pH-Induced physicochemical modifications of native phosphocaseinate suspensions: Influence of aqueous phase

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Summary — pH-Induced physicochemical changes and rennet coagulation times of casein micelle suspensions have been studied with native phosphocaseinate powder dissolved in water, 0.1 mol L\(^{-1}\) NaCl and milk ultrafiltrate. An increase in ionic strength between water and NaCl led to higher diffusible calcium and phosphorus, to greater micelle voluminosity and to greater casein and mineral solubilization during acidification. Dissolution of the phosphocaseinate in ultrafiltrate resulted in physicochemical properties very close to milk. The acidification process can be divided into five successive steps. Relations between physicochemical changes are discussed, together with the reduction of rennet coagulation time with pH and ionic strength reduction.

micelle / pH / ionic strength / calcium / phosphorus / native phosphocaseinate / rennet coagulation / acid coagulation

Résumé — Modifications physicochimiques des suspensions de phosphocaséinate natif induites par le pH : influence de la phase aqueuse. L’effet du pH sur les évolutions physicochimiques de suspensions de micelles de caséine, ainsi que sur les temps de coagulation pressure, a été étudié sur des poudres de phosphocaséinate natif reconstituées dans de l’eau, une solution 0,1 mol L\(^{-1}\) en NaCl et un ultrafiltrat de lait. L’augmentation de force ionique entre l’eau et NaCl provoque une augmentation des teneurs en calcium et en phosphore diffusibles, une plus grande voluminosité des micelles, et une augmentation de la solubilisation des caséines et des minéraux intervenant lors de l’acidification. La solubilisation du phosphocaséinate natif dans l’ultrafiltrat conduit à un comportement proche du lait. L’acidification peut être divisée en cinq phases successives. Les relations entre les évolutions physicochimiques sont discutées, ainsi que la diminution des temps de coagulation pressure avec la baisse du pH et de la force ionique.

micelle / pH / force ionique / calcium / phosphore / phosphocaséinate natif / coagulation pressure / coagulation acide
INTRODUCTION

Acidification of milk is fundamental to many industrial dairy processes such as casein, cheese and yoghurt manufacture. Previous work established that during the decrease of pH, calcium and inorganic phosphate were solubilized from the micelles (Brulé et al, 1974; Van Hooydonk et al, 1986a; Visser et al, 1986; Dalgleish and Law, 1989; le Graet and Brulé, 1993), caseins dissociate from the micelles near pH 5.4 at 20°C, and then reassociate (Roefs et al, 1985; Van Hooydonk et al, 1986a; Dalgleish and Law, 1988), micelle size decreases (Heertje et al, 1985; Roefs et al, 1985; Banon and Hardy, 1991, 1992), zeta potential decreases (Darling and Dickson, 1979; Banon and Hardy, 1992) or reaches a minimum at pH 5.4 and a maximum at pH 5.1 (Schmidt and Poll, 1986) and solvation decreases (Tarodo de la Fuente and Alais, 1975; Snoeren et al, 1984; Creamer, 1985; Visser et al, 1986). Finally, at its isoelectric pH, casein becomes aggregated in a network.

However, the understanding of the role and the sequence of each of these changes is still limited. Heertje et al (1985) suggested that the particles aggregating at pH 5.0 are different from the native ones, ie, those at physiological pH. The complex change of zeta potential with pH reduction (Schmidt and Poll, 1986) can be explained by the \( \beta \)-casein acting as the starting point for the casein aggregation (Heertje et al, 1985; Visser et al, 1986) or by the pH-dependent affinity of calcium for micellar casein or inorganic phosphate (Schmidt and Poll, 1986).

In the present paper, we describe the behaviour of casein micelles during acidification. Tangential membrane microfiltration of milk by the Bactocatch procedure as proposed by Fauquant et al (1988), followed by spray drying of the diafiltered retentate (Schuck et al, 1994) results in an enriched micellar caseinate called native phosphocaseinate powder. This product is close in protein composition to a commercial calcium caseinate powder (Schuck et al, 1994) and it exhibits reduced rennet clotting time and increased gel development kinetics (Pierre et al, 1992). Hence, this powder may be regarded as a suitable way to standardise milk in cheese making processes as described by Salles et al (1995), because it enhances technological properties by increasing the protein concentration, without increasing the lactose content. Moreover, it may be regarded as an attractive way to produce various micellar suspensions.

The aim is to establish the effect of the aqueous phase and of the ionic strength on casein and mineral dissociation during acidification and to show relationships between the various physicochemical changes. The native phosphocaseinate powder was dissolved in water, 0.1 mol L\(^{-1}\) NaCl solution, and milk ultrafiltrate. Acidification was induced by adding glucono-delta-lactone (GDL), because that is a simple system to reach desired and stable pH values in quiescent conditions, without using bacteria.

