Use of casein micelles to improve the solubility of hydrophobic pea proteins in aqueous solutions via low-temperature homogenization

Abigail Krentz, Israel García-Cano, Joana Ortega-Anaya, and Rafael Jiménez-Flores*
Department of Food Science and Technology, The Ohio State University, Parker Food Science and Technology Building, Columbus 43210

ABSTRACT

The dairy industry struggles to maintain consumer attention in the midst of declining fluid milk sales. Current trends create an opportunity to incorporate plant-based proteins with milk to produce a high-protein, multisourced, functional food product. Plant-based proteins, such as those in peas, can be challenging to use in food systems because of their low solubility and undesirable off-flavors. Casein micelles have unique structural properties that allow for interactions with small ions and larger macromolecules that aid in their noteworthy ability as a nanovehicle for hydrophobic compounds. The objective of this study was to use the inherent structure of the casein micelle along with common dairy processing equipment to create a stable colloidal dispersion of casein micelles with pea protein to improve its solubility in aqueous solutions. We created 3 blends with varying ratios of casein-to-pea protein (90:10, 80:20, 50:50). We subjected the mixtures to 3 cycles of homogenization using a bench-top GEA 2-stage homogenizer at 27,580 kPa maintained at 4°C, followed by pasteurization at 63°C for 30 min. The resulting blends were homogeneous liquids with increased stability due to the lack of protein precipitation. Further protein analysis by HPLC and AA sequencing revealed that vicilin, an insoluble storage protein, was the main pea protein incorporated within the casein micelle structure. These results supported our hypothesis that low-temperature homogenization can successfully be used to create a colloidal dispersion with increased solubility, in which insoluble plant-based proteins may be incorporated with casein micelles in an aqueous solution. Additionally, 3-dimensional microscope images of the blends indicated a noticeable difference between the surface roughness upon addition of pea protein to the casein micelle matrix. This research highlights a promising application for other plant-based proteins to be used within the dairy industry to help drive future product innovation while also meeting current processing conditions and consumer demands.

Key words: casein micelle, pea protein, homogenization, colloidal dispersion

INTRODUCTION

Caseins account for 80% of the total proteins in milk and self-assemble into casein micelles (CM). The CM contain the 4 major CN subgroups (αS1, αS2, β, κ) along with colloidal calcium phosphate, held together by noncovalent interactions including electrostatic, hydrophobic, and hydrogen bonding (Holt and Horne, 1996; Horne, 1998; Walstra, 1999). The structure of CM can be disrupted by the addition of chelating agents, NaCl, and cations. Additionally, changes in the solution temperature and pH can also disrupt the micelle structure. For example, calcium phosphate exhibits greater solubility at lower temperatures, and thus is released from the micelle upon cooling to 4°C (Koutina et al., 2014). Similarly, hydrophobic β-CN is also released from the micelle at lower temperatures (Pierre and Brule, 1981; Liu et al., 2013). The release of colloidal calcium phosphate and β-CN creates a looser, dissociated micelle structure. The same effect is observed upon the addition of chelating agents, such as sodium citrate. Upon the addition of sodium citrate 0.02 M, the resulting CM structure is larger in size due to the complexing of sodium citrate to calcium phosphate, ultimately leading to the release of β-CN (Heertje et al., 1985).

The unique structure and hydrophobic core of CM allows them to serve as carriers of essential nutrients native to CM, such as calcium, phosphate, and protein, along with the potential to serve as encapsulation vehicles for hydrophobic compounds. In general, encapsulation is carried out by first disassociating the CM, then facilitating interactions between the hydrophobic compounds and the hydrophobic CN regions, and finally reassembling the CM structure (Glab and Boratyński, 2017). For example, vitamin D2 and D3 have been incorporated within CM via ultra-high-pressure homogenization and restoration of mineral content to enrich low-fat foods (Menéndez-Aguirre et al., 2011;
Menéndez-Aguirre et al., 2014; Cohen et al., 2017). Additionally, soybean oil, fish oil, and rapeseed oil have been successfully encapsulated within the hydrophobic regions of CM through pH shifts and ultrasound treatments (Ghasemi and Abbasi, 2014).

