

Preparation of antigen: *M. bovis* (Egy Bu6-Dak-12) was grown in PPLO medium for 48 hours and harvested by centrifugation at 14000 r.p.m. for 20 minutes. Pellet was washed for three times with phosphate buffer saline (PBS) pH 7.2. The pellet is re-suspended in PBS and the protein concentration was estimated as described by [15]

ProteoJET Membrane Protein Extraction Kit : Fermentas, Cat.no. K0321,EU was used according to the manufacturer

Vaccine preparation: according to [14].

Vaccine evaluation:

Determination of Hydrophilic-Lipophilic Balance (HLB) value of the oil emulsion: -[16] and [17].

a-Physical evaluation

- Emulsion stability test: according to Specification WHO/M/13.R4 [18]
- Viscosity testing according to [19].
- Each 100 ml of vaccine containing 9.7 ml span 80 (Sigma), 59 ml paraffin oil (Sigma) as oil phase and 1.8 ml tween 80 (sigma) and 29.5 ml PBS (1mg/ml membrane protein) as aqueous phase.

b-Sterility testing (according to the Code of Federal Regulations "9 CFR"): For detection of Bacteria, Fungi and Mycoplasma contamination.

c-Safety and Potency test: Challenge test according to [11]:

Design of the laboratory experiment:

Four groups of New Zealand rabbits (9-10 weeks old) were weighing about 1.5 kg were housed separately and vaccinated 0.5 ml S/C. These groups represented as:

- 1) Group A (vaccinated/challenged): 4 rabbits were inoculated with membrane vaccine then boosting after 2 weeks and challenged 2 weeks later with 0.2 ml (10^8 cfu / ml) S/C of field *M. bovis* (Egy-Bu-6-DK-12).
- 2) Group B (vaccinated / not challenged): 4 rabbits were vaccinated 2 weeks later S/C with 0.5 ml *M. bovis* Vaccine then boosting after 2 weeks and not challenged. These were monitored for adverse effects (Safety test) and antibody response.
- 3) Group C (challenged): 4 rabbits were inoculated with 0.2 ml of *M. bovis* field isolate S/C after 2 weeks.
- 4) Group D (unvaccinated / not challenged): 4 rabbits as control negative group.

The membrane immune response in the vaccinated groups was detected using ELISA (Mycoplasma bovis ELISA kit, Bio-X Diagnostics, Belgium)

Field study and sample collection: Milk samples were collected every two weeks from all vaccinated and contact animals, which were subjected to ELISA, culture and PCR.

3. Results

3.1 Laboratory Experiment

For evaluation the potency and safety of the prepared membrane *M. bovis* vaccine, a laboratory experiment was carried out in rabbits. Group (A) showed the highest antibodies titer after challenge and extended to the end of the experiment, followed by groups B and C. While the control negative (group D) was the least.

Concerning culture and PCR, group C gave positive results one week after challenge and extended to the end of the experiment. While groups A, B and D were negative for culture and PCR. (Fig.1).

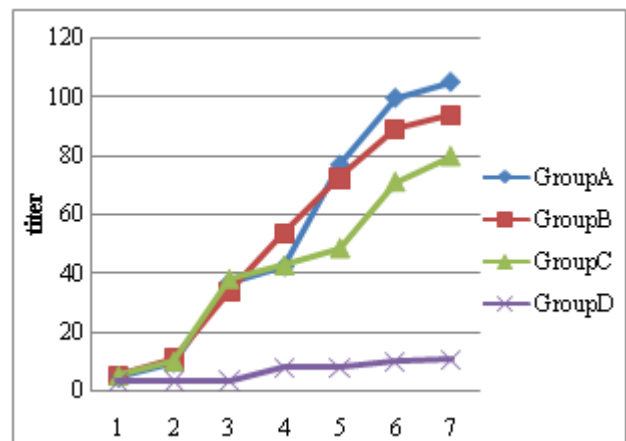


Figure 1: ELISA results of *M. bovis* membrane vaccine in rabbits

3.2 Field evaluation of the prepared *M. bovis* membrane vaccine

Two localities in Fayoum Governorate (farm 1 represented by 56 animals, farm 2 represented by 80 animals) three herds were chosen from each farm, and in Behera Governorate, one farm represented by one herd (32 animals) was chosen. All animals were tested for the detection of *Mycoplasma bovis* infection using ELISA, isolation, and PCR.

