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Formation of heat-induced protein aggregates in milk as a means to recover the whey protein fraction in cheese manufacture, and potential of heat-treating milk at alkaline pH values in order to keep its rennet coagulation properties. A review

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Abstract – The heat-treatment of cheese milk, or whey, to denature the whey proteins has long been the most applied means of recovering these proteins, either directly in the cheese curd, or as added to the cheese milk prior to renneting. In heat-treated milk, the interaction of the denatured whey proteins with the casein micelles, however, limits the primary phase of the enzymatic reaction, and prevents fusion of the casein micelles. Cheeses containing heat-denatured whey proteins also exhibit excessive moisture, a crumbly and soft texture, and poor meltability. Various technological means to reduce these drawbacks are reviewed. Especially, it is generally accepted that the heat-treatment of skim milk at alkaline pH generates aggregates of denatured whey proteins and κ-casein in the serum phase of milk, rather than on the surface of the casein micelles. As a consequence, the casein micelles are depleted in κ-casein, and free of denatured whey proteins. However, the attempts made to exploit this interesting protein distribution in cheese-making remain scarce, despite their promising results.

rennet coagulation / heat-treatment / whey protein / cheese

摘 要 – 在干酪生产过程中通过热处理回收乳清蛋白, 以及在碱性 pH 下加热牛乳以保持凝乳酶的凝固特性 （文献综述）。长期以来在干酪的生产过程中一直采用热处理牛乳或乳清使乳清蛋白变性, 通过这种方法来回收这些蛋白。有时直接将这些蛋白加入凝块或在凝乳之前将这些蛋白加入牛乳中。在牛奶的热处理过程中, 由于变性的乳清蛋白和酪蛋白胶束之间相互作用限制了初期的酶反应, 进而阻止了酪蛋白胶束的融合（聚合）。干酪中含有变性的乳清蛋白会表现出水分含量过高、质地脆、软、融融性非常差。关于如何克服干酪生产中这种缺陷的技术手段已有许多文献报道。通常比较容易接受的方法是在碱性条件下热处理脱脂乳, 此时变性的乳清蛋白和 κ-酪蛋白的凝胶物存在于牛乳的乳清相中, 不是在酪蛋白胶束的表面。其结果是酪蛋白胶束中既不存在 κ-酪蛋白也不含有变性的乳清蛋白。尽管这种处理方法可以获得令人满意的结果, 但对这种蛋白分布在干酪生产中的应用还很少。

凝乳酶凝乳 / 热处理 / 乳清蛋白 / 干酪

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Résumé – Récupération des protéines sériques du lait dans le caillé fromager au moyen de la formation d’agrégats thermo-induits, et intérêt de chauffer le lait à pH alcalin dans le but de conserver son aptitude à la coagulation prèseuse. Revue. Le traitement thermique du lait ou de lactosérum est un des moyens les plus utilisés pour incorporer les protéines sériques ainsi dénaturées, soit directement dans le caillé, soit sous forme d’ingrédient ajouté au lait fromager. Cependant, l’interaction entre les protéines sériques dénaturées et les micelles de caséines, due au chauffage du lait, a pour effet de limiter la phase primaire de la coagulation par la prèseuse, ainsi que la fusion des micelles déséquilibrées. Les fromages obtenus à partir de lait chauffé sont aussi plus humides, plus mous, plus granuleux et moins aptes à la fonte que les fromages standards. Les moyens technologiques de pallier ces inconvénients sont rappelés. En particulier, il est connu que le traitement thermique du lait à pH alcalin favorise la formation d’agrégats de protéines sériques dénaturées et de caséine \( \kappa \) dans la phase soluble du lait, plutôt qu’en surface des micelles de caséines. En conséquence, la teneur en caséine \( \kappa \) des micelles est réduite et celles-ci ne sont pas recouvertes de protéines sériques dénaturées. Cependant, peu d’études ont tenté d’exploiter ces caractéristiques du lait chauffé à pH alcalin en technologie fromagère, malgré des résultats prometteurs.

coaulation prèseuse / traitement thermique / protéine sérique / fromage

1. INTRODUCTION

In traditional cheese-making processes, a large proportion of the whey proteins (\( \beta \)-lactoglobulin, \( \alpha \)-lactalumlin, bovine serum albumin, immunoglobulins, lactoferrin, enzymes, etc.) are eliminated in the whey during drainage. These whey proteins have excellent nutritional value, due to their high content in cystein, and to the high content of \( \alpha \)-lactalumlin in tryptophan residues [91]. They also have very good foaming and emulsion properties in their native state, or when partially denatured [34, 79, 167], and high water retention and gelation properties in their denatured state [32, 33, 79, 155, 160].

The recovery of the whey protein fraction in cheese technology has long been extensively investigated. Heat-treatment of the milk to denature the whey proteins and aggregate them with the casein fraction is one of the major means to reach this goal; however, this causes severe inhibition of gel formation, and texture defects of the curd [75, 87, 142, 168]. The present review stresses that heat-treatment of skim milk at pH > 6.7 induces changes in the distribution of the heat-induced protein aggregates between the serum and the micelle phases of milk, related to the higher dissociation rate of \( \kappa \)-casein upon heating at alkaline pH values, and suggests that new technological approaches may be investigated in order to optimise the recovery of the whey proteins in the cheese curd, and improve the rennet coagulation properties of the heated milk.

2. PRINCIPAL MEANS FOR THE RECOVERY OF THE WHEY PROTEIN FRACTION IN CHEESE-MAKING

2.1. Thermal coprecipitation of the whey proteins and casein

Heat-treatment of milk at temperatures exceeding ~ 60 °C leads to denaturation (unfolding) of the whey proteins, followed by aggregation through hydrophobic interaction and disulphide-thiol interchanges to form heat-induced aggregates, either on the surface of the casein micelles (micelle-bound aggregates), or in the serum phase of the milk (serum aggregates) [3–5, 116]. These aggregates essentially contain whey proteins and \( \kappa \)-casein; micelle-bound aggregates may also contain traces of \( \alpha s_2 \)-casein [26, 62, 121, 139, 144]. They can reach a diameter of 30 to 100 nm [76, 157].

