% v/v solution. Then, the sample was analyzed by using Zetasizer 3000HSA (Malvern Instruments, Worcestershire, UK). pH of the system was around 6–6.5, near to pKa of chitosan. Three different synthesis batches (N = 3) were used in this study to obtain the average particle size, polydispersity index (PDI) and zeta potential.

2.6.3. Surface Morphology of PCNP Samples by Field-Emission Scanning Electron Microscopy (FESEM)

The surface morphologies of PCNP, PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA were observed using FESEM analysis. Samples were prepared as for DLS study, which were then sonicated for five minutes. Small volumes of samples were dropped on cleaned stubs and left to dry for three days in a 50 °C oven. The dried samples were coated with platinum by using JEOL JEC-3000FC auto fine coater (Tokyo, Japan). Finally, the samples were analyzed under an electron microscope, JEOL JSM 7600F (Tokyo, Japan). The NP diameters and particle size distributions were calculated using Image J software by National Institutes of Health (version 1.52a) from the FESEM image analysis of 50 individual particles [55].

2.6.4. Surface Morphology of PCNP Samples by High-Resolution Transmission Electron Microscopy (HRTEM)

HRTEM was used to determine morphology and clarify functions of detected structures; also, to measure particle size and check its uniformity. Firstly, samples were prepared as for DLS study, followed by sonication for fifteen minutes. Then, a tiny droplet of each sample was applied onto a formvar/carbon film, 400 mesh copper TEM grids. After that, the samples were letft to dry for 7 h under a lamp, followed by the analysis process by JEOL JEM 2100F Field Emission TEM (Tokyo, Japan).

2.6.5. Encapsulation Efficiency Study of Thymoquinone and L-Ascorbic Acid into PCNP

The encapsulation efficiency (EE) was calculated by comparing the difference in absorbance of a total compound and free compound. Total compound refers to compound solution only, while free compound refers to the compound that is unencapsulated in PCNP-TQ, PCNP-LAA or PCNP-TQ-LAA. Both solutions contained the same concentration of the compound. The % EE indicates the percentage of compound successfully encapsulated in the carrier; it was calculated using the following formula [56]:

$$Encapsulation \ efficiency, \ \% = \frac{Abs. \ of \ total \ compound - Abs. \ of \ free \ compound}{Abs. \ of \ total \ compound} \times 100$$
(2)

The absorbance was measured using Lambda 35 UV–Vis spectrophotometer (PerkinElmer, Waltham, MA, USA) at wavelengths of 257 nm and 267 nm for LAA and TQ, respectively. The wavelengths were determined based on the highest peak detected. Triplicate test (N = 3) analysis of single and dual compounds loaded in PCNP were tested.

2.6.6. Preliminary Study of Thymoquinone and L-Ascorbic Acid In Vitro Release from PCNP

The in vitro release study of thymoquinone (TQ) and L-ascorbic acid (LAA) loaded PCNP formulations were conducted first by freeze-drying all samples using a Coolsafe 95-15 PRO freeze drier (ScanVac, Lynge, Denmark) for 48 h. Then, the pellet was loaded into a quartz cuvette containing phosphate-buffered saline (PBS) pH 7.4. The cuvette was later inserted into a UV–Vis spectrophotometer (PerkinElmer, Waltham, MA, USA) and set to wavelengths of 257 nm and 267 nm for LAA and TQ, respectively. The sample was left in the UV–Vis spectrophotometer for 48 h, with auto data recording set for every 1 min (time drive option).

A calibration curve for each compound is needed in order to convert the absorbance reading to the concentration of the compound released. After considering the dilution factor, the percentage of compound released (%) could then be measured using the following formula:

Compound released,
$$\% = \frac{Released \ compound \ concentration}{Encapsulated \ compound \ concentration} \times 100$$
 (3)

Obtained release profiles were analyzed with zero order, first order, Higuchi, Hixson–Crowell and Korsmeyer–Peppas models.

Zero-order kinetic model is as follows [57]:

$$m_t = m_b + k_0 t, \tag{4}$$

where m_t is the amount of compound released at time t, m_b is the amount of compound in solution before release (usually 0) and k_0 is the zero-order rate constant.

First-order kinetic model is as follows [57]:

$$\ln(m_0 - m_t) = \ln(m_0) - k_1 t, \tag{5}$$

where m_0 is the amount of compound in the formulation before the dissolution and k_1 is the first-order release rate constant.