The animals proved to be negative for all used tests were chosen for vaccination. The dose was 4 ml injected S/C, followed by booster dose after two weeks. Milk samples were collected every two weeks for six months from all vaccinated and contact animals were subjected to ELISA, culture and PCR. At Fayoum (Fayoum 1&2) ELISA geometric mean titer started with (13.4-18.9) after two weeks post vaccination and increased to reach (153.24-195.2) at the end of experiment (24 weeks). While at Behera GMT was (23.56) after two weeks and increased to be (145.69) Table(1)

3.3 Isolation and PCR results of vaccinated herds:

The collected milk samples of the vaccinated herds were negative for isolation and PCR continued to give negative results till the end of the experiment, whereas the contact animals were positive for *Mycoplasma bovis* infection by isolation and PCR, indicating that the immune response due

to vaccination (vaccinated cattle were protected against the infection), Photo (1).

Table 1: Geometric mean titer of the herds from Fayoum and Behera Governorates vaccinated with local *M. bovis* vaccine

Governorate	Fayoum Governorate						Behera
	Farm1			Farm2			
	Herd1	Herd2	Herd3	Herd1	Herd2	Herd3	
Weeks							
2	14.98	14.98	13.4	18.92	15.03	18.5	23.56
4	46.34	38.9	44.19	22.2	25.32	27.2	32.64
6	62.11	58.49	53.29	26.5	30.03	44.4	45.66
8	71.93	62.11	77.14	36.66	36.17	50.15	52.51
10	83.9	71.93	95.58	50.4	49.21	59.21	64.66
12	105.84	105.84	115	70.66	68.37	61.73	75.68
14	110.66	110.66	130	75.34	72.15	72.74	98.20
16	113.27	115.27	134.5	80.53	81.65	83.61	95.56
18	132.45	133.29	160.21	105.23	113.32	105.2	107.71
20	164.3	165.2	175.2	132.1	123.52	121.5	119.36
22	180.85	171.32	190.42	140.1	139.2	135.35	128.35
24	189.27	175.2	195.2	155.21	162.47	153.24	145.69

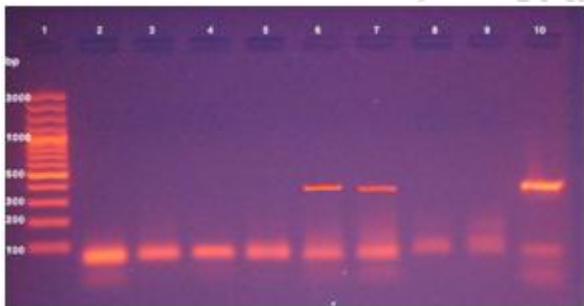


Photo 1: PCR results of vaccinated and non-vaccinated infected cows with *Mycoplasma bovis* mastitis
 Lane 1: 100 bp DNA ladder Lanes 2, 3, 4, 5 and 8: vaccinated cows
 Lanes 6 and 7: non-vaccinated infected cows with *M. bovis* mastitis (contact)
 Lane 9: control negative Lane 10: control positive

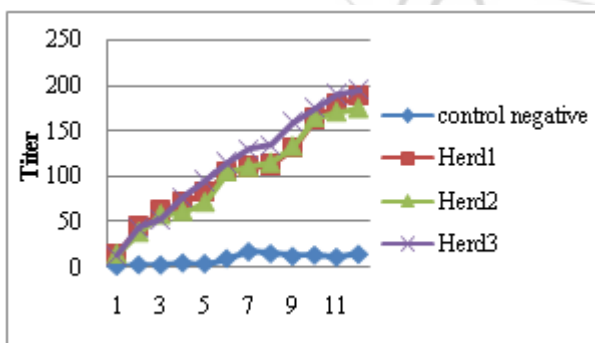


Figure 2: ELISA results of three herds at Fayoum Governorate (Farm 1) vaccinated with *M. bovis* vaccine