Serpelloni [136], Banks [10], Banks et al. [12] and Law et al. [86] have taken advantage of the heat-induced binding of denatured whey protein to the casein micelles to recover the whey proteins in Cheddar curds. Another application of this approach in cheese-making can be found in Sanchelima [130], where a continuous flow process is described. The heat-treatments used in these studies range from 80 to 95 °C for a few seconds to several minutes, so that significant heat-denaturation of the whey proteins can be obtained [30, 31]. Ultrafiltration (UF) may be used to increase whey
Rennet coagulation of milk heated at alkaline pH

protein concentration, and hence increase the rate of whey protein denaturation during heating [122]. For fresh cheese, the so-called Thermo-Quark process can be applied, where the rennet coagulum of acidified, heated (90–95 °C, 2–3 min) skim milk is further heated at 50–60 °C for 1–2 min to complete the denaturation of α-lactalbumin prior to drainage by centrifugation [118].

2.2. Addition of heat-denatured whey protein to cheese milk

To avoid the consequences of the heat-induced interaction between denatured whey proteins and κ-casein on the rennet coagulation properties of the milk (see Sect. 3), the whey proteins may be recovered by heating the lactoserum itself, prior to addition to cheese milk [109]. During heating, the whey proteins denature and aggregate through hydrophobic interactions and thiol/disulphide exchanges. Increased protein concentration prior to heating allows increased denaturation rates [66, 68], e.g. using UF concentration of whey [21], as opposed to concentration of total dry matter using evaporation where increasing lactose content exerts a protective effect on the native structure of β-lactoglobulin [123, 124]. As opposed to the milk medium, where the reported sizes for heat-induced protein aggregates are < 1 μm, the whey proteins in heated lactoserum readily form large, visible flocs that can reach up to the mm scale. The heat-aggregated whey protein can then be easily concentrated by UF [21, 119] or centrifugation, e.g. by the Centri-Whey process [52, 53, 161]. The size of the aggregates can be controlled by physical means, including homogenisation [110] or shear of the heated whey [40], or the control of the whey protein denaturation through the temperature of heating [8, 147]. At constant temperature, decreasing lactose concentration reduces the protective effect of the sugar on denaturation of the whey proteins [21, 124], while a combination of increased heating regimen and lactose concentration produces large aggregates with a sandy texture [70, 147]. Adjusting the pH of the whey, generally in the range 3.5 to 7, prior to heat-treatment, is also a common method to control the size and hydration of the aggregates, although the reported optimum pH values differ significantly [40, 84, 148]. Other chemical means used to control the size of the aggregates are the variation in concentration of calcium or sodium salts [23, 148], or the use of transglutaminase enzyme [145].

Commercial applications of microparticulated whey products have found outlets under trademarks such as Dairy-Lo™ (Pfi zer Food Science Group, New York, NY, USA) or Simplesse® (NutraSweet Co., St Louis, MO, USA). These products are largely used as fat substitutes in the manufacture of low-fat products, including cheeses.

As an alternative to the use of commercial ingredients, the addition of particulated whey protein to cheese milk can also be performed in a single flow process. This involves the separation of the casein and whey protein fractions using membrane filtration techniques [126] and separate heating of the two fractions, followed by re-combination [105].

2.3. Use of membrane technologies to pre-concentrate cheese milk

As opposed to the above approaches, concentration of the total protein fraction of milk by UF allows the retention of native whey proteins into either intermediate concentrated retentates, or “pre-cheese” retentates which have essentially the composition of the targeted cheese [103, 112]. The pre-cheese can be coagulated using acidification and/or renneting, and processed up to moulding without extended drainage [38, 103]. Prior heat-treatment of the milk, resulting in formation of aggregates of denatured whey proteins and κ-casein prior to concentration, may be used to prevent the loss of the concentrated whey proteins during syneresis [49]. Pre-concentration of whey using UF, prior to its addition to milk and concentration into liquid pre-cheese [29] or to UF-concentrated milk [151] have also been proposed to increase whey protein recovery in the cheese curd.

A major drawback of the UF concentration of cheese milk is the increase in the
buffering capacity of the retentates, because of the concentration of colloidal calcium phosphate. Satisfactory corrections are possible through acidification of the milk prior to concentration, or increase in ionic strength by addition of sodium chloride during or after concentration [75, 111]. Both methods induce the solubilisation of calcium ions, either through reducing ionisation of the phosphoseryl groups of the casein molecules, or displacement of calcium by sodium within the micelles [20, 88]. Addition of sodium chloride to the retentate will also prevent coagulation during the UF process, especially in the case of heated milk that is acidified prior to or during UF. Replacement of calcium by sodium induces an increase in the surface charge, voluminosity and hydration of the casein micelles; and the increased ionic strength of the medium will lower their apparent isoelectric pH [39]. These effects of NaCl addition therefore ensure a higher stability of the milk system to environmental changes.

For the above reasons, the most successful development of UF cheeses, in particular made with heated milk, have been high-moisture, low-pH white cheeses such as Quarg and Cast Feta. In the latter case, UF concentration of heated milk improves smoothness of the curd [87] and processing of the curd directly in its retail package prevents fine losses [112]. Alternatively, ultrafiltration can be performed after the complete acidification and renneting of the milk, to make fresh cheeses or cream cheeses. Positive relationships have been reported between the amount of heat-denatured whey protein in the starting milk, and increased viscosity of the fresh cheeses [81, 82, 101].

