

Characterisation of the thermostable protease AprX in strains of *Pseudomonas fluorescens* and impact on the shelf-life of dairy products: preliminary results

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Abstract

Bacterial proteases are involved in food spoilage and shelf-life reduction. Among the bacterial proteases, a predominant role in spoilage of dairy products seems to be played by the thermostable metallo-protease AprX, which is produced by various strains of Pseudomonas fluorescens. Differences in AprX enzyme activity among different strains were highlighted, but the most proteolytic strains were not identified. In this study, the presence of the aprX gene was evaluated in 69 strains isolated from food matrices and 18 reference strains belonging to the P. fluorescens group, which had been previously typed by the multi locus sequence typing method. Subsequently, a subset of reference strains was inoculated in ultra-high temperature milk, and the expression of the aprX gene was evaluated at 22 and 6°C. On the same milk samples, the proteolytic activity was then evaluated through Azocasein and trinitrobenzenesulfonic acid solution assays. Finally, to assess the applicability of the former assay directly on dairy products the proteolityc activity was tested on industrial ricotta samples using the Azocasein assay. These results demonstrate the spread of aprX gene in most strains tested and the applicability of Azocasein assay to monitor the proteolytic activity in dairy products.

Introduction

Thermostable protease activity was indicated as one of the main factors responsible of dairy products spoilage, such as gelation of ultra-high temperature (UHT) milk, with consequences on the shelf-life and significant economic losses for the food industry. Proteases are predominantly active against the casein fraction, which causes gelation of UHT milk and/or the formation of bitter off-flavors. The gelation of UHT milk derives from the formation of complexes between the k-casein and β-lactoglobulin that are denatured as a result of the heat treatment. Gelation is triggered by enzymatic processes of protein degradation (Rauh et al., 2014; Datta and Deeth, 2001). The proteolysis during storage of UHT milk seems to be a consequence of the interaction between the plasminogen-plasmin system (endogenous protease of the milk) with proteases of bacterial origin. Among the bacterial proteases involved in spoilage processes a predominant role seems to be played by the thermostable metallo-protease AprX, which is produced by various strains of the species Pseudomonas fluorescens. The AprX protease, produced and released by bacteria in milk, is resistant to heat and is able to maintain unaltered its activity even after heat treatments that milk may undergo during its processing such as pasteurization, UHT treatment, and cheese-making (Ismail and Nielsen, 2010; Frohbieter et al., 2005). Such enzyme activities could also have a role during cheese ripening for the formation of flavors in seasoned cheeses, and aromatic characteristics of milk cultures of the P. fluorescens group are strain-dependent (Morales et al., 2005; Carraro et al., 2011).

To evaluate proteolytic activity, several assays were developed and applied in *P. aerugi*nosa (Kessler and Safrin, 2014) and some of them tested for AprX activity on P. fluorescens strains (Dufour et al., 2008; Marchand et al., 2009a). The AprX protein and its encoding gene have been studied extensively (Liu et al., 2007; Maunsell et al., 2006; Nicodeme et al., 2005; Woods et al., 2001; Liao and McCallus, 1998). Differences in AprX enzyme activities were highlighted, but the identification of the most proteolytic strains was not completely elucidated (Dufour et al., 2008; Marchand et al., 2009a, 2009b). This is due partially to the unreliability of methods currently used for the identification of P. fluorescens strains. Recently, P. fluorescens was recognized as a species group (Mulet et al., 2010) and a molecular typing approach was developed and applied on several P. fluorescens reference and field strains (Andreani et al., 2014). The determination of the genetic diversity among species and strains belonging to the P. fluorescens group provides an accurate method for strain identification.

In the present study the presence of the *aprX* gene was evaluated in 69 strains isolated from food matrices and 18 reference strains belonging to the *P. fluorescens* group. All strains were previously typed by multi locus sequence typing (MLST) method and allocated in a *subgroup* as genetically related to a specific reference type strain.

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In a selected group of reference strains, the activity and gene expression of AprX were tested in milk. Finally, the applicability of Azocasein assay directly on dairy products was evaluated on industrial *ricotta* samples. The results demonstrated the spread of *aprX* gene in most of the strains tested and the applicability of trinitrobenzenesulfonic acid solution (TNBS) and Azocasein tests to monitor the proteolytic activity in dairy products.

