Using real-time polymerase chain reaction as an alternative rapid method for enumeration of colony count in live Brucella vaccines

Waleed S. Shell1, Mahmoud L. Sayed1, A. A. Samy2, Ghada Mohamed Al-Sadek1, Gina Mohamed Mohamed Abd El-Hamid1 and Abdel Hakam M. Ali1

1. Central Laboratory for Evaluation of Veterinary Biologics, Cairo, Abbassia, Egypt; 2. Department of Microbiology and Immunology, National Research Center, Egypt.

Corresponding author: Waleed S. Shell, e-mail: tarikwaleedshell@hotmail.com,
Co-authors: MLS: m_lotfi8@hotmail.com, AAS: ayman_samy@hotmail.com, GMA: ghada.elsadek@yahoo.com, GMMAE: gina_mohammed@msn.com, AMA: hakam2060@gmail.com
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Abstract

Aim: Brucellosis is a major bacterial zoonosis of global importance affecting a range of animal species and man worldwide. It has economic, public health, and bio-risk importance. Control and prevention of animal brucellosis mainly depend on accurate diagnostic tools and implementation of effective and safe animal vaccination program. There are three types of animal Brucella live vaccines - Brucella melitensis Rev-1 vaccine, Brucella abortus S19, and B. abortus RB51. Evaluation of these vaccines depends mainly on enumeration of Brucella viable count. At present, used colony count method is time consuming, costly and requires especial skills. Hence, the aim of this study is to use and standardize real-time polymerase chain reaction (RT-PCR) as an alternative, quantitative, sensitive, and rapid method to detect the colony count of Brucella in live Brucella vaccine.

Materials and Methods: Four batches of different live Brucella vaccines were evaluated using of conventional bacterial count and RT-quantitative PCR (RT-qPCR) using BSCP31 gene specific primers and probe. Standard curve was generated from DNA template extracted from 10-fold serial dilution of living B. abortus RB51 vaccine to evaluate the sensitivity of RT-qPCR.

Results: Results revealed that three batches of living Brucella vaccines were acceptable for Brucella colony count when traditional bacterial enumeration method was used. Results of RT-qPCR were identical to that of conventional bacterial count.

Conclusions: Results concluded that RT-qPCR was relatively sensitive compared to traditional bacterial colony count of these vaccines.

Keywords: Brucella, colony count, RB51, Rev-1, real-time polymerase chain reaction, S19, vaccines.

Introduction

Brucellosis is a major bacterial zoonosis of global importance affecting a range of different mammals including cattle, sheep, goats, swine, rodents, marine mammals, and man worldwide. In food animals, the disease primarily affects the reproductive system with concomitant loss in fertility and productivity of affected animals. In man, infection is characterized by recurrent febrile episodes that lead to the description of this disease as undulant fever (economic and public health importance) [1]. The severity of this disease and lack of vaccines suitable for use in man has led to the investigation of Brucella as agents for bioterrorism (bio-risk importance) [2]. Vaccines to be used for human are not yet available, and so eradication of human brucellosis largely depend on the eradication of the disease in animals. Eradication of brucellosis in animals has been a goal for many countries. To control brucellosis, comprehensive vaccination, surveillance, and quarantine programs should be implemented. Both control and prevention procedures are highly dependent on accurate diagnostic tools and implementation of effective and safe animal vaccination programs [3].

There are three types of animal Brucella live vaccines - Brucella melitensis Rev-1 vaccine (0.5-2×10⁵ colony forming unit [CFU]/dose) for vaccination of sheep and goats, Brucella abortus S19 (0.5-5×10⁹ CFU/dose) for vaccination of cattle and bufaloes, and B. abortus RB51 (1-3.4×10⁹ CFU/dose) for vaccination of cattle and bufaloes. Evaluation of these vaccines depends mainly on enumeration of viable count, smoothness or roughness, safety test and potency test [4]. European Pharmacopoeia [5] reviewed that the dose of Rev-1 vaccine in sheep and goats should contain not fewer than 0.5×10⁹ and not more than 4×10⁹ live bacteria per dose.

At present, practiced colony count method is time consuming, costly and requires especial skills.
Hence, the aim of this study was to use and standardize real-time polymerase chain reaction (RT-PCR) as a quantitative, sensitive, and rapid method to detect the colony count of live *Brucella* vaccine.

