Physicochemical stability of lycopene-loaded emulsions stabilized by plant or dairy proteins

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Highlights
• Lycopene-loaded emulsions were prepared with plant or dairy protein emulsifiers.
• Caseinate and pea protein-stabilized emulsions were physically stable for 14 days.
• After 14 days of incubation >65% of the lycopene remained encapsulated.
• Pea protein is an interesting alternative for dairy protein in emulsion production.
Abstract

Lycopene is a lipophilic bioactive compound that can be challenging to deliver in vivo. To mediate this, delivery strategies, such as protein-stabilized oil-in-water (O/W) emulsions, have been suggested to improve the physicochemical stability and bioavailability of lycopene. In this research, the effects of plant (soy and pea) and dairy (whey and sodium caseinate) proteins on physical stability (droplet size, charge, interfacial rheology) and lycopene retention in canola O/W emulsions (pH = 7.0, 10% oil) were compared.

Particle size distribution for sodium caseinate and pea protein-stabilized emulsions remained unchanged after 14 days of refrigerated storage, while whey and soy protein isolate-stabilized emulsions became unstable. Zeta potential was largely negative (−45 to −60 mV) for all emulsions and the lycopene concentration in plant protein-stabilized emulsions at 14 days of storage was similar to that in sodium caseinate-stabilized emulsions. Sodium caseinate formed relatively viscous films at the oil-water interface, while the other proteins formed more elastic layers. Despite this difference, both the caseinate and pea protein-stabilized emulsions were promising delivery vehicles, indicating that plant-derived proteins can be feasible alternatives to dairy emulsifiers.

![Diagram showing stability and lycopene content for whey, soy, casein, and pea proteins in O/W emulsions](image-url)
1. Introduction

Recently, there has been growing interest in enriching food products with bioactive ingredients (e.g., flavors, vitamins, antioxidants or phytochemicals) to produce a desired functionality. Lycopene is the most potent singlet oxygen quencher amongst carotenoids (Di Mascio, Kaiser, & Sies, 1989; Rao, Waseem, & Agarwal, 1998) that could be used as a naturally derived antioxidant or as a health-promoting ingredient. However, lycopene is largely insoluble in water and chemically labile. Therefore, encapsulation strategies should be considered, such as using emulsion-based delivery systems.

Dairy proteins have been extensively used for food applications, and in particular to stabilize the interface in oil-in-water (O/W) emulsions. Compared to other emulsifiers (e.g., surfactants or modified starch), dairy proteins, such as whey protein isolate (WPI) and sodium caseinate (SC), can improve the physical and chemical stability of carotenoid-loaded emulsions (Mao et al., 2009; Mao, Yang, Yuan, & Gao, 2010). The high colloidal stability is attributed to the ability of dairy proteins to form thick and sterically-stabilized interfacial layers (Dickinson, 2001). In emulsion stability, the interfacial protein layer plays a critical role in the physical stabilization process (Wilde, 2000). Amongst dairy proteins, whey proteins (mostly represented by the globular protein β-lactoglobulin) have a rigid structure, which is known to lead to different interfacial organization compared to SC (primarily β-casein), which has a flexible structure (Dickinson, 2013) and in turn may lead to different effects on the physical and perhaps chemical stability of emulsions. Besides, Cornacchia and Roos (2011) found that the different protein chemistries of WPI and SC affected β-carotene retention in O/W emulsions, with the latter protein providing a better oxidative barrier. Dairy protein emulsifiers have also proved to promote the bioavailability of bioactives. Ribeiro et al. (2006) found that interfacial WPI combined with Tween 20 or sucrose laurate improved cellular uptake of lycopene and astaxanthin, compared to emulsions stabilized with only Tween 20 in colon carcinoma cells (lines HT-29 and Caco-2). Although the mechanism of enhanced bioavailability was not elucidated, the authors alluded to potential interactions between the carotenoids and β-lactoglobulin as a possible explanation.

