Evaluation of the bacteriological quality of raw cow’s milk at various stages of the milk production chain on farms in Algeria

M. Hamiroune (1, *) , A. Berber (3) & S. Boubekeur (3)

(1) Department of Agronomic and Veterinary Sciences, Faculty of Natural and Life Sciences, Ziane Achour University, B.P. 3117, Moudjbara road, Djelfa, Algeria
(2) High National Veterinary School, Issad Abbes road, Oued Smar, Algiers, Algeria
(3) Institute of Veterinary Sciences, Blida 1 University, B.P. 270, Soumâa road, Blida, Algeria

* Corresponding author: mouradhamiroune@gmail.com

Summary

This study evaluates hygiene practices on 53 dairy farms in the Jijel and Blida regions of Algeria. A survey questionnaire was drawn up covering milking conditions and cleaning of the equipment. In parallel, bacteriological analyses were carried out to estimate the rate, source and development of bacterial contamination in raw milk produced on the farm. In addition, screening was performed to detect the presence of inhibitor residues.

The results of the survey revealed poor livestock conditions and milking practices that could explain the presence of bacteria in cow’s milk.

The bacteriological results showed that 76.1% of milk samples taken from cow udders complied with legal standards, compared with only 35.8% of milk samples taken from storage tanks. Moreover, bacterial inhibitors were detected in 28.8% of milk samples. These results showed that the hands of milkers, udders, teat cups, utensils, the water used during milking and the milking environment were all potential sources of milk contamination by the bacteria under investigation.

These results suggest that, to improve the bacteriological quality of milk, there is a need to introduce a quality policy which places a premium on milk of high bacteriological quality and aims to generalise good hygiene practices throughout the dairy production chain.

Keywords

Algeria – Bacterial contamination – Bacteriological quality – Cows – Hygiene practices – Milking – Raw milk.

Introduction

Milk figures prominently in the diet of Algerians, which explains why the dairy sector has seen annual growth of 8% (1). Algeria is the leading consumer of raw milk in the Maghreb region, with almost 3 billion litres a year. The hygiene quality of raw milk is therefore vital (2).

To maintain hygiene conditions on farms and up to the arrival of milk in dairies, the bacteriological quality of the milk must be monitored (3).

There are several risk factors for milk contamination at different stages of production on the farm, prompting the authors to conduct this study with the principal aim of evaluating the bacteriological quality of raw milk in the Jijel and Blida regions and identifying raw milk contamination risk factors on the farm.

Materials and methods

Farm selection

This study was conducted on 53 dairy farms covering 360 milking cows in the Jijel and Blida regions in Algeria, during the period from March 2013 to July 2014.

A non-random convenience sampling plan was defined to include herds that differed in terms of size, milking method
and equipment or in preparation of the udders for milking (washing and disinfection).

Epidemiological survey

On each dairy farm, a survey was carried out and the milking process was monitored on the sampling day. The survey questionnaire form indicates the cows sampled, milk production systems and milking hygiene.

Sampling

In order to assess the bacteriological quality and sources of contamination of milk produced on the farm, raw milk samples were taken, as well as samples from the environment and the milking equipment.

Before milking, a 100 ml sample of the water used for milking was taken, as well as 100 ml of the water used for rinsing the milking utensils. Swab samples were taken from the milkers’ hands, teat cups and skin of the udders of each of the cows.

During milking, a flask containing sterile water was exposed for 30 minutes to assess environmental contamination (environmental sample).

The samples were taken aseptically and placed in labelled sterile vials. The authors used the individual milk sampling technique described by Mialot (4).

The samples were then placed in a cooler and transported directly to the testing laboratories (the laboratory of the Algerian centre for quality control and packaging [CACQE] and the laboratory for veterinary testing, quality control, compliance and applied research [AVCQ-LAB] in Baraki), where they were refrigerated at +4°C. The time between sampling and the first analyses barely exceeded 24 hours.

Table I shows the number of samples taken by sampling site.

Detection and enumeration of contaminating microorganisms

Different dilutions with a tryptone salt broth (TSB) were used depending on the nature of the sample; they varied between $10^{-1}$ and $10^{-8}$.

In each sample, a search was made for five groups of bacteria: total aerobic mesophilic flora, faecal streptococci, faecal coliforms, Staphylococcus aureus and Clostridium sulphite-reducers (CSR) (5).