**Materials and Methods**

**Vaccines**

Eight lyophilized living *Brucella* vaccines of different batches (two *B. abortus* S19, four *B. abortus* RB51, and two *B. melitensis* Rev-1). The lyophilized vaccines were reconstituted in vaccine diluents and were used for bacteriological colony count and genomic DNA extraction.

**Bacterial colony count of living *Brucella* vaccines**

About 0.1 ml of expected countable dilutions of different live *Brucella* vaccines were inoculated in five plates of tryptone soya agar and spread with a sterile glass. CFU per vaccine dose were enumerated according to protocols described previously [4,6].

**Extraction of genomic DNA from *Brucella* strains**

Genomic DNA extraction from single dose of live *Brucella* vaccines for evaluation of *Brucella* viable count and from 10-fold serial dilutions of RB51 vaccine from $10^{5}$ to $10^{10}$ CFU/ml for generation of standard curve (RB51 vaccine vial of 5 doses $2 \times 10^{10}$ to $10^{10}$ CFU/ml for serial dilution). Genomic DNA extraction was performed using G-spin Total DNA Extraction Kit (*iNtRON*) following the kit manufacturer’s protocol.

**Oligonucleotide primers and probes used in RT-PCR**

Real-time PCR on tested samples was done using the primers and probe [7,8] identifying and targeting the bscp31 gene (GenBank accession number M20404) [7] (Table-1 and Figure-1).

**RT-PCR**

RT-PCR assay was standardized and performed in Stratagene MX3005P quantitative PCR (qPCR) system. The PCR Master Mix and PCR cycling conditions used are given in Tables-2 and 3.

**RT-PCR standard curves**

Standard curves were generated by plotting the cycle threshold values (CT) of the RT-qPCR performed on 10-fold serial dilutions of purified DNA from 10-fold serial dilutions of *B. abortus* RB51 vaccine ($10^{5}$ to $10^{10}$ CFU/ml) against the log input cells/ml [9]. *Brucella* species concentrations were determined by the viable cell plate count method as mentioned above [4,6].

**Results and Discussion**

In the absence of effective and safe human vaccine against brucellosis, animal vaccination against brucellosis is an important issue in control and eradication of brucellosis in animals and human. For more than 60 years, *B. abortus* S19 vaccine for buffaloes and cattle and *B. melitensis* Rev-1 vaccine for goats and sheep remain as the most efficient *Brucella* vaccines, and their use is of a great impact on the control and incidence of brucellosis in domestic ruminants and humans [10]. S19 and Rev-1 vaccines are used in vaccination of calves and ewes, respectively, in a dose of $0.5 \times 10^{6}$ CFU/dose and $0.5 \times 10^{6}$ CFU/dose [4]. Rev-1 vaccine can be used in a dose of $0.5 \times 10^{5}$ CFU/dose [5]. RB51 vaccine strain was developed in 1982 by Prof. Gerhardt Schurig’s group and is derived from a virulent smooth *B. abortus* biovar 1 strain 2308. RB51 vaccine is used in vaccination of cows in a dose of $1 \times 10^{6}$ CFU/dose [4,11]. Evaluation of these vaccines depends mainly on identification of vaccinal strains, enumeration of *Brucella* viable count, safety and potency. Enumeration of *Brucella* viable count is time consuming and needs special skills [4]. This study was designed to use a RT-qPCR as alternative, sensitive, and rapid method to detect colony count in *Brucella* vaccines. Eight batches of *Brucella* vaccines, two *B. abortus* S19, four *B. abortus* RB51, and two *B. melitensis* Rev-1 were evaluated by conventional bacterial colony count and RT-qPCR.

By using conventional colony count, seven batches of living *Brucella* vaccines used in this study

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**Table-1:** RT-PCR oligonucleotides primers and probe of BCSF31 for *Brucella* species.

| Primer                  | Sequence (5’-3’)                              | Amplicon size (bp) |
|------------------------|-----------------------------------------------|-------------------|
| BSCP31 Forward primer  | GCTCGGTTGCGCAATATCATGC                      | 151 bp            |
| BSCP31 Reverse primer  | GGATAAAGCGTGCCAGAGA                       |                  |
| RT-PCR probe           | AAATCTTCACCTGGCCATCA-FAM/BHQ1               |                  |

RT-PCR=Real-time polymerase chain reaction
were with satisfactory results and within the standard international range of acceptable dose for animal’s vaccination. On the other hand, one of the RB51 vaccine batches was unacceptable with colony count of $6\times10^9$ CFU/dose as shown in Table-4.