The drawback of using dairy proteins for producing functional food emulsions is their low sustainability and impact on the environment (Erb et al., 2016, VandeHaar and St-Pierre, 2006). Plant proteins represent a large and relatively underutilized resource that is more sustainable and requires less energy for production compared to their animal-derived counterpart (de Boer, Helms, & Aiking, 2006; O’Kane, Vereijken, Happe, Gruppen, & J S Van Boekel, 2004). Recent reviews (Shi & Dumont, 2014; Song, Tang, Wang, & Wang, 2011) have also highlighted functional properties of different biobased films from plant proteins as the utilization of such renewable proteins has gained popularity. Despite the growing interest for plant-derived proteins as emulsifiers (Chihi, Mession, Sok, & Saurel, 2016), the link with stabilization of bioactive components in O/W delivery systems is hardly ever made. Many plant proteins, including soy protein isolate (SPI) and pea protein isolate (PPI) have been reported as promising functional emulsifiers (Aoki, Taneyama, & Inami, 1980; Bengoechea, Cordobés, & Guerrero, 2006; Lam & Nickerson, 2013; Pelgrom, Berghout, Van Der Goot, Boom, & Schutyser, 2014; Phoon, San Martin-Gonzalez, & Narsimhan, 2014), yet it is still arguable whether they perform as well as dairy proteins, or even outperform them (Chove, Grandison, & Lewis, 2001). SPI and PPI are
both from commonly consumed plant sources and exhibit good emulsifying properties as they have been shown to form stable O/W droplets that were not drastically bigger compared to β-lactoglobulin-stabilized droplets (Benjamin, Silcock, Beauchamp, Buettner, & Everett, 2014). Interfacial properties of SPI and PPI have also been studied and demonstrate potential to physically stabilize O/W emulsions by forming strong viscoelastic films (Chang et al., 2015). Despite the numerous studies characterizing soy and pea protein functionality, limited work (Fernandez-Avila, Arranz, Guri, Trujillo, & Corredig, 2016; Tapal & Tiku, 2012) has been conducted specifically on SPI, consisting primarily of globular proteins glycinin and conglycinin (Chronakis, 1996), and PPI, consisting primarily of legumin and vicilin/convicilin (O’Kane et al., 2004), for improving bioactive delivery. Tapal and Tiku (2012) conducted research on curcumin and SPI complexation and found that >80% of the bioactive was retained during simulated gastric conditions. Fernandez-Avila et al. (2016) also found promising results for plant protein (SPI and PPI)-stabilized emulsions, as conjugated linoleic acid (CLA) delivery was enhanced compared against non-emulsified CLA for both proteins in a Caco-2 cell model. Despite these promising first results, it is still unknown whether plant proteins could be a valuable alternative to dairy proteins for the production of functional emulsions loaded with bioactives, such as lycopene. In fact, direct comparisons between plant and dairy proteins and the link between interfacial properties and bioactive encapsulation have hardly been touched upon.

For the design of emulsion-based encapsulation systems, we believe it is necessary to connect the physicochemical stability of emulsions with the structural organization of the oil-water interface. Consequently, the aims of this study were to determine the effect of interfacial dairy or plant protein on the: 1) physical stability (particle size and zeta potential) and 2) chemical stability (lycopene retention) of emulsions, and 3) interfacial organization (adsorption kinetics and dilatational rheology). Ultimately, we have attempted to relate these findings and provide guidelines for the design of sustainable protein-stabilized emulsion-based delivery systems.

2. Materials and methods

2.1. Materials

Canola oil and tomato paste for lycopene extraction were purchased from local supermarkets (Wageningen, Netherlands). MP Alumina N-Super I (MP Biomedicals, France) was mixed with canola oil overnight as previously described (Berton, Genot, & Ropers, 2011) to strip the oil of tocopherols and surface-active impurities. All-trans-lycopene standard, all solvents (analytical grade) and other reagents were purchased from Sigma Aldrich (Zwijndrecht, the Netherlands). Proteins were generously donated by the suppliers as follows: 97.5% purity WPI (BIPRO, Davisco, Switzerland), 80% purity SC (Sodium Caseinate S, DMV International, Amersfoort, Netherlands), and 90% purity SPI (soy protein isolate SUPRO EX 37, Solae Europe SA, Switzerland) and 80–90% PPI (pea protein isolate NUTRALYS F85, Roquette, France). Ultrapure water (Millipore Milli-Q water purification system) was used for all experiments.

2.2. Methods

2.2.1. Preparation of lycopene oil stock

Approximately 250 g of tomato paste were combined with 10 g of celite, 10 g of sodium bicarbonate, and 500 mL of an extraction solvent (1:1 v/v hexane (0.1% butylated
hydroxytoluene w/v) − ethyl acetate). The mixture was held under a stream of nitrogen and in an ice-bath while stirring at 250 rpm with an overhead IKA mixer for 1.5 hours. The mixture was then vacuum filtered with No. 1 filter paper (Whatman, United Kingdom) to separate solids from liquids, transferred to a separatory funnel, and washed with a saturated solution of sodium chloride in water. The lower aqueous phase was drained and the upper hexane layer was collected, flushed with nitrogen and rotary evaporated almost to dryness. Stripped oil (~80 g) was added to solubilize the lycopene crystals prior to transferring to a borosilicate screw top bottle. The resulting lycopene-in-oil mixture was held under a stream of nitrogen to remove residual solvent until constant weight was achieved. This entire process was repeated 10 times and individual batches of lycopene oil were pooled, prior to aliquoting into 35 mL batches, flushing with nitrogen, and storing at −20° C. The lycopene content of the stock oil was determined after dilution in hexane spectrophotometrically at 471 nm, analyzed using high-performance liquid chromatography (Kean, Hamaker, & Ferruzzi, 2008), and then compared against an all-trans-lycopene standard to identify cis- and trans- isomers (Ho, Ferruzzi, Liceaga, & San Martin-González, 2015). The resulting stock oil had a total lycopene concentration of 0.236 mg/g of oil and consisted primarily of all-trans-lycopene (~90%).