MATERIALS AND METHODS

**Preparation of native phosphocaseinate suspensions (NPCS)**

The native phosphocaseinate powder P1 was prepared according to Pierre et al (1992) and Schuck et al (1994). Chemical composition of this powder is reported in table I. The powder was reconstituted at 50°C to a concentration of 25 g L\(^{-1}\) of casein (ie, 31.94 g L\(^{-1}\) of powder); 0.2 g L\(^{-1}\) sodium azide was added to the aqueous phase used for the reconstitution, ie, deionized water or 0.1 mol L\(^{-1}\) NaCl solution prepared in deionized water or milk ultrafiltrate (UF). Raw whole milk (Besnier Bridel, l’Hermitage, France) used to prepare UF was skimmed twice at 43°C with Westfalia Separator DD100Z. UF was obtained by ultrafiltration on a spiral membrane of 3000 Da molecular mass cut-off.
Acidification of native phosphocaseinate

Table 1. Chemical composition of the native phosphocaseinate powder.

<table>
<thead>
<tr>
<th>Component</th>
<th>g kg⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dry matter</td>
<td>926.20</td>
</tr>
<tr>
<td>Total nitrogen</td>
<td>836.16</td>
</tr>
<tr>
<td>NCN</td>
<td>53.40</td>
</tr>
<tr>
<td>NPN</td>
<td>4.33</td>
</tr>
<tr>
<td>Lactose</td>
<td>6.7</td>
</tr>
<tr>
<td>Ashes</td>
<td>81.50</td>
</tr>
<tr>
<td>Ca</td>
<td>28.49</td>
</tr>
<tr>
<td>Na</td>
<td>0.48</td>
</tr>
<tr>
<td>K</td>
<td>0.89</td>
</tr>
<tr>
<td>Mg</td>
<td>1.00</td>
</tr>
<tr>
<td>Cl</td>
<td>1.57</td>
</tr>
<tr>
<td>Citrate</td>
<td>1.32</td>
</tr>
</tbody>
</table>

*Using 6.38 as converting factor; NCN, non-casein nitrogen; NPN, non-protein nitrogen.

After calculation with the factor 6.38; NCN: azote non caséinique; NPN: azote non protéique.

(MMCO) (S10Y3, Amicon, Epernay, France) at 50 °C. Preparation of native phosphocaseinate suspensions (NPCS) in each aqueous phase and subsequent analyses were carried out in duplicate.

**Acidification of NPCS**

Defined amounts of GDL (Lysactone, Roquette Frères, Lestrem, France) were added at 20 °C to NPCS to reach pH values between 4.0 and 7.5, after 15 h at 20 °C. After GDL addition, NPCS were stirred and left at 20 °C in a water bath. The pH obtained in these conditions was related in advance by a polynomial function (maximal degree five) to the amount of GDL. This equation allows to compute the content of GDL to reach precise values of pH. One mol L⁻¹ NaOH was added to micelle suspensions in UF to reach pH 7.0.

**Fractionation**

Each sample was ultracentrifuged at 20 °C during 2 h at 75 000 g in a L8-55 ultracentrifuge (Beckman Instrument France, Gagny, France), with a 50.2 Ti rotor. Supernatants were carefully removed by syringe, and micellar pellets were drained for 5 min.

1 h centrifugation at 1000 g at 20 °C was carried out in a Cryofuge M 7000 (Heraeus Sepsitech, Les Ulis, France).

Ultrafiltration was carried out at 20 °C on a Centriflo CF 25 (MMCO 25000 Da, Amicon, Epernay, France) at 500 g during 1 h. The first 2 mL of ultrafiltrate and the concentrated suspensions were discarded and 7 mL of each sample were then ultrafiltered.

**Physicochemical analysis**

Calcium and phosphorus (P) contents were determined by atomic absorption spectrometry (Brulé et al, 1974) and International Dairy Federation standard method (1987), respectively, on micelle suspensions and Centriflo ultrafiltrates. Concentrations of minerals in ultrafiltrate were converted into diffusible mineral concentrations in NPCS by multiplying by a correction factor of 0.96, according to Pierre and Brulé (1981). This correction accounted for excluded volume effect, but neglected the Donnan effect (Holt, 1985). This probably caused a systematic error for diffusible amounts. Micellar mineral contents were calculated by subtracting the diffusible concentrations from the total one.

Total nitrogen contents (TN) of 1000-g supernatants were estimated by Kjeldahl with 6.38 as conversion factor.

Non-micellar caseins were defined as caseins which do not sediment after 2 h centrifugation at 75 000 g. They were obtained by optical density measurements (OD) at 280 nm (Uvikon 810, Kontron SA, St-Quentin-en-Yvelines, France) of the supernatants diluted in 10 mmol L⁻¹ EOTA (pH 10) (adapted from Oriehuis and Teernstra, 1992). ODs of suspensions and supernatants were calculated by multiplying the OD measured by the dilution factor. For NPCS dissolved in water and NaCl, the percent of OD in supernatant (OD%) at each pH was calculated as:

\[
\text{OD} \% = \frac{\text{OD of supernatant at pH}_i \times 100}{\text{OD of NPCS at pH}_i}
\]

while for the NPCS dissolved in UF the calculation was:

\[
\text{OD} \% = \frac{\text{OD of supernatant at pH}_i - \text{OD of UF used}}{\text{OD of NPCS at pH}_i - \text{OD of UF used}} \times 100
\]

to compare the solubilization of caseins with the pH in the various media. OD values of the two
UF were 5.00 and 6.04. Supernatants which gave the higher OD% were analysed by HPLC on reversed-phase C4 column according to Jaubert and Martin (1992). β-casein (Eurial, Nantes, France) was used as standard for calibration, and absorbivity values of the major caseins were used for quantification (Swaisgood, 1992).