The detection and enumeration of total aerobic mesophilic flora (TAMF) was carried out on glucose agar with yeast extract (plate count agar [PCA]) after incubation at 30°C for 72 hours (6).

Faecal coliforms (FC) were detected and enumerated on violet red bile lactose agar (VRBL), incubated for 24 hours at 44°C. All red colonies (lactose+) that appeared with a minimum diameter of 0.5 mm were considered to be faecal coliforms (7).

Staphylococcus aureus (SA) was detected and enumerated on Baird Parker agar with egg yolk and tellurite incubated at 37°C for 24–48 hours. Black, shiny convex colonies appeared surrounded by a clear halo of 2–5 mm in diameter. This was confirmed by Gram stain test (+), catalase test (+) and coagulase test (+) (8).

Faecal streptococci (FS) were enumerated on Rothe broth (Pasteur Institute, Algeria). A millilitre of each sample to be analysed was added to 9 ml of TSB. In this way, the authors obtained a mother dilution of $10^{-1}$ from which the decimal dilutions were made. A millilitre of each dilution was then placed in three tubes of Rothe broth. Following incubation for 48 hours at 37°C, the contents of the positive tubes (those that were cloudy) were then seeded on bile agar and bile esculin azide (BEA) for confirmation and allowed to incubate at 37°C for 24 and 48 hours (9).

For CSR at 46°C, an aliquot of milk placed in a sterile test tube was preheated for 10 minutes at 80°C to destroy vegetative forms and to activate the spores. Using a sterile pipette, 1 ml of the test sample (milk heated for 10 minutes at 80°C) was then injected deep into the tryptose-sulfite-cycloserine agar (TSC) (Pasteur Institute, Algeria) and the inoculum was mixed gently into the culture medium, without forming bubbles to avoid oxygenation of the medium. The tubes were then plunged into cold water to solidify the mixture. Following incubation at 46°C for 20 ± 2 hours, only the characteristic colonies, i.e. those surrounded by a black halo, were counted (10).
The enumeration results obtained for the different floras were interpreted according to the standards laid down in interministerial decree No. 35-1998 of January 1998 on the microbiological specifications of certain foodstuffs (5) (Table II).

**Detection of bacterial inhibitors in milk**

The DelvoTest® SP-NT (DSM Food, the Netherlands) was used to detect bacterial inhibitors in raw milk. It is based on inhibiting the growth of *Bacillus stearothermophilus* var. *calidolactis*, a bacterium which is very sensitive to a wide range of antibiotics and sulphonamides. It takes the form of ampoules containing an agar medium seeded with spores of *B. stearothermophilus* and enriched with growth nutrients. In each previously identified ampoule, 100 µl of a milk sample were introduced using a micropipette fitted with a disposable tip. The ampoules were placed in a water bath at 64 ± 1°C for three hours. On removal, the colour of the medium was examined by the naked eye. If a sample had clearly changed from violet to yellow, it indicated that the sample contained no bacterial inhibitors. In the presence of bacterial inhibitors the sample remained a violet colour.

**Statistical analyses**

Geometric mean calculations were performed for the enumeration of bacteria isolated at different points of the raw milk production chain on the farm.

The chi-square ($\chi^2$) test was used to test the relationships between the bacterial composition of milk and milk production points on the farm, as well as between the presence of bacterial inhibitors and milk production points on the farm.

**Results**

**Description of milking practices**

The results of the survey on milk production systems are presented in Table III. It transpires that:

- on almost all farms (86.8%), the milking machine was cleaned using only water
- very few milkers washed their hands (17%)
- on 83% of farms, the cows’ udders and teats were washed before milking using the same washing cloth for all the cows
- 88.7% of farmers neglected to wipe down the teats
- only 26.4% of farmers soaked the teats in a disinfectant solution, the remaining 73.6% failed to do this
- 73.6% of farmers neglected to discard the foremilk on the ground, compared with only 26.4% who did use this practice.