2.2.2. Preparation of the aqueous phase

WPI and SC were added to 0.01 M phosphate buffer (pH = 7) and stirred with a magnetic stir bar overnight at room temperature at 100 rpm prior to emulsification the following morning. Preliminary experiments showed SPI and PPI contained an insoluble fraction (Supplementary data, Table A.1), which negatively affected the emulsifying properties. Hence, the nonsoluble fraction of the plant protein was removed prior to emulsification. SPI or PPI were combined with 0.01 M phosphate buffer (pH = 7) and stirred for 48 h at 200 rpm at 4° C. The resulting mixtures were centrifuged at 10,000 × g for 10 min at 20° C. The supernatant was collected and centrifuged again under the same conditions. The resulting supernatant, containing the soluble protein fraction, was carefully collected and stored at 4° C prior to use. The soluble protein concentration was estimated following a standard protocol for BCA Protein Assay (Thermoscientific, 2015). The removed pellet, ~70 wt% (SPI) and ~75 wt% (PPI) of the protein isolate mass, likely contained insoluble proteins and polysaccharides. The day prior to emulsification, soluble plant protein solutions were diluted with 0.01 M phosphate buffer (pH = 7) to obtain 5 or 7 g/L of protein for SPI and PPI, respectively. The diluted solutions were stirred with a magnetic stir bar overnight at room temperature at 100 rpm prior to emulsification the next morning.

2.2.3. Preparation of lycopene-loaded emulsions

Preliminary experiments were conducted to determine the optimal concentration of protein (WPI, SC, SPI, & PPI) to use that would allow for small, physically stable emulsion droplets while limiting excess emulsifier using an adapted protocol from Berton, Genot et al. (2011). An excess protein amount ( < 30%) was selected for all systems to obtain less than one-third of protein excess in the aqueous phase, while ensuring similar emulsion droplet sizes for all samples after emulsification. The aqueous phase of emulsions made with varying concentrations (5–20 g/L) of protein was collected after centrifugation at 1840 x g for 1.5 h. The amount of protein in the aqueous phase was then estimated as previously described (Thermoscientific, 2015) at 562 nm
using a DU 720 UV–vis spectrophotometer (Beckman Coulter, Woerden, Netherlands). Selected concentrations of proteins for emulsions were determined to be 5 g/L for WPI, SC, and SPI and 7 g/L for PPI as these allowed for a small droplet size (0.1–0.2 μm) while limiting the excess protein to <30% of soluble protein (Supplementary data, Figure A.1).

Aliquots of lycopene stock oil were removed from freezer storage and placed in an ultrasonic water bath for 30 min to solubilize lycopene crystals in the oil. A coarse emulsion was prepared by mixing the lycopene oil stock (10% wt) with aqueous protein solution (90% wt) via an Ultra Turrax at 11,000 rpm for 30 s. The coarse emulsion was then immediately passed through a high pressure M-110Y Microfluidizer (Microfluidics, Massachusetts, USA) for five times at 800 bar. The freshly prepared emulsions were flushed with nitrogen and stored in borosilicate screw top vials at 4 °C, in the dark. The resulting emulsions were sampled and measured (for physical stability) and aliquoted and stored (for chemical stability) at 0, 3, 7, and 14 days. Aliquots for lycopene quantification were stored in glass vials, covered with aluminium foil, flushed with nitrogen, and stored at −20°C until tested.

2.2.4. Physical stability of emulsions
2.2.4.1. Particle size

Emulsion droplet size was measured using a static light scattering instrument (Mastersizer 2000, Malvern Instruments Ltd.; Worcestershire, UK). Non-diluted emulsion samples were directly added to an attached Hydro SM small volume sampling unit for measurement. In order to assess if samples flocculated, 1 mL of emulsion was added to 4 mL of 10% sodium dodecyl sulfate (SDS) solution in water, gently vortexed for a few seconds, and then the droplet size was measured again.

All samples were measured within an obscuration range of 12–16%. Particle size of emulsion droplets is reported as the mean (d_{3,2}) and represents the average of three independent emulsion measurements, each of which were the average of three measurements.

2.2.4.2. Zeta potential

The zeta potential of emulsions was determined by measuring the electrophoretic mobility of droplets via laser Doppler velocimetry using a Zetasizer Nano ZS (Malvern Instruments Ltd.; Worcestershire, UK). Measurements were conducted with a backscatter detection angle of 173° C and calculated following the Smoluchowski model with refractive indices of 1.330 and 1.475 for water and canola oil, respectively. Samples were diluted with ultrapure water to 1.25% (v/v) and measured after 2 min of equilibration at 25 °C with 3 measurements per sample. The zeta potential values were expressed as the average from three independent samples.