In this study, bcsp31 gene was selected to be used in RT-qPCR for evaluation of colony count of living *Brucella* vaccines which is highly conserved gene among *Brucella* species and also used frequently as a gene target for diagnosis of human brucellosis [12-14], and therefore could potentially detect *B. melitensis* and *B. abortus* strains which were included in this study [15,16]. Moreover, it is specific method as it did not amplify DNA from any non-*Brucella* templates. The bcsp31 PCR was found to be 100% specific and was the most sensitive assay when compared with to omp2 and the 16S rRNA PCR [17]. BSCP31 PCR was used by many researchers for specific identification of genus *Brucella* from seropositive, active, relapsing, chronic cases in humans [18-20]. Furthermore, this gene target has been used specifically to detect *Brucella* in human cerebrospinal fluid, blood, and serum [21-23], in clinical tissues from seals [24] and in buffalo milk [25]. Many reports have been published on the diagnostic efficiency of qPCR assays using bcsp31 gene for diagnosis of brucellosis in human samples [26] and also used for screening of brucellosis from camel serum [27].

TaqMan technology determines the PCR cycle at which the increase in fluorescence of the reporter dye reaches a CT is proportional to the log of the amount of target DNA and hence the log of the number of bacteria in the sample. Standard graph was based on *B. abortus* RB51 DNA extracted from tenfold serial dilution of RB51 vaccine (Figure-2). The RT-PCR assay with primers and probe specific for the *Brucella* BCSP31 gene was positive for all vaccine samples. The CT values were clearly inversely related to the quantity of organisms, especially during standard curve generation. These CT values corresponded to $10^{11}-10^9$ CFU (positive at CT=16) to $10^9$ CFU (positive at CT=32) of *Brucella* organism when the values were fit into the standard curve generated by using the results for serial dilutions of RB51 vaccine. Colonies count $10^{11}-10^9$ CFU represents the range of acceptable colonies count of all type of *Brucella* vaccines ($1\times10^9$ CFU/dose in case of Rev-1 vaccine to $3.4\times10^9$ CFU/dose in case of RB51 vaccine) (Figure-3).

As shown in Figures-2 and 3 and Table-4, results of RT-qPCR were in agreement with results of traditional bacterial colony count except with one batch of Rev-1 vaccine where results of traditional colony count and RT-qPCR were $3\times10^9$/CFU/dose and $5.163\times10^9$/CFU/dose, respectively, but still results within the same log. Results of qPCR were with sensitivity of 87.5%. Findings confirmed that the unaccepted batch of RB51 vaccine by traditional colony count was out of standard international range of *B. abortus* RB51 vaccine.

Results agree with Angel et al. [28] who used RT-qPCR for enumeration of acetic acid bacteria with 100% sensitivity when compared with plating and microscope counting also was in agreement with Chaloemnon et al. [29] who enumerated the gastrointestinal microbiota (*Lactobacilli, Bifidobacteria, and Escherichia coli*) in weaning pigs by conventional culture and RT-PCR. Aline et al. [30] and Susan et al. [9]. Found high agreement with the results of traditional colony count and RT-qPCR when used to

| Table-2: Preparation of PCR master mix. | Component | Volume/reaction |
|----------------------------------------|-----------|----------------|
| 2x QuantiTect Probe RT-PCR master mix  | 12.5 µl   |
| Forward primer                        | 0.2 µl (200 mm) |
| Reverse primer                        | 0.2 µl (200 mm) |
| Probe                                 | 0.1 µl (100 µm) |
| DNase free water                      | 6.8 µl   |
| Template DNA                          | 5 µl     |

RT-PCR=Real-time polymerase chain reaction

| Table-3: RT-PCR cycling conditions. | Stage                | Temperature | Time | Cycles |
|-----------------------------------|----------------------|-------------|------|--------|
| Primary denaturation              | 95°C                 | 10 min      | 1    |
| Amplification                     | 95°C                 | 30 s        | 40   |
| Secondary denaturation            | 60°C                 | 90 s (optics on) |