Materials and Methods

Bacterial strains

P. fluorescens group strains are listed in Table 1. For each strain, the sequence type (ST) and the *subgroup* obtained by MLST analysis (Andreani *et al.*, 2014) are reported. Strains were conserved at -80°C in Tryptic Soy Broth [TSB; Oxoid, Basingstoke, UK; with 50% v/v glycerol (Sigma-Aldrich, Saint Louis, MO, USA)].

DNA and RNA extraction

For DNA extraction, a single colony from a fresh culture on CFC *Pseudomonas* Agar Base (CFC PAB; Oxoid) was re-suspended in 100 μ L of nuclease-free water, vortexed at high speed for 5 seconds and incubated at 95°C for 10 minutes. The tube was vortexed again and centrifuged for 2 minutes at 14,000 rpm. The supernatant was transferred to a fresh tube and stored at -20°C (Martino *et al.*, 2011).

For RNA extraction, a single pure colony of each strain was inoculated in triplicate (giving 3 biological replicates for each strain) in 3 mL MBM Broth $[0.7\% K_2 HPO_4, 0.3\% KH_2 PO_4, 0.05\%$ trisodium citrate, $0.01\% MgSO_4$, $0.1\% (NH_4)_2 SO_4$, 0.2% glucose] and kept at 22°C for



Table 1. Species, subgroups, food origin, sequence type, presence of AprX gene and proteolytic activity in plate of Pseudomo	mas fluo-
rescens strains.	