### 2.4. Whey cheeses

The whey proteins present in by-products of traditional cheese-making may also be recovered through the preparation of specific traditional whey cheeses made from the concentration of whey, or of a mix of whey and milk, and moulding of the concentrate [42]. Recent reviews have made a distinction between two processes for making whey cheeses [45]. A first group of whey cheeses may be prepared by the combined heating and evapo-concentration of a mixture of whey and milk, until Maillard browning and a high viscosity are reached, then moulding of the concentrated product. The sugars crystallise on further cooling, to give the appropriate texture of the products. Theses cheeses are essentially produced in Scandinavian countries, e.g. Mysost in Norway, and in some Mediterranean countries like Greece, which produces Myzithra [56, 135].

The second group of whey cheeses, that includes Ricotta in Italy, Sérac or Brousse in France, and Brocciu in Corsica, are produced by the heat-treatment of sweet whey, with or without added milk, concentrated or not, at temperatures of 80–95 °C for 30 to 45 min [56, 127]. The whey proteins extensively denature and aggregate until flocs are formed that can be easily collected by filtration or centrifugation. The precipitate can be further acidified by addition of an acidulant or of lactic acid bacteria.

Whey cheeses comply with different standards than those that apply to cheeses. The distinction is essentially based on the casein to whey protein ratio of the starting dairy material, that should be equal to or higher than that of milk in standard cheeses [41], and lower in that of whey cheeses [42]. Also, the manufacture of standard cheeses involves the coagulation of the milk protein, in particular, the casein fraction, by rennet or another coagulating enzyme to form a gel or into any other end-product having similar textural and chemical properties [41].

### 3. THE USE OF pH AS A MEANS TO CONTROL THE FORMATION OF HEAT-INDUCED MILK PROTEIN AGGREGATES

#### 3.1. Disulphide bonds, number and size of the heat-induced protein aggregates

Various studies have reported the strong impact of pH on the denaturation and aggregation of the whey proteins. The denaturation rates of α-lactalbumin and β-lactoglobulin in milk heated at 80 °C both increased with pH from 5.5 to 8.8, although the two proteins
Ren net coagulation of milk heated at alkaline pH had different profiles of sensitivity against pH [85]. In model solutions of β-lactoglobulin [66, 68, 163] or in skim milk [57], heated at pH values in the maximum range 3 to 10, heat-induced protein aggregates were formed at all pH values. The aggregation rate increased with pH values in the range 6.5 to 8.0, so that more aggregates were formed at pH 7.0 or 8.0 than at pH 6.5 [66, 68]. The reactivity and accessibility of the free thiol of β-lactoglobulin was higher at pH 8.0 than at 7.0, as a result of the proximity of the pK value of the thiol (~ 8.2) [66]. As a result, the conversion of thiols into disulphide bonds increased with pH up to at least 6.9 [163] or up to pH 11 [113], as did the contribution of intermolecular disulphide bonds in the formation of heat-induced protein aggregates of β-lactoglobulin [67] or of κ-casein and whey protein in milk [57]. Conversely, the contribution of electrostatic interactions decreased as pH increased, as a result of the increased negative charge of the β-lactoglobulin (isoelectric pH ~ 5.4) [67].

The effect of highly alkaline pH values on the denaturation and sulphydryl-mediated polymerisation of whey proteins has been reported to occur even in the absence of heating, so that high molecular-weight aggregates were formed at 22 °C in solutions of whey protein isolate at pH 9 and 11 [113]. This property was exploited by Connolly [25] in the manufacture of co-precipitates of casein and whey proteins after a short exposure of milk materials to pH 9 to 11, and subsequent precipitation at acidic pH values.

The analysis of heat-induced aggregates of β-lactoglobulin by size-exclusion chromatography, light scattering, or gel electrophoresis, showed that the molecular mass of the aggregates was about ten-fold smaller (decrease from 10^6 to 10^5, or from 10^7 to 10^6 g·mol–1, depending on conditions) as the pH of heat-treatment was increased from 6.5 to 8.0 [66–68]. A negative relationship between the pH of heat-treatment and the size of the heat-induced aggregates was also reported for the aggregates formed on the surface of casein micelles, in milk at pH 6.35 to 6.90 [156] or 6.5 to 6.7 [4], and for those formed in the serum phase of milk at pH 6.5 to 7.2 [128], but not at pH 6.7 to 6.9 [156]. Hoffmann and Van Mil [67] suggested that conditions favourable to disulphide bonding at high pH values, and therefore to termination reaction (e.g. SH/SH oxidation, [113]), and to over-aggregation through hydrophobic interactions at low pH values, could account for the relationship of the size of the heat-induced whey protein aggregates with pH.

These results, summarised in Table I, therefore indicate that the heat-induced aggregates of whey proteins form at a faster rate, and in greater amounts, as the pH of

<table>
<thead>
<tr>
<th>Property of the aggregates</th>
<th>Variation as pH increases</th>
<th>pH range studied</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentration</td>
<td>Increases</td>
<td>≥ 6.5</td>
<td>[57] (skim milk), [66, 68, 163] (β-lg)</td>
</tr>
<tr>
<td>Proportion of covalent bonds in the aggregates</td>
<td>Increases</td>
<td>≥ 6.5</td>
<td>[67] (β-lg), [113] (WPI)</td>
</tr>
<tr>
<td>Size</td>
<td>Decreases</td>
<td>[6.3–8.0]</td>
<td>[66–68] (β-lg), [4, 128], [156] (skim milk)</td>
</tr>
<tr>
<td>Proportion of aggregates bound to the casein micelles</td>
<td>Decreases</td>
<td>[6.5–7.5]</td>
<td>[4–6, 57, 117, 139–141], [156] (skim milk)</td>
</tr>
<tr>
<td>Proportion of κ-casein in the aggregates</td>
<td>Increases?</td>
<td>[6.3–6.9]</td>
<td>[156]</td>
</tr>
</tbody>
</table>
heat-treatment increases. They are also smaller, and richer in covalent disulphide bonds, as pH increases. Further information, e.g. on the effect of the pH of heat-treatment on the initial stages of denaturation of the whey proteins, can be found elsewhere [35].