Drained pellets of 75 000 g centrifugation were dried at 103 °C during 7 h and the solvation in g of water per g of dry pellet were deduced. The total sediment volume, \( V_s \) (mL g\(^{-1}\)) was calculated as:

\[
V_s = C_{\text{cas}} \times V_{\text{cas}} + C_w \times V_w
\]

where \( C_{\text{cas}}, C_w \) are the casein and water concentrations in the wet pellet (g\(^{-1}\)), respectively, and \( V_{\text{cas}}, V_w \) are the specific volumes of caseins (0.7 mL g\(^{-1}\)) and water (1.0 mL g\(^{-1}\)) (Van Hooydonk et al, 1986a). We assumed whey proteins, lactose and mineral contributions to be negligible, because their concentrations in the pellet were almost constant or low. Casein voluminosity obtained by pellet solvation measurements, \( V_{\text{pellet}} \), is \( V_s/C_{\text{cas}} \) (Van Hooydonk et al, 1986a).

Apparent dynamic viscosity of micelle suspensions was measured at 20 °C using a coaxial cylinder viscometer LS30 (Contraves, Zurich, Switzerland) at 94.5 s\(^{-1}\) strain rate. Relative viscosity was calculated as the ratio of dynamic viscosity of NPCS and of the measured values for dynamic viscosity of water (1.00 mPa s), NaCl (0.95 mPa s) or UF (1.09 mPa s). Voluminosity was calculated according to the equation of Eilers, cited in Van Hooydonk et al, 1986a):

\[
\eta_{\text{rel}} = \left(1 + \frac{1.25 \Phi}{\Phi_{\text{max}}}\right)^2 \left(1 - \frac{\Phi}{\Phi_{\text{max}}}\right)
\]

where \( \eta_{\text{rel}} \) is the relative viscosity of the NPCS, \( \Phi \) is the volume fraction of casein and \( \Phi_{\text{max}} \) is a constant of value 0.79. \( \Phi \) is equal to \( C_{\text{cas}} \times V_{\text{visc}} \), where \( C_{\text{cas}} \) is the casein concentration of caseinate (25.10\(^{-3}\) g mL\(^{-1}\)) and \( V_{\text{visc}} \), the voluminosity obtained by viscosity measurements. Whey proteins were neglected because their concentrations were very low.

Average micelle diameters were estimated by a dynamic light-scattering method on a Coulter N4MD apparatus (Coultronics, Margency, France) at 20 °C after 1:300 dilution in Dalgleish buffer (1984) (Pierre et al, 1995). The computed autocorrelation function of the scattered light is related to the intensity-weighted average diffusion coefficient and to the particle diameter, assuming a log-Gaussian distribution. Diluted suspensions were stabilised for 10 min inside the cell holder, and a measurement of light intensity scattered at 90° during 300 s was carried out. Each suspension was acidified in duplicate and means were calculated so that each curve represents four experiments.

**Rheological and scanning electron microscopy of acid gels**

A constant speed cone penetrometer (Stevens LFRA) was used to evaluate firmness of acid gels with a cone of 90° moving 10 mm inside the gel at 2 mm s\(^{-1}\), as described by Korolczuk and Mahaut (1988). GDL was added to NPCS to reach pH 4.4, as described above. Just after addition, NPCS was placed in the test cup of 25 mm height and 62 mm diameter. The gel height was 20 mm. Measurements of firmness in Pa were performed at room temperature.

Gel samples at pH 4.4 (3 x 3 x 2 mm) were fixed at room temperature for 48 h in a solution of 2.5% glutaraldehyde prepared in 0.1 mol L\(^{-1}\) (pH 7.2), sodium cacodylate, dehydrated in alcohol series and critical point dried from carbon dioxide. They were fractured by hand, mounted on stubs and sputter coated with gold (about 70 nm). A Philips scanning electron microscope (XL 20) was operated at 12 kV. Particle sizes were determined on screen.

**Renneting**

Hansen rennet powder (Boll, Arpajon, France) was prepared at 50 g kg\(^{-1}\) as described by Famelart (1994) and used immediately after thawing at 1 mL per L of NPCS. Coagulation time of non-acid-aggregated samples prewarmed during 30 min was evaluated with a Formagraph at 30 °C.

**RESULTS**

**Initial NPCS characterization**

The pHs of NPCS were 7.45, 7.36 and 6.73 in water, NaCl and UF, respectively. Diffusible calcium and phosphorus contents were higher for NPCS in NaCl than in water (figs 2, 3). At pH 6.7, for NPCS dissolved in water, diffusible calcium and
Acidification of native phosphocaseinate

Phosphorus were 2.862 and 1.677 mmol kg⁻¹, respectively. In NaCl, these values became 5.525 and 2.435 mmol kg⁻¹ (table II). Mineral diffusible contents of NPCS in UF were 8.337 and 13.339 mmol kg⁻¹ for calcium and phosphorus, respectively. They are increased by the mineral contents of the ultrafiltrate used for dissolution. However, it appears that diffusible minerals of NPCS in UF were less than the diffusible content of NPCS in water plus the UF content (table II).