Higuchi model is as follows [58]:

$$m_t = k_H \sqrt{t},\tag{6}$$

where k_H is Higuchi rate constant.

Hixson–Crowell model is as follows [59]:

$$\sqrt[3]{m_0} - \sqrt[3]{m_{left}} = k_{H-C}t,$$
 (7)

where m_{left} is the amount of compound left in the formulation over time *t* and k_{H-C} is Hixson–Crowell rate constant.

Korsmeyer–Peppas model is as follows [60]:

$$\log\left(\frac{m_t}{m_{\infty}}\right) = \log k_{K-P} + n\log t \tag{8}$$

where m_{∞} is the amount of compound released after an infinitive time, k_{K-P} is Korsmeyer–Peppas rate constant and n is the parameter indicative of the release mechanism.

3. Results and Discussion

3.1. Quantification of Hydrophobic Palmitic Acid Functionalization on Chitosan by (TNBS) Assay

The amount of primary amine content in chitosan (CS) before and after modified with palmitic acid are shown in Table 1 and Figure 3. At optimum TPP volume of 250 μ L, 25% of free amine content decreased from 88.35 ± 5.78% (CNP) to 62.99 ± 2.18% (PCNP). Decreased in the free primary amine groups of CS with TPP indicates increased in the crosslinking reactions between cationic amine groups of CS and anionic TPP. In other words, functionalization of the amine group of CS with hydrophobic palmitic acid NHS was a success and this outcome is similarly seen in previous studies [30,41]. This is essential to ensure adequate sites for hydrophobic thymoquinone encapsulation later. As shown in Figure 4, the palmitic acid only conjugated the amine sites of CS partially and that is crucial because some of the amine sites will be utilized by TPP crosslinker to interact during the PCNPs synthesis later [61].

Table 1. Percentage of free amine and significancy test (*t*-test) between chitosan nanoparticles (CNP) and PCNP at different sodium tripolyphosphate (TPP) volume. (a) significant **** with a *p* value of <0.0001, (b) significant *** with a *p* value of 0.0004, (c) significant **** with a *p* value of <0.0001, (d) significant **** with a *p* value of <0.0001, (e) significant **** with a *p* value of <0.0001, (f) significant *** with a *p* value of 0.0004, and (g) significant *** with a *p* value of 0.0004.

		Percentage of Free Amine (%)				<i>t</i> -Test		
Label Volume of TPP (µL)	CNP	Standard Deviation	PCNP	Standard Deviation	<i>p</i> -Value	Significance between CNP and PCNP Reading at Different TPP Volume		
а	0	100.00	0	84.76	0.77	< 0.0001	****	
b	50	97.41	1.15	77.11	2.93	0.0004	***	
с	100	94.05	0.94	74.04	0.79	< 0.0001	****	
d	150	93.47	1.59	71.61	1.78	< 0.0001	****	
e	200	92.54	2.16	64.72	2.14	< 0.0001	****	
f	250	84.87	2.81	62.99	2.18	0.0004	***	
g	300	92.38	4.70	58.17	2.43	0.0004	***	

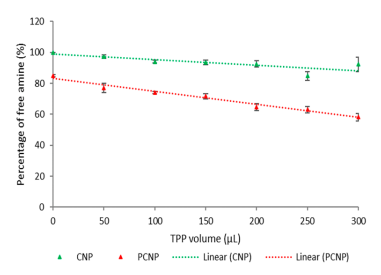


Figure 3. Percentage of free amine content in chitosan before (CNP) and after (PCNP) modification across a range of TPP volume. The free amine content in PCNP was lesser than in CNP. The loss of primary amine in CNP was due to the conjugation of palmitic acid during modification.

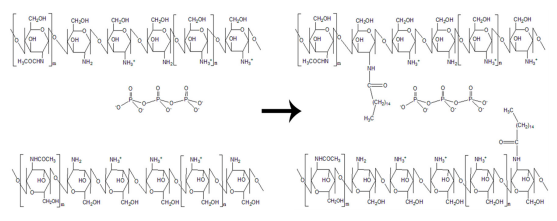


Figure 4. Conjugation of palmitic acid to amine group of chitosan. (**left**) Structure refers to unmodified CNP, and (**right**) structure refers to modified PCNP. "m" and "n" denote repetition of acetylated and deacetylated group of chitosan, respectively.