The aim of the presented research was to use CM to encapsulate another type of hydrophobic compound, plant-based protein. The demand for plant-based protein is growing due to consumers’ desires for high-protein health benefits and environmentally conscious products (Boland et al., 2013). Pea protein (PP), chosen as the plant-based protein for this study, has become fairly popular due to its balanced AA composition and branched chain AA that assist in muscle synthesis and expansion (Babault et al., 2015). Pea protein is a non-allergenic, gluten-free, and cost-effective plant protein. However, it can be a challenge to use in food systems because of its low solubility and undesirable bitterness. Pea protein has a high percentage of hydrophobic proteins (65–80%), making it a good hydrophobic model to represent plant-based proteins (Nehete et al., 2013; Lu et al., 2020). Other highly hydrophobic plant-based proteins such as soy, wheat, corn, and sunflower have potential to be applied in a CM encapsulation system (Nehete et al., 2013).

The objective of this study was to create a stable colloidal dispersion of CM with PP to improve the solubility of PP in aqueous solution. We hypothesized that low-temperature homogenization could be applied to an unstable dispersion of disrupted CM and PP, and the resulting blend would be a stable colloidal dispersion in which the insoluble PP associate with CM in suspension. Unlike previous research studies, this protocol used readily available dairy processing equipment that could be easily reproduced in any dairy facility. This research highlighted a promising application for other plant-based proteins to be used within the dairy industry to help drive future product innovation.

**MATERIALS AND METHODS**

**Sample Preparation**

Commercial fat-free milk (Dean Foods Company) was purchased at local grocery stores. The skim milk sample was prepared by first dissociating the CM with 5.16 g/L food-grade trisodium citrate (It’s Just!, 138 Foods Inc.; Heertje et al., 1985). The skim milk was equilibrated at 4°C for 24 h. The commercial skim milk used for this study had a nutrition label claiming 3.3% protein, of which 2.6% represented CM (Linn, 1988). For this reason, a PP slurry with 2.6% protein was developed to have a 1:1 protein ratio between the CN in the milk sample and the protein in the PP slurry. The 2.6% PP slurry was created by combining PP isolate (Mettle Nutrition LLC) with distilled water and continuous agitation at 25°C for 24 h. The milk sample and the PP slurry were combined at different ratios of CM:PP (90:10, 80:20, and 50:50). We wanted the major protein component to be contributed by the CM; therefore, we did not create blends with less than 50% CM. The skim milk with trisodium citrate and the 2.6% PP slurry were the control samples used in this study to understand how each protein component behaved on its own.

**Homogenization and Pasteurization of CM:PP Blends**

Each protein blend underwent 3 cycles of low-temperature (4°C) homogenization using a bench-top GEA 2-stage homogenizer (NS2002 H, GEA Niro Soavi S.p.A.) at 24,130 kPa (first stage) and 3,447 kPa (second stage). After homogenization, they were batch pasteurized in a water bath at 63°C and held for 30 min. The sample blends were stored at −20°C until further analysis.

**Particle Size Analysis**

The hydrodynamic diameter of the dispersed particles in the liquid CM:PP blends and control samples were measured by dynamic light scattering using NanoBrook ZetaPALS Potential Analyzer (Brookhaven Instruments Corporation). We added 3 mL of distilled water to a clean cuvette, and then 100 μL of liquid sample was added. Backscattering angle 173° was used for weakly scattered particles such as proteins. The temperature was maintained at 25°C with an equilibration time of 1 min. The measurements were performed over a range of 1.0 to 10,000 nm over 5 min, and the sample specifications were set to be 0.133 refractive index, 0.890 mPa-s viscosity, and pH = 7.0. Results are reported as averages from 5 replicates.

**Soluble Protein Analysis**

The soluble protein content of each CM:PP blend and control sample were measured pre- and post-homogenization via Quick Start Bradford Protein Assay (Bio-Rad). We combined 20 μL of 1:1,000 diluted sample with 140 μL of Tris-HCl 50 mM (pH = 8) buffer and 40 μL of Bradford dye (Bio-Rad). The mix reaction was transferred to a 96-well plate and mixed thor-
oughly. The protein concentration was measured with a spectrophotometer (acuSkan GO 1510, Thermo Fisher Scientific) at a wavelength of 595 nm. The standard curve was prepared using BSA (Bio-Rad).

