Development of oil-in-water microemulsion encapsulating Vitamin E for thermal and hydrolysis degradation study

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Abstract

Vitamin E is widely used for medicinal and cosmeceuticals purposes. However, it is easy to degrade by the environment. In this study, the degradation of Gold Tri.E™ was studied and determined. Gold Tri.E™ is a mixture of Vitamin E homologs (tocotrienol and tocopherol) extracted from palm oil (Elais Guineensis). A nanocarrier system has been optimized to encapsulate Gold Tri.E™ from degrading and increasing its stability as a bioactive compound. An oil-in-water (o/w) microemulsion was formulated and optimized as the best carrier to encapsulate Gold Tri.E™ with the mean particle size of 32.60 ± 3.60 nm and 99.99 ± 0.01% encapsulation efficiency (EE). Degradation of the Gold Tri.E™ in o/w microemulsion was significantly reduced as compared to the bare Gold Tri.E™. This suggested that the system could protect Gold Tri.E™ from thermal and hydrolysis degradation. Thus, the ease of preparation, low-cost production, and small particle size obtained when Gold Tri.E™ encapsulated in this system give promising drug delivery system to encapsulate, protect, and increase the shelf life of Gold Tri.E™.

Keywords: Vitamin E, tocopherol, tocotrienol, degradation study, microemulsion

INTRODUCTION

Vitamin is an essential micronutrient that an organism needs in small quantities for the proper functioning of its metabolism. Vitamins can be water-soluble or lipid-soluble. Most vitamins are not single molecules, but groups of related molecules called vitamers. Vitamin E is a lipid-soluble vitamin that consists of tocopherols and tocotrienol (Sarbandi et al., 2018). The chemical structures of vitamin E homologs, which are tocotrienol and tocopherol, were depicted in Fig. 1 (a) and (b), respectively.

Fig. 1(a) Chemical structure of vitamin E homologs, which is tocopherol.

where,
- α-Tocopherol: R'≡R''≡CH3
- β-Tocopherol: R'≡CH3, R''≡H
- γ-Tocopherol: R'≡CH3, R''≡H
- δ-Tocopherol: R'≡R''≡H

Fig. 1(b) Chemical structure of Vitamin E homologs, which is tocotrienol.

where,
- α-Tocotrienol: R'≡R''≡CH3
- β-Tocotrienol: R'≡CH3, R''≡H
- γ-Tocotrienol: R'≡CH3, R''≡H
- δ-Tocotrienol: R'≡R''≡H

Primary dietary sources of vitamin E include vegetable oils, seeds, and cereal grains (Combs & McClung, 2017). It has a fundamental role as an antioxidant in the metabolism of all cells, which scavenges free radicals and protects membrane lipid from oxidation (Combs & McClung, 2017; Devereux et al., 2019). However, Vitamin E is highly susceptible to degradation in the ambient environments, such as temperature, light, and humidity (Hategekimana & Zhong, 2015).

Hence, the colloidal carrier system is one of the best alternatives that can be applied in the aim to protect and deliver
the lipophilic bioactive component to the targeted site (Aggarwal et al., 2010; Hategekimana & Zhong, 2015). Various nanocarriers have been explored to encapsulate vitamin E, such as liposome (Amiri et al., 2018), nanostructured lipid (Ying & Misran, 2017), polymeric nanoparticles (Zhang et al., 2017) and emulsion (Alayoubi et al., 2013; Yang & McClements, 2013; Raikos, 2017). In this study, we explored the formulation of microemulsion as a protective carrier for vitamin E from degradation due to the simpler laboratory preparation, which giving more advantages to scale up at the industrial level and involving the low cost of production (McClements, 2012).

Gold Tri.E™, which is a trademark product of Sime Darby Plantation Berhad, would be used as a vitamin E model. Gold Tri.E™ contains a mixture of tocotrienol and tocopherol that extracted from *Elais Guineensis* palm fruit (Heng et al., 2013). Gold Tri.E™ is yellow and commonly used as a dietary supplement, nutraceutical, functional food ingredient, and cosmeceutical (Muhammad et al., 2012). By incorporating the Gold Tri.E™ into oil-in-water (o/w) microemulsion, the Gold Tri.E™ would be expected to be protected from degradation. Furthermore, its small average particle size could enhance the delivery of Gold Tri.E™ to the targeted site and improve the efficacy of Gold Tri.E™.

**EXPERIMENTAL**

**Materials**

The Gold Tri.E™ 20P, 30P, 50, and 70 were generously provided by Sime Darby Food & Beverage Marketing Sdn. Bhd. (Selangor, Malaysia). Oleic acid and Tween 80 were purchased from Fluka (Buchs, Switzerland). All samples were prepared by using deionised water with the resistivity of 18.2 Ω cm⁻¹, collected from a Barnstead Diamond Nanopure Water Purification unit that was pre-filtered by Barnstead Diamond™ RO unit (Barnstead International, USA).