3.2. Functional Groups Determination by Fourier-Transform Infrared Spectroscopy (FTIR)

ATR-FTIR spectra of the NPs samples; PCNP, PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA are shown in Figure 5 in the range of 4000–650 cm⁻¹. A number of moieties were recognized based on the peaks and the numerical values of transmitted beams are reported in Table 2. Based on Table 2, PCNP, PCNP-LAA and PCNP-TQ-LAA have the same set of peaks. The FTIR interpretation was done by referring to the Virtual Textbook of Organic Chemistry, Infrared Spectroscopy topic by William Reusch [62].

	Functional Group	Wavenumber (cm ⁻¹)	Percent Transmittance (%)			
Peak			PCNP	PCNP-TQ	PCNP-LAA	PCNP-TQ-LAA
а	Alcohol OH (H bonded) and NH stretch (2° amine)	3399	7.47	4.15	3.17	6.18
b	Amine NH ₂ scissor (1° amine), carboxylic acid C=O (amide) and alkene C=C stretch	1647	24.20	20.24	12.46	21.96
с	Amine NH ₂ scissor (1° amine)	1563	26.91	-	10.89	18.99
d	Carboxylic acid NH (amide) bend	1542	-	62.19	-	-
e	Alkane CH ₂ and CH ₃ deformation and carboxylic acid C-O-H bend	1414	42.41	-	24.14	36.45
f	Alkane CH ₂ and CH ₃ deformation	1379	-	56.40	-	-
g	Amine C-N and carboxylic acid C-O stretch, P=O stretch	1106	3.19	9.01	8.52	7.02
h	Alkene =C-H and =CH ₂ bend	903	12.77	28.42	26.88	18.32
i	Amine NH ₂ and N-H wagging (shifts on H-bonding)	718	64.79	63.52	77.46	86.57

Table 2. Percent transmittance of functional groups existed in PCNP, PCNP-TQ, PCNP-LAAand PCNP-TQ-LAA.

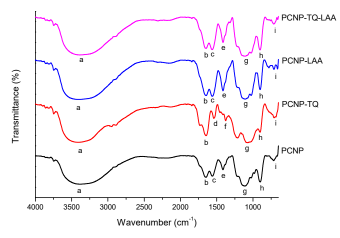


Figure 5. Fourier-transform infrared spectroscopy (FTIR) spectra of PCNP, PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA. The labelled peaks that represent certain functional groups are stated in Table 2.

PCNP-TQ, on the other hand, has extra peak d (1542 cm⁻¹) and peak f (1379 cm⁻¹) that represent carboxylic acid NH (amide) bend and alkane CH_2 and CH_3 deformation, respectively. The emergence of these peaks could be supported by the chemical structure of TQ itself (refer to Supplementary Materials, Figure S1). TQ has many C=O components that might bind with the amine of palmitic acid-NHS and CS. This answers the emergence of peak d and indirectly proves that hydrophobic TQ had successfully interacted with hydrophobic palmitic acid-NHS, which enhanced its encapsulation. Peak f in PCNP-TQ, on the other hand, emerged to resemble CH_2 and CH_3 deformation in conjunction with TQ association with palmitic acid NHS.

Peak a at 3399 cm⁻¹ appeared in all samples and it suggests an overlay of alcohol OH (H bonded) with secondary amine NH stretch. The strong and broad peak a was majorly contributed by OH (H bonded) stretch, while NH stretch was weak. For peak a, encapsulated PCNPs had stronger bands as compared to PCNP because PCNP has the least source of OH. PCNP-LAA showed the strongest band because LAA has the highest OH sites that could have bonded with the OH from TPP (refer to Supplementary Materials, Figure S1 for LAA chemical structure). On the other hand, NH contributed to the emergence of the peak a and CS played an important role in supplying NH for all PCNPs conditions.

Peak b at 1647 cm⁻¹ appeared in all samples and they are primary amine, C=O amide and alkene C=C stretch. It also represents N–C=O group resulted in the addition of palmitic acid to chitosan during the hydrophobic modification. This N–C=O group was less significant in unmodified chitosan samples as reported by Othman et al., 2018 in Table 1 [42]. Therefore, this justifies the successful insertion of palmitic acid to chitosan. Besides, similar palmitoyl-chitosan nanoparticles have been reported to exhibit palmitic acid in this range [41,45].