**Overall bacteriological quality of milk**

With regard to the criteria required by interministerial decree No. 35-1998 of 24 January 1998 on the microbiological specifications of raw milk (5), the results obtained from the 466 samples can be summarised as follows:

- 8.6% (31/360) of the individual milk samples, 15.1% (8/53) of milk samples from the milking machine and 47.2% (25/53) of milk samples from storage tanks did not comply with the legal criteria for all the enumerated bacteria

**Table II**

Microbiological specifications of raw milk (acceptability thresholds) in force in Algeria at the time of the study (5)

<table>
<thead>
<tr>
<th>Microbiological parameter</th>
<th>Acceptability threshold in raw milk</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total aerobic mesophilic flora at 30°C</td>
<td>$10^2$ cfu/ml</td>
</tr>
<tr>
<td>Faecal streptococci</td>
<td>Absence/0.1 ml</td>
</tr>
<tr>
<td>Faecal coliforms</td>
<td>$10^2$ cfu/ml</td>
</tr>
<tr>
<td><em>Staphylococcus aureus</em></td>
<td>Absence</td>
</tr>
<tr>
<td><em>Clostridium</em> sulphite-reducers at 46°C</td>
<td>50 cfu/ml</td>
</tr>
<tr>
<td>Bacterial inhibitors</td>
<td>Absence</td>
</tr>
</tbody>
</table>

**Table III**

Characteristics of milking practices on the 53 Algerian dairy cattle farms that participated in the study

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Proportion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Milking machine cleaning</td>
<td>Water + cleaning product 13.2% (7/53) Water only 86.8% (46/53)</td>
</tr>
<tr>
<td>Hand washing by milkers</td>
<td>Practised 17% (9/53) Not practised 83% (44/53)</td>
</tr>
<tr>
<td>Udder and teat washing before milking</td>
<td>Practised (collective washing) 83% (44/53) Not practised 17% (9/53)</td>
</tr>
<tr>
<td>Teat washing</td>
<td>Practised 11.3% (6/53) Not practised 88.7% (47/53)</td>
</tr>
<tr>
<td>Disinfection of teats after milking</td>
<td>Practised 26.4% (14/53) Not practised 73.6% (39/53)</td>
</tr>
<tr>
<td>Discarding the foremilk</td>
<td>Practised (discarding on the ground) 26.4% (14/53) Not practised 73.6% (39/53)</td>
</tr>
</tbody>
</table>
– 15.3% (55/360) of the individual milk samples, 24.5% (13/53) of milk samples from the milking machine and 17% (9/53) of milk samples from storage tanks complied with the legal criteria for only some enumerated bacteria
– 76.1% (274/360) of the individual milk samples, 60.4% (32/53) of milk samples from the milking machine and 35.8% (19/53) of milk samples from storage tanks complied with the legal criteria for all the enumerated bacteria.

A deterioration in milk quality was observed between the udder and the storage tank. The proportion of good quality milk fell from 76.1% to 35.8%, while poor quality milk rose from 8.6% to 47.2% (Fig. 1).

**Frequency of bacterial inhibitors in milk on the farm**

Bacterial inhibitors were detected in 28.8% (134/466) of all the raw milk samples. The 134 positive samples were distributed as follows:

– 30.6% (110/360) were individual milk samples
– 26.4% (14/53) were milk samples from the milking machine
– 18.9% (10/53) were milk samples from the storage tanks.

The rate of bacterial inhibitors in raw milk was particularly high in the individual milk samples (30.6%) (Table IV). The frequencies varied significantly depending on the sampling site \( (p < 0.05) \).

**Sources of milk contamination**

The proportion of samples contaminated by the bacteria studied (TAMF, FS, FC, SA and CSR) varied from 0% for samples from the hands of milkers (FS, FC and CSR), the milking environment (SA and CSR) and the milking water (FS and FC) to 98.1% for samples of water used during milking (TAMF) (Table V).

TAMF was found in all sample types at levels varying from 79.2% for samples taken from the hands of milkers to 98.1% for those from milking water.

While FS and FC were not detected in the samples taken from the hands of milkers or in the samples of water used during milking, they were often found in the samples taken from utensils (60.4% and 66% respectively), from udders (51.9% and 57.8% respectively) and from teat cups (41.5% and 45.3% respectively).

While CSR were detected in the samples taken from udders (10.8%), from utensils (9.4%), from teat cups (5.7%) and from the water used at different stages of milking (18.9%), they were not found in the samples taken from the hands of milkers or in the environmental samples.