2.2.5. Chemical stability of emulsions
2.2.5.1. Lycopene extraction and quantification in emulsions

Lycopene was extracted from emulsion samples using a method previously described (Ax, Mayer-Miebach, Link, Schuchmann, & Schubert, 2003) with modifications. Precisely 3 mL of ethanol, 1 mL of saturated sodium chloride in water, and 4 mL of solvent (0.1% BHT in hexane...
w/v) were added to 1 mL of emulsion sample. The samples were then vortexed and flushed with nitrogen prior to sonication for 5 min. Following this, a Pasteur pipette was used to carefully collect the upper hexane phase. Extraction with additional solvent was repeated until the hexane phase was colorless (4 repetitions). Extracts were diluted with hexane to achieve absorbance values between 0.1–0.8, sonicated for 10 s to ensure solubilisation of lycopene crystals, and measured with a UV–vis spectrophotometer at 471 nm. The total lycopene content was calculated using a molar extinction coefficient of $1.85 \times 10^5 \text{ M}^{-1} \text{ cm}^{-1}$, which was calculated as described previously (Britton, Liaaen-Jensen, & Pfander, 2004). The chemical stability of lycopene was expressed as the relative retention of lycopene ($C_{\text{relative}}$) and the absolute lycopene content. The relative lycopene retention and the encapsulation efficiency (EE) are defined as follows (Eqs. (1) and (2)):

\[
(1) C_{\text{relative}} (\%) = \left( \frac{C_t}{C_0} \right) \times 100
\]

\[
(2) \text{EE} = \left( \frac{C_0}{C_i} \right) \times 100
\]

Where $C_t$ is the lycopene content (mg/100 g of emulsion) in the lycopene at time $t$ and $C_0$ is the lycopene present in the emulsion on day 0 of storage. $C_i$ represents the amount of lycopene initially added to 100 g of emulsion. The absolute lycopene content was determined as the lycopene content (mg lycopene/100 g of emulsion) measured at each time point. Lycopene stability was determined in triplicate from three independent emulsions. To prevent lycopene degradation, all extractions were conducted in the dark. Hexane extracts containing lycopene were flushed with nitrogen, sealed in glass vials, wrapped in foil, and held on ice up until measurement (within 30 min). Samples handled in this manner did not show differences in lycopene content when measured within this time frame.

2.3.2. Properties of protein films at the oil-water interface
2.3.2.1. Adsorption kinetics

The interfacial tension at the interface between stripped oil and aqueous protein solutions was measured using an automated drop tensiometer (Teclis, Longessaigne, France). Preliminary experiments (data not shown) indicated that there was no observable change in the adsorption kinetics of whey proteins when lycopene was present in the oil, compared to pure stripped oil (for a lycopene-to-whey protein ratio similar to that in emulsion systems). Therefore, stripped canola oil was used as the oil phase for this series of experiments. It was used to fill a 0.5 mL glass syringe, connected to a 16-gauge stainless steel needle to form a model oil droplet (surface area of 60 mm$^2$). The continuous phase was protein solutions (0.1 g/L) in 10 mM phosphate buffer (pH 7) in a 40 x 23.6 x 15 mm glass cuvette (Hellma Analytics, Jena, Germany). Protein adsorption kinetics was measured during 2-h runs to ensure equilibrium and was run in, at least, duplicate to ensure repeatability. Interfacial tension was determined by fitting the experimental data to the Young-Laplace equation. Following each experiment, needles and syringes were cleaned with a 1% detergent solution (Hellmanex, Hellma Analytics, Jena, Germany) using an ultrasonic bath. Prior to use, needles and syringes were rinsed with ethanol and copious amounts of ultrapure water.
2.3.2.2. Interfacial rheology

Following the 2-h equilibration period used to allow for protein adsorption at the oil-water interface, oscillation cycles were applied to the model drop to investigate the viscoelastic response of the protein interfacial film to dilatational deformation. The drop was subjected to amplitude sweeps (2–35%) under a constant frequency of 0.01 Hz. The dilatational elastic modulus \( E'_{d} \) and the dilatational viscous modulus \( E''_{d} \) were determined from the intensity and phase of the first harmonic of a Fourier transform of the oscillating surface tension signal, and are defined as follows (Eqs. (3) and (4)):

\[
(3) E'_{d} = \Delta g \frac{A_0}{\Delta A} \cos \delta
\]
\[
(4) E''_{d} = \Delta \gamma \frac{A_0}{\Delta A} \sin \delta
\]

Where \( \Delta \gamma \) is the change in surface tension, \( A_0 \) is the initial drop surface, \( \Delta A \) is the change in drop surface during the oscillations and \( \delta \) is the phase shift.