A higher content of non-micellar casein (fig 5) was observed for NPCS dissolved in water (18% of OD), than in NaCl (11.5%) and in UF (9%), and micelle sizes (fig 7) were 210, 220 and 240 nm, respectively. Initial solvation of NPCS (fig 8) dissolved in NaCl was 25% higher than the others.

**Acidification**

Figure 1 shows pH values of NPCS obtained with increasing amounts of GDL. The acidification curves were similar. The initial pH values were lower for NPCS in UF than in water or NaCl. GDL amounts needed to reach pH 4 were higher for NPCS in UF than in water or NaCl.

**Mineral solubilization with pH**

Between pH 7.0 and 5.2, diffusible calcium and phosphorus contents increased continuously, while below pH 5.2, diffusible calcium still increased and diffusible phosphorus decreased (figs 2, 3).

The extent of solubilization during successive steps of pH (pH 6.7–6.0; pH 6.0–5.6; pH 5.6–5.2; pH 5.2–4.0) as was done by Le Graet and Brulé (1993) showed a substantial increase of diffusible minerals during the step pH 6.7–6.0 and pH 6.0–5.6 for NPCS dissolved in high ionic strength. It is obvious from figures 2 and 3 that diffusible minerals of NPCS dissolved in NaCl reached the final, low values at higher pH.

Linear relations were found between micellar calcium and micellar inorganic phosphorus from pH 7.0 to pH 5.3 (fig 4). Slopes were higher for NPCS in water than in the other solvents.

**Table II. Calcium (Ca) and phosphorus (P) contents (mmol kg⁻¹) of UF (milk ultrafiltrate used to dissolve the powder) and diffusible mineral contents of the native phosphocaseinate suspensions (NPCS) dissolved in water, 0.1 mol L⁻¹ NaCl and UF at pH 6.7 and 20 °C.**

<table>
<thead>
<tr>
<th>Total content of UF (mmol kg⁻¹)</th>
<th>Diffusible content (mmol kg⁻¹) of:</th>
</tr>
</thead>
<tbody>
<tr>
<td>NPCS in water</td>
<td>NPCS in 0.1 mol L⁻¹ NaCl</td>
</tr>
<tr>
<td>Ca exp 1**</td>
<td>7.100</td>
</tr>
<tr>
<td>exp 2**</td>
<td>7.775</td>
</tr>
<tr>
<td>P exp 1**</td>
<td>13.677</td>
</tr>
<tr>
<td>exp 2**</td>
<td>12.677</td>
</tr>
</tbody>
</table>

* On the assumption that diffusible content of NPCS dissolved in water added up to the UF content: 9.800 mmol kg⁻¹ = total content of UF (7.100) + diffusible content of NPCS in water (2.700). ** Two replicates are presented.

**En supposant qu'il y a addition des concentrations diffusibles du NPCS dissout dans l'eau et des concentrations de l'UF: 9.800 mmol kg⁻¹ = teneur en minéraux totaux de l'UF (7.100) + teneur en minéraux diffusibles des suspensions de phosphocaséinate natif (NPCS) dans l'eau (2.700). ** Deux répétitions sont présentées.
Fig 1. Values of pH after glucono-delta-lactone (GDL) addition in native phosphocaseinate suspensions in water (Δ), 0.1 mol L\(^{-1}\) NaCl (■) and milk ultrafiltrate (□). Contents of GDL are expressed in g added for 100 g of final solution. GDL (lysoctone) was added at 20 °C and pH were measured after 15 h incubation at 20 °C.

Évolution du pH après l'addition de glucono-delta-lactone (GDL) aux suspensions de phosphocaséinate natif dans l'eau (Δ), NaCl 0,1 mol L\(^{-1}\) (■) et l'ultrafiltrat de lait (□). Les contenus en GDL sont exprimés en g ajoutés pour 100 g de suspension finale. La GDL (lysoactone) était ajoutée à 20 °C et le pH était mesuré après 15 heures d'incubation à 20 °C.

Fig 2. Solubilization with the pH at 20 °C of calcium from suspensions of native phosphocaseinate. Concentrations of calcium in permeates obtained by ultrafiltration on Centriflo CF25 are presented. The pH values were obtained after the addition of GDL during 15 h at 20 °C. Two replicates for each medium are presented. Same symbols as in the legend to figure 1.

Solubilisation du calcium des suspensions de phosphocaséinate natif avec le pH à 20 °C. Les concentrations en calcium dans les perméats obtenus sur Centriflo CF25 sont présentées. Les différentes valeurs du pH étaient obtenues 15 heures après l'addition de la GDL à 20 °C. Deux répétitions de chaque suspension sont présentées. Mêmes symboles que pour la figure 1.

**Casein dissociation**

Figure 5 shows the non-micellar caseins as the fraction of optical density present in 2h, 75 000 g supernatants. Non-micellar caseins at 20 °C appeared to have a peak at pH 5.4–5.5 in every medium. The fraction of non-micellar caseins at pH 5.4–5.5 determined by HPLC was 7%, 30% and 12% in water, NaCl and UF respectively. \(\kappa\)-Casein was the most important casein in supernatants of NPCS in water and UF (38 and 51% of total non-micellar caseins, respectively) and \(\beta\)-casein was present at the highest percentage (43%) in the supernatant of NPCS in NaCl. A decrease of non-micellar caseins with decreasing pH was observed between pH 7.4 and 6.0, particularly for the NPCS in water (fig 5).