Strain	Species	Subgroup	Source	ST	AprX gene	Prot 6°C	teolitic ac 22°C	tivity 31°C
DSM 17152T	Pseudomonas gessardii	P. fluorescens subgroup	Mineral water	7	+	+	++	-
DSM 15294T	Pseudomonas brenneri	P. fluorescens subgroup	Natural mineral water	8	+	+	++	-
DSM 17967T	Pseudomonas mandelii	P. mandelii subgroup	Mineral water	9	+	+	+	-
DSM 17150T	Pseudomonas jessenii	P. jessenii subgroup	Mineral water	10	+	-	+	-
DSM 16610T	Pseudomonas koreensis	P. koreensis subgroup	Agricultural soil	11	+	+++	+	++
DSM 17489T	Pseudomonas orientalis	P. fluorescens subgroup	Spring water	12	+	+++	++	-
DSM 18928T	Pseudomonas synxantha	P. fluorescens subgroup	Cream	13	+	++	+	-
DSM 18862T	Pseudomonas azotoformans	P. fluorescens subgroup	Paddies	14	+	+++	+++	-
DSM 6252T	Pseudomonas lundensis	P. fragi subgroup	Prepacked beef	15	+	-	-	-
DSM 14020T	Pseudomonas rhodesiae	P. fluorescens subgroup	Natural mineral water	16	+	++	+++	-
DSM 113201	Pseudomonas veronii	P. fluorescens subgroup	Mineral water	17	+	++	++	-
DSM 17149T	Pseudomonas libanensis	P. fluorescens subgroup	Spring water	18	+	+++	++	
DSM50415	Pseudomonas fluorescens	P. fluorescens subgroup	Soil	19	+	++	++	+
CECT124T	Pseudomonas corrugata	P. corrugata subgroup	Tomato: pith necrosis	20	+	тт	++	++
CECT1241 CECT229T	Pseudomonas marginalis	P. fluorescens subgroup	<i>Cichorium intybus</i> (endive)	20	+ +	-	+++	++
CECT225T CECT378T	Pseudomonas fluorescens	P. fluorescens subgroup	Pre-filter tanks, town water works	22				-
CECT3781 CECT446T	Pseudomonas fragi		Unknown	22	+	++	++	-
	Ŭ	P. fragi subgroup			+	+++	+++	-
CECT4470T	Pseudomonas chlororaphis	P. clororaphis group	Plate contaminant	24	+	++	++	++
ps_1		P. fluorescens subgroup	Mozzarella cheese	25	+	+	++	-
ps_2		P. fluorescens subgroup	Blue mozzarella cheese	26	+	++	++	-
ps_3		P. fragi subgroup	Mozzarella cheese	27	+	-	-	-
os_4		P. fluorescens subgroup	Mozzarella cheese	28	+	-	+++	-
ps_5		P. fluorescens subgroup	Mozzarella cheese	29	+	-	++	-
ps_6		P. fluorescens subgroup	Blue mozzarella cheese	30	+	++	++	
ps_7		P. koreensis subgroup	Mozzarella cheese	31	+	+++	+++	-
ps_8		P. fluorescens subgroup	Mozzarella cheese	32	+	+++	+++	-
ps_9		P. fluorescens subgroup	Mozzarella cheese	33	-	++	++	-
ps_10		P. fluorescens subgroup	Mozzarella cheese	34	+	++	++	-
ps_11		P. fluorescens subgroup	Mixed salad	35	+	+	++	-
ps_12		P. fragi subgroup	Butter	36	+	-	-	-
ps_13		P. fluorescens subgroup	Blue mozzarella cheese	37	+	++	++	-
ps_14		P. koreensis subgroup	Pork	38	+	-	+	+
ps_15		P. fluorescens subgroup	Salmo trutta marmoratus (trout)	39	+	++	++	-
ps_16		P. fluorescens subgroup	Salmo trutta fario (trout)	40	+	+++	+++	++
ps_17		P. fragi subgroup	Ricotta	41	+	++	+	-
os_18		P. fluorescens subgroup	UHT milk	42	+	++	++	++
os_19		P. fragi subgroup	UHT milk	43	-	-	+	++
os_20		P. fluorescens subgroup	UHT milk	44	+	+++	++	+
os_21		P. koreensis subgroup	Mozzarella cheese	45	+	+	+	-
ps_22		P. fluorescens subgroup	Blue mozzarella cheese	46	+	+	++	-
ps_23		P. koreensis subgroup	Mozzarella cheese	47	+	-	++	-
ps_24		P. fluorescens subgroup	Cheese	48	+	-	++	-
os_25		P. fluorescens subgroup	UHT milk	49	+	-	+	-
os_26		P. koreensis subgroup	Human	50	+	++	++	++
ps_20 ps_27		P. fluorescens subgroup	Mixed salad	51	+	++	++	-
os_28		P. koreensis subgroup	Mixed salad	52	+	++	+++	+++
ps_20 ps_29		P. fluorescens subgroup	Mixed salad	53	+	+++	+++	
ps_23 ps_30		P. fluorescens subgroup	Mixed salad	54	+	+++	+++	
ps_30 ps_31		P. fluorescens subgroup	Mixed salad	55	+ +	++	+++	_
		P. clororaphis group	Mixed salad	55 56		TT	+++	-
ps_32		1 0 1			+	-	-	-
ps_33		P. fluorescens subgroup	Mixed salad	57 E 0	+	++	++	-
ps_34		P. fluorescens subgroup	Mixed salad	58	+	+++	+++	-

Continued on next page.



24 h to reach 108 CFU/mL. Then, cultures were diluted five times in 3 mL of milk. After 24h of incubation at 22°C or five days at 5°C, 1 mL of culture was then extracted using the RNAeasy Mini Kit (Qiagen, Hilden, Germany) following the manufacturer's protocol. DNase treatment was performed using the Qiagen RNase-Free DNase Set (Oiagen). RNA was eluted in 30 µL of RNAse-free H₂O. Purified DNA and RNA were quantified using a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA). One microgram of total RNA for each sample was reverse transcribed to cDNA using SuperScript® II Reverse Transcriptase (Invitrogen, Carlsbad, CA, USA). To evaluate DNA contamination in RNA samples treated with DNAse, no-RT (reverse-transcriptase) samples were also checked in PCR amplification.

Polymerase chain reaction amplification

Primers for *aprX* gene amplification were designed using PriFi software (http://cgiwww.daimi.au.dk/cgi-chili/PriFi; Fredslund *et al.*, 2005) using as template the alignment of *aprX* sequences from complete genomes of *P. fluorescens* group already available in Genbank (http://www.ncbi.nlm.nih.gov/genome). The sequence of primers was Pse_AprXF1 CAGACCCTGACCCACGARATCGG and Pse_AprXR1 TGAGGTTGATCTTCTGGTTCTGGG. The *rpoD* housekeeping gene was used as positive control for DNA and RNA extraction using primers reported in Andreani *et al.* (2014).