**Staphylococcus aureus** was isolated mainly from the water used at the different stages of milking (50.9%), from samples taken from the hands of milkers (39.6%) and from udders (28.9%). The lowest levels were found on utensils (5.7%) and teat cups (7.5%).

**Critical points and the presence of contaminating bacteria**

A comparison of the bacterial counts in milk at different sampling points on the farm (from the cow's udder to the storage tank) showed a significant difference \( (p < 0.05) \) for each group of bacteria identified (Table VI).

In individual milk samples, 78.9% contained TAMF, 23.6% contained FS, 32.8% contained FC, 16.1% contained SA and 3.3% contained CSR.

**Table IV**

Proportion of milk samples containing bacterial inhibitors taken from the 53 Algerian dairy cattle farms that participated in the study

<table>
<thead>
<tr>
<th>Source of the sample</th>
<th>Proportion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Milk from the udder</td>
<td>30.6% (110/360)</td>
</tr>
<tr>
<td>Milk from the milking machine</td>
<td>26.4% (14/53)</td>
</tr>
<tr>
<td>Milk from storage tanks</td>
<td>18.9% (10/53)</td>
</tr>
</tbody>
</table>
In the milk in storage tanks, the proportions were respectively 96.2% (TAMF), 64.2% (FS), 75.5% (FC), 58.5% (SA) and 5.7% (CSR).

The bacterial load in raw milk samples rose progressively along the farm production chain (Fig. 2).

**Discussion**

Samples from raw cow’s milk and the environment, as well as from milking equipment, were taken on several dairy farms in the Jijel and Blida regions of Algeria. Convenience sampling was employed in order to include farms of different sizes using different methods and different milking equipment. Milk quality was assessed according to the Algerian standards in force for the microbiological specifications of raw milk.

Poor hygiene conditions during milking and milk storage, as well as lack of hygiene among milkers (dirty hands, poor-quality milking water, etc.) and in the equipment used for milk production, were the causes of the poor hygiene quality of the milk produced. In fact, bacterial contamination of milk worsened as it progressed along the production chain.

**Hygiene assessment of milking practices**

The survey conducted on these dairy farms revealed that, in general, neither milking conditions, nor equipment cleaning, nor milk storage were optimal. On all of the farms covered by the study, milking was carried out under poor hygiene conditions and cleaning products were rarely used for udder preparation or for equipment cleaning.

**Cleaning milking machines**

The majority of milkers (86.8%) rinsed the milking machine with water only, compared with 13.2% who used a mixture of water and a cleaning product. These results are similar to those of the study conducted in Monastir (11), where 10% of farmers alternated the use of acid and alkaline detergents during cleaning operations.

**Hand washing by milkers**

The level of hygiene among the majority of milkers was unacceptable: only 17% of them washed their hands before each milking, while most (83%) did not.

According to Thomelin (12), it is vital to ensure the best possible hygiene conditions in order to reduce

**Table V**

<table>
<thead>
<tr>
<th>Source of the sample</th>
<th>TAMF (52/53)</th>
<th>FS (0/53)</th>
<th>FC (0/53)</th>
<th>SA (50.9/27)</th>
<th>CSR (18.9/10)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water used during milking</td>
<td>98.1%</td>
<td>0%</td>
<td>0%</td>
<td>50.9%</td>
<td>18.9%</td>
</tr>
<tr>
<td>Utensils</td>
<td>94.3%</td>
<td>60.4%</td>
<td>66%</td>
<td>5.7%</td>
<td>9.4%</td>
</tr>
<tr>
<td>Milking environment</td>
<td>81.1%</td>
<td>13.2%</td>
<td>18.9%</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Udders</td>
<td>83.9%</td>
<td>51.9%</td>
<td>57.6%</td>
<td>28.9%</td>
<td>10.8%</td>
</tr>
<tr>
<td>Treat cups</td>
<td>96.2%</td>
<td>41.5%</td>
<td>45.3%</td>
<td>7.5%</td>
<td>5.7%</td>
</tr>
<tr>
<td>Hands of milkers</td>
<td>79.2%</td>
<td>0%</td>
<td>0%</td>
<td>39.6%</td>
<td>0%</td>
</tr>
</tbody>
</table>