**Compositional Analysis**

Moisture (%), solids (%), and fat (%) of the CM:PP blends and homogenized milk and pea slurry controls were measured using manufacturer’s instructions for Smart 6 CEM Milk 0T01-T6 method (CEM, Smart 6). The protein (%) of the CM:PP blends was measured via Dumas combustion using manufacturer’s instructions on a Rapid MAX N-Exceed nitrogen and protein analyzer (Rapid MAX N-Exceed, Elementar). The conversion factor of 6.38 (milk) was used to convert elemental nitrogen to percent protein. The combined carbohydrates and ash (%) were measured via difference in solids. Results are reported as averages from 3 replicates.

**Coagulation of CM:PP Blends**

We transferred 50 mL of each CM:PP blend to a clean test tube and placed in a 37°C incubator. Then, 1 mL of CaCl₂ 1% was added to the sample and set for 10 min at 37°C. To coagulate the CN in each blend, chymosin (Chy-Max M 127223, Chr. Hansen) was first diluted with distilled water to a 1:100 ratio, and 0.25 mL of this dilution was added to each sample and shaken gently back and forth for 30 s. The samples were incubated at 37°C for 24 h to achieve full curd formation. Afterward, the curd was removed from the whey using a cheese cloth. Control experiments were carried out under the same conditions using homogenized skim milk (positive control) and homogenized PP slurry, which did not coagulate under these conditions (negative control). Each CM:PP blend was prepared in triplicate.

**SDS-PAGE**

To determine the profile of the proteins incorporated in the CM:PP blends (curd and whey components) compared with the control samples, 10% SDS-PAGE gels were prepared in not-reducing conditions. A mixed solution containing the diluted sample with 45 μg of protein and 2× Laemmli sample buffer (65.8 mM Tris-HCl, pH 6.8, 2.1% SDS, 26.3% glycerol, 0.01% bromophenol blue; Bio-Rad) was loaded into each well. The gel ran for 70 min at 120 V. After the running time, the gels were stained with Coomassie blue (Bio-Rad), and an image of each gel was taken using Chemidoc-Touch Imaging System (Bio-Rad) and analyzed using Image Lab software 6.1 (Bio-Rad).

**3-Dimensional Laser Microscope**

An aliquot of each coagulated CM:PP blend was placed on a 3-mm microscope slide and gently dried at 37°C in an oven. Afterward, it was analyzed with a 3-dimensional laser microscope (VK-X200 series, Keyence) under different magnifications (10×, 20×, 50×, and 150× lenses). Four images were analyzed for each sample, and the most representative was chosen and depicted for analysis. Five areas of each image were analyzed to determine surface roughness (Sa; as the arithmetical mean height in μm) using Multi File Analyzer software (version 1.3.1.120; Keyence). The averages of each CM:PP blend Sa values were compared with Sa values of the positive control (coagulated skim milk).

**Reversed-Phase HPLC**

The proteins of the CM:PP blends and control samples were separated and identified via reversed-phase (RP)-HPLC (LC-20AD, Shimadzu USA Manufacturing Inc.) coupled to an Aeras 3.6-μm Widepore XB-C8 (LC column 250 × 4.6 mm) and a fluorescence detector (RF-10AXL, Shimadzu). Solvent A was 0.1% trifluoroacetic acid (TFA) in HPLC grade water and solvent B was 0.1% TFA in HPLC grade acetonitrile. Both solvents were degassed for 10 min before being vacuum filtered through a 0.22-μm nylon filter. The flow rate was 1 mL/min. From 0 to 20 min, solvent B was 25 to 50%. From 20 to 30 min, solvent B was 50 to 80%. From 30 to 32 min, solvent B was 80 to 25%. The excitation wavelength (λex) and emission wavelength (λem) in the detector were 280 and 348 nm, respectively, as these are the optimal wavelengths to detect tryptophan. The column oven was set at 40°C, and the sample injection volume was 5 μL. The light emitted from the excited atoms from the analyte was measured in μV. The samples were prepared by using the method described by Ma et al. (2017) with some modifications. In brief, 800 μL of protein sample were combined with 800 μL of solution A (5.37 mM sodium citrate, 19.5 mM d-L-dithiothreitol, 6 M guanidine hydrochloride in 0.1 M BisTris buffer, pH = 7). The samples were then vortexed for 10 s and incubated at room temperature for 1 h. An aliquot of 500 μL was further mixed with 1,000 μL of solution B (4.5 M guanidine hydrochloride in a solution of water, acetonitrile, and TFA in a ratio
of 89.9:10:0.1). The 1.5-mL sample mixture was filtered through a 0.22-μm filter into an amber HPLC vial and stored at 4°C until the day of analysis.