3.2. Dissociation of κ-casein and distribution of the heat-induced protein aggregates

Early studies on the stability of heat-treated or UHT-treated skim milk showed that κ-casein dissociated more extensively from casein micelles during heating at pH > 6.9, than at lower pH values [140, 141]. Because they interact with κ-casein upon heating, the denatured whey proteins remain in the serum phase when the heat-treatment is performed at pH > 6.9, whereas a high proportion of them can be sedimented with the casein micelles by ultracentrifugation when the heat-treatment is performed at lower pH values [57, 117, 139–141]. Virtually no aggregates could be found in the serum phase of milk at pH < 6.7 [156]. Other studies showed a strong relationship between the pH of milk and the proportion of β-lactoglobulin attached to the casein micelles upon heating. In skim milk heated at 90 °C, 70–80% w/w of the β-lactoglobulin was bound to the micelles at pH 6.5, versus 30% w/w at pH 6.7, and 10% w/w at pH 7.1 [4–6]. In skim milk heated at 80 °C, 60% w/w of the β-lactoglobulin was bound to the micelles at pH 6.55, versus 30% w/w at pH 6.9 [156]. In concentrated milk heated at 120 °C, 40% w/w of the whey protein was bound to the micelles at pH 6.55, versus 25% at pH 6.85 [139]. At a defined extent of denaturation of the whey protein, the hydrodynamic diameters of the casein micelles after heating decreased with increasing pH of heat-treatment, which was attributed to the binding of less numerous and/or smaller whey protein/κ-casein aggregates to the surface of the micelles [4, 7].

All these results clearly show that the higher the pH of heat-treatment, the higher the dissociation rate of κ-casein, and the more heat-induced aggregates found in the serum phase, rather than on the surface of the casein micelles (Fig. 1, Tab. 1).

However, it is not yet established whether the formation of heat-induced aggregates in the serum phase of milk is a cause, or a consequence, of the heat-induced dissociation of κ-casein. On the basis that the dissociation rate of κ-casein did not increase as fast as the proportion of heat-induced serum aggregates as a function of pH, Anema and Li [5] suggested that the increased negative charge of the casein micelles at higher pH values prevented the denatured whey protein from interacting with the surface of the casein micelles, so that dissociation of κ-casein should precede its interaction with the whey protein in the serum phase. Conversely, other studies have reported a positive relationship between the amount of β-lactoglobulin in the milk,
and the dissociation rate of κ-casein, to the extent that blocking the free thiol group of β-lactoglobulin with N-ethylmaleimide (NEM) prevented the extensive dissociation of κ-casein upon heating [3, 141]. Furthermore, a positive relationship has been demonstrated between the pH of heat-treatment, and the heat-induced dissociation of the caseins, especially κ-casein [2, 139]. κ-Casein dissociated to a greater extent as the pH and the temperature of heat-treatment increased, especially > 60 °C [2]. The current hypothesis that would best explain most of these results is that the κ-casein dissociated upon heating is prevented from returning to the surface of the micelles because it aggregates with denatured whey proteins in the serum phase. However, the possibility that the denatured whey proteins that attach on the surface of casein micelles actively contribute to dissociation of the κ-casein cannot be ruled out. It is also possible that differently glycosylated forms of κ-casein dissociate differently [50].

Vasbinder and De Kruif [156] have demonstrated that the coating of the casein micelles by heat-induced aggregates of denatured whey protein and micellar κ-casein become less and less homogenous as pH decreases from 6.90 to 6.35, with larger aggregates binding to less numerous κ-casein sites (Fig. 1). At a defined extent of denaturation of the whey proteins, these results suggest that larger heat-induced aggregates have a higher whey protein to κ-casein ratio, in agreement with the compositions of heat-induced aggregates found in the serum of heated control and casein-depleted milk [62].

4. DETRIMENTAL EFFECTS OF HEAT-TREATMENT ON THE CHEESE-MAKING PROPERTIES OF MILK AND THE POTENTIAL OF HEATING MILK AT ALKALINE pH VALUES

4.1. Rennet coagulation properties of heated milk

The heat-treatment of cheese milk for reasons of food safety or for the recovery of the whey proteins generally induces an increase in the rennet coagulation time (RCT); this is also true when a mixture of already denatured and aggregated whey protein and milk is heated [60]. The formation of heat-induced aggregates of denatured whey proteins and κ-casein on the surface of the casein micelles seems responsible for the inhibition of the renneting process in heated cheese milk [114, 120, 164]. Displacement of these heat-induced aggregates into the serum phase of milk upon heating milk at pH > 6.9 may therefore be an interesting means to reduce the detrimental effect of heating on the RCT of milk. The effect of such distribution of the heat-induced aggregates on the RCT may compare with that of the addition of “particulated” heat-denatured whey proteins to cheese milk on the RCT [48, 96] and would reduce the decrease in the curd firming rate and final strength of the gels observed with such addition as a consequence of the replacement of part of the casein, as the effective gel-forming protein fraction, by whey proteins [48, 149, 150]. Heat-treatment of milk at alkaline pH would also generate particles with sizes well below the 8 to 10 µm limit [76, 128] at which particles physically hinder formation of the rennet gel instead of improving its texture through being “inert fillers” in the network [74, 149].