The loss tangent (\( \tan \theta \)) was calculated by the following equation:

\[
(5) \tan \theta = \frac{E''_{d}}{E'_{d}}
\]

2.3.3.1. Statistical analysis

All emulsions were prepared in triplicate with physical and chemical stability measurements reported as the mean and standard deviation of all measurements per emulsion type. Statistical analysis was conducted with JMP version 11 (SAS Institute Inc.; Cary NC, USA). Data were subjected to one-way analysis of variance (ANOVA) with \( \alpha = 0.05 \). The Tukey-Kramer method was conducted post-hoc for mean comparisons (\( \alpha = 0.05 \)).

3. Results and discussion

3.1. Physical stability of lycopene-loaded emulsions

All emulsions exhibited small droplet size (d_{3,2}: 0.13–0.29 \( \mu \)m) between 0 and 14 days of storage (Fig. 1) with span values between 2.17-3.16 (Supplementary data, Figure A.2). At day 0, all emulsion droplet sizes had a d_{3,2} between 0.14 and 0.24 \( \mu \)m, yet plant protein-stabilized emulsion droplets were significantly larger compared to dairy-protein stabilized emulsions. SC-, WPI-, and PPI-stabilized emulsions remained physically stable with a similar droplet size at day 14 compared to day 0 (Fig. 2), in contrast to SPI-stabilized emulsions, which exhibited a larger d_{3,2} value at day 14. In order to understand what caused this, all emulsion samples were measured with and without SDS to check for flocculation. The SC and PPI-stabilized emulsions exhibited similar particle size distributions at day 0 and day 14, with and without SDS (Fig. 2), indicating they were not subjected to flocculation. Conversely, the SPI-stabilized emulsion showed particle size distributions that exhibited a left-shift when diluted in SDS solution, both at t = 0 and 14 days indicating that some flocculation occurred. Yet, after addition of SDS the particle size distribution of SPI-stabilized emulsions was similar at day 0 and day 14, indicating that the emulsion was stable to coalescence. The particle size distribution of the WPI-stabilized emulsion...
shifted to higher values after 14 days compared to the initial measurement, which remained unchanged after SDS treatment, indicating that coalescence occurred to a limited extent.

Fig. 1. Particle size ($d_{3,2}$, left y-axis) of lycopene-loaded emulsions over time. Response values shown represent the mean ± standard deviation (n = 3), with letters denoting samples that are significantly different at a given storage time ($\alpha = 0.05$).

Fig. 2. Comparison of particle size distributions of lycopene-loaded emulsions stabilized with WPI (A), SC (B), SPI (C), and PPI (D) at day 0 (solid line, black), day 0 with 1% SDS (solid line, gray), day 14 (dotted line, black), and day 14 with 1% SDS (dotted line, blue). Identical distributions with and without SDS dilution suggest that flocculation did not occur in such samples. When Day 0 and Day 14 distributions are identical the emulsions are stable.
All samples exhibited negative initial zeta potentials between −45 and −60 mV, which did not change over the course of 14 days (Supplementary data, Fig. A.3). Large negative zeta potential values were expected as emulsions were prepared at a pH above the isoelectric point of all proteins tested (Supplementary data, Table A.1). Although zeta potential can give an indication of electrostatic stabilization, proteins are mostly known for the formation of thick, viscoelastic layers at the oil-water interface that are directly linked to their efficiency at preventing emulsion droplet coalescence (Dickinson, Owusu, Tan, & Williams, 1993), as will be discussed in more detail in the interfacial rheology section.

3.2. Encapsulation stability of lycopene-loaded emulsions

All emulsions contained around 1.4 mg lycopene/100 g emulsion directly after their preparation, and by the end of storage after 14 days they all had a relative lycopene retention >65% (Fig. 3) corresponding to >0.8 mg/100 g emulsion. The highest relative lycopene retention amongst emulsions was with SC at ~87%, closely followed by PPI, with a retention of ~81%. Both values were significantly higher than found for the WPI- and SPI-stabilized emulsions. Although the lycopene stabilities were statistically different depending on the chosen protein, we should consider that differences in droplet surface area could also have affected the physicochemical stability. SC has been reported to better protect emulsions against lipid oxidation compared to WPI, and also better than SPI (Hu, McClements, & Decker, 2003), which is in accordance with our findings. The relatively low stability obtained with WPI compared to the work of Hu may be the result of the difference in pH that was applied, 3.0 versus 7.0 used for this study: isoelectric points are ~5.1 for WPI (Alting, Hamer, de Kruif, & Visschers, 2000) and ~5.6 for SPI and PPI (Chove et al., 2001; Liu, Elmer, Low, & Nickerson, 2010). Antioxidant properties of the legume proteins may have also influenced the lycopene stability of the PPI and SPI-stabilized emulsions. Han and Baik (2008) reported that both the free (soluble) and bound (insoluble) portion of soy and peas exhibit antioxidant activity.
Fig. 3. Relative retention of lycopene, as a function of time for lycopene-loaded emulsions. Response values shown represent the mean ± standard deviation (n = 3), with same letters denoting values that are not significantly different (α = 0.05).

