Evaluation of the General Organization of Veterinary Services control program of animal brucellosis in Egypt: An outbreak investigation of brucellosis in buffalo

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Abstract

Background and Aim: Brucellosis is a major constraint to livestock production in Egypt as well as many developing countries worldwide. Bovine brucellosis is an economically important disease with reproductive failure as a principal manifestation resulting in abortion, premature birth and decreased milk production in females, and orchitis and epididymitis in males. In spite of the efforts of Egyptian veterinary services to overcome brucellosis, the disease is still prevalent in both animals and humans and represents one of the most important public health hazards in Egypt. The aim of the present work was to investigate the efficacy of the control program implemented by the General Organization of Veterinary Services in Brucella infected buffalo farm on serological, molecular, cultural, and histopathological basis. Brucella melitensis biovar 3 was recovered from 6 buffalo-cows.

Materials and Methods: Blood samples were collected from a total of 750 non-vaccinated lactating buffalo-cows. These animals were proved positive for Brucella by the Egyptian brucellosis national program. Sera were tested using buffered acidified plate antigen test and rose Bengal test as screening tests and complement fixation test as a confirmatory test. Positive animals were separated for slaughter under the supervision of the Egyptian veterinary authorities. Remaining animals were tested every 3 weeks with slaughtering of positive cases and this continued until the remaining animals revealed three successive negative serological tests. Different lymph nodes (prescapular, prefemoral, mediastinal, retropharyngeal, and supramammary) were collected from 11 Brucella seropositive buffalo-cows slaughtered after being confirmed serologically as Brucella infected cases. Samples were collected and processed for bacterial isolation and nucleic acid detection using polymerase chain reaction (PCR). Parts of these specimens were fixed in 10% neutral buffered formalin for 48 h then processed by paraffin embedding technique.

Results: “Test and slaughter” policy was applied on Brucella infected dairy buffalo farm. The program continued for 6 months with slaughtering of positive cases until the herd was proved Brucella free. B. melitensis biovar 3 could be recovered from six buffalo-cows. Universal PCR confirmed Brucella on genus level and Bruce-ladder multiplex, PCR confirmed the presence of B. melitensis on the species level. Histopathological examination of Brucella-infected lymph nodes revealed massive rarified and depleted lymphoid areas of both sub-capsular and deep cortical lymphoid follicles, macrophage cells granulomatous reaction, as well as fat, infiltrates, and chronic vasculitis. The chronic nature of Brucella lesions has been confirmed in this study as indicated by the chronic vasculitis and collagen deposition.

Conclusion: Freedom status from brucellosis in this study required 6 months which are considered long time allowing the spread of infection to other localities especially under unhygienic conditions, husbandry system favoring mixed populations of different ages, sex, aborted and pregnant, and lack of controlled movement of animals. Therefore, effective control of animal brucellosis requires surveillance to identify infected animal herds, elimination of the reservoirs, and vaccination of young heifers. B. melitensis biovar 3 is the cause of the Brucella outbreak in buffalo which still remains the prevalent type of Brucella in Egypt. The disease runs a chronic course allowing further spread of infection.

Keywords: bruce-ladder, brucellosis, buffalo, histopathology, polymerase chain reaction.

Introduction

Brucellosis is a major constraint to livestock production in Egypt as well as many developing countries worldwide. Bovine brucellosis is an economically important disease with reproductive failure as a principal manifestation resulting in abortion, premature birth and decreased milk production in females, and
Brucellosis in ruminants is most symptomatic only in primary infections where animals do not continually abort, however, aborted material, uterine, and vaginal discharges, as well as milk, contain a huge number of bacteria [9].

Bovine abortion is less common with an infection caused by Brucella melitensis than in Brucella abortus infections [10]. Interestingly, in Egypt B. melitensis is the most predominant type isolated from animals and humans [11-15].

Different Brucella species are incriminated as highly pathogenic in humans, with B. melitensis classically reported as the agent responsible for most of the human infections [16].