Both the primary and secondary phases of the rennet coagulation process are affected by heat-treatment. In milk at its natural pH value, Van Hooydonk et al. [154] and Ferron-Baumy et al. [49] have shown that the initial velocity, Vi, of the primary phase of the action of rennet was 20 to 25% slower when the whey proteins were extensively denatured (heat-treatment exceeding 90 °C for 5 min), compared with unheated milk. Van Hooydonk et al. [154] also showed that the amount of hydrolysable κ-casein was reduced by 10% w/w after heat-treating skim milk at 120 °C for 5 min. Partial inhibition of the primary phase of the action of rennet on milk can be accounted for by the steric hindrance and the changes in electronegativity of the surroundings of the Phe105-Met106 bond of κ-casein, induced by the binding of heat-denatured whey proteins on the casein molecule [49].
The increased extent of heat-induced dissociation of κ-casein at alkaline pH, in combination with the increased formation of soluble heat-induced whey protein/κ-casein aggregates described earlier, may allow a better access of the enzyme to the Phe₁₀₅-Met₁₀₆ bond, despite the occurrence of denatured whey proteins, although this may depend on the structure of the aggregates. It is also a possibility that micelles depleted in κ-casein after heat-treating the milk at pH > 6.9 may be destabilised at a lower level of proteolysis than required for control micelles.

Van Hooydonk et al. [154] and Vasbinder et al. [158] also reported a large inhibitory effect of the heat-treatment of milk on the secondary phase of the rennet action, i.e., on the aggregation and gelation of the casein micelles. As a consequence of their coating by denatured whey proteins, the casein micelles of heated milk have a lower surface hydrophobicity [92], thus lowering the potential for interaction. Their spherical shape may also be retained longer than for untreated micelles, therefore preventing fusion, as suggested by the microstructure of rennet gels of heated and control milks observed by scanning electron microscopy [102, 120]. It is possible that milk heated at alkaline pH values, where less (hydrophilic) κ-casein and less heat-induced protein aggregates remain on the surface of the micelles, would show a reduced inhibition of the secondary phase of rennet coagulation.

An alternative explanation for the heat-induced increase in the rennet coagulation time of heated milk is the precipitation onto the casein micelles of the soluble calcium and phosphate ions during heating, to create a quasi-irreversibly insoluble tricalcium phosphate form, as suggested by Ustunol and Brown [152]. These authors reported an increase in the RCT of milk submitted to prolonged heating at 25 or 50 °C, i.e., at temperatures lower than those required to induce the denaturation of, e.g., β-lactoglobulin. This calcium-dependent inhibition of the renneting process is accounted for by the lack of screening effect of soluble calcium on the casein micelle, and therefore the increase in their net repulsive charge and related stability. Van Hooydonk et al. [154] further indicated that the relative insolubility of the heat-induced form of calcium phosphate would induce changes in the mineral equilibrium of milk, promoting the depletion of the casein micelle in colloidal calcium phosphate and therefore the formation of softer rennet casein gels. These changes in mineral equilibrium of heated milk could be at the origin of the so-called “rennet hysteresis” of heated milk, where the RCT of heat-treated milk is further increased by a prolonged storage in the cold [114, 154]. Heat-treatment [162] and alkalinisation of the milk [153] both decrease the proportion of soluble calcium (and phosphate) ions, thus increasing the detrimental effects of these changes on the formation of rennet gels, although one application of alkaline heat-treatment in order to reduce milkstone can be mentioned [46]. However, the effects of calcium and phosphate changes on the inhibition of the rennet action in heated milk are generally regarded as lower than those induced by the formation of heat-induced protein aggregates on the surface of the casein micelles [95, 154, 158], albeit this is still debated [133].

Controversial results have been reported in the literature on the effect of heat-treating milk at pH 7.5, rather than 6.7, on the rennet coagulation time. Absence of rennet coagulation of a milk sample heated at 90 °C for 10 min at pH 7.5, then neutralised to pH 6.7 for 24 h at 5 °C, was reported [154]. The heat-treatment of skim milk at 90 °C for 30 s, at pH values ranging from 6.0 to 9.0, induced little variations in the kinetics of the primary phase of the renneting reaction, as long as pH was below 7.3 [89]. At heat-treatment pH ≥ 7.3, Leaver et al. [89] reported a decrease in the rate of rennet action up to pH 7.8, followed by an increase at higher pH values.

On the other hand, the RCT of skim milk heated at 72 °C for 16 s and at pH 7.2, then acidified to pH 6.4, was only ~ 6 min, versus ~ 22 min for skim milk heated, and renneted, at pH 6.7 [71]. Rennet coagulation times < 6 min were obtained after pH-cycling milk heated at pH 7.5, to pH 5.5 at 4 °C, then to pH 6.2 or 6.4 prior to renneting [71]. Among various samples of heated
skim milk, those heated at pH 7.3, then pH-cycled, showed maximum rennet coagulation properties, although not as good as those of raw milk [142]. These studies tend to show a positive effect of heat-treating milk at pH > 6.7 on the RCT, albeit in most cases the effect of such alkaline heat-treatment could not be clearly distinguished from the effect of post-heating treatments, such as acidification or pH-cycling. When renneting was performed directly on the milk samples, without particular treatment following heating at 80 °C for 10 min at pH values ranging from 6.35 to 6.90, it was, however, found that the milk samples heated at pH 6.90 gelled earlier and gave stronger rennet gels than milk heated at pH 6.70, as a result of the larger proportion of aggregates to be found in the serum phase rather than on micellar κ-casein [156].