3.3. Properties of protein layers at the oil-water interface

3.3.1. Adsorption kinetics

Interfacial tension at the oil-water interface with proteins initially dissolved in the aqueous phase was determined and expressed as a function of time (log scale) as shown in Fig. 4. In the absence of protein, the stripped oil-water interface exhibited a constant interfacial tension at ∼36 mN/m (data not shown) and was in accordance with values previously obtained in our laboratory for stripped vegetable oil, whereas a decrease in interfacial tension over time was observed when proteins were present. SC, SPI, and PPI led to roughly similar equilibrium interfacial tensions of approximately 15.8 mN/m, 15.6 mN/m, and 15.9 mN/m, respectively, by the end of the two hour run while WPI led to a higher value at roughly 18.3 mN/m, indicating that it is less surface active in comparison to the other proteins.
Fig. 4. Adsorption kinetics of WPI (A), SC (B), SPI (C), and PPI (D) at the O/W interface as a function of time (log scale). The slope of the line correlates with the rate of adsorption to the interface. The dashed line represents the interfacial tension of the stripped oil-water interface in the absence of protein at \( \sim 36 \) mN/m.

SC appeared to have the fastest rate of adsorption, followed by the plant proteins—PPI being faster than SPI—with WPI exhibiting the slowest rate of adsorption at the oil-water interface. SC adsorbs quickly to the interface due to a relatively higher amount of nonpolar groups compared to proteins such as WPI (Dickinson, 2011, Nakai and Li-Chan, 1988). SC differs from WPI, SPI, and PPI in its structure; specifically, \( \beta \)-casein consists of flexible, random coil proteins with little secondary structure due to the number and distribution of prolyl residues, and to a lack of covalent intramolecular bonding (Dickinson, 2001), which makes caseins flexible, amphiphilic proteins. Conversely, disulfide bridges and cysteine residues in \( \beta \)-lactoglobulin, the main component of WPI, stabilize the protein’s globular tertiary structure (McClements, Monahan, & Kinsella, 1993). WPI may show the slowest rate of adsorption because of its rigidity and relatively slow conformational changes at the interface compared to SC. In particular, the random coil structure of SC allows for rapid conformational changes at the interface, and therefore faster decrease in the interfacial tension (Freer et al., 2004), whereas WPI and the globular plant proteins (SPI and PPI) have stronger intra-protein interactions that limit the rate of adsorption at the oil-water interface. Although the adsorption kinetics provide insights into potential interfacial behaviour of the different proteins, it does not predict emulsion stability. Besides, protein flexibility affects the structure of the formed interfacial films, which is investigated in more detail in the next section.

3.3.2. Interfacial rheology

In emulsions, coalescence can happen if a hole is created in the interfacial film that separates two colliding droplets. Such a rupture can be seen as a dilatational deformation, thus we tested the dilatational properties of protein-stabilized interfaces (Bos and van Vliet, 2001, Murray, 2011). With the exception of WPI, the elastic and viscous moduli of the protein layers did not have a
large dependence on the applied deformation (Fig. 5), implying that the measurements were conducted within the linear viscoelastic regime. Compared to all other samples, the SC layer exhibited substantially lower elastic moduli (Fig. 5A), and thus higher loss tangent (Fig. 5B), while the elastic moduli for WPI, SPI, and PPI all appear to be substantially higher (> 15 mN/m) than their corresponding loss moduli. This indicates that the SC layer exhibited more viscous behaviour compared to the other protein layers, which is likely due to the random coil and lack of secondary structure characteristic of SC (Dickinson, 1992). Our findings are in agreement with other studies in which SC was also reported to form viscous layers at the oil-water interface (Erni, Windhab, & Fischer, 2011) due to loose packing and weak interactions between interfacial casein proteins (Dickinson, 2001). A viscous interface, which is characteristically less dense and compact compared to an elastic one, is formed with SC primarily due to its flexibility as a protein, but also due to its hydrophobicity as SC preferentially orients along the oil phase as opposed to building adsorbed layers at the oil-water interface (Maldonado-Valderrama et al., 2005).

Fig. 5. Elastic (filled shapes) and loss (open shapes) moduli (A) and loss tangent (B) of proteins at deformations between 0.03 and 0.35. Higher loss tangent values indicate a more viscous response, while lower values indicate a more elastic behavior. Response values shown represent the mean ± standard deviation (n = 3). Statistical differences amongst protein films are shown (B) with same letters denoting values that are not significantly different (α = 0.05).

