A field study evaluating the humoral immune response in Mongolian sheep vaccinated against sheeppox virus

Petra Fay1 | Georgina Limon1 | Gerelmaa Ulziibat3 | Buyantogtokh Khanui3 | Odonchimeg Myagmarsuren2 | Gessica Tore1 | Bodisaikhan Khishgee3 | John Flannery1 | Batkhuyag Sandag4 | Batchuluun Damdinjav3 | Philippa M. Beard1,2

1 The Pirbright Institute, Pirbright, UK  
2 The Roslin Institute, Easter Bush, University of Edinburgh, UK  
3 State Central Veterinary Laboratory, Zaisan, Khan-Uul District, Ulaanbaatar, Mongolia  
4 The Mongolian General Authority for Veterinary Services, Ulaanbaatar, Mongolia

Correspondence  
Philippa M. Beard, The Pirbright Institute, Ash Road, Pirbright, GU24 0NF, UK. Email: Pip.Beard@pirbright.ac.uk

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Abstract  
Sheeppox is a transboundary disease of small ruminants caused by infection with the capripoxvirus sheeppox virus. Sheeppox is found in Africa, the Middle East and Asia and is characterized by fever, multifocal cutaneous raised lesions and death. Vaccination with live attenuated capripoxvirus (CPPV) strains is an effective and widely used strategy to control sheeppox outbreaks; however, there are few reports of post-vaccination field surveillance studies. This study used a commercially available enzyme-linked immunosorbent assay (ELISA) to examine quantitative and temporal features of the humoral response of sheep vaccinated with a live-attenuated CPPV strain in Mongolia. Four hundred samples were tested using the ELISA commercial kit, and a subset of 45 samples were also tested with a virus neutralization test (VNT). There was substantial agreement between the VNT and ELISA tests. Antibodies to CPPV were detected between 40 and 262 days post-vaccination. There was no significant difference between serological status (positive/negative) and sex or age; however, an inverse correlation was found between the length of time since vaccination and serological status. Animals between 90 and 180 days post-vaccination were more likely to be positive than animals greater than 180 days post-vaccination. Our results show that a commercial CPPV ELISA kit is a robust and reliable assay for post-CPPV vaccination surveillance in resource-restricted settings and provide temporal parameters to be considered when planning sheeppox post-vaccination monitoring programmes.

Keywords  
capripoxvirus, humoral immunity, mongolia, post-vaccination monitoring, poxvirus, sheeppox, sheeppox virus, virus neutralization assay

1 | BACKGROUND

Sheeppox (SP), goatpox (GP) and lumpy skin disease (LSD) are transboundary diseases caused by infection with viruses of the genus capripoxvirus (CPPV), namely sheeppox virus (SPPV), goatpox virus (GTPV) and lumpy skin disease virus (LSDV). All three viruses cause systemic disease in ruminants characterized by fever, multifocal cutaneous raised lesions and death. LSDV causes disease only in cattle,
while SPPV and GTPV cause disease in sheep and goats. The host preference of SPPV and GTPV varies with some isolates of SPPV causing disease only in sheep, some isolates of GTPV causing disease only in goats and some isolates of SPPV and GTPV causing disease in both sheep and goats (Babiuk et al., 2009). SP and GP are high consequence diseases. They reduce production of meat, milk, wool and cashmere and decrease the value of affected animals, therefore having a substantial negative effect on farmers’ livelihoods (Babiuk et al., 2008; Bolajoko et al., 2019; Garner et al., 2000; Limon et al., 2020). Furthermore, countries with endemic SP or GP face restrictions on trade of live animals and animal products (Tuppurainen et al., 2017).

Mongolia is a landlocked country located in central Asia bordered by Russia to the north and China to the south, east and west. Mongolia has one of the highest livestock per capita ratios in the world, with a human population of 3.2 million compared to 4.3 million cattle, 30.1 million sheep and 27.3 million goats (Mongolian Statistical Information Service, 2017). Since 1977, there have been three SP outbreaks in Mongolia: the 2006–2007 outbreak affecting five provinces (Beard et al., 2010), the 2013 outbreak affecting two provinces and the extensive 2015–2017 outbreak affecting eight provinces. During this latest, the Mongolian General Authority for Veterinary Services started a risk-based vaccination campaign as a control strategy. In 2016, a targeted post-vaccination surveillance programme was employed.

Vaccination is considered one of the most effective methods for control of CPPVs (Tuppurainen et al., 2017). Live-attenuated strains of SPPV and GTPV are the most common type of vaccine used against SP and GP; however, the duration of the humoral immune response following vaccination with live attenuated CPPV vaccines is poorly understood. Manufacturers often recommend annual vaccination regimes to maintain herd immunity; however, these recommendations are often based on research conducted for LSD in cattle and/or under controlled conditions (Boumart et al., 2016; Kali et al., 2019; Klement et al., 2018; Milovanovic et al., 2019). There is very little published research describing the humoral immune response following regional or national vaccination programmes to protect against SP or GP.