Fig 3. Solubilization of phosphorus from native phosphocaseinate suspensions dissolved in water, 0.1 mol L\(^{-1}\) NaCl and UF with the decrease of pH at 20 °C. Concentrations of phosphorus in permeates obtained by ultrafiltration on Centriflo CF25 are presented. The pH values were obtained after the addition of GDL during 15 h at 20 °C. Two replicates for each medium are presented. Same symbols as in the legend to figure 1.

Solubilisation du phosphore des suspensions de phosphocaséinate natif dans l'eau, NaCl 0,1 mol L\(^{-1}\) et l'UF avec le pH à 20 °C. Les concentrations en phosphore dans les perméats obtenus sur Centriflo CF25 sont présentées. Les différentes valeurs du pH étaient obtenues 15 heures après l'addition de la GDL à 20 °C. Deux répétitions de chaque suspension sont présentées. Mêmes symboles que pour la figure 1.
Acidification of native phosphocaseinate

Casein precipitation
Nitrogen contents of 1000 g-supernatant decreased from 95% to 10% of total nitrogen around pH 5.0 (fig 6). The pH value at half precipitation was 5.2, 4.7 and 4.8 for NPCS in water, 0.1 mol L⁻¹ NaCl and UF, respectively. Precipitation appeared less abrupt at higher ionic strength.

Particle size analysis
Figure 7 shows the decrease of average micelle diameter from pH 7.0 to 5.4-5.5.

The reduction was about 10% in water and UF and 17% in 0.1 mol L⁻¹ NaCl, pH values lower than 5.5 caused the micelles to aggregate, and their size can no longer be determined.

Micelle solvation
Similar shapes for micelle solvation as a function of pH were observed in the different media (fig 8). Minimum solvation was found at pH 4.6, and a maximum peak appeared at pH 5.4. The first minimum for solvation was observed at pH 6.6 in water, at pH 6.3 in NaCl and at pH 6.1 in UF. Smaller
variations were obtained for phosphocaseinate dissolved in water than in UF or NaCl.

**Viscosity measurements**

The relative viscosity decreased from pH 7.5 to pH 6.4–6.5 for NPCS in water or NaCl, and from pH 7.0 to pH 6.0 for NPCS in UF (fig 9). The reduction amounted to 10%, 7% and 12% for water, NaCl and UF, respectively.

Voluminosity values obtained by viscosimetry and pellet solvation measurement showed the same tendencies: a decrease from pH 7.0 to 6.3 followed by an increase (table III). Values obtained by the two methods exhibit great differences in magnitude. Decreases in voluminosity were stronger for NPCS in UF than in the other medium.

**Structure and texture of acid gels**

Gel firmness obtained after acidification were significantly different (at least at 0.04%) in the three aqueous phases and decreased as ionic strength increased. Values of 239 ± 35 Pa, 40 ± 9 Pa and 52 ± 10 Pa were calculated, in water, NaCl and UF, respectively. It is evident from figure 10 that particles in gel made with water were larger (305 ± 141 nm) than with NaCl (128 ± 61 nm), and with UF (150 ± 84 nm). NPCS dissolved in water appeared to produce a fibrous gel at pH 4.4, with chains made of several particles, entrapping the water in well defined void spaces, while NPCS dissolved in higher ionic strength led to smaller particles aggregated in clusters.

**NPCS renneting**

Decreasing the pH led to reduction of rennet coagulation times as generally observed by others on milk. The pH effect was stronger between pH 7.0 and 6.5. Between pH 5.9 and 5.5, coagulation times were only slightly dependent on pH, specially for NPCS in UF. Decreasing the ionic strength from NaCl to water led to high reduction of coagulation times from 23 min to 8 min at pH 6.7 and to greater differences at increasing pH values. From pH 7.0 to 6.7, rennet coagulation times decreased from UF to NaCl medium, while for pH values lower than 6.7, higher coagulation times were observed for NaCl suspensions.

Table III. Effect of pH on casein voluminosity at 20 °C. \( V_{\text{pellet}} \) and \( V_{\text{visc}} \) are the casein voluminosity obtained from pellet hydration measurements and viscosity measurements on native phosphocaseinate suspensions.

<table>
<thead>
<tr>
<th>Aqueous phase</th>
<th>( V_{\text{pellet}} ) at pH 7.0</th>
<th>( V_{\text{pellet}} ) at pH 6.3</th>
<th>( V_{\text{pellet}} ) at pH 5.6</th>
<th>( V_{\text{visc}} ) at pH 7.0</th>
<th>( V_{\text{visc}} ) at pH 6.3</th>
<th>( V_{\text{visc}} ) at pH 5.6</th>
<th>( \Delta_{\text{max}} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>2.67</td>
<td>2.57</td>
<td>2.73</td>
<td>3.39</td>
<td>2.90</td>
<td>3.22</td>
<td>0.10</td>
</tr>
<tr>
<td>0.1 mol L(^{-1}) NaC</td>
<td>3.19</td>
<td>3.03</td>
<td>3.18</td>
<td>4.90</td>
<td>4.09</td>
<td>5.06</td>
<td>0.27</td>
</tr>
<tr>
<td>UF</td>
<td>2.82</td>
<td>2.42</td>
<td>2.51</td>
<td>5.03</td>
<td>3.59</td>
<td>3.74</td>
<td>0.43</td>
</tr>
</tbody>
</table>

\( \Delta_{\text{max}} \) Maximal reduction of voluminosity.