**Fourier transform infrared spectroscopy (FTIR)**

Functional group analysis of Gold Tri.E™ 20P, 30P, 50, and 70 was characterized by using Spectrum 400 Fourier Transform-Infrared (FT-IR) Spectrometer (Perkin Elmer, USA) equipped with attenuated total reflectance (ATR) accessory (GladiATR-PIKE Technologies, USA) in the Department of Chemistry, UM. Five mg of sample was placed on the clean, residue-free ATR crystal surface and scanned at 4 cm⁻¹ within the wavenumbers of 450 to 4000 cm⁻¹.

**Differential scanning calorimetry (DSC)**

Thermal behaviour of Gold Tri.E™ 20P, 30P, 50, and 70 was analyzed by using a Shimadzu DSC-60 Differential Scanning Calorimeter (Shimadzu, Kyoto, Japan). The DSC has been precalibrated with indium (Aboulfazli et al., 2014). About 10 mg of the sample was carefully transferred into Tzero Aluminum Hermetic DSC pans and placed in the DSC cell by using the empty pan as a standard pan. Both sample pans were heated to 180 °C with the scan rate of 10 °C min⁻¹ in the same single furnace. The heating run was performed with a continuous flow of nitrogen at 50.0 ml min⁻¹.

**Preparation of oil-in-water (o/w) microemulsion**

Five g of oleic acid was mixed with 10 g of deionized water with the presence of 2, 5, 10, 20, or 25% w/w of Tween 80. The mixtures were undergone mild stirring at ambient temperature (24.8±0.1°C). The o/w microemulsion obtained was left for about 30 minutes prior to analysis. The formulation of the o/w microemulsion was as depicted in Table 1. The optimum formulation was then further incorporated with 0.05% w/w of Gold Tri.E 30P. The o/w microemulsion was then incubated at room temperature for further analysis.

**Table 1 Formulation of the o/w microemulsion.**

| Formulation | Microemulsion composition (% w/w) |
|-------------|----------------------------------|
|             | Tween 80 | Oleic Acid | Water   |
| M1          | 0        | 5          | 95      |
| M2          | 2        | 5          | 93      |
| M3          | 5        | 5          | 90      |
| M4          | 10       | 4          | 86      |
| M5          | 20       | 4          | 76      |
| M6          | 25       | 4          | 41      |

Mean particle size and polydispersity index (PDI) measurement

The mean particle size and polydispersity index (PDI) of microemulsion were measured by employing the Zetasizer Nano ZS (Malvern Instruments, U.K.). A 1 cm path length quartz cuvette was used for the measurement of the mean particle size and PDI measurement. The sample was equilibrated for 2 minutes at 25.0±0.1 °C and the measurement was done in triplicates.

**Encapsulation efficiency of Gold Tri.E™**

The supernatant at the bottom of the vials was carefully collected and analyzed by a UV-Vis spectrophotometer. The amount of free Gold Tri.E™ 30P was determined from the standard calibration curve of Gold Tri.E™ 30P at 295 nm with the coefficient of determination, $R^2$, of 0.9940. Standard calibration curve of Gold Tri.E™ 30P is available upon request. Encapsulation efficiency (%EE) for each formulation was determined by using equation (1).

$$\%EE = \left( \frac{W_f - W_p}{W_f} \right) \times 100\%$$

Degradation study by UV-Vis spectroscopic method

Degradation of Gold Tri.E™ 30P was analysed by using Cary 50 UV-Vis Spectrophotometer (Agilent Technologies, USA). The o/w microemulsion, Gold Tri.E™ 30P, and o/w microemulsion encapsulating Gold Tri.E™ 30P were carefully transferred into glass vials and incubated at 8.0, 25.0, or 45.0±0.1 °C in the absence of light for 1 hour. The amount of Gold Tri.E™ 30P presented after incubation was quantified from the standard calibration curve at 295 nm.

**RESULTS AND DISCUSSION**

**Characterization of Gold Tri.E™**

Fig. 2 shows the FTIR spectra of Gold Tri.E™ 20P, 30P, 50, and 70% containing the mixture of vitamin E homologs. The broad peak at 3331 to 3433 cm⁻¹ was attributed to the OH stretching bend. The peaks within 2925 to 2854 cm⁻¹ showed the presence of asymmetric and symmetric stretching vibrations of the -CH₂ and -CH₃ (Silva et al., 2009). The characteristic peak at 1450 cm⁻¹ showed the presence of phenyl skeletal, while methyl asymmetric bending at 1440 cm⁻¹ and symmetric bending at 1377 cm⁻¹. The phenyl skeletal stretch and methyl asymmetric bending were overlapped with each other with respect to the presence of shoulder shown in the spectra. Peaks at 1240 to 1220 cm⁻¹ were corresponded to the -CH₂ while peak
1086 cm\(^{-1}\) was for the plane bending of phenyl and peak 928 cm\(^{-1}\) was represented the trans =CH\(_2\) stretching (Man et al., 2005). There was also a strong peak shown at 1087 cm\(^{-1}\), indicating the presence of C-O stretch. An overtone peak at 1850 to 2340 cm\(^{-1}\) was a signal from the combination of bands and assigned as the ring substitution pattern of the phenyl ring. The peaks from the vibration motions, as showed in Fig. 2, reconfirmed the structure of Tri.E™ that contained Vitamin E homologs, which are tocopherol and tocotrienol.