3.3. Particle Size Distribution by Zetasizer

The particle size distribution of all samples (PCNP, PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA) were analyzed by zetasizer. The size, polydispersity index (PDI) and zeta potential of particles were successfully determined as shown in Table 3 and Figure 6. Particle size resembles the size of nanoparticles, while PDI resembles homogeneity of particles distribution; lower PDI samples are made up of more uniform particles size and therefore, they are more monodisperse [30]. Zeta potential, on the other hand, indicates the surface charge of nanoparticles that develops at the particle–liquid interface (slipping plane).

Overall, PCNP had the smallest particle size average, 92.6 ± 8.6 nm because it has not been encapsulated with any antioxidant yet. Therefore, the crosslinked PCS with TPP resulted in the smallest particles, with a size less than 100 nm. PNCP also had the lowest PDI value, 0.277 ± 0.103 ; this indicates that it had the most even dispersion of particles as compared to other samples. Studies reported that incorporation of compounds or drugs into nanoparticles resulted in increased of particle size [45,63]. Likewise, in this study, the encapsulated nanoparticle samples, PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA showed an increment in size. PCNP-LAA had a slightly larger particles, 165.8 ± 12.9 nm than PCNP-TQ, 158.3 ± 13.9 nm because LAA has a higher molecular weight, 176.12 g/mol, while TQ is 164.20 g/mol. Furthermore, the final concentration of LAA used in the PCNP-LAA and PCNP-TQ-LAA formulation was 160 µM; which is 10 µM more than TQ encapsulated in PCNP-TQ and PCNP-TQ-LAA. Additionally, TQ also has more double bonds as compared to LAA; hence, it may have a smaller radius as it is more compact (refer to Supplementary Materials, Figure S1). For PCNP-TQ-LAA, the average particle size was the biggest, 247.7 ± 24.0 nm, and it signifies that LAA and TQ had been efficaciously encapsulated. Besides that, the PDI value of PCNP-TQ-LAA was 0.348 ± 0.043 and it is lower than PCNP-TQ, 0.374 ± 0.052 and PCNP-LAA, 0.392 ± 0.087 . This remarks that dual loaded PCNP had more unvarying particles size against single loaded PCNP-TQ and PCNP-LAA due to more complex crosslinking formation.

For zeta potential, the values among samples did not differ much. It can be concluded that the encapsulation of thymoquinone, TQ and L-ascorbic acid, LAA did not contribute to a significant change of the nanoparticle surface charge. Therefore, it proves that they did not conjugate to the nanoparticle, instead they were encapsulated. In addition, the zeta potentials of PCNPs were almost similar with reported studies, which were in the range of +20 mV to +30 mV [46,64,65]. Moreover, the positively charged PCNPs could provide better interaction with negatively charged mucosal membrane; this could later facilitate the PCNPs delivery and cellular uptake [46,66].

Table 3. Particle size, polydispersity index and zeta potential of all samples; PCNP, PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA.

Sample	Particle Size (nm)	PDI	Zeta Potential (mV)
PCNP	92.6 ± 8.6	0.277 ± 0.103	22.35 ± 1.48
PCNP-TQ	158.3 ± 13.9	0.374 ± 0.052	19.45 ± 1.20
PCNP-LAA	165.8 ± 12.9	0.392 ± 0.087	20.60 ± 1.13
PCNP-TQ-LAA	247.7 ± 24.0	0.348 ± 0.043	19.60 ± 1.27