**AA Sequencing**

The AA sequence of the PP of interest was analyzed via capillary-liquid chromatography-nanospray tandem mass spectrometry (LC/MS/MS). The protein sample was loaded into a 10% SDS-PAGE stained in Coomassie blue. The single band was excised from the gel and was analyzed at The Mass Spectrometry and Proteomics Facility (Ohio State University, Columbus). They performed an in-gel digestion with sequencing grade-modified trypsin and extracted the peptides. The peptides were separated via capillary-LC/MS/MS and the data collected was searched in Mascot Daemon by Matrix Science version 2.7.0. Proteins identified with at least 2 unique peptides were considered to be a reliable identification.

**Statistical Analysis**

All experiments were conducted in quintet (particle size analysis, Sa) or triplicate (compositional analysis, soluble protein). One-way ANOVA analyses followed by Tukey-Kramer HSD multiple-comparison were conducted to determine statistical differences \( (P < 0.05) \) between pairs using JMP software (JMP 14.0.0, SAS Institute Inc.).

**RESULTS AND DISCUSSION**

**Disruption of CM**

Successful encapsulations within CM first involve the disruption of the micelle to expose the hydrophobic core to facilitate greater interactions with the added hydrophobic compound (Głęb and Boratyński, 2017). The proposed protocol suggests using a temperature decrease and an addition of sodium citrate to disrupt the CM structure. For the purpose of the study, it was assumed that the soluble particles in milk were all CM, due to the low percentage and small size of serum proteins, in addition to the low percentage of fat present (Holt, 1985; Beliciu and Moraru, 2009). We observed significantly larger sized CM in the 4°C milk compared with the 60°C milk, with hydrodynamic diameters (mean ± SD) of 197.11 ± 2.07 nm and 163.15 ± 5.28 nm, respectively \( (P < 0.0001; \text{data not shown}) \). Upon the addition of sodium citrate to the milk at 4°C, the hydrodynamic diameter increased to 220.06 ± 2.34 nm, which was significantly greater than the milk with no sodium citrate addition \( (P < 0.0001; \text{data not shown}) \). In addition to the larger soluble particle size, the milk at 4°C + sodium citrate appeared less turbid and less white, indicating a successful disruption of the micelle structure (Fox et al., 2015). These results support the proposed protocol to use low-temperature (4°C) processing methods in combination with sodium citrate addition to successfully disrupt the CM structure.

**Soluble Protein Analysis**

Three CM:PP blends at different ratios pre- and posthomogenization are shown in Figure 1A. The pre-homogenized blends at all CM:PP ratios showed clear separation between the 2 protein segments, an indication of an unstable colloidal dispersion. The insoluble PP settled out of solution and formed a solid pellet. On the other hand, the posthomogenized blends at all CM:PP ratios were homogeneous with no visible sedimentation. All 3 posthomogenized blends remained visibly stable at 4°C for up to 3 wk with no formation of an insoluble pellet. The soluble protein content (Figure 1B) of the 90:10 and 50:50 blends significantly increased upon homogenization \( (90:10, P = 0.0268; 50:50, P = 0.0015) \). The soluble protein content of the 80:20 blend also increased upon homogenization, but not by a significant value \( (80:20 P = 0.0924) \). The process of homogenization has been shown to disrupt the 3-dimensional globular PP bodies into smaller, more soluble aggregates suspended in the aqueous phase (Moll et al., 2021). In agreement with Moll et al. (2021), the present results supported the disruption of globular PP bodies by the fact that the PP slurry control prehomogenization had a soluble protein content of 8.24 ± 1.38 mg/mL, which significantly increased posthomogenization to 66.47 ± 17.06 mg/mL \( (P = 0.0041; \text{Figure 1B}) \).