PCR amplifications were performed in an Applied Biosystems 2720 Thermal Cycler

(Thermo Fisher Scientific, Waltham, MA, USA) in a final volume of 20 µL of amplification mix containing 1U of GoTaq polymerase (Promega, Madison, WI, USA), 1X GoTaq Buffer, 1.5 mM MgCl₂, 0.2 mM each deoxynucleotide triphosphate (dNTP), 250 mM each primer and 5 ng of genomic DNA as template. The reaction mixture was subjected to the following thermal cycle: an initial step at 94°C for 2 min to activate the polymerase and 35 cvcles each of denaturation at 94°C for 20 seconds, annealing of the primers at 60°C for 30 seconds and extension at 72°C for 1 minute and a final step of extension at 72°C for 7 min. Amplified products were analysed by electrophoresis on 1.8% agarose-Tris-acetate-EDTA (TAE) gels, stained with SYBR[®] Safe DNA Gel Stain (Invitrogen) and visualized on a UV transilluminator (Gel Doc XR[™]; Biorad, Hercules, CA, USA).

Table 1. Continued from previous page.

Strain	Species	Subgroup	Source	ST	<i>AprX</i> gene	Proteolitic 6°C 22°C			
ps_35		P. fluorescens subgroup	Mixed salad	59	+	+++	++	-	
ps_36		P. fluorescens subgroup	Mixed salad	60	+	+++	++	-	
ps_37		P. clororaphis subgroup	Mixed salad	61	+	+	++	+++	
ps_38		P. clororaphis subgroup	Mixed salad	62	+	+	++	+++	
ps_39		P. fluorescens subgroup	Mixed salad	63	+	++	++	+	
ps_40		P. fluorescens subgroup	Ricotta	64	+	+++	++	-	
 ps_48		P. fragi subgroup	Pork	72	+	++	+++	+	
ps_50		P. fragi subgroup	Pork	74	+	++	+++	+++	
ps_51		P. clororaphis group	Pork	75	+	++	+++	+++	
ps_54		P. fragi subgroup	Pork	78	+	++	++	+	
ps_55		P. koreensis subgroup	Pork	79	+	-	++	+++	
ps_56		P. koreensis subgroup	Sashimi	80	+	+	++	+++	
ps_57		P. fluorescens subgroup	Sashimi	81	+	++	+++	++	
ps_58		P. corrugata subgroup	Sashimi	82	+	-	++	++	
ps_59		P. fluorescens subgroup	Sashimi	83	+	+++	+++	++	
ps_60		nd	Sashimi	nd	+	+++	+++	+++	
ps_61		P. fluorescens subgroup	Sashimi	84	+	+	+++	++	
ps_62		nd	Sashimi	nd	+	+	+++	+++	
ps_63		nd	Sashimi	nd	+	++	+++	++	
ps_64		P. fluorescens subgroup	Sashimi	85	+	-	+++	+	
ps_65		P. koreensis subgroup	Rocket	86	+	++	+++	+++	
ps_66		P. koreensis subgroup	Rocket	87	+	++	+++	+++	
ps_67		P. koreensis subgroup	Valerian	88	+	++	+++	+++	
ps_68		P. corrugata subgroup	Valerian	89	+	-	+++	+++	
ps_69		P. mandelii subgroup	Dairy-product	90	+	-	+	+	
ps_70		P. fragi subgroup	Dairy-product	91	+	+	++	-	
ps_71		P. fragi subgroup	UHT milk	92	-	+	+	+	
ps_72		P. fragi subgroup	Dairy-product	93	+	-	++	+	
ps_73		P. fluorescens subgroup	Dairy-product	94	-	-	+	-	
ps_74		P. fragi subgroup	Dairy-product	95	+	++	+	+	
ps_75		P. fluorescens subgroup	Blue mozzarella cheese	29	+	-	++	-	
ps_76		P. fluorescens subgroup	Mozzarella cheese	26	+	++	++	-	
ps_77		P. fluorescens subgroup	Meat	96	+	-	++	-	
		J							
ps_78		P. fluorescens subgroup	Mozzarella cheese	97	+	-	+	-	

ST, sequence type; UHT, ultra-high temperature; nd, not determined.