4.2. Quality of cheeses made with heated milk

The heat-treatment of milk at temperatures allowing the denaturation of the whey proteins significantly increases the cheese yield, through the increases in the protein yield and moisture content of the curd. In Cheddar cheeses, the studies reported a slight increase in fat yield, a +2.5 to +8% w/w increase in protein yield, and a +2.5 to +5% w/w increase in humidity after heat-treating the cheese milk at ~ 90 °C for up to 1 min [10, 72, 142]. In Havarti cheese, heat-treating the milk at 85 °C for 17 s led to an increase in protein yield (+2% w/w) and in humidity (+3% w/w) of the curd [93]. Heat-treating the milk at 91 °C for 16 s at an alkaline pH value of 7.5 also helped increase the protein yield of Cheddar cheeses, despite a slightly higher loss of fines during drainage, compared with control milk at either natural pH or pH 7.5 [72]. In Cheddar cheeses made with milk heated at 90 °C for 30 s, more whey proteins were denatured, and more were recovered in the rennet curd, whose total protein content increased as the pH of heat-treatment increased in the range 6.5 to 8.7 [12]. In the light of the work of, e.g., Vasbinder and De Knuij [156] or Renan et al. [128], these results suggested that despite the higher proportion of heat-induced whey protein/κ-casein aggregates formed in the serum phase of heated milk at alkaline pH values, these aggregates were retained in the cheese curd upon renneting, cutting and drainage. A slight tendency for increased moisture content could be noticed in cheeses made from milk heated at pH > 7.5, compared with the lower pH of heat-treatment [12].

However, the organoleptic qualities of the cheeses, especially texture and flavour, may be affected by the heat-treatment of the milk. The curds of Cheddar cheeses made with heated milk are generally crumbly, and can be difficult to process mechanically [10, 12, 102]. The firmness of fresh curds, obtained by renneting heated milk at pH 6.6, was reported to be inversely correlated to the rate of whey protein denaturation [60], although increased heating temperature from 80 to 130 °C prior to the manufacture of Mozzarella cheese (renneting pH ~ 5.6) resulted in greater firmness of the cheeses, possibly as a result of a bias with the solid-not-fat content [132]. The firmness of ripened Cheddar cheeses made with heated milk generally compared with cheeses made with pasteurised milk [12, 13]. Moderate high temperature-short time heat-treatment was also reported to smoothen the texture of white soft cheese made from UF-concentrated whole milk; longer heating times resulting in mealmess [137]. The increased retention of water in the cheese curd due to the high water-binding capacity of denatured whey proteins may be detrimental to drainage, curd fusion and firmness, and therefore to the manufacture of hard cheeses such as Cheddar cheese [129] and model semi-hard cheeses similar to Gouda, Provolone, Havarti or Halloumi [58]. Conversely, satisfactory textures were reported when manufacturing semi-hard cheese such as Cheshire using milk heated at 97 °C for 15 s, providing adjustments of the standard process [102]; or soft cheeses such as Camembert using milk heated at temperatures up to 80 °C for 1 to 3 min [9, 54] or UHT-treated milk [17]. A high water-binding capacity, a low proteolysis and strong covalent bonds as introduced by denatured, aggregated whey proteins during the heat-treatment of milk, are especially
beneficial to the yield, taste and texture of fresh acid cheeses such as quarg, or fromage frais [65, 74, 81, 82, 87, 95, 99, 101]. The functional properties of cheeses made with heated milk may also be adversely affected by the occurrence of whey proteins in the curd. Cheddar [10, 12] or Mozzarella cheese [64] made with cheese milk that included heat-denatured whey proteins showed impaired melting and stretching capacity, and reduced oiling-off as a result of an increased interaction between fat and protein [10], changes in the protein matrix of the curd, or the binding of water and calcium by the denatured whey proteins [74]. Similar effects have been reported in the presence of native whey proteins in UF-Mozzarella cheese [100], and may be explained by the denaturation of these proteins during cooking [112].

Heat-treatment of cheese milk may also induce sensible changes in the flavour profiles of cheeses. Reduction in free fatty acid production can be observed in milk lacking their endogenous microbial flora [58, 80], despite the fact that high heat-treated milk generally yields better acid production by lactic acid bacteria than raw milk, as reviewed by Feldstein and Westhoff [47]. Flavour changes also include modification of the free amino acid and peptidic profiles released during ripening of semi-hard cheeses [58] or of a range of soft to hard cheeses [83]. The release of peptides after 3 months of ripening, in Cheddar cheeses made with heated milk, was reported to benefit from adjustment of the milk to alkaline pH values prior to heating. As the pH of heat-treatment increased in the range 6.0 to 8.7, the peptide profile of the cheeses resembled more closely that of a control Cheddar cheese, made from simply pasteurised milk; as a result, Cheddar cheeses made from milk heated at 90 °C for 30 s at alkaline pH were more acceptable than when heated at lower pH values [12]. The development of the typical Cheddar flavour was also higher in cheeses made with milk heated at pH 7.1 and 7.5, compared with milk heated at pH 6.7 or even with control pasteurised milk [12]. The cheeses were more crumbly when prepared with heated milk, but this character was reduced at heat-treatment pH of 7.1 or 7.5. Poor melting properties were reported, whatever the pH of heat-treatment. It was also reported that the flavour defects of the Cheddar cheeses made from heated, pH-adjusted milk developed faster than their qualities, so that the same cheeses, aged 9 months, were less acceptable than at 3 months of age [12]. Heating milk at alkaline pH also generated less proteinaceous fouling of the heating vessel, and less cooked flavour of the cheese milk, than at lower pH values [72].

The key effects of the recovery of whey protein in cheese using heat-denatured whey proteins, on the rennet coagulation properties of cheese milk and on cheese quality, are summarised in Table II. The increase in moisture content of cheeses containing denatured whey proteins is omitted because its qualification as advantage or drawback depends on the variety of cheese produced. Table II stresses that only a small number of studies have investigated the effects of alkaline pH of heat-treatment in cheese technology.