\( \Delta_{\text{max}} \) Réduction maximale de la voluminosité.
Acidification of native phosphocaseinate

The effect of the aqueous phase on the initial NPCS

Diffusible salts being responsible for about 50% of the buffering of milk (Lucey et al, 1993), it was not surprising that the buffering of micelle suspension was greater in UF (fig 1).

According to Brulé et al (1974), Van Hooydonk et al (1986b) and Le Graet and Brulé (1993), the addition of 0.1 mol L^{-1} NaCl in milk at pH 6.7 leads to about a 0.75–1.00 mmol kg^{-1} diffusible calcium increase (11% increase) and no diffusible phosphorus increase. In this study, the solubilization of phosphocaseinate powder in NaCl, compared to solubilization in water led to a solubilization of 2.662 mmol kg^{-1} of calcium and of 0.758 mmol kg^{-1} of phosphorus from the colloidal calcium phosphate (CCP) (mean values from table II). According to a hypothetical tricalcium-phosphate composition (Ca/P = 1.5 mol mol^{-1}), the increase of ionic strength due to NaCl caused solubilization of 0.758 mmol kg^{-1} phosphorus and this would correspond to solubilization of 1.137 mmol kg^{-1} of calcium bound to phosphorus. It follows that 43% of NaCl-solubilized-calcium was bound to phosphorus and 57% likely bound to the phosphoserine residues. This solubilization of the CCP occurred because, first, the diffusible phase of the NPCS dissolved in NaCl was not saturated with calcium phosphate, as in milk and, second, because an increase in ionic strength is known to produce a decrease of the free ion activity coefficient. This increases the stochiometric solubility product of salts and consequently, the concentration of ionic...
Fig 8. Pellet solvation in g of water per g of dry pellet as a function of pH, at 20 °C. Drained pellet obtained after 75 000 g, 2 h centrifugation of native phosphocaseinate suspensions were dried at 103 °C for 7 h. Two repetitions are shown for each medium. Same symbols as in the legend to figure 1.

Fig 9. Relative viscosity of native phosphocaseinate suspensions obtained from viscosity measured in coaxial cylinder at 94.5 s⁻¹ strain rate and 20 °C. Relative viscosity (suspension viscosity/solvent phase viscosity) was obtained using 1.00, 0.95 and 1.09 measured values for viscosity of water, 0.1 mol L⁻¹ NaCl and UF, respectively. Two repetitions are shown for each medium. Same symbols as in the legend to figure 1.

species such as calcium phosphate. Anyway, it is well known that sodium added tends to partly displace calcium from the CCP to the aqueous phase. According to Grufferty and Fox (1985) and Le Graet and Brulé (1993), calcium directly bound to phosphoseryl residues is affected by added NaCl.

The effect of the aqueous phase on the acidification process

The diffusible mineral values with pH decrease obtained for NPCS in UF are within the range of values published elsewhere (Dalgleish and Law, 1989; Le Graet and Brulé, 1993). The two ultrafiltrates used were not exactly the same (table II), which would explain the discrepancy seen in figures 2 and 3 for NPCS dissolved in UF.

The differences in mineral solubilization with acidification between water, NaCl and UF are difficult to explain, because the dissolution of the phosphocaseinate powder in the aqueous phase (notably in water and NaCl) can lead to a mineral solubilization (table II). Total mineral solubilization obtained with the pH reduction was observed at higher pH values for NPCS dissolved in NaCl (figs 2, 3). On the one hand, the increased ionic strength led to a higher salt solubility (Walstra and Jenness, 1984). On the other hand, the increase in ionic strength caused a lower apparent pK value for ionic groups of caseins and consequently a retardation of the mineral solubilization (Baumy et al, 1989). This explained the lower pH values obtained for nitrogen precipitation when micelles were suspended in a higher ionic strength medium (fig 6). Slopes and intercepts for linear relations between micellar calcium and mi-
Acidification of native phosphocaseinate

Fig 10. SEM micrographs of gel at pH 4.4 obtained with GDL for native phosphocaseinate suspensions (NPCS) in water (a, d), 0.1 mol L\(^{-1}\) NaCl (b, e) and UF (c, f), at two different magnifications. In water, casein particles were larger, and gel appeared as a fibrous network of chained particles, with well-defined void spaces. The gel of NPCS in NaCl or UF was composed of clusters of smaller particles with large, and diffuse void spaces.