Proteolytic activity in plate

The reference and field strains were tested for their proteolytic activity by agar diffusion assays at 6°C for 10 days, at 22 and 31°C for 7 days. All the strains were revitalized through a 72-hours-preinoculum in TSB at 22°C. A dilution of 10⁵ CFU/mL was applied for the subsequent tests. The extracellular protease activity evaluation was conducted on Nutrient Agar (NA; Biokar diagnostics, Paris, France) with 2% UHT milk observing a clear zone around the colonies. The presence of a clear zone around the colonies after incubation was indicative for proteolysis.

Quantification of extracellular proteolytic activity (Azocasein assay)

Proteolytic activity of bacterial strains was quantified using Azocasein (Sigma-Aldrich) as substrate. One hundred µL of a 3% (w/v) Azocasein stock solution were added to 100 µL of cell free supernatant fluid and 300 uL of 50 mM Na₂HPO₄ pH 7.5. The negative control was set up with 100 µL of not inoculated milk. The mixture was incubated at 37°C for 1 h and the reaction was stopped by adding 500 µL of 20% (w/v) trichloroacetic acid (TCA). The sample was centrifuged at 12,000 g for 10 min and absorbance of the supernatant was measured at 366 nm using Multiskan GO UV/Vis spectrophotometer (Thermo Fisher Scientific). Absorbance of the blank (500 µL not incubated sample plus 500 µL of TCA 20%), and of the negative control were subtracted from sample absorbance. The results are reported as OD₃₆₆.

Quantification of proteolysis (trinitrobenzenesulfonic acid solution test)

The experimental set-up made it possible to calculate the net proteolytic activity produced during 2 weeks of storage at 37°C after a heat treatment simulating UHT process and storage (as described in Marchand *et al.*, 2008, 2009a). Hydrolysis of proteins was measured by the determination of the release of a-amino

groups by the trinitrobenzenesulfonic acid (TNBS) method. The free amino groups react with the TNBS reagent (Sigma-Aldrich) at pH 9.2 in the dark. A yellow-orange colour develops and its intensity is determined by absorption measurements at 420 nm. The degree of proteolysis is calculated from the increase in absorption after 2 weeks of storage at 37°C and expressed as mmol of glycine equivalents mL-1 milk, using glycine (2.5, 2.25, 2, 1.75, 1.5, 1.25, 1, 0.75, 0.5, 0.25 e 0 mM; Sigma-Aldrich) to create a standard curve. The experiment was repeated twice, first in macro method (experiment A using a 7800 UV/VIS spectrophotometer; JASCO, Easton, MD, USA) and second time in micro method (experiment B using a Multiskan GO UV/Vis spectrophotometer (Thermo Fisher Scientific).

Proteolytic activity in industrial *ricotta* samples

Of ricotta samples, 10 g were inoculated with 1 mL of a 10⁸ cell/mL of a fresh culture of P. fluorescens group strains growth in TSB in sterile 50 mL tubes and were maintained at refrigeration temperature for seven days. The Azocasein assay was carried out in order to evaluate the proteolytic activity of the inoculated ricotta in comparison to three industrial ricotta samples, belonging to the same lot, collected and analyzed 24 h from production. One gr of ricotta was sampled in duplicate from each of the four inoculated samples and the three fresh ricotta samples and diluted in 5 mL of Phosphate-buffered saline (PBS). After mixing, 2 mL were centrifuged at 12,000 rpm for 5 minutes, then 100 µL of the supernatant was used for the Azocasein assay as previously described.

Results

Distribution of the aprX gene in



Pseudomonas fluorescens group

The distribution of the *aprX* gene was evaluated on 18 reference and 69 field strains and the result is reported in Table 1. Only for four strains (4.6%) the PCR amplification of *aprX* gene, repeated twice, gave a negative result. All these four strains resulted positive to the amplification of the *rpoD* gene. The four strains belonged to the *P. fluorescens* subgroup (ps_9 and ps_73) or to the *P. fragi* subgroup (ps_19 and ps_71).