Although the heating conditions of milk may be easy factors to control, it should be borne in mind that heat-treatment induces intricate and complex changes not only in the form and distribution of the denatured whey proteins, but also in the pH and buffering properties of the milk, respectively due to the transformation of lactose through the Maillard reaction and formation of formic acid [24], and to calcium phosphate precipitation [97]. These environmental changes are in turn very likely to influence the protein system, and hence the formation and properties of the whey protein/κ-casein heat-induced aggregates (Sect. 3, this review) and properties of the curd. Guinee et al. [58] have, for instance, reported that the higher the heat-treatment of cheese milk, the lower the pH measured in semi-hard cheeses from day 1 and throughout ripening. This could account for some heat-induced changes in the cheese texture, e.g. coarseness and mealiness, otherwise attributed to the interaction of denatured whey proteins with casein micelles. Heat-treatment of the milk prior to cheese-making, and/or the subsequent pH, structural mineral and protein changes of the milk and
Rennet coagulation of milk heated at alkaline pH 11 curd, may also influence growth of the micro-organisms [47, 58] and activity of the enzymes, both with potentially important consequences on texture and flavour of the cheeses. Depending on conditions, proteolysis of the αs- and β-caseins by plasmin may be reduced [14, 15, 146] or increased [16, 22, 77]. Various factors have been proposed to account for these changes, including the heat-denaturation of plasmin or plasminogen [14, 15, 146], or of an inhibitor thereof [16, 22, 77]. Native [63, 90, 112] or denatured whey proteins in cheese curds are thought to inhibit proteolysis of β- and/or αs-caseins by plasmin and/or chymosin [74, 93], e.g. through plasmin/β-lactoglobulin heat-aggregation [78]. Mistry and Kasperson [111] suggested that a higher moisture content of the cheeses, as induced by the incorporation of denatured whey protein, enhances the activity of plasmin and residual chymosin in the curd. The whey proteins themselves

Table II. Key advantages and drawbacks of the three approaches presented in this review that use heat-denatured whey proteins in order to increase cheese yield. RCT: rennet coagulation time, WP: whey proteins, ○: casein micelles, ●: whey protein, □: κ-casein and ▲: “particulated” whey proteins.

<table>
<thead>
<tr>
<th>Method applied to the milk</th>
<th>Representation of the dairy system</th>
<th>Advantages</th>
<th>Drawbacks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heat-treatment at natural pH</td>
<td>– Increased cheese yield [10, 72, 93, 142]</td>
<td>– Increased RCT and decreased curd fusion due to the coating of the casein micelles by WP [114, 120, 164]</td>
<td>– Crumbly, soft texture of the curd [10, 12, 60, 102]</td>
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<td></td>
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<td>– Poor melting properties [10, 12, 64]</td>
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<td></td>
<td></td>
<td>– Changes in proteolysis and flavour profile [14–16, 58, 74]</td>
</tr>
<tr>
<td>Addition of “particulated” whey protein</td>
<td>– Minimal increase of the RCT [48, 96]</td>
<td>– Decreased gel firming rate and final strength due to the decrease of the concentration of casein [48, 149, 150]</td>
<td>– Crumbly, soft texture of the curd [48, 96]</td>
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<td>– Poor melting properties [107]</td>
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<td>– Decreased gel firming rate and final strength due to the decrease of the concentration of casein [48, 149, 150]</td>
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<td>– Crumbly, soft texture of the curd [48, 96]</td>
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<td>– Poor melting properties [107]</td>
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<td></td>
<td>– Reduced flavoured and texture drawbacks of milk heated at natural pH [12]</td>
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</tbody>
</table>
did not appear to be hydrolysed during renneting or ripening of the cheeses, whether they were in native or denatured form [13, 74, 100, 131]. Promotion or inhibition of primary proteolysis will affect the development of bitterness in cheeses made from heated milk, but also curd fusion, ripening, and melting properties of the cheeses, based on yet debated mechanisms that do not always involve the heat-denatured whey proteins. UF concentration of cheese milk is eventually known to increase the buffering capacity of the cheese curd and to influence growth and lysis of the ripening microorganisms [55]. Its use in combination with heat-treatment to recover the whey proteins in cheese may generate effects on texture and flavour that are independent of properties of the denatured whey proteins. Care should therefore be taken as to interpret the direct effects of heat-treatment and pH changes on cheese quality.

4.3. Technological means used to restore the cheese-making properties of heated milk

Various approaches have been proposed in the literature to attempt to restore the cheese-making properties of heated milk. It is, for instance, a common practice to increase the level of ionic calcium, usually by addition of up to 0.3–0.6 g·L\(^{-1}\) calcium chloride, depending on the intensity of heat-treatment [10, 143]. Ionic calcium screens the charges on the casein micelles and reduces the interparticle repulsion, thus favouring flocculation; it also helps increase the gel strength by forming calcium bonds between phosphocasein molecules in the rennet gel. In the case of milk heated at alkaline pH, where significant precipitation of calcium phosphate is expected, it is essential to maintain an adequate concentration of soluble calcium.