**Table VI**

<table>
<thead>
<tr>
<th>Source of the sample</th>
<th>TAMF (284/360)</th>
<th>FS (85/360)</th>
<th>FC (118/360)</th>
<th>SA (50/360)</th>
<th>CSR (12/360)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Milk from the udder (individual)</td>
<td>78.9%</td>
<td>23.6%</td>
<td>32.8%</td>
<td>16.1%</td>
<td>3.3%</td>
</tr>
<tr>
<td>Milk from the milking machine</td>
<td>83%</td>
<td>26.4%</td>
<td>41.5%</td>
<td>22.6%</td>
<td>3.8%</td>
</tr>
<tr>
<td>Milk from storage tanks</td>
<td>96.2%</td>
<td>64.2%</td>
<td>75.5%</td>
<td>58.5%</td>
<td>5.7%</td>
</tr>
</tbody>
</table>
contamination of udders and milk by bacteria that can enter when a cow's sphincters are open.

**Udder and teat washing before milking**

Most milkers (83%) performed collective washing of teats and udders before milking. These results are similar to those of M'Sadak et al. (13), who found that the majority of farmers (93%) prepared the udder by pre-washing with water using the same cloth for all the cows.

According to Noireterre (14), this udder preparation method increases the risk of transmitting pathogens from an infected quarter to a healthy quarter of the udder with the subsequent onset of mastitis.

**Teat washing**

This stage can minimise the risk of mastitis, improve milk quality and prevent slippage of teat cups and the entry of air (vacuum fluctuation) into milking units (15).

This study found a teat-washing frequency (11.3%) well below that reported by M'Sadak et al. (67%) in a study of the effect of milking conditions on the udder health of dairy cows in the Mahdia region of Tunisia (16).

**Disinfection of teats**

This study found that very few milkers (26.4%) disinfected the teats after milking. These results are significantly lower than those reported by M'Sadak et al. (11) in a recent study in the Monastir region, where 59% of farmers applied this practice.

According to Bareille and Lemarchand (17), teat disinfection after milking can reduce the rate of new intra-mammary staphylococcus infections by 50–95%. According to Hanzen (18), teat disinfection after milking can reduce the number of microorganisms transferred via the teat canal during milking, which may have developed at the tips between milking sessions. It can also be used to treat any injuries to the teats.

**Discarding the foremilk**

The authors found that only 26.4% of milkers discarded the foremilk, even though doing so has advantages over early detection of clinical mastitis and the elimination of microorganisms in the teat canal (18). This corroborates the observations of M'Sadak et al. (11) in the Monastir region, where 28% of farmers used this practice.

**Bacterial inhibitors in milk**

According to the interministerial decree on the microbiological specifications of certain foodstuffs (5), good quality milk should not contain bacterial inhibitors. However, 28.8% of the 466 raw milk samples analysed in this study contained bacterial inhibitors and the majority of farmers treated mastitis with penicillin and/or tetracycline (according to the results of the authors' survey questionnaire). These figures show the scale of bacterial inhibitor use on the dairy farms studied.

These results are in line with those of Srairi et al. (19) and Kouame et al. (20), who reported levels of around 25% and 24.7% respectively.

Heavy contamination of milk samples with tetracycline and/or penicillin was also confirmed by a study by Ben Mahdi and Ouslimani (21), who reported a contamination rate of around 97.3%.

In addition, these results revealed differences in the proportion of bacterial inhibitors, depending on the site. They were found in 30.6% of individual samples of milk (on leaving the udder), in 26.4% of milk samples taken from milking machines and in 18.9% of samples from storage tanks. These figures indicate the scale of antibiotic use on dairy farms where the milk is collected and hence the extent of the resulting risk to consumer health.

High levels of bacterial inhibitors in individual raw milk samples are probably caused by the widespread, uncontrolled use of intra-mammary pharmaceutical
Faecal coliforms and streptococci were found on utensils (39.6%) and on the udders (28.9%). Faecal contamination was found in the water used at the different milking stages (50.9%), on the hands of milkers (39.6%) and on the udders (28.9%). The presence of faecal coliforms and faecal streptococci in raw milk is strongly associated with faeces-soiled udder skin and poorly designed feed (that had been in contact with the ground), which contaminates the milk either directly or via faeces. These are pathogenic bacteria and their presence indicates recent or older faecal contamination of the ground (27).