Direct laboratory methods of diagnosis of brucellosis in animals such as bacterial isolation have high specificity but are time-consuming and require an appropriate degree of biosafety. Polymerase chain reaction (PCR) has been introduced as an effective tool for rapid detection and confirmation of Brucella infection as well as differentiating Brucella species. Only, both culture and molecular techniques can definitively establish the presence of infection and determine the responsible Brucella species. On the other hand, serological procedures are used more often as quick and less expensive diagnostic tools. However, serological tests have several limitations concerning specificity and sensitivity, especially when testing individual animals [17].

Control of brucellosis presents considerable difficulties due to its contagiousness, uncertainties about detection [18], wide host range, significant numbers of carriers [19], and latently infected animals [9]. National high committee of zoonoses of the General Organization of Veterinary Services (GOVS) established the brucellosis control program which depends mainly on the test-and-slaughter policy. Herds containing even one positive animal were kept under quarantine, and all animals were to be subjected to periodical testing every 21 days. Quarantine measures were released if the animals pass three consecutive negative tests at 21 days intervals [20].

In spite of the efforts of Egyptian veterinary services to overcome brucellosis since 1988, the disease is still prevalent in both animals and humans and represents one of the most important public health hazards in Egypt as the prevalence of animal brucellosis in Egypt in the past 20 years ranged from 0.33% to 1.32% according to the official reports of the GOVS, Egypt [21].

The aim of the present work was to investigate the efficacy of the control program implemented by the GOVS in Brucella infected buffalo farm on serological, molecular, cultural, and histopathological basis.

Materials and Methods

Ethical approval

All clinical samples in this study were collected as per standard sample collection procedure without giving any stress or harm to the animals. The present work was approved by the Ethical Committee for Medical Research at the College of Veterinary Medicine, Beni-Suef University and Animal Care Guidelines of the General Organization for Veterinary Services, Egypt.

Study area

The study was conducted on a buffalo farm of 952 buffaloes at Cairo - Ismailia desert road, Ismailia Governorate, Egypt from September 2016 to April 2017. This farm was located under quarantine by the veterinary authorities due to Brucella infection.

Study population and samples

- This study was carried out on a Brucella infected dairy buffalo farm of 952 animals under quarantine of the GOVS. Animals were kept overcrowded and reared in open system in which animals in different ages; aborted and pregnant ones; males and females were housed together.
- Blood samples were collected from a total of 750 non-vaccinated adult lactating buffalo-cows out of 952 animals. About 20 ml of blood were aseptically drawn from jugular vein into a sterile evacuated test tube using a jack. The test tubes were left at room temperature in a sloping position to allow clotting for 2 h. The collected samples were labeled, identified and transferred to the laboratory where they held in refrigerator till the next day to give a chance for serum to separate. The serum was siphoned off by Pasteur Pipette after centrifugation at 3000 rpm for 10 min. Clear sera were stored in Cryotubes at −20°C until its use for serological studies.
- Sera were screened using buffered acidified plate antigen test (BPAT) and rose Bengal test (RBT). Positive samples were further tested with complement fixation test (CFT) as recommended by the OIE [22]. Positive animals were separated for...
slaughtering under the supervision of the Egyptian veterinary authorities. Animals were tested every 3 weeks for detection of animals which are still incubating the disease according to the roles of the GOVS with slaughtering of positive animals after each test. This continued until obtaining three successive negative serological tests.

- Different lymph nodes (prescapular, prefemoral, mediastinal, retropharyngeal, and supramammary) were collected from 11 slaughtered seropositive buffalo-cows. Samples were collected and processed for bacterial isolation according to Alton et al. [23]. Parts of these specimens were fixed in 10% neutral buffered formalin for 48 h then processed by paraffin embedding technique.

**Brucella strains**

- Reference *Brucella* strains; *B. melitensis* Ether, *B. abortus* 544, and *Brucella suis* 1330 were supplied by the Central Veterinary Laboratory, Weybridge, Surrey KT15 3NB, UK.
- Field isolate of *B. melitensis* biovar 3 isolated from cattle (Collection culture of the Department of Veterinary Medicine, Beni-Suef University) was also used.
- *B. abortus* S19 vaccine strain was obtained from Veterinary Serum and Vaccine Research Institute, Abbasia, Cairo, Egypt. Reference *Brucella* strains and the field *Brucella* isolate were used as control positives for the identification and typing of *Brucella* isolates recovered in this study.