Acidification of the milk prior to renneting is another means to increase the concentration of ionic calcium by dissociating both colloidal, and heat-induced, insoluble calcium phosphate complexes. As the pH approaches 5.5, i.e., the pH of highest activity of chymosin, the rates of coagulation and firming of the rennet gel increase [19]. Banks et al. [11] and Banks [10] thus lowered the renneting pH of milk heated at 90 °C for 30 s or 1 min from pH ~ 6.5 to 5.8 or 6.2 to make Cheddar cheese. However, acidification also increases association of the enzyme with the casein molecules [36], and therefore retention of the enzyme in the curd, where its strong activity leads to increased proteolysis [27, 51, 69, 159]. This effect is more significant in rennet-type curds such as Cheddar [69], where acidification in conventional processes essentially follows a fast renneting step and is usually limited, than in soft cheeses where acidification starts prior to rennet addition and largely proceeds throughout the renneting process, curd development and drainage [51]. In Cheddar cheese, acidification of the milk prior to renneting also slows down lactic acid production by the lactic acid bacteria, resulting in inhibited flavour development [11]. Cycling the pH of heated milk has proven a most efficient alternative to increase the concentration of ionic calcium by the use of pH adjustment of the milk, without leading to excessive chymosin retention and proteolysis [71, 73, 143]. In this process, the heated milk is acidified to pH 5.5 to 6.2, stored at 5 °C for 24 h and eventually neutralised at pH 6.5 to 6.7 prior to renneting [154]. Maximum rennet coagulation properties of heated milk were, for instance, reported through combining alkaline pH of heat-treatment, and pH-cycling of the milk prior to enzyme addition [11]. Long storage in the cold may be avoided by acidification to pH values below 5.5 [97] or by using injection and depressurisation of CO\(_2\) to adjust the pH [61]. Lucey et al. [98] reported that milk that had been pH-cycled to values lower than 5.5 and restored to 6.6, showed a decreased buffering property at pH ~ 5.1 and higher calcium activity, indicating an increase in soluble ionic calcium. Furthermore, a buffering peak appeared at pH 6.3 on back titration of pH-cycled milk, that the authors attributed to the precipitation of part of the dissociated calcium and phosphate ions into dicalcium phosphate dihydrate or octacalcium phosphate. This agreed with the suggestion by Van Hooydonk et al. [154] that the neutralisation
step of the pH cycle may induce the re-formation of calcium phosphate complexes that are different in nature to those naturally found in (heated) milk. Lucey [95] and Guillaume et al. [61] further hypothesised that the destructuration of the casein micelles upon pH-cycling make previously buried κ-casein molecules more accessible to hydrolysis by rennet.

Alternatively to exploiting the mineral balances of milk to restore the cheese-making properties of heated milk, other approaches make use of the protein fraction. Zoon [168] proposed only heating a fraction of a milk bulk or blending heated milk with raw milk so that unaffected κ-casein could take the lead in the renneting reaction kinetics. Vasbinder and De Kruif [156] have reported that milk heated at pH 6.35 exhibited a rennet coagulation behaviour comparable with that of unheated milk. This would result from the more heterogeneous distribution of the denatured whey proteins on the surface of casein micelles as the pH of heat-treatment decreases, which leaves more κ-casein free of denatured whey proteins, and therefore accessible to chymosin.

More specifically, Delespaul and Remars [37], Quiblier et al. [126], Schreiber and Hinrichs [134] and Nelson and Barbano [115] extracted a large proportion of the whey proteins from the milk by either chromatography or membrane technology, respectively, so that the depleted milk could be heated without having the extensive interaction of heat-denatured whey protein with the κ-casein molecules, i.e., without increase in the RCT and quality decrease in the rennet gel. The co-product of such separation can also be heated to denature the whey protein, concentrated using UF, and added back to the casein fraction to make cheese with a high protein recovery and little modification of the rennet coagulation properties [105]. Concentration of heated milk by ultrafiltration techniques is another useful approach to restore the cheese-making properties of heated milk [49, 60, 65, 104, 126, 138, 165, 166]. It is postulated that concentration reduces the distance between casein micelles, and hence increases the probability for interaction and bridge formation during renneting [59, 106, 164]. This practice is especially recommended to make fresh or soft cheeses where high whey protein recovery is beneficial because it leads to a relatively high moisture content and a thick texture.

4.4. Of the use of alkalis in cheese-making

The use of alkalis, like any other food additive, is strictly controlled by policies (at national and supra-national levels, such as in the European Union) generally inspired by the norms released by the Codex Alimentarius. Alkalis generally proposed in patents using pH manipulation of dairy products to make dairy ingredients and dairy products, such as sodium, calcium or magnesium salts of hydroxide, oxide, carbonate and other substances [18, 145] fall under the Codex Alimentarius general standard for food additives [44]. However, the general standard for cheese [41] only authorises glucono-δ-lactone, calcium carbonate and magnesium carbonate as acidity regulators, within the limits of the Good Manufacturing Practice (GMP). The addition of alkalis to milk in order to make cheese may therefore induce important restrictions of, e.g. designation of the product, unless otherwise enforced by national policies.

The addition of alkalis to milk, especially sodium salts, might also have important consequences on the nutritional value of the product, hence on its recommendation by the bodies in charge of food safety [1] and finally on its acceptance by today’s health-concerned consumers. Calcium and magnesium salts may meet nutritional recommendations better, although uses out of the standard regulation for cheese may fall under a category of enriched dairy products and should hence meet the specific standards for such products [143] for the Codex Alimentarius, sometimes through a time-consuming agreement procedure.

5. CONCLUSION

As outlined in the present review, the changes in the distribution of the heat-induced
whey protein/κ-casein aggregates, and in the dissociation rate of κ-casein, upon heating milk at alkaline pH, may be an interesting, yet insufficiently investigated factor to use to incorporate the whey protein into the cheese curd with minimum deleterious effects on the rennet coagulation properties of the milk. Combination of alkaline heat-treatment with other means such as UF concentration or calcium addition may improve some of the processes presently used to reach these goals, as demonstrated by the results on combined alkaline treatment and pH-cycling of milk [72, 142]. At present, the heat-treatment of milk at alkaline pH values has found little application in the restoration of the rennet coagulation properties of heated milk. In most cases, such a process was used to induce whey protein aggregates or protein-calcium complexes not otherwise present in the unheated milk, giving ingredients with modified viscosity, calcium content, or lactose content [18], to increase the yield of isolation of whey protein/casein complexes [25] or to increase the solubility of the isolates upon reconstitution [57]. In some other cases, alkaline heat-treatment has been used as a preliminary to the use of transglutaminase [145], sometimes in combination with rennet [28] to manufacture dairy ingredients. In yoghurt-making, heating at pH 7.1 improves the acid gelation properties of the milk, and final firmness of the gel [5, 94]. This review therefore demonstrates the need for further investigations to make use of the potential of alkaline heat-treatment of milk in order to increase retention of the whey proteins into cheese curds, as intended by Ménard et al. [108].

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