Compared to SC-based interfaces, WPI-based ones exhibited a more elastic behaviour, which can be attributed to strong intermolecular interactions and a high two-dimensional packing efficiency at the interface (Dickinson, 2001). SPI- and PPI-based layers exhibited loss tangents more similar to that of the WPI-based layer, which was expected since plant proteins are globular (Boye et al., 2010) and known to produce an interconnected, viscoelastic monolayer at the oil-water interface (Chang et al., 2015).
3.4. Comparison and design considerations for protein-stabilized emulsions

All our emulsions had similar and small droplet size, therefore, effects of interfacial area, that are reported to potentially influence chemical stability (Lethuaut, Métro, & Genot, 2002) or not (Berton-Carabin, Ropers, & Genot, 2014; Hu, McClements, & Decker, 2003; Osborn & Akoh, 2004) can rather safely be disregarded in the interpretation of the results. Besides, we designed our emulsions in such a way that the fraction and concentration of non-adsorbed proteins was low, so that the contribution of this non-adsorbed fraction to their physicochemical stability was presumably limited (Berton, Genot et al., 2011, Berton, Ropers et al., 2011, Faraji et al., 2004).

Most probably, the protein properties and the resulting interfacial layers affect lycopene stability. Steric forces influence emulsion physical stability, particularly for SC-stabilized emulsions, as electrostatic forces are expected to play a lesser role in stabilization for flexible proteins (Dickinson, 2010), while for the other less flexible proteins, thicker layers are expected to stabilize the interfaces. Hu et al., 2003a, Hu et al., 2003b discussed the amino acid composition of SC, which contains relatively high amounts of antioxidative tyrosine, proline, and methionine, as a potential explanation for improved oxidative stability of emulsions stabilized with SC compared to SPI and WPI, although they express that this relationship is not clear. In another study, high-pressure processing at 1379 bar vs. 345 bar was reported to induce a tighter packing in the cross-linked interfacial layer of SC-stabilized emulsions, which was related to a higher oxidative stability (Phoon, Paul, Burgner, Fernanda San Martin-Gonzalez, & Narsimhan, 2014). Other studies have reported that increasing processing temperature of protein-stabilized emulsions results in further unfolding of proteins and potential alteration of conformation (Let, Jacobsen, Sørensen, & Meyer, 2007). In particular, whey proteins have been reported to exhibit antioxidant properties post-homogenization due to the unfolding and exposure of sulfhydryl groups, which can either repel (Hu et al., 2003a) or scavenge free radicals (Let et al., 2007; Tong, Sasaki, Mcclements, & Decker, 2000). Aside from the antioxidant effects of protein, other impurities (e.g., polysaccharides and starches) existing in the commercial plant protein isolates could contribute to the lycopene stability. In particular, phenolic compounds in peas, which exist primarily in the hulls (Shahidi et al., 2001), may have contributed to the chemical stability of the PPI-stabilized emulsions.

From the above it is clear that interfacial properties are related to the physicochemical stability of an emulsion, which is mostly linked to providing a denser barrier against oxidizing agents and coalescence (Georgieva, Schmitt, Leal-Calderon, & Langevin, 2009), however it is difficult to find clear experimental evidence for this. As discussed previously, elastic interfaces are the result of an interconnected protein network. The gel-like viscoelastic interface observed in this study amongst WPI, SPI, and PPI-stabilized emulsions would be expected to form a rigid layer, which in theory could better physically stabilize the system and limit contact between the lipid phase and oxidizing agents. However, globular proteins may exhibit localized empty patches due to depletion (Bos & van Vliet, 2001), which potentially has detrimental consequences for lycopene stability.

Despite the mechanical and structural properties of the interface, chemical properties, such as oxygen permeability through a protein layer, should also be taken into consideration. β-casein films at the air-water interface were found to have a higher oxygen permeability compared to
that of β-lactoglobulin (Toikkanen et al., 2014), while β-casein-stabilized emulsions have been found to exhibit better oxidative stability (based on oxygen uptake and formation of conjugated dienes, hexane, and propanal) in various conditions compared to β-lactoglobulin-stabilized emulsions (Berton, Ropers, Bertrand, Viau, & Genot, 2012; Berton, Ropers, Viau, & Genot, 2011), and this is most probably caused by the fact that caseins are better at scavenging free radicals (Clausen, Skibsted, & Stagsted, 2009) and binding iron (Faraji, Mcclements, & Decker, 2004; Sugiarto, Ye, Taylor, Singh, & Singh, 2010) compared to whey proteins.

Yet, protein flexibility and interfacial elasticity alone cannot be used to simply explain the stability of lycopene-loaded emulsions. It is likely that chemical properties of the proteins aided in lycopene stability, although future work could be done to directly assess this. Especially pea protein is of great interest; given its relatively high stability and encapsulation capacity, it is expected to serve as a genuine alternative for animal-based proteins in emulsion formulations.

4. Conclusions

This work systematically investigated the physical and chemical stability of lycopene-loaded emulsions prepared using various proteins as emulsifiers. Especially emulsions stabilized with casein and pea protein exhibited both high chemical (encapsulation% > 80%) and physical stability (no change in particle size) after 14 days. Interestingly, no correlation could be found between the elasticity of the protein layers at model oil-water interfaces, and the physicochemical stability of the corresponding emulsions. This is most probably due to the fact that adsorbed casein molecules induced strong steric repulsion, resulting in an additional emulsion stabilization effect, and lycopene protection effects due to the protein ability to chelate metals ions and scavenge free radicals.