**Primers**

Primer sequences used for universal and multiplex Bruce-ladder PCR are listed in Table-1.

**Serological examination**

BPAT using buffered acidified plate test antigen (killed *B. abortus* strain 99 antigen, at concentration of 11% in lactate buffer, pH 3.7±0.03), RBT using RBT antigen (Rose Bengal stained, 8% cells killed *B. abortus* strain 99 antigens in lactate buffer, pH 3.65±0.05) and CFT, warm micro technique using CFT antigen (*B. abortus* biovar 1 strain 1119-3 cells in phenol saline, at a concentration of 4.5%, pH 6.8) were conducted according to Alton et al. [23] and OIE [22].

**Bacteriological examination**

Lymph nodes of 11 seropositive buffalo-cows that were slaughtered under supervision of the GOVS were cultured on tryptose agar medium (Oxoid) with antibiotics selective antibiotic supplement [24] according to Alton et al. [23]. Plates were incubated at 37°C in an atmosphere of 10% CO₂ and examined daily for 10 days for growth. The criteria used for identification were the colonial morphology, biochemical tests, requirement for additional atmospheric 10% CO₂, production of hydrogen sulfide gas, production of urease, growth on media containing the inhibitory dyes thionin and fuchsin, agglutination with polyclonal monospecific antisera A and M and R, and phage typing using Tbilisi (Tb) and Izatnagar (Iz,) according to the methods recommended by Alton et al. [23] and OIE [22].

**DNA extraction**

*From Brucella culture*

Few colonies were harvested and suspended in 200 μl of sterile, DNase, RNase-free deionized water in Eppendorf tubes. Bacterial cells were inactivated by heating the tubes at 100°C for 10 min. Killed bacterial cells were centrifuged at 15,700 g for 10 min. The supernatant containing crude DNA template was pipetted into new sterile Eppendorf tubes discarding these Diment, Ouahran-Bettache et al. [25].

*From tissue specimens*

Twenty-five mg of fresh animal tissue was transferred into a microcentrifuge tube and 200 μl of GST Buffer and 20 μl of Proteinase K were added and then vortex thoroughly then incubated at 60°C overnight or until the sample lysate becomes clear. The tubes were centrifuged for 2 min at 12,000 rpm. the supernatant was carefully transferred to a new 1.5 ml microcentrifuge tube. Extraction was carried out according to the instruction manual of gSyn™ DNA extraction kit, Geneaid (New Taipei City, 22180 Taiwan, Cat. No. GS 100).

**Table-1:** Primer sets used for PCR.

| Primer          | Sequence (5’–3’)          | Amplicon size (bp) | DNA targets                                      |
|-----------------|---------------------------|---------------------|--------------------------------------------------|
| BMEI0535f       | GGC-CAT-TCT-TGG-ATG-AA    | 450                 | Immunodominant antigen, gene bp26                |
| BMEI0535r       | CGC-AGG-CGA-AAA-CAG-CTA-TAA | 587             | Erythritol catabolism, gene eryC (D-erythrulose-1-phosphate dehydrogenase) |
| BMEI0428f       | GCC-GCT-ATT-ATG-TGG-ACT-GG | 1682                | Glycosyltransferase, gene wboA                   |
| BMEI0428r       | AAT-GAC-CTT-ACG-GTC-ATT-CG | 1071                | Outer membrane protein, gene omp31               |
| BMEI0998f       | ATC-CTA-TGG-CCC-CGA-TAA-AGG | 272                 | ABC transporter binding protein                  |
| BMEI0997r       | GCT-TGG-CAT-TCT-TGG-ATG-AGC | 218                 | Ribosomal protein S12, gene rpsL                |