Proteolytic activity of

Pseudomonas fluorescens strains in plate

Proteolytic activity was observed at least at one temperature condition in 17 reference strains and 65 fields strains (for a total of 94.2%), indicating that proteolysis is a common spoilage mechanism for *P. fluorescens* group strains. The non-proteolytic strains are *P. lundensis* DSM6252T, ps_3, ps_12, ps_32, ps_79. The complete data are reported in Table 1.

Proteolytic activity and *aprX* gene expression in milk samples inoculated with *Pseudomonas fluo-rescens*

The experiment was set up to simulate the conditions during UHT milk production (as described in Marchand et al., 2008, 2009a) and a schematic representation of the analyses is reported in Figure 1. Briefly, 100 µL (10⁷ cells) of a fresh culture of each strain was inoculated in 10 mL of UHT milk and grown for 24 hours. An aliquot of 100 µL of the culture was then inoculated in 10 mL of UHT milk and grown for 24 hours. Double growth in milk until exponential phase was done to adapt bacterial strains to milk. The culture was then diluted to 103 cfu/mL in UHT milk and incubated at 6°C for five days. At the end of the incubation, the Azocasein test was performed as described in Materials and Methods to measure the global proteolytic activity (thermoresistent and non-

Table 2. Proteolityc activity and AprX gene expression in Pseudomonas fluorescens group references strains.

Tuble 21 Troteonty and There expression in Tromosmonic junorescent group reserves strains											
Type strains	Subgroups	Proteolitic activity in plate*		<i>Apr</i> X gene expression°		Azocasein OD ₃₆₆		TNBS [#] mM glicine		Azocasein [§] <i>Ricotta</i> OD ₃₆	
		6°C	22°C	31°C	22°C	6°C	Pre-HT	Post-HT	Experiment A	Experiment B	
DSM 17489T	P. orientalis	+++	++	-	+++	+-+	0.28	0.22	3.69	11.85	1.02
DSM 6252T	P. lundensis	-	-	-			0.02	0.01	1.92	6.72	nd
DSM 14020T	P. rhodesiae	++	+++	-	+++		0.07	0,04	1.71	24.09	0.34
DSM 17149T	P. libanensis	+++	++	-	+++		0.37	0.17	-	-	1.38
CECT378T	P. fluorescens	++	++	-	++-		0.1	0.03	3.15	7.24	1.01
CECT446T	P. fragi	+++	+++	-	+++		0.28	0.18	-	-	nd
CECT229T	P. marginalis	-	+++	-			0.01	0.02	-	-	nd

TNBS, trinitrobenzenesulfonic acid solution; HT, high-temperature. *+, ++ or +++ depending on the size of the clear zone around the colonies; °+ or – is positive or negative result in the amplification of each triplicate; texperiment A was performed in macromethods, Experiment B in micromethods; seported the average between the OD of the duplicates. nd, not determined.



This preliminary experiment was performed with six reference strains, including a negative control (not inoculated milk) and the results are summarized in Table 2. The gene expression study demonstrated that, in the condition tested, the *aprX* gene is not expressed in *P. lundensis* and *P. marginalis*. The Azocasein assay gave OD₃₆₆ values ranging from 0.01 to 0.37. The TNBS assays gave values in mM Glycin from 0 (negative results for three strains) to 24.09 in micro method and from 0 to 3.69 in macro method. Proteolytic activity data is only partially in agreement with expression data.

Proteolytic activity in industrial *ricotta* samples

The results of proteolytic activity measured on *ricotta* samples are reported in Table 2 and Figure 2. The data evidenced variable activity in the freshly produced *ricotta* samples despite the three samples belonged to the same lot of production.

Discussion

The analysis of 87 *P. fluorescens* strains for the presence of the *aprX* gene demonstrated that the gene is widespread in this bacterial group. The *aprX* negative strains are not strongly genetically related (Andreani *et al.*, 2014) suggesting that the lack of *aprX* amplification, could be due to loss of the gene or mismatches in primer sites that occurred independently in each strain. However, the *aprX* gene expression study demonstrated strong variability across strains, which might explain the large variability in proteolytic activity reported in previous studies (Dufour *et al.*, 2008; Marchand *et al.*, 2009a, 2009b).