**Compositional Analysis**

The compositional analysis of the 90:10, 80:20, and 50:50 CM:PP blends were compared with commercial skim milk in Table 1. As expected, the 90:10 CM:PP blend had the most similar composition to skim milk. There was no significant difference between the 90:10 CM:PP blend and skim milk moisture content \( (P = 0.3397) \), solids content \( (P = 0.3366) \), and protein content \( (P = 1.0) \). In general, as the amount of PP in the blend increased, the moisture content increased, and the solids content decreased. This result was expected because the initial PP slurry only had a solids content of 3.3%, compared with the initial skim milk with a solids content of 9.05%, due to the additional components in milk. As the PP content of the blend
increased, the protein content significantly decreased compared with skim milk. The 80:20 and 50:50 CM:PP blends had statistically similar protein contents to each other \( (P = 0.6088) \), but statistically different protein contents to the 90:10 CM:PP blend \( (P = 0.0231 \) and \( P = 0.0045 \), respectively) and skim milk \( (P = 0.0246 \) and \( P = 0.0048 \), respectively). Protein content in milk is affected by the genetics, age, stage of lactation, and nutrition of the cow and is not standardized in the industry (Linn, 1988). In the case of this study, the skim milk protein content was 4.72 ± 0.03%, which was greater than what was expected based on the nutritional label (3.3%). Therefore, the protein content inevitably decreased upon the replacement of skim milk with PP slurry in the blend. All CM:PP blends and skim milk have statistically similar fat contents \( (P = 0.1348) \). There was relatively little fat to begin with in the skim milk (0.07%), and the PP contributed negligible fat to the blend.

**Particle Size Analysis**

We measured the hydrodynamic diameter of the dispersed particles in pre- and posthomogenized skim milk and PP slurry controls to understand the effect of low-temperature processing on the individual protein components. The particle size of CM in skim milk did not change significantly between pre-homogenization and posthomogenization \( (220.06 \pm 2.34 \text{ nm and } 216.00 \pm 8.35 \text{ nm, respectively}; P = 0.8721; \text{Figure 2}) \). This result was expected because the size and structure of CM is not affected at pressures less than 200 MPa (Huppertz et al., 2004, 2006). On the other hand, the soluble particle size of PP slurry pre-homogenization was significantly greater than the PP slurry posthomogenization \( (956.80 \pm 10.30 \text{ nm and } 323.18 \pm 5.85 \text{ nm, respectively}; P < 0.0001; \text{Figure 2}) \). Smaller protein aggregates tend to have improved solubility due to the increased surface area and ele-

**Table 1. Compositional data (mean ± SD) of CM:PP blends and skim milk**

<table>
<thead>
<tr>
<th>Sample (CM:PP)</th>
<th>Moisture (%)</th>
<th>Solids (%)</th>
<th>Protein (%)</th>
<th>Fat (%)</th>
<th>CHO + ash2 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>90:10</td>
<td>91.31 ± 0.01a</td>
<td>8.69 ± 0.01a</td>
<td>4.73 ± 0.07a</td>
<td>0.09 ± 0.02a</td>
<td>4.82</td>
</tr>
<tr>
<td>80:20</td>
<td>92.00 ± 0.08a</td>
<td>7.99 ± 0.06a</td>
<td>4.29 ± 0.10a</td>
<td>0.08 ± 0.01a</td>
<td>4.36</td>
</tr>
<tr>
<td>50:50</td>
<td>94.31 ± 0.47a</td>
<td>5.69 ± 0.47a</td>
<td>4.14 ± 0.25a</td>
<td>0.10 ± 0.01a</td>
<td>4.24</td>
</tr>
<tr>
<td>Skim milk</td>
<td>90.95 ± 0.10a</td>
<td>9.05 ± 0.10a</td>
<td>4.72 ± 0.03a</td>
<td>0.07 ± 0.02a</td>
<td>4.57</td>
</tr>
</tbody>
</table>

*Different lowercase letters indicate a significant difference \( (P < 0.05) \).
1CM:PP = ratio of casein micelle to pea protein.
2By difference; CHO = carbohydrates.
todynamic forces between protein and water molecules (Jambrak et al., 2008; Jiang et al., 2017). This improved solubility correlated to the increased soluble protein and smaller dispersed particle sizes identified in the PP slurry posthomogenization. Based on the hydrodynamic diameter results, it was inferred that any change in soluble particle size in the blends was due to the disruption of globular PP.