PCR=Polymerase chain reaction

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PCR assay

Universal PCR was performed for molecular identification of Brucella spp. in DNA extracts from Brucella cultures and lymph nodes at the genus level according to Garcia-Yoldi et al. [26] using Primer sequences targeting Erythritol catabolism, gene eryC (D-erythulose-1-phosphate dehydrogenase), and Primer sequences for amplification of target gene (Immunodominant antigen, gene bp26) (Table-1). On the other hand, Bruce-ladder multiplex PCR was carried out for molecular identification of Brucella in DNA extracts from Brucella cultures at the species level using five primers, table according to Garcia-Yoldi et al. [26], using IN gene Bruce-ladder, IN gene Bruce-ladder VR: Batch No 180515, Ingenasa, Madrid, Spain (Table-1). The PCR amplification was carried out using Labnet® Multigene Gradient thermal cycler, Catalog TC9600-G- 230V (Labnet International, Inc. Edison, NJ, USA). The cycling conditions were 4 min at 95°C for initial heating, 35 cycles of 45 s at 94°C, 45 s at 60°C, 60 s at 72°C, and final extension for 7 min at 72°C. The PCR amplicons were analyzed by running 10 μl of the PCR products in 1% agarose gel stained with ethidium bromide (0.5μg/mL). Thereafter, gels were photographed under UV illumination using gel documentation and analysis system. Brucella species was determined according to the molecular size of the amplified products using DNA ladder (100 bp and 1 kb), (Biomatik R Code No. M7123 and M7508), Biomatik Corporation, Ontario, Canada.

B. melitensis positive control was used as control positive in universal PCR. B. abortus RB51 Bruce-ladder kit control, B. suis Bruce-ladder kit control, B. melitensis Rev1 Bruce-ladder kit control, and B. melitensis positive control were used as positive control in Bruce ladder PCR.

Pathological examination

Lymph nodes of all the bacteriologically positive six buffalo-cows from which Brucella isolated were fixed in 10% neutral buffered formalin for about 2 days. The specimens were processed by paraffin embedding method, sectioned 5-7 μm and stained with hematoxylin and eosin according to Bancroft and Gamble [27] for histopathological examination. In addition, Masson’s trichrome stain was used to confirm the presence of collagen fibers.

Statistical analysis

Statistical analysis was carried out using the 95% confidence interval for the population proportion according to Petrie and Watson [28].

Results

Serological examination using BPAT, RBT, and CFT of 750 buffalo-cows revealed a prevalence of 79 (10.53%) in the first examination followed by 11 (1.64%), 52 (7.88%), 8 (1.32%), and 3 (0.50%) at the 2nd, 3rd, 4th, and 5th examinations, respectively (Table-2). Successive serological testing continued every 3 weeks for detection of animals which are still incubating the disease with slaughtering of positive animals detected after each testing according to the roles of the GOVS. The first negative test for the remaining animals (N 597) was at the 6th examination.

Table-2: Successive serological examinations of Brucella-infected buffalo farm.

| Examination       | Number of examined buffaloes | Positive (%) | 95% CI |
|-------------------|-------------------------------|--------------|--------|
| 1st (September 24)| 750                           | 79 (10.53)   | (0.083, 0.126) |
| 2nd (October 16) | 671                           | 11 (1.64)    | (0.006, 0.025) |
| 3rd (November 8) | 660                           | 52 (7.88)    | (0.058, 0.098) |
| 4th (November 30)| 608                           | 8 (1.32)     | (0.004, 0.022) |
| 5th (December 23)| 600                           | 3 (0.50)     | (-0.004, 0.010) |
| 6th (January 16) | 597                           | 0 (0.00)     | 0      |
| 7th (February 8) | 597                           | 0 (0.00)     | 0      |
| 8th (March 4)    | 597                           | 0 (0.00)     | 0      |

Total serologically positive animals 153 (20.4%). CI=Confidence interval

Table-3: Typing of six Brucella isolates recovered from buffaloes at species and biovar level.