RT-PCR=Real-time polymerase chain reaction

| Table-4: *Brucella* count by traditional methods and RT-qPCR based on a standard graph generated by *brucella* RB51 DNA within the range. | *Brucella* vaccines samples | CT | Estimation of *brucella* vaccines by rt-PCR | *Brucella* viable count by traditional methods | Acceptance |
|---------------------------------------------------------------------------------------------------------------------------------|-----------------------------|----|--------------------------------------------|-----------------------------------------------|-----------|
| RB51                                                                                                                              | 21.69                       | 2.89×10^9 CFU/dose                     | 3.4×10^10 CFU/dose                             | Accepted  |
| RB51                                                                                                                              | 23.00                       | 1.25×10^10 CFU/dose                     | 3.4×10^10 CFU/dose                             | Accepted  |
| RB51                                                                                                                              | 22.14                       | 2.325×10^10 CFU/dose                    | 1.2×10^11 CFU/dose                             | Accepted  |
| RB51                                                                                                                              | 28.65                       | 4.19×10^9 CFU/dose                      | 6×10^10 CFU/dose                               | Not accepted |
| S19                                                                                                                               | 27.98                       | 5.025×10^9 CFU/dose                     | 4×10^9 CFU/dose                                | Accepted  |
| S19                                                                                                                               | 31.52                       | 1.6×10^9 CFU/dose                       | 4.8×10^9 CFU/dose                              | Accepted  |
| Rev-1                                                                                                                             | 27.87                       | 5.163×10^9 CFU/dose                     | 3×10^10 CFU/dose                               | Accepted  |
| Rev-1                                                                                                                             | 32.00                       | 1×10^9 CFU/dose                         | 1.5×10^9 CFU/dose                              | Accepted  |

RT-qPCR=Real-time quantitative polymerase chain reaction, CFU=Colony forming unit, CT=Cycle threshold
enumerate *Lactobacillus helveticus* in dairy products and *Streptococcus pneumoniae*, respectively.

In disagreement with these results, Botaro et al. [31] reviewed that the qPCR protocol can be used as a rapid diagnostic assay to accurately detect *Staphylococcus aureus* from bovine milk, but this protocol is not accurate for counting of *S. aureus* in bronopol-preserved milk samples from naturally infected mammary glands. Same findings were revealed from enumeration of living *E. coli* O157:H7 on plants [32].

A major drawback of qPCR is its inability to differentiate the DNA from viable and dead cells, and this is a critical factor for many researchers’ especially in the food industry, water pollution researches, so to remedy this shortcoming, researchers have used biological dyes such as ethidium monoazide and propidium monoazide to pre-treat samples before DNA extraction which is important issue especially in food industry [32,33].

Results of this study may be more applicable than other studies which used RT-PCR for identification of organisms from tissues, water, etc., which may give false results due to nonspecific reactions which especially occurs when RT-PCR used for identification of multiple organisms using universal primers sets. However, in this study, we evaluated vaccines which contain one organism type (*Brucella*) as these vaccines were tested for sterility before counting process. Furthermore, although RT-PCR measure the total number of living and dead (dead cells as a results of freeze drying process) *Brucella* cells, but due to all these vaccines are subjected to the same factors as freeze drying program so ratio of living to dead cells were nearly constant and so it would not have an effect on the sensitivity of the RT-PCR.

**Conclusions**

In this study, RT-qPCR assay was developed to enumerate colony count in live *Brucella* vaccines using DNA template extracted from tenfold serial dilutions of different living *Brucella* vaccines. The
The authors have conducted PCR work on different live bacterial vaccines such as Brucella, E. coli (Poulvac_poultry vaccine), Salmonella vaccine (Megan VAC-1, poultry vaccine), and Streptococcus equi vaccine (PINNACLE® I.N., horse vaccine) and evaluated the sensitivity of standard curves generated from DNA template extracted from tenfold serial dilutions of different living bacterial vaccines and from 10-fold serial dilution of template DNA.

**Authors’ Contributions**

All authors designed and planned this research work. DNA extraction from different live Brucella vaccine batches and from serial dilution of control positive live Brucella vaccine batch were carried out by AAS, GMA, GMMAE and AMA. Traditional Colony count of different live Brucella vaccines batch was performed by all authors. RT-PCR on different live Brucella vaccines batch and construction of standard curve to estimate bacterial count in vaccine batches were carried out by WSS, MLS, AAS and AMA. All authors contributed equally in preparation and revision of the manuscript and collection of scientific papers related to the subject of this research. All authors read and approved the final manuscript.

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**Competing Interests**

The authors declare that they have no competing interests.

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