Images de microscopie à balayage des gels à pH 4,4 obtenus avec la GDL à partir des suspensions de phosphocaseinate natif (NPCS) dans l'eau (a et d), NaCl 0,1 mol L\(^{-1}\) (b et e) et l'UF (c et f), à deux grandissements différents. Dans l'eau, les particules caséiniques étaient plus grandes, et le gel apparaissait comme un réseau fibreux de particules en chaînettes, avec des pores bien définis. Le gel obtenu pour le NPCS dans NaCl ou l'UF était composé de paquets agglomérés de particules plus petites, avec des pores de taille moins bien définie.
cellular inorganic phosphorus for NPCS in UF were in total agreement with the range of values cited in Dalgleish and Law (1989) (1.61–1.98 mol Ca/mole P). The slopes were higher for NPCS dissolved in water, suggesting that micellar calcium directly bound to casein was solubilized together with the colloidal calcium phosphate. This suggests that the calcium dissociation with pH can be differently affected by the ionic strength.

In the changes occurring during lowering of the pH some successive stages can be conveniently distinguished. The various changes occurring are dependent on each other. Dissociation of casein from the micelles and changes in solvation are involved.

Phase I

The pH ranges of this step were from pH 7.0–7.5 to 6.6, to pH 6.3 and to 6.1 for NPCS dissolved in water, NaCl and UF, respectively. Micelle size, micelle solvation, and viscosity decreased during this step as described by Roefs et al (1985) on milk at 8 °C, by Van Hooydonk et al (1986a) and by Banon and Hardy (1991, 1992). This is in accordance with a voluminosity decrease due to the ionisation regression as the pH decreased. This led, first, to less negatively charged proteins and hence to the reduction of repulsive forces between adjacent chains. This results in the onset of the progressive collapse of the outer-hairy-layer described by Banon and Hardy (1992) between pH 6.6 and 5.8 in milk at 20 °C. Second, and well established, the protonation of mineral phosphate led to the dissociation of calcium and phosphorus.

During this phase, less mineral matter dissociated from micelles in water, and more was dissolved from micelles in NaCl, compared with micelles in UF. It can be concluded that it was not directly the mineral solubilization that governed the physicochemical changes, and that the increase in ionic strength led to increased mineral solubilization in these ranges of pH, probably because solubility of calcium phosphate is increased by higher ionic strength.

It is noteworthy that, during this step, a decrease of non-micellar caseins was observed, particularly for the NPCS in water (fig 5). A similar decrease was observed in the study of Dalgleish and Law (1988), but only at 30 °C. It can be suggested that some non-micellar caseins present in NPCS dissolved in water may return to the micelles, due to the reduction of repulsive forces between charged amino acid residues.

Phase II

This step ended at pH 5.5–5.6, at which the casein dissociation was maximal, as in Dalgleish and Law (1988) on milk at 20 °C. As the mineral content of micelles further decreased, viscosity, casein solvation and casein dissociation from micelles seemed to increase. This can be explained by the reduction of electrostatic interactions between caseins, and the subsequent increase of water-casein interactions. Micelles placed in NaCl exhibited a greater voluminosity (fig 8), and more casein dissociation (fig 5). Replacement of calcium by sodium or addition of NaCl in milk is known to increase the milk viscosity and the water-holding properties of casein micelles. This is explained, on the one hand, by changes in the structure of micelles probably due to the displacement of calcium from the CCP by sodium and, on the other hand, by the increase of the ionisation of caseins due to the decrease of activity coefficients. The latter led to the increase of the micelle voluminosity and also of the functional properties of casein material (Burgess, 1982; Snoeren et al, 1984; Creamer, 1985).

Casein solubilization values at pH 5.4–5.5 for NPCS in UF (12%) were smaller than the values obtained by Dalgleish and Law (1988) for milk at 20 °C (30%). Release of soluble casein during microfiltration of the phosphocaseinate (TN-NCN in
Acidification of native phosphocaseinate microfiltrates was less than 5%) prior to its drying process might account for this discrepancy. \( \kappa \) - and \( \beta \)-caseins were mainly released from the micelles, because \( \kappa \)-casein is supposed to be mainly located at the micellar surface and \( \beta \)-casein is able to diffuse out of the micelle. Supernatants of NPCS in NaCl were enriched in \( \beta \)-casein, though ionic strength increase is known to have a negative effect on \( \beta \)-casein extraction (Famelart et al, 1989).

According to Van Hooydonk et al (1986a), the swelling of micelles between pH 6.0 and 5.6 is only explained by the solubilization of the CCP. An increase in ionisation or in hydrophilic characteristics of molecules leads to an increase in amphiphilic properties, ie an increase of the tendency of hydrophobic part to cluster. Experiments on the effect of heating on milk proteins (Haque and Kinsella, 1988) show that hydrophobic side chains residues were withdrawn within proteins, while ionisable residues were more exposed. When the charge of proteins is reduced by pH changes of the eluant in high-performance hydrophobic interaction chromatography, hydrophobic interactions increase, and vice versa, which means that hydrophobic residues are less withdrawn in the core of the protein when less ionised (Fausnaugh and Regnier, 1986; Hjerten et al, 1986). Hydrophilicity of proteins would affect the exposure of hydrophobic residues, ie, the contribution of hydrophobicity to the formation and stability of protein complexes. It might be possible that during the decrease of ionic interactions between casein and between casein and water, a substantial reduction of hydrophobic interactions took place, leading to the swelling of micelles or even of submicellar units.