| Brucella isolates CO₂ H₂S Urease Growth on dyes Thionin Fuchsin Lysis by phage Monospecific sera Conclusion |  |
|------------------------------------------------|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| Field isolates (6)                               | - | - | + in 20 h | + | + | + | - | - | + | + | + | - | B. melitensis 3 |
| Reference strains                                |  |
| B. melitensis Ether                              | - | - | + in 18-24 h | + | + | + | - | - | + | + | + | - | B. melitensis 3 |
| B. abortus544                                    | - | + | + in 2 h | - | - | + | + | + | + | + | - | B. abortus 1 |
| B. suis1330                                      | - | +++ + + in <15 min | + | + | - | - | + | - | + | + | - | B. suis 1 |

RTD=Routine test dilution, Tp=Tbilisi, Iz1=Izatnagar, a=1:50000, b=1:100000, A=Anti Brucella abortus, M=Anti Brucella melitensis, R=Rough Brucella antiserum
followed by the 7th and 8th negative examinations where the herd was considered free from *Brucella* infection. Quarantine measures were released because animals passed three consecutive negative tests at 21 days intervals. A total of 153 (20.4%) serologically positive animals were detected in this study. This program continued for about 6 months until all positive cases were detected and eliminated by slaughtering.

Bacteriological examination of different lymph nodes of 11 seropositive buffalo-cows revealed isolation of 6 (54.55%) *Brucella* isolates. Cultures were smooth, and colonies were elevated, transparent, convex, with intact borders, and brilliant surface and have a honey color under transmitted light. According to the results of biochemical tests, requirement for additional atmospheric 10% CO₂, production of hydrogen sulfide gas, production of urease, growth on media containing the inhibitory dyes thionin and fuchsin, agglutination with polyclonal monospecific antisera A and M and R, and phage typing using Tb and Izatnagar (Iz) summarized in Table-3, all the six isolates were typed as *B. melitensis* biovar 3 (Table- 3).

**PCR assays**

Using primer sequences targeting Erythritol catabolism, gene *eryC* (D-erythrulose-1-phosphate dehydrogenase), PCR has amplified the fragment 587 bp, Figure-1a confirming the presence of *Brucella* on genus level. Using primer sequences targeting immunodominant antigen, gene *bp26PCR* has amplified the fragment 450 bp, Figure-1b confirming the presence of *Brucella* on genus level. Bruce-ladder multiplex PCR has amplified three fragments of 587 bp, 1071 bp, and 1682 bp, sizes confirming the presence of *B. melitensis* (Figure-1c). On the other hand, Bruce-ladder PCR of *B. melitensis* Rev1 vaccine strain (reference Kit) amplified four fragments of 218 bp, 587 bp, 1071 bp, and 1682 bp sizes (Figure-1c). Rev1 strain only revealed the 218 bp, produced by the BMEI0752 primer pair. This primer pair detects a unique point mutation in the rpsL gene, coding for the ribosomal protein S12 of the vaccine strain *B. melitensis* Rev-1 and responsible for streptomycin resistance of Rev-1[29]. *B. suis* (reference Kit) amplified fragments of 272 bp, 587 bp, 1071 bp, and 1682 bp sizes (Figure-1c) and *B. abortus* Bruce-ladder kit control amplified fragments of 587 bp and 1682 bp sizes (Figure-1c). *B. abortus* RB51 vaccine strain (reference Kit) showed two amplicons of 587 bp and the specific additional band 2524 bp (Figure-1c) which confirms the results reported by López–Goni et al. [30] who stated that *B. abortus* RB51 can be distinguished by a specific additional 2,524-bp fragment.

Histopathological examination of *Brucella*-infected lymph nodes revealed massive rarified and depleted lymphoid areas of both sub-capsular and deep cortical lymphoid follicles (Figure-2a-e). Few numbers of neutrophils, large number of lymphocytes associated with histiocytes or macrophages in the form of diffuse loose granulomatous reaction with lack of giant cells were observed (Figure-3a and b). Fatty infiltrates detected in *Brucella* infected lymph nodes, either discrete multiple variable sized fat globules or