The hydrodynamic diameter of each CM:PP blend was analyzed pre- and posthomogenization (Figure 2). The soluble particle size of the 80:20 and 50:50 blends increased significantly posthomogenization ($P = 0.0006$ and $P < 0.0001$, respectively). Before homogenization, CM were the major soluble particle represented in the blend. However, after homogenization, insoluble globular PP were disrupted into smaller soluble aggregates that contributed to the increase in soluble particle size and content. However, these results do not show how the PP interact in the soluble phase or whether or not they associate with any milk component. On the other hand, the soluble particle size of the 90:10 blend decreased upon homogenization. The authors believe that the PP was not in sufficient quantity (10%) to have a significant effect on increasing the overall soluble particle size or that the 90:10 ratio created smaller complexes of casein and PP (Figure 2).

**Protein Profile Analysis**

Sodium dodecyl sulfate-PAGE analysis was used to understand qualitatively how the PP incorporate within the soluble phase. We analyzed homogenized skim milk and PP slurry via SDS-PAGE (Figure 3). The major milk proteins included immunoglobulins, lactoferrin, BSA, CN, β-LG, and α-LA, which have molecular weights of 160, 80, 66.5, 20 to 25, 18.3, and 14.2 kDa, respectively (El-Loly, 2007; Edwards and Jameson, 2014; Vincent et al., 2016). About 80% of PP represents the following insoluble storage globulin proteins: legumin, vicilin, and convicilin (Barac et al., 2010). Legumin contains various subunits of 40- and 20-kDa molecular weights. Vicilin is composed of subunits with molecular weights of ~50, ~47, ~34, and ~30 kDa subunits. Convicilin contains subunits of ~71 kDa (Barac et al., 2010). Figure 3 shows an intense band around 47 kDa in the PP slurry (lane 1); however, there is no band in the skim milk (lane 2) at this similar molecular weight. This PP at 47 kDa is easily identifiable and will not be confused with any milk proteins, and thus can be used as an indicator to determine where the PP incorporates within the blend (whey proteins, CM, or self-association). To assess what fraction was enriched in PP, the CM:PP blends and controls were coagulated with rennin Chy-Max M 127223 to separate the whey proteins (soluble fraction) from CM (coagulated precipitate). The CM:PP blends and milk coagulated under these conditions; however, the PP slurry did not. We analyzed the total uncoagulated blend, the liquid whey component, and the solid CN curd component through SDS-PAGE (Figure 4).

All CM:PP blends contained the 47 kDa band (Figure 4 lanes 1, 4, and 7), which remained in all the CM fractions (Figure 4 lanes 3, 6 and 9) rather than the whey fractions (Figure 4 lanes 2, 5, and 8), indicating that the homogenization process incorporated this PP into CM. To identify the 47-kDa protein, we conducted AA sequence analysis. We identified 44 peptides that represented 77% of coverage with a pea vicilin sequence using the NCBI BLAST database (https://blast.ncbi.nlm.nih.gov/Blast.cgi; data not shown). These observations confirmed the incorporation of the insoluble storage globulin PP vicilin with CM via low-temperature homogenization. The method limitation of SDS-PAGE as applied to this experiment was the inability to differentiate PP from milk proteins through characteristics other than molecular weight. For this reason, only the 47 kDa vicilin was identified as incorporating within the CM; however, it is possible that other PP also incorporate within the CM. Additional methods such as electron microscopy that can better

![Figure 2](image_url)

**Figure 2.** Hydrodynamic diameter (nm) of dispersed particles of casein micelle:pea protein (CM:PP) blends and controls, pre- and posthomogenization. Values are means of data from triplicate experiments with standard deviation indicated by vertical bars. Different lowercase letters (a-f) indicate a significant difference ($P < 0.05$).
differentiate between PP and CM may be used in future experiments (Jiang et al., 2017).

**RP-HPLC Analysis**

We used RP-HPLC with fluorescence detection to further understand the interaction between PP and CM in the blend based on their intrinsic protein fluorescence due to aromatic side chains such as Trp, Tyr, and Phe (Lu et al., 2020). Figure 5 shows the chromatogram of the CM:PP blends and their respective whey components after coagulation compared with skim milk and PP slurry. There were 2 fluorescence peaks produced by PP at 6.7 and 12.8 min, whereas skim milk did not have a signal at these retention times, indicating that PP was the cause of this fluorescence. In contrast, the PP fluorescence was detected in all 3 CM:PP blends at these specific retention times, but not in their whey components, indicating again that PP were interacting with the CM. The results from RP-HPLC further demonstrated that the proposed protocol described in this paper effectively created a stable colloidal dispersion between insoluble PP and CM. It should be noted that the RP-HPLC analysis was only used for qualitative results due to the lack of available commercial standards for pea vicilin.