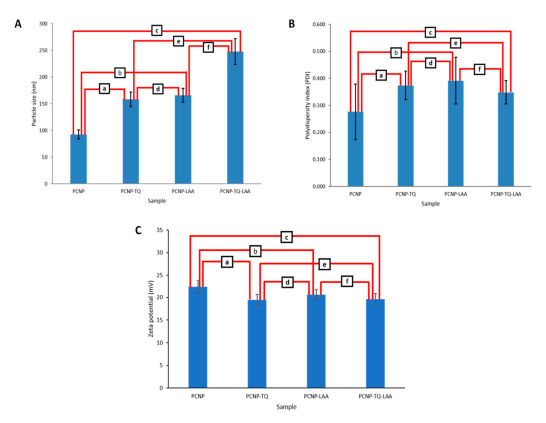


Figure 6. (**A**) Particle size of PCNP, PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA, with *t*-test conducted to see the significance of particle size changes between samples. The confidence level was set to 95% and the results are labelled as a to f. (a) Significant ** with a *p*-value of 0.0022, (b) significant ** with a *p*-value of 0.0012, (c) significant *** with a *p*-value of 0.0005, (d) not significant with a *p*-value of 0.5320, (e) significant ** with a *p*-value of 0.0051, and (f) significant ** with a *p*-value of 0.0065. (**B**) Polydispersity index and significance test (*t*-test) of PCNP, PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA. (a) Not significant with a *p*-value of 0.2168, (b) not significant with a *p*-value of 0.2129, (c) not significant with a *p*-value of 0.5437, and (f) not significant with a *p*-value of 0.4818. (**C**) Zeta potential and significance test (*t*-test) of PCNP, PCNP-TQ, PCNP-TQ, PCNP-LAA and PCNP, PCNP, PCNP-TQ, PCNP-LAA, The differences between all samples were insignificant with (a–f) *p*-values of 0.1649, 0.3161, 0.1851, 0.4284, 0.9146 and 0.4936, respectively.

3.4. Surface Morphologies of Nanoparticles

Morphologies of empty PCNP, single loaded PCNP-TQ and PCNP-LAA and dual loaded PCNP-TQ-LAA were performed by FESEM and HRTEM. FESEM images were taken at 50,000 magnification as shown in Figure 7A, while HRTEM images were taken at 10,000 magnification as shown in Figure 7C. All samples were performed at least thrice.

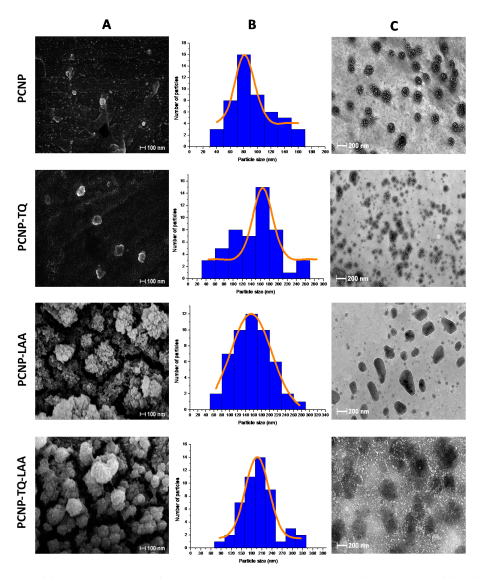


Figure 7. Field-emission scanning electron microscopy (FESEM) images (column **A**), normal distribution of particles based on FESEM images (column **B**) and high-resolution transmission electron microscopy (HRTEM) images (column **C**) of PCNP, PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA samples (from top to bottom row). Particle sizes increased as the encapsulated number of antioxidant increased (from empty PCNP to PCNP-TQ-LAA).

Based on FESEM images Figure 7A, the shape of nanoparticles is spherical with uneven surface texture. As reported by a few researchers, the spherical shape of nanoparticles establishes more efficacious therapeutics delivery process by having large surface areas [11,16,17]. The formation of particles was clearly seen and the particle sizes were in range if compared with PSD study. Figure 7B shows a histogram and normal distribution curves of particle size based on the FESEM images. The bin range for PCNP is 20 while the other bin range is 30. The average particle size of PCNP was 81.49 ± 14.33 nm. PCNP had the smallest particle size average in comparison with PCNP-TQ, 165.32 ± 21.80 nm, PCNP-LAA, 153.50 ± 51.89 nm and PCNP-TQ-LAA, 195.90 ± 34.58 nm. For PCNP-TQ particle size distribution (refer to Figure 7B, PCNP-TQ), the second most measured particles have a size range of 90–110 nm. This clarifies that there are few particles unencapsulated and it matches with encapsulation efficiency percentage of TQ as stated in Table 4, which is only $41.3 \pm 0.6\%$. On the other hand, PCNP-LAA has better particle dispersion as compared to PCNP-TQ because of the selectivity of chitosan that readily interacted with LAA which is of the same nature, hydrophilic [67].