![Figure-1: (a) Universal polymerase chain reaction (PCR) using primer sequences targeting Erythritol catabolism, gene *eryC* (Lane 1: 100 bp DNA ladder, Lane 2, 3, 4, 5: *Brucella* cultures (587 bp), Lane 6: *Brucella* lymph node DNA extract (587 bp), Lanes 7, 8: *Brucella melitensis* positive control (587 bp), Lane 9: *Brucella abortus* S19 control). (b): Universal PCR using primer sequences targeting immunodominant antigen, gene *bp26* (Lane 1: 100 bp DNA ladder, Lane 2, 3, 4, 5: *Brucella* cultures (587 bp), Lane 6: *Brucella* lymph node DNA extract (587 bp), Lanes 7, 8, 9: *Brucella melitensis* positive control (587 bp), Lane 9: *Brucella abortus* S19 control). (c): Multiplex Bruce-ladder PCR of *Brucella* isolates (Lane 1: DNA ladder, Lane 2: *B. abortus* Bruce-ladder kit control (587 bp and 1682 bp), Lane 3: *B. abortus* RB51 Bruce-ladder kit control (587 bp and 2524 bp), Lane 4: *Brucella suis* Bruce-ladder kit control (272 bp, 587 bp, 1071 bp, and 1682 bp), Lane 5: *B. melitensis* Rev1 Bruce-ladder kit control (218 bp, 587 bp, 1071 bp, and 1682 bp), Lane 6: *B. melitensis* positive control (587 bp, 1071 bp, and 1682 bp), Lanes 7, 8: *B. melitensis* cultures (587 bp, 1071 bp, and 1682 bp).]
single large globule (Figure-4a-c). Endothelial hyperplasia and hypertrophy of tunica media and adventitia of lymphoid medullary blood capillaries with diffuse collagen deposition either in a diffuse manner or localized around blood capillaries as well as few numbers of neutrophils were observed (Figure-5a). Massive areas of necrosis were replaced by single and/or multiple eosinophilic structures less masses using hematoxylin and eosin and green in color using Masson’s trichrome stain till the total lack of the germinal center (Figure-5b). Perivascular edema (Figure-6) could be detected.

Discussion

The seroprevalence of anti-Brucella antibodies in 750 buffalo-cows using BPAT, RBT, and CFT was 10.53% (n=79) in the first examination. Such high prevalence may be due to exposure of the highly susceptible non-vaccinated buffalo-cows to Brucella infection through the introduction of an infected animal which is considered an extremely important risk
factor that can lead to spread of the infection to the whole herd. Removal of infected animals for slaughtering failed to stop the spreading of the disease, as the sero testing continued to detect positive animals in the subsequent testing for over 6 months where at last 597 buffalo-cows proved serologically negative for three successive tests with 3 weeks intervals. This can be attributed to a husbandry system favoring mixed populations of different ages, sex, aborted and pregnant, and lack of calf hood vaccination. Animals incubating the disease and latent infections as it has been estimated by Nielsen and Duncan [3] that about 20% of calves born by infected heifers could be found persistently infected with brucellae may play a role in the failure of the control program. In addition, exposure of susceptible animals to uterine and vaginal discharges of infected animals that contain a huge number of bacteria is an important risk factor for disease transmission between infected and susceptible animals as reported by Holt et al. [31]. Infected tissues and milk are the main sources of environmental contamination that spreads the disease horizontally [9]. Of extremely important consideration is the possible long interval between *Brucella* infection and development of clinical signs. It is reasonable to say that the current results indicate an outbreak of *Brucella* infection. Efforts of the veterinary authorities to control brucellosis in Egypt through the organized “test and slaughter program” failed to eliminate the disease or reduce the prevalence of the disease as the prevalence of animal brucellosis in Egypt in the past 20 years ranged from 0.33%-1.32% according to the official reports of the GOVS, Egypt [21]. Nearly similar prevalence was recorded by Abd-El Halim et al. [19] in Assiut, Egypt who recorded the prevalence of the disease in buffaloes as 12.9% with failure of control program of the GOVS. Lack of surveillance program to identify infected animals/herds and absence of control of the movement of infected animals to prevent re-introduction of the disease which is considered the major risk factor, especially in Egypt [26], are contributing factors.