To evaluate the spoilage activity of AprX in dairy products, milk and *ricotta* cheese were used as template to inoculate strains positive to *aprX* gene. The experimental protocol for milk, reported in Figure 1, was designed to simulate UHT milk productive process, with milk samples incubated first at refrigeration temperature, and after heat treatment, incubated at 37°C. Similar experimental design to simulate UHT milk production was proposed and applied in previous studies (Marchand *et al.*, 2008, 2009a).

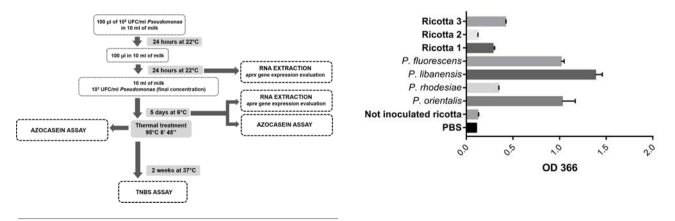
The proteolytic activity was measured with two different assays, the first, the Azocasein assay, measures directly the activity with a colorimetric reaction, the second, the TNBS assay, measures free amino acids (aminogroups), as products of the proteolysis. The two different assays are both easy and fast to be carried out, inexpensive and require a common spectrophotometer. In the present study, these assays were applied to compare their sensibility on dairy products inoculated with a bacterial culture. However, the results of the two assays are only partially comparable. The Azocasein assay results are in good agreement with *aprX* expression data, if considering value less to 0.05 OD₃₆₆ as negative. On the contrary, TNBS assay results are discordant with aprX expression and Azocasein assay data for P. lundensis and P. libanensis. On the basis of these results the Azocasein assay might be more reliable as a direct measure of the enzyme activity, however some concerns remain. First, the Azocasein assay had not a standard curve to be used to compare the results. Secondly, protease activity immediately after the heat

treatment might be slower as a consequence of the treatment itself and this effect might be different in the different strains. This effect could be due to differences in protein structure despite the nucleotidic sequence was reported to be very conserved among *P. fluorescens* strains (Marchand *et al.*, 2009b) or to different resistance of strains to heat treatment. The gelation of UHT milk, if occurs, usually takes place after weeks from production. This long time might depend on the time required by proteases to reactivate after heat treatment or to some chemical modification that might occur in milk and activate proteases.

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Anyway, this preliminary study strongly confirms strain variability in protease activity. P. lundensis and P. fragi were indicated as the species mostly involved in spoilage of milk (Marchand et al., 2009a). For these two strains, contrasting results were obtained in the present study. Regarding P. lundensis type strain, the *aprX* expression was negative in milk as it was proteolytic activity in plate and at Azocasein assay, even if the TNBS assay yielded a positive result. On the contrary, P. fragi type strain expressed aprX and was positive for proteolytic assay both in plate and with Azocasein assay, but was negative with TNBS assay. Thanks to MLST molecular typing, the P. fluorescens group fields strains were accurately identified and the application of the experiment in milk using fields strains belonging to P. fragi subgroup (that include P. lundensis) is interesting to confirm if the spoilage phenotype is related to this taxonomic group.

Finally, with the aim to test the applicability of these assays to analyze dairy products, in which spoilage activity might be due to thermoresistent proteases, industrial *ricotta* was chosen as a study case. In fact, industrial *ricotta* is produced at high temperature (90°C) and successively pasteurized (80°C) starting from milk whey that usually presents high bacterial loads (among these bacteria, *Pseudomonas* is



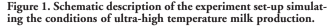


Figure 2. Proteolytic activity in *ricotta* samples.

often present). Industrial *ricotta* often shows premature spoilage that reduces product shelflife. The application of Azocasein assay on freshly produced industrial *ricotta* samples showed protease activity that might be one of the responsible factors for premature spoilage. This result highlights the importance of good quality raw material to extend shelf-life also for heat-treated products.

Conclusions

These preliminary data highlight the interest, but also the complexity of studying the proteolytic activity of the *aprX* gene in *P. fluorescens*. The availability of MLST-typed strains might help to identify the major *aprX* producing strains and evaluate if such spoilage activity is a phenotypic trait linked to specific lineages in the *P. fluorescens* group. Azocasein and TNBS assay should be improved, but might be suitable (in particular the Azocasein assay) to evaluate proteolytic activity in dairy products such as industrial *ricotta* or UHT milk to assess in advance the spoilage potential during shelf-life.

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