**Overall bacterial quality of milk and sources of contamination**

These results show that milk becomes increasingly contaminated as it progresses through the different stages of milking. Between the udder and the milk storage tank, the proportion of good quality milk samples fell from 76.1% to 35.8%.

This rapid decline in the bacteriological quality of milk as it passes along the farm production chain is the result of successive instances of contamination from utensils, the udders, the teat cups, the milking environment and the hands of milkers. Milk becomes contaminated during milking operations and the more it is handled, the greater the risk of bacterial contamination.

Bacteria detection revealed that the udder skin, utensils and teat cups carried all the bacteria under investigation (TAMF, FS, FC, SA and CSR). The udders of some cows were more contaminated than others and the mixing of raw milk from several cows contributed to the drop in milk quality in storage tanks.

In addition, *S. aureus* was found in the water used at the different milking stages (50.9%), on the hands of milkers (39.6%) and on the udders (28.9%). Faecal streptococci and faecal coliforms were found on utensils (60.4% and 66% respectively), on the udders (51.9% and 57.8% respectively), on teat cups (41.5% and 45.3% respectively) and in the milking environment (13.2% and 18.9% respectively). *Clostridium* sulphite-reducers were found at low levels in the water used at the different milking stages (18.9%), on the udders (10.8%), on utensils (9.4%) and on teat cups (5.7%). All these elements are therefore sources of contamination of raw milk.

The presence of TAMF in raw milk is an indicator of the overall level of hygiene on farms. TAMF includes microorganisms that cause spoilage or contamination, acidifying lactic flora and sometimes pathogenic bacteria. Enumeration of these flora is the method most commonly used by dairy processing plants to assess the bacterial quality of milk and it is therefore an important indicator of hygiene conditions during milking (25). The high levels of these flora found in samples from milk cooling tanks is probably the result of intensive bacterial growth arising from failure to control hygiene conditions during milking and milk storage.

The presence of faecal coliforms and faecal streptococci in raw milk indicates an environmental source of contamination. Their proliferation in raw milk reflects a failure to observe the required hygiene measures during milking, and probably contamination during milk storage. Faecal coliforms and streptococci in raw milk are strongly associated with faeces-soiled udder skin and poorly designed and improperly cleaned milking equipment (25). Bonfoh et al. (26) report that poor cleaning of recipients in contact with milk on the farm left residual levels of contamination of around $4.1 \log_{10}$ colony-forming units (cfu)/ml.

*Clostridium* sulphite-reducers were found in animal feed (that had been in contact with the ground), which contaminates the milk either directly or via faeces. These are pathogenic bacteria and their presence indicates recent or older faecal contamination of the ground (27).

*Staphylococcus aureus* is a contagious agent living on cow udders that can be transmitted from one cow to another (28). This bacterium can enter milk either directly, by excretion from udders infected with clinical or subclinical staphylococcal mastitis, or by environmental contamination during the handling and processing of raw milk (29, 30). When the udder is infected, *S. aureus* is excreted in the milk in highly variable quantities from $0$ to $10^6$ cfu/ml (31). These results, which support those of Kouame et al. (20), show that this bacterium came mainly from the water used at the different stages of milking (50.9%), from the hands of milkers (39.6%) and from udders (28.9%).
Conclusion

This study shows that the increasing bacterial load in milk as it passes along the farm production chain is the result of successive instances of contamination associated with poor hygiene practices during milking.

The search for sources of contamination along the entire raw milk chain showed that udders, milkers’ hands, teat cups, utensils, the milking environment and the water used during milking were all sources of milk contamination by the bacteria under investigation. In addition, bacterial inhibitors were detected in the milk samples analysed.

To improve the quality of raw milk, farmers need to implement a range of hygiene measures in cowsheds and during milking, all the more rigorously and systematically because the animals’ environment is highly contaminated. This environmental contamination could be reduced by introducing manure storage and spreading practices to prevent the recycling and spread of bacteria. This will be difficult to achieve without the effective participation of farmers following information campaigns targeted at them.

Acknowledgements

The authors would like to thank all the staff of the laboratory of the Algerian centre for quality control and packaging (CACQE) and the laboratory for veterinary testing, quality control, compliance and applied research (AVCQ-LAB).

References