Even at low temperatures, Gold Tri.E™ 50 and 70 were in liquid form, resulting in the absence of melting point, \(T_m\), within the tested temperatures. Gold Tri.E™ 20P and 30P displayed thermal stability with melting points at 148 and 150 °C, respectively. \(T_m\), \(T_p\), and \(\Delta T_{1/2}\) were reported in Table 2. Gold Tri.E™ 30P was selected to be encapsulated in the o/w microemulsion based on their highest \(T_m\), as it will stable at a higher temperature (Hategekimana & Zhong, 2015).

| Sample | \(T_p\) (°C) | \(T_m\) (°C) | \(\Delta T_{1/2}\) (°C) |
|--------|-------------|-------------|-----------------------|
| Tri.E 20 | 133 | 148 | 140 |
| Tri.E 30 | 134 | 150 | 141 |
| Tri.E 50 | NA | NA | NA |
| Tri.E 70 | NA | NA | NA |

Particle size and Polydispersity Index (PDI) of o/w microemulsion

Fig. 4 shows the average particle size and PDI of o/w microemulsion at 25.0±0.1 °C. The incorporation of Tween 80 was found to affect both mean particle size and PDI of the formulation. The amount of Tween 80 in the formulation was optimized to produce the small average particle size of o/w microemulsion and low PDI. Low particle size is needed to deliver deeper to the targeted site, while low PDI indicated the homogenous size of the particle (McClements, 2012). Generally, two trends were observed, the decrease in average particle size from M1, M2, to M3 (a) and the increase in average particle size from M4, M5, to M6 (b). These trends were happened due to the amount of Tween 80 in the formulation. Their particle size was depended on the Tween 80 composition, showing that the average particle size and PDI of o/w microemulsion can be engineered through the composition variation of Tween 80 in the formulations lipid (Ying & Misran, 2017).

The o/w microemulsion M3 has a transparent appearance and a mean particle size of 32.6±3.6 nm. This was agreeable with the previous report on the colloidal dispersion that tended to become transparent when the particle size fell around 30 nm (McClements, 2012). This formulation was selected to encapsulate Gold Tri.E™ 30P.
Encapsulation efficiency (%EE)

Encapsulation of Gold Tri.E™ 30P in o/w emulsion was analyzed by measuring the free Gold Tri.E™ 30P in the formulation and quantified from the standard calibration curve. The %EE of Gold Tri.E™ 30P in o/w microemulsion was calculated to be 99.99%, which may due to the optimized amount of surfactant in the formulation. The %EE is improved as compared to the previous study using different formulation (Ying & Misran, 2017). This excellent %EE was resulted from the nature of Gold Tri.E™ 30P, which is hydrophobic. This characteristic would spontaneously force the Gold Tri.E™ 30P to be encapsulated in the core of o/w microemulsion, as depicted in the graphical abstract.

Degradation study of Gold Tri.E™ in o/w microemulsion

Vitamin E has a high tendency to experience physical instability and degradation (Hategekimana & Zhong, 2015). Therefore, the degradation of Gold Tri.E™ 30P in the colloidal system was studied to provide the precaution steps during processing and storage.

Fig. 5 shows the variation in the amount of Gold Tri.E™ 30P as it was incubated in various ambient temperatures. The bare Gold Tri.E™ 30P indicated a more significant reduction in thermal degradation as compared to hydrolysis degradation. In the colloidal system, Gold Tri.E™ 30P would simultaneously undergo both thermal and hydrolysis degradations. However, M3 showed both thermal and hydrolysis stabilities against the incubated ambient temperatures. Encapsulation of Gold Tri.E™ 30P in o/w microemulsion showed protection against thermal and hydrolysis degradations, where there were small changes in the amount of Gold Tri.E™ 30P at the incubated temperature.

Thus, it emphasized that the o/w microemulsion system could be a promising system to be used as a carrier system Gold Tri.E™ 30P as it was able to protect the Gold Tri.E™ 30P from degradation.

CONCLUSION

Gold Tri.E™, which contains a mixture of Vitamin E homologs, was encapsulated in the o/w microemulsion. The encapsulation efficiency yielded 99.99%. The average particle size was 32.6±3.6 nm, which could possibly increase the surface area and penetrate deeper into the skin. The experiments showed that o/w microemulsion could successfully protect Gold Tri.E™ 30P from thermal and hydrolysis degradations.

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