HRTEM images, on the other hand, show layers of compounds by looking at the different color tones. Black particles represent the nanoparticles that have an average size that is almost the same as FESEM and PSD results. Besides, there are lots of tiny white pods in PCNP and PCNP-TQ which are palmitic acid micelles, the chemical used to modify chitosan to hydrophobic. The existence of palmitic acid in the system could be proven by looking at the FTIR interpretation. However, once PCNP was loaded with LAA, the white pods were less obvious and solid lines outlining those particles appeared to separate them from the surrounding. Meanwhile in PCNP-TQ-LAA, the image of particles became opaque, which indicated that they were fully occupied.

In addition, small white pods were also seen outside of the black particles. They are globular unreacted palmitic acid micelles. The amount of palmitic acid loaded might be excessive; therefore, they leached out from the NPs. It was proposed that the palmitic acid will attach to chitosan as an anchor on the surface of it; instead, it formed circular shapes. The palmitic acid anchor might be too long, which caused the formation of a circle as a result of joint ends.

3.5. Encapsulation Efficiency (EE)

This study was conducted to quantify how much of the loaded compound was successfully encapsulated. The concentration of TQ and LAA loaded in each sample PCNP-TQ, PCNP-LAA and PCNP-TQ-LAA were 150 μ M and 160 μ M, respectively. These optimized concentrations were used according to the previous study on unmodified chitosan nanoparticles done by Othman et al. (2018) in Section 3.5 [42]. According to Table 4 and Figure 8, LAA showed better EE as compared to TQ in both single and dual system PCNP with EE values of 73.0 ± 2.6% (116.8 μ M) and 90.0 ± 0.0% (143.9 μ M), respectively. These EE of LAA were much higher as compared to its EE before the modification by palmitic acid. By looking at Section 3.5, Table 4 by Othman et al., 2018, the EE of LAA in unmodified CNP-LAA and CNP-TQ-LAA were 69.3 ± 1.8% (110.9 μ M) and 22.8 ± 3.2% (36.5 μ M), respectively [42].

Table 4. The concentration of TQ and LAA loaded and encapsulated in PCNP samples.

Compound	Loaded Compound Concentration (µM)	EE (%)	Standard Deviation	Encapsulated Compound Concentration (µM)
TQ in PCNP-TQ	150	41.3	0.6	62.0
TQ in PCNP-TQ-LAA		64.9	5.3	97.3
LAA in PCNP-LAA	160	73.0	2.6	116.8
LAA in PCNP-TQ-LAA		90.0	0.0	143.9

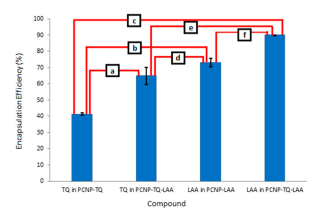


Figure 8. The percentage encapsulation of TQ in PCNP-TQ, TQ in PCNP-TQ-LAA, LAA in PCNP-LAA, and LAA in PCNP-TQ-LAA. A *t*-test was conducted to see the significance of the percentage encapsulation changes between samples. The confidence level was set to 95% and the results are labelled as a to f. (a) Significant ** with a *p*-value of 0.0097, (b) significant *** with a *p*-value of 0.0005, (c) significant *** with *p*-value of <0.0001, (d) not significant with a *p*-value of 0.0762, (e) significant ** with a *p*-value of 0.0080, and (f) significant ** with *p*-value of 0.0031.

On the other hand, only $41.3 \pm 0.6\%$ (62.0 µM) and $64.9 \pm 5.3\%$ (97.3 µM) of TQ were encapsulated in PCNP-TQ and PCNP-TQ-LAA, respectively. Lower EE of TQ compared with LAA in PCNPs was contributed by the hydrophobic nature of the compound itself, which requires stronger energy to interact with the main polymer, hydrophilic chitosan [33,46,68]. However, by having the palmitic acid modifier that holds TQ in place, the EE of TQ in the modified PCNPs were found to be higher compared to EE of TQ in the unmodified CNPs. As stated by Othman et al., 2018 in Section 3.5, Table 4, the EE of TQ in unmodified CNP-TQ and CNP-TQ-LAA were 68.7 ± 4.8% (103.1 µM) and 35.6 ± 3.6% (53.4 µM), respectively [42].