**Microstructure Analysis**

The surface of the curd component of the CM:PP blends was further analyzed by 3-dimensional laser microscopy. Figure 6 shows the 90:10, 80:20, and 50:50 CM:PP curds under 150× lens compared with a skim milk curd. Through visual comparison, the skim milk curd had fewer, but more pronounced, CN coagulates on the curd surface. As the amount of PP increased, the size of the surface ridges decreased, and the ridges became more numerous. The 90:10 curd had very pronounced surface ridges in the lower part of the image that were similar to the milk CN coagulate. However, in the upper right area of the 90:10 curd surface, the texture started to look more similar to the 80:20 curd surface. The 50:50 curd surface looked different than the skim milk curd, with unrecognizable coagulates compared with the intact CN coagulates. The 50:50 curd had less pronounced ridges, but they were more numerous. The arithmetical mean height was measured to evaluate the Sa. As the amount of PP increased, the surface roughness (μm) decreased. The ranking of the surface roughness of each sample was as follows: skim milk (3.99 ± 2.63 μm) > 90:10 blend (2.70 ± 1.70 μm) > 80:20 blend (1.70 ± 0.58 μm) > 50:50 blend (0.79 ± 0.38 μm). The surface roughness of the skim milk was significantly different than the surface roughness of the 80:20 CM:PP blend (P = 0.0002) and the 50:50 CM:PP blend (P < 0.0001). We hypothesized that this change in surface texture was due to the incorporation of PP in the CN structure, which disrupted the CN coagulum.

**CONCLUSIONS**

In this work, we were able to take advantage of the conformational changes that CM undergo upon exposure to cold temperatures and sodium citrate addition to prepare a codispersion with PP that significantly increased the stability of this hydrophobic protein fraction in colloidal suspensions. Our method to create a stable colloidal dispersion used the unique CM structure optimal for protein blending and current dairy processing equipment, such as homogenizers, to allow for easy reproducibility in any dairy facility. The CM:PP blend showed promising results for the use of mixed protein,
functional food product in the form of liquid, gel, or powder applications. Future experiments should deal with the amount of PP that associates within the CM, the nature of these associations, and sensory analysis. There is great potential to apply this research to other plant-based proteins or nutraceuticals with low solubil-

**Figure 4.** Sodium dodecyl sulfate-PAGE (10%) stained in Coomassie blue of casein micelle:pea protein (CM:PP) blends (90:10, 80:20, 50:50) and their respective whey and CN components. M = molecular weight marker; lane 1 = 90:10 total blend; lane 2 = 90:10 whey component; lane 3 = 90:10 CN component; lane 4 = 80:20 total blend; lane 5 = 80:20 whey component; lane 6 = 80:20 CN component; lane 7 = 50:50 total blend; lane 8 = 50:50 whey component; lane 9 = 50:50 CN component. Forty-five micrograms of protein was loaded in each well.

**Figure 5.** Reversed-phase HPLC with fluorescence detection chromatogram of casein micelle:pea protein (CM:PP) blends and their respective whey components compared with skim milk and homogenized PP slurry.
ity to be further used in an innovative way that allows the dairy industry to provide highly nutritious protein products, while also capitalizing on current consumer trends.

**ACKNOWLEDGMENTS**

This project was funded by the Parker Endowment (Department of Food Science and Technology, The Ohio State University, Columbus). The project was supported by NIH (Bethesda, MD) award number grant P30 CA016058 for the use of the CCIC Mass Spectrometry and Proteomics (MSandP) Facility (Columbus, OH) services and resources. The Fusion Orbitrap instrument was supported by NIH award number grant S10 OD018056. The authors thank Molly Davis (Department of Food Science and Technology, The Ohio State University, Columbus) for her editing. The authors have not stated any conflicts of interest.

**REFERENCES**


Beliciu, C. M., and C. I. Moraru. 2009. Effect of solvent and temperature on the size distribution of casein micelles measured by...