Eradication of brucellosis by test and slaughter seems unfeasible in developing countries because of limited resources to compensate farmers whose animals are slaughtered during such screening programs [32].

*Brucella* organisms were isolated from different lymph nodes of six seropositive buffalo-cows out of 11. Failure of isolation from the other 5 positive serological cases may be due to a low number of viable *Brucella* organisms in the tissue sample or due to contamination with other bacteria and the fastidious nature of *Brucella* organisms [33]. Typing of the six *Brucella* isolates involved a combination of growth characteristics including colonial morphology, oxidase, urease, CO2 requirement, H2S production, growth in the presence of fuchsin and thionin, lysis by Tbilisi and Izatnagar, Iz, bacteriophages and agglutination with monospecific A and M anti-sera that confirmed the presence of *B. melitensis* biovar 3.

*B. melitensis* biovar 3 was considered as the prevalent *Brucella* type in Egypt [11-15]. In addition, Abd-El Halim et al. [19] in Assiut, Egypt, isolated the same brucella species and biovar isolated in this study; *B. melitensis* biovar 3 from 3 aborted buffaloes out of 87 (3.45%). Primarily, *B. melitensis* infects small ruminants. Transmission to large ruminants may be the outcome of close contact between sheep, goats, cattle, and buffalo which is considered by Holt et al. [31] as the main risk factor of transmission of *B. melitensis* biovar 3 to cattle and buffalo in Egypt where *B. melitensis* biovar 3 is the prevalent type in both large and small ruminants.

Using primer sequences targeting Erythritol catabolism, gene *ervC* (Derythulose-1-phosphate dehydrogenase), PCR has amplified the fragment 587 bp, Figure-1a confirming the presence of *Brucella* on genus level. Interestingly, the PCR testing *B. abortus* S19 DNA did not produce the 587-bp fragment common to all *Brucella* strains as it is the only *Brucella* strain unable to oxidize erythritol [34]. The preferential utilization of erythritol rather than glucose is characteristic for pathogenic *Brucella* strains [35]. Using primer sequences targeting immunodominant antigen, gene *hp26PCR* has amplified the fragment 450 bp, Figure-1b confirming the presence of *Brucella* on genus level. These assays are designed to exploit a single unique genetic locus. They are useful for identification when species is not critical [29].

Concerning molecular characterization of *Brucella* isolates on the species level, Bruce-ladder multiplex PCR has amplified three fragments of 587 bp, 1071 bp, and 1682 bp, sizes Figure-1c confirming the presence of *B. melitensis*. On the other hand, Bruce-ladder PCR of *B. melitensis* Rev1 vaccine strain has amplified four fragments of 218 bp, 587 bp, 1071 bp, and 1682 bp sizes. Rev1 strain only revealed the 218 bp, produced by the BMEI0752 primer pair. This primer pair detects a unique point mutation in the *rpsL* gene, coding for the ribosomal protein S12 of the vaccine strain *B. melitensis* Rev-1 and responsible for streptomycin resistance of Rev-1, [30]. *B. suis* has amplified fragments of 272 bp, 587 bp, 1071 bp, and 1682 bp sizes and *B. abortus* Bruce-ladder kit control has amplified fragments of 587 bp and 1682 bp sizes. *B. abortus* RB51 vaccine strain (Figure-1c) showed two amplicons of 587 bp and the specific additional band 2524 bp which confirms the results reported by López–Goni et al. [36] who stated that *B. abortus* RB51 can be distinguished by a specific additional 2,524-bp fragment. This can be explained after Venmulapalli et al. [37] who demonstrated that the *whoA* gene encoding a glycosyltransferase, an enzyme essential for the synthesis of O antigen, is disrupted by an IS711 element (an insertion sequence) in *B. abortus* RB51 and computer analysis of the nucleotide sequence from RB51 revealed that the *whoA* gene

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was interrupted by an 842-bp fragment that resulted in development of the 2524 fragment (1682+842=2524).