Performance of each protein could be ranked for each property measured, however, it is perhaps more valuable to consider the collective characteristics for each of the protein-stabilized emulsions. Although SC appeared to perform optimally, PPI was a strong plant contender and demonstrated comparably good properties as it stabilized emulsions against flocculation and coalescence, exhibited relatively rapid protein adsorption, and stabilized lycopene to a similar extent as SC. Overall, SC and PPI both exhibited relatively good physical and chemical stabilization for lycopene-loaded emulsions, while SPI and WPI exhibited better stabilization for either physical or chemical stabilization, rather than both (Table 1). This research demonstrates that selected plant proteins can perform well compared to dairy proteins for lycopene encapsulation and have potential as dairy alternatives for chemical protection against oxidation in colloidal systems.
Table 1. Summary comparison of physical and chemical properties lycopene-loaded emulsions stabilized with WPI, SC, SPI, or PPI. Proteins that strongly demonstrated relatively high (++++) values for a given characteristic are compared against those with intermediate (+++ or ++) and lower (+) values.

<table>
<thead>
<tr>
<th></th>
<th>Small Droplet Size</th>
<th>Physical Stability</th>
<th>Fast Adsorption</th>
<th>Highly Elastic Interface</th>
<th>Lycopene Retention</th>
</tr>
</thead>
<tbody>
<tr>
<td>WPI</td>
<td>++</td>
<td>++</td>
<td>+</td>
<td>+++</td>
<td>+</td>
</tr>
<tr>
<td>SC</td>
<td>+++</td>
<td>+++</td>
<td>++++</td>
<td>+</td>
<td>+++</td>
</tr>
<tr>
<td>SPI</td>
<td>+</td>
<td>+</td>
<td>++</td>
<td>+++</td>
<td>++</td>
</tr>
<tr>
<td>PPI</td>
<td>++</td>
<td>+++</td>
<td>+++</td>
<td>++</td>
<td>+++</td>
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</tbody>
</table>

Industrial relevance text

This work investigated the physical and chemical stability of lycopene-loaded emulsions prepared with dairy or plant proteins as emulsifiers. The outcomes are highly relevant to industry, as they constitute a basis to formulate food emulsions that not only encapsulate highly lipophilic and labile bioactives, but that are also made of sustainable ingredients (plant proteins). This addresses the current priorities of the food sector, i.e., making foods that are both healthy and sustainable.

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Appendix A. Supplementary Data

Figure A.1. Determination of optimal protein concentration. Particle size (left y-axis) and correlating percent of excess protein (right y-axis) versus protein concentration added to the emulsion for WPI (A), SC (B), SPI (C), and PPI (D). Dashed line denotes the selected protein concentration.
Figure A.2. Span of lycopene-loaded emulsions over time. Response values shown represent the mean + standard deviation (n=3), with same letters denoting values that are not significantly different (α=0.05).

Figure A.3. Initial zeta potential of lycopene-loaded emulsions fabricated with proteins and protein blends. Response values shown represent the mean + standard deviation (n=3), with same letters denoting values that are not significantly different (α=0.05).
Figure A.4. Encapsulation efficiency of lycopene in protein stabilized emulsions at t=14 days. Response values shown represent the mean ± standard deviation (n=3), with same letters denoting values that are not significantly different (α=0.05).

Table A.1. Physical properties of the different protein samples in phosphate buffer (10 mM, pH 7.0).

<table>
<thead>
<tr>
<th></th>
<th>Solubility (wt%) a</th>
<th>Particle size (nm) b</th>
<th>Zeta potential (mV) b</th>
</tr>
</thead>
<tbody>
<tr>
<td>WPI</td>
<td>~ 100</td>
<td>5.0 ± 0.1</td>
<td>-16.4 ± 1.7</td>
</tr>
<tr>
<td>SC</td>
<td>~ 100</td>
<td>13.7 ± 1.9</td>
<td>-25.8 ± 1.5</td>
</tr>
<tr>
<td>SPI</td>
<td>30.2 ± 0.4</td>
<td>112.2 ± 33.4</td>
<td>-23.2 ± 1.3</td>
</tr>
<tr>
<td>PPI*</td>
<td>25.1 ± 2.4</td>
<td>Main peak: 37.3 ± 7.0</td>
<td>-22.9 ± 1.3</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Second peak: 248.6 ± 31.8</td>
<td></td>
</tr>
</tbody>
</table>

a Percentage of soluble proteins starting with dispersing 6 wt% powdered sample in buffer. See text for details regarding the separation and determination of soluble proteins.

b Determined by dynamic light scattering (refractive indices for dispersed material and continuous phase, 1.45 and 1.33, respectively; for the Zeta potential determination the Smoluchowski model was used).

* Extensive characterization for this sample is available in the work of Pelgrom et al. (2015).
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