Micelle voluminosity increased during this step, while the micelle diameter showed only a further decrease. The weakening of internal bonds of the micelle led to its swelling. This disaggregation led to the decrease in diameter because free submicelles are leaving the micelle, despite an increase in voluminosity.

**Phase III**

Between pH 5.5–5.6 and pH 5.3–5.4, casein dissociation decreased, while the average micelle size and solvation increased. According to Snoeren et al (1984), the peak in solvation may be partly due to the dissociation of \( \beta \)-casein from the micelles. This unexpected observation of solvation increase and casein dissociation decrease was already mentioned by Van Hooydonk et al (1986a), while according to Roefs et al (1985), this discrepancy is absent in the cold. The solvation changes being mainly due to the changes of the micellar structure, it is then possible to observe a decrease in 'soluble' casein, despite a solvation increase. As the pH decreased, new interactions, probably hydrophobic and electrostatic, occurred between these caseins in free submicelles, which explained the reduction in 'soluble' caseins. The caseins are held together by these new interactions and we assumed that this new structure is able to retain more water, because less deformable during the centrifugation stress. Interactions being reduced at 4 °C, the less rigid structure did not show this behaviour: casein solvation and dissociation decreased simultaneously in the cold.

**Phase IV**

The decrease of solvation and of soluble total nitrogen from pH 5.3–5.4 to 4.4–4.6 were due to casein precipitation at the isoelectric pH, caused by the general charge neutralisation. The effect of ionic strength on the isoelectric pH values of the four caseins is substantially different. This is expected from the comparison of isoionic pH values calculated in Swaisgood (1992) and isoelectric pH values. This explained that the decrease of soluble nitrogen with pH was less abrupt for high ionic strength suspensions (fig 6).
Phase V

Below the pH of minimal solvation, i.e., the isoelectric pH (pH 4.4–4.6), casein began to be positively charged. Water-holding properties of proteins increased, as the 'solute' casein contents. Negative ionisation of phosphate groups at these pH values (pK 3; Walstra and Jenness, 1984) would explain adsorption of inorganic phosphate on acid casein (fig 3) as described by Le Graet and Brulé (1993).

The effect of the aqueous phase on acid gels

Reliable correlations between microstructure and firmness in acid gels were already mentioned, depending on the temperature, the acidulant used, and the pH values (Harwalkar and Kalab, 1981, 1986; Kalab et al, 1983). Gels in the form of coarse clusters, with large pores result in low firmness, while a more continuous protein network with chained particles exhibit greater firmness. Increasing ionic strength of Na-caseinate with increasing amounts of NaCl is known to produce acid gels with lower modules (Roefs and Van Vliet, 1990). In this study, the particles in acid gels of higher ionic strength were smaller and the gels appeared as coarse clusters of particles. Gels of NPCS in water at pH 4.4 are composed of particles with a higher net positive charge than in higher ionic strength. This is clearly demonstrated in figure 6, were the pH at half precipitation was 5.2 for micelles in water and about 4.7 for high ionic strength. The higher positive charge of particles in water may first prevent interparticle interactions, resulting in a finer and chained gel, and second, resulted in a swelling of casein particles.

The effect of the aqueous phase on rennet coagulation time

The reduction of ionic strength of the NPCS markedly decreased the rennet coagulation time. Reduced rennet clotting time for native phosphocaseinate compared with milk was already mentioned by Pierre et al (1992). We observed (Famelart, unpublished work) that the reconstitution of the phosphocaseinate used in this study in 0 to 0.12 mol L$^{-1}$ NaCl solutions resulted in the decrease of both the enzymatic and the aggregation rates. An increase in rennet clotting time with ionic strength increase was previously observed by Visser et al (1980) on model substrates, and by Grufferty and Fox (1985) and Van Hooydonk et al (1986b) on milk, provided that the pH after the salt addition has been corrected. Ionic strength increase is supposed to screen the negative charge on chymosin and the positive clusters in κ-casein near the Phe105-Met106 peptide linkage, and to lower the enzyme-substrate attraction (Visser et al, 1980).

Curves of relative viscosity versus pH—reflecting micelle voluminosity—and of rennet coagulation times from pH 7.0 to 6.3 were very close, with substantially lower voluminosity in water. In our opinion, and with the suggestion that rennet-induced aggregation is diffusion-controlled (Van Hooydonk et al, 1984; Famelart, 1994), steric repulsions due to increased voluminosity in NaCl and UF were mainly responsible for long coagulation times and for the changes in rennet coagulation time from pH 7 to 6.3. At lower pH values, the reduction of negative charge would have played the major part.

CONCLUSION

Native phosphocaseinate used in this study might be regarded as a relevant model of milk micelles. It raised the possibility of studying the effect of the aqueous phase on the mineral exchanges between diffusible and colloidal phases and the behaviour of the micelles toward acidification or rennet processes in the absence of whey proteins. It is relevant to mention that it con-
sists of an attractive substrate for technological milk processing such as yoghurt and cheese manufacture. NPCS dissolved in UF behave like native milk. Moreover, when dissolved in water, the micelle suspension is able to form a strong gel at acidic pH and to coagulate with rennet at shorter times than milk, as already shown by Pierre et al (1992).

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REFERENCES

MH Famelart et al