Following the modification, both TQ and LAA showed an increase in the EE for the dual loaded system. The EE of TQ in modified PCNP-TQ-LAA increased to about 29% compared to unmodified CNP-TQ-LAA. Meanwhile, the EE of LAA in modified PCNP-TQ-LAA increased to about 67% compared to unmodified CNP-TQ-LAA. This proves that the modification of chitosan by palmitic acid NHS was able to augment the EE of antioxidants. Same observations were seen in the increment of drugs EE when palmitic acid was used as a modifier in chitosan [30,41]. This clarifies that the palmitic acid is a highly potential modifier in polymeric nanocarrier for encapsulation enhancement of pharmaceuticals.

3.6. Preliminary Study of Thymoquinone and L-Ascorbic Acid In Vitro Release from PCNP

Release study was conducted to quantify how much of the encapsulated compound was successfully released from the PCNP carrier. The percentage and concentration of compounds released were included in Table 5. Release of TQ from the dual loaded system, PCNP-TQ-LAA was 37% more than the release of TQ from single loaded PCNP-TQ. LAA also showed higher release percentage from the dual loaded system as compared to a single loaded system, PCNP-LAA by around 19%. This trend indicates that when TQ and LAA were encapsulated together in a carrier, they could be released more efficiently.

Label	Release of	Released				
		Percentage (%)	Concentration (µM)	Total Time (h)		
А	TQ from PCNP-TQ	60.6	37.6	43.9		
В	TQ from PCNP-TQ-LAA	97.5	94.9	43.9		
С	LAA from PCNP-LAA	17.0	19.8	33.7		
D	LAA from PCNP-TQ-LAA	36.1	52.0	33.7		

Table 5. Release percentage and concentration of thymoquinone (TQ) and L-ascorbic acid (LAA) from PCNP samples.

The zero-order, first-order, Higuchi, Hixson–Crowell and Korsmeyer–Peppas plots were made for each release sample (A, B, C and D as labelled in Table 5). Release kinetic of a sample was determined by selecting the highest correlation value, R² from the best fit line of all models. Figure 9 summarizes the best release kinetic model for all samples. Only sample C follows the Higuchi model, while the rest follow the zero-order model. The zero-order explains the constant release rate of a pharmaceutical product. It means TQ and LAA were able to be released controllably from the dual loaded system, PCNP-TQ-LAA. This indirectly tells us that the system could help people predict how long to expect a pharmaceutical product to work.

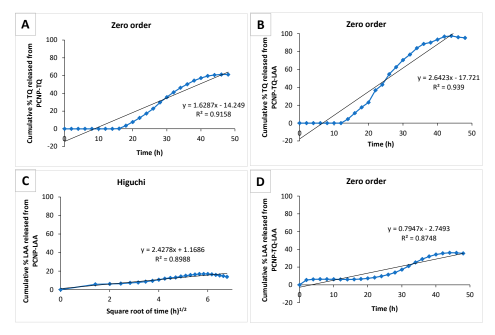


Figure 9. Release kinetics summary of TQ and LAA from all samples. (**A**) Release of TQ from PCNP-TQ, (**B**) release of TQ from PCNP-TQ-LAA, (**C**) release of LAA from PCNP-TQ-LAA and (**D**) release of LAA from PCNP-TQ-LAA. (**A**,**B**,**D**) follow zero-order release kinetics, while (**C**) follows the Higuchi release kinetic.

4. Conclusions

In this study, modification of biodegradable and biocompatible chitosan with palmitic acid NHS prior to nanoparticles formation had resulted in average particle size of around 250 nm for dual loaded thymoquinone and L-ascorbic acid palmitoyl-chitosan nanoparticles, PCNP-TQ-LAA. The robust and easy modification had also successfully increased the encapsulation efficiency of thymoquinone and L-ascorbic acid in the dual loaded system by about 29% and 67%, respectively; therefore, less loaded antioxidants were unencapsulated. This is supported by FTIR spectra of particular peaks to resemble related functional groups and also by morphology images using FESEM and HRTEM. It could also be proven that a 2:1 volume ratio of chitosan to palmitic acid NHS is remarkably appropriate to obtain partial hydrophobic sites in chitosan for more efficient encapsulation of antioxidants. The achievement of modifying this carrier with better capability of encapsulating dual classes hydrophilic and hydrophobic antioxidants for more release kinetics, the polymeric nanoparticles carrier could potentially be reused with different sets of drugs for more effective delivery.

Supplementary Materials: The following are available online at http://www.mdpi.com/2227-9717/8/9/1040/s1, Figure S1: Chemical structure of thymoquinone, TQ (left) and L-ascorbic acid, LAA (right).

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