Such multiplex Bruce-ladder PCR assay proved capable to differentiate Brucella species as it is directed toward genetic loci that are variable among species [29]. Differential PCR-based assays are particularly useful for epidemiological traceback. The major advantage of PCR assays is the time taken compared to conventional methods which require several days to isolate and identify Brucella organisms [38,39].

Histopathological examination of Brucella-infected lymph nodes revealed massive rafified and depleted lymphoid areas of both subcapsular and deep cortical lymphoid follicles. Similar findings were reported by Ahmed et al. [40] who reported lymphoid depletion in the white pulp of spleen in buffaloes naturally infected with B. melitensis. Moreover, Maria-Jesus et al. [41] described mild lymphoid depletion in the splenic nodules in mice experimentally infected with virulent Brucella organisms. On the other hand, this finding comes in contrast with Barrionuevo et al. [42], Jubb et al. [7], and Pollak et al. [43] who found that lymphoid hyperplasia was the predominant feature in case of brucellosis. In addition, Xavier et al. [44] reported variable degrees of lymphoid hyperplasia associated with neutrophilic and histiocytic infiltrates in the medulla and paracortex in Brucella infected bovine internal iliac lymph nodes and Meador et al. [45] described lymphofollicular hyperplasia, sinusoidal histiocytosis, and medullary plasmacytosis in bovine supramammary lymph nodes. Interestingly, Neta et al. [46] reported that Brucella infection resulted in lymphoid hyperplasia in bovine lymph nodes and spleen, and lymphoid depletion in the thymus and Hosein et al. [47] reported hyperplasia in the white pulp of the spleen of most examined Brucella infected cows and depletion in the white pulp of spleen and the cortex of lymph nodes of some cows. Cheville et al. [48] explained lymphoid depletion as a result of immunodeficiency due to the multiplication of Brucella in the lymphocytes and macrophages of infected animals. The obtained results revealed very few numbers of neutrophils, a large number of lymphocytes associated with histiocytes or macrophages in the form of a diffuse loose granulomatous reaction with lack of giant cells. Such results have been supported by those of Barrionuevo et al. [42] and Pollak et al. [43] who attributed the presence of a population of monocytes to the secondary internalization of the bacteria.

The chronic nature of brucellosis has been confirmed in this study as indicated by the chronic vasculitis that was characterized by endothelial hyperplasia and hypertrophy of tunica media and adventitia of lymphoid medullary blood capillaries with diffuse collagen deposition either in a diffuse manner or localized around blood capillaries, as well as few numbers of neutrophils. Chronic granulomatous lymphadenitis observed in this study, coincide with the results obtained by Benitez et al. [49] who mentioned that collagen fibrotic proliferation could contribute to granuloma formation as the compact structure of granuloma successfully prevents the dissemination of the microorganisms. Perivascular edema may suggest the second episode of bacterial invasion to the tissue especially when associated with chronic lesions. Fatty infiltrates detected in Brucella infected lymph nodes, either discrete multiple variables sized fat globules or single large globule, may be attributed to gradual replacement of the lymph nodes parenchyma with fat. Similar findings were reported by Gonzales-Peramato et al. [50].

Conclusion

Freedom status from brucellosis in this study required 6 months which are considered long time allowing the spread of infection to other localities especially under unhygienic conditions, husbandry system favoring mixed populations of different ages, sex, aborted and pregnant, and the lack of controlled movement of animals. Therefore, effective control of animal brucellosis requires surveillance to identify infected animal herds, elimination of the reservoirs and vaccination of young heifers. B. melitensis biovar 3 is the cause of the Brucella outbreak in buffalo which still remains the prevalent type of Brucella in Egypt. The disease runs a chronic course allowing further spread of infection.

Authors’ Contributions

HIH designed the study, interpreted the results and reviewed the manuscript. AM performed the fieldwork and collected the samples. HMZ was responsible for the bacteriological identification of the Brucella isolates. AMSM, SR and BEM carried out the PCR assays and drafted the manuscript. NMS was responsible for the histopathological examination. All authors read and approved the manuscript.

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Competing Interests

The authors declare that they have no competing interests.

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