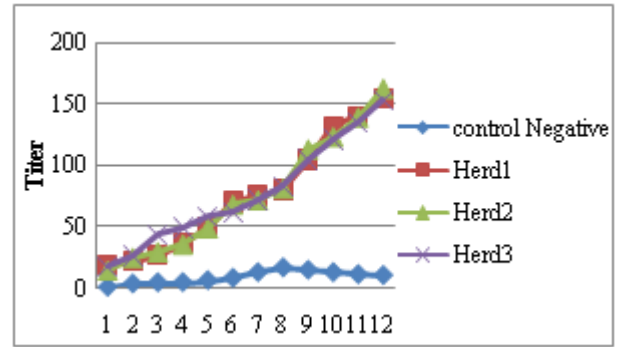


Figure 3: ELISA results of three herds at Fayoum Governorate (Farm 2) vaccinated with *M. bovis* vaccine

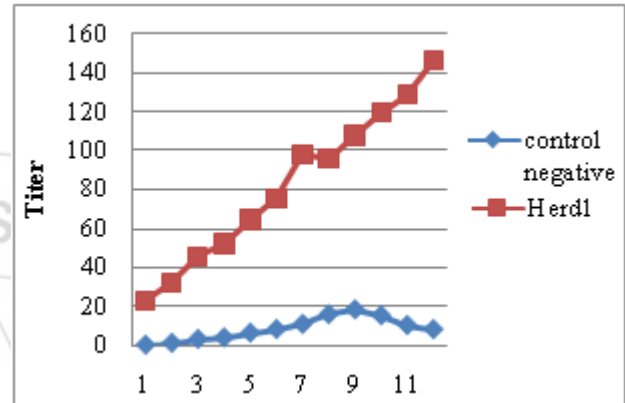


Figure 4: ELISA results of one herd at Behera Governorate vaccinated with *M. bovis* vaccine

4. Discussion

Mycoplasma bovis mastitis is a highly contagious disease that results in milk loss and culling of infected animals. Several species of *Mycoplasma* have been associated with mastitis (*M. bovis*, *M. californicum*, *M. canadense*, *M. bovigenitalium*, *M. alkalescens*, *M. arginini*, *M. bovis* and *M. dispar*) [20].

In Egypt *M. bovis* was isolated from clinical, subclinical and tank milk of bovine dairy herds at different Governorates [4], [5] and [6]. *M. bovis* strain isolated from clinical mastitis cases in buffaloes at Dakahlia Governorate identified using PCR, sequenced and submitted to GenBank. This isolate was used for preparation of *M. bovis* membrane vaccine which tested for safety and potency in rabbits showed the highest immune response detected by ELISA in Group A (vaccinated), followed by group B (vaccinated / challenged) and group C (challenged). These results agreed with [21] who mentioned that rabbits vaccinated with *M. bovis* and *M. bovigenitalium* saponized vaccine had the highest antibody titers. Also [10] concluded that calves vaccinated s/c with *M. bovis* bacterin and received two boosters at three week intervals showed lower respiratory disease (16.3%) when compared with control calves (27%).

In the present work, the prepared vaccine was evaluated in field study on bovine dairy herds in two governorates (Fayoum and Behera). Milk samples were collected from vaccinated and contact animals every two weeks and continued for 6 months.

M. bovis specific antibody levels detected by the indirect ELISA gradually increased after the booster dose to reach (198.2) in herd 3 (Farm1, Fayoum), while herds 2 (Farm1, Fayoum) gave a GM titer of (162.47), followed by the dairy farm at Behera Governorate (145.69) after 6 months. Culture and PCR were negative in the all vaccinated animals till the end of study. Also milk production was normal in quantity and quality without any pathological changes. On the other hand, the contact animals (infected with *M. bovis* mastitis) were positive for culture and PCR with pathological change in milk (secretion containing visible particles, seropurulent or aqueous suspension) and drop of milk yield.

[22] mentioned that vaccination with killed *M. bovis* systemically and in the mammary gland elicited local and systemic antibody. [23] indicated that saponin in combined vaccine can produce a specific humeral immune response to *M. agalactiae* over 6 months. Also [24] detected a high immune status with GMT 40.3, 3 weeks post-vaccination and has risen to 80.6 after 4 week and was maintained to 64 at the end of experiment.

Immunization of animals with high quality vaccines is the primary means of control for many animal diseases. In other cases, vaccines are used in conjunction with national disease control or eradication programs.

5. Conclusion

The present work aimed to prepare and evaluate *M. bovis* membrane vaccine prepared from local field isolate. Three farms at Fayoum and Behera Governorates were choice for the field study which extended to six months. The vaccine proves to be safe, efficacious and immunogenic.

6. Acknowledgment

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