In this study, we used a subset of serum samples collected from sheep during post-vaccination surveillance in Mongolia to investigate the humoral immune response of vaccinated sheep following a risk-based SP vaccination campaign. We used the results to identify potential factors that might play a role in the detection of seroconversion and to assess the suitability of a commercial enzyme-linked immunosorbent assay (ELISA) test for post-vaccination monitoring in a non-endemic resource restricted setting.

2 METHODS

2.1 Vaccination

All animals were vaccinated with a live-attenuated capripoxvirus vaccine produced by Biocombinat SOI, in Mongolia. The sheeppox vaccine was the live-attenuated SPPV Perego strain (original date 1978), grown in primary lamb testis cell cultures. The vaccine was administered to sheep subcutaneously into a hairless skin area (fore-flank or tail patch) in compliance with Mongolian hygiene regulations. Mongolia does not carry out sheeppox vaccination programmes routinely; therefore, to the best of the authors’ knowledge this was the first sheeppox vaccine administered to the animals.

2.2 Sample collection

Blood samples from sheep were collected as part of the post-vaccination surveillance programme for SP implemented by the Mongolian General Authority for Veterinary Services in 2016. All provinces that were part of the vaccination programme in the country (n = 8; Figure 1) were part of the post-vaccination monitoring evaluation.

FIGURE 1 Geographical location of Mongolia (dark brown – top left map) and provinces where vaccination was conducted (in alphabetical order) (1) Dornold, (2) Dornogovi, (3) Dungovi, (4) Govisumber, (5) Khenthi, (6) Sukhbaatar, (7) Tuv and (8) Ulaanbaatar
Multistage sampling was used to select animals for testing. Briefly, in each province between one and three soums (or districts) were randomly selected. Within each soum, herds were randomly selected and 20 sheep in each herd were randomly selected for sampling. A total of 2000 samples were collected.

For each sample collected, herder location (province and soum name), name of the herder/owner, age and sex of the animal, date when the animal was last vaccinated for SP (according to the vaccination records) and date of sampling were recorded.

### 2.3 Sample processing

Blood samples were stored at 4°C to clot after which the separated serum was collected. Samples were stored at 4°C and transported to the State Central Veterinary Laboratory (SCVL) of Mongolia, in Ulaanbaatar, Mongolia, within two days post-collection. On receipt, serum samples were centrifuged at 3000 rpm for 10 min, serum supernatant collected, aliquoted, and stored at −20°C.

### 2.4 ELISA

From the 2000 samples collected as part of the post-vaccination monitoring, 400 samples were randomly selected for comparative serological testing by both SCVL in Mongolia and The Pirbright Institute in the UK using a commercially available ELISA (ID Screen® Capripox Double Antigen Multi-species ELISA kit; IDvet, Grabels, France) following manufacturer instructions. Aliquots of the same serum samples were tested singly at both the SCVL and Pirbright. The optical density readings were used to calculate the percentage of seropositivity (%SP) for each sample. A sample with a %SP value of ≥30% was considered positive.

### 2.5 Fluorescent virus neutralization test

The recombinant EGFP-095-LSDV Neethling virus was generated through insertion of the enhanced green fluorescent protein (EGFP) marker at the N terminus of the LSDV ORF 095, encoding for a putative core protein of the LSDV Neethling strain (GenBank: AF409138.1). BS-C-1 (ATCC® CCL-26™) cells were infected with the LSDV Neethling strain at a multiplicity of infection of 0.1. Two hours after infection, cells were transfected with a recombinant transfer plasmid carrying the EGFP sequence fused to the LSDV095 gene and flanked by an upstream and a downstream LSDV homology region of, respectively, 427 and 338 bp. Infected/transfected cells were incubated for about 5 days during which homologous recombination occurred between the LSDV DNA genome and the recombinant construct. After the incubation period, single EGFP positive BS-C-1 cells were sorted into pre-seeded 96-well plates twice using a BD FACSARia™ III sorter (BD Biosciences). The recombinant fluorescent virus was recovered from supernatants of infected cell lysates and separated from the parental virus through fivefold limiting dilutions of the viral suspension. The isolated recombinant EGFP-095-LSDV Neethling virus was then purified through a sucrose cushion to obtain a working stock to be used in the development of the fluorescent virus neutralization test (FVNT). Full genome sequencing of EGFP-095-LSDV Neethling was carried out to confirm the genetic modification (data not shown).

Forty-five serum samples were selected and tested for neutralizing antibodies by the FVNT. Samples were randomly selected based on the ELISA test results spread over time. Time between vaccination and sampling was re-categorized into (i) short (<98 days), (ii) medium (99–132 days) and (iii) long (>133 days), and %SP values were re-categorized into (i) low (%SP ≤ 0.50), (ii) medium (%SP 0.51 to 74.38) and (iii) high (%SP ≥ 74.39), using the first and the third quartile as cutoffs in both cases. Samples were then grouped into the nine categories and five samples selected from each category for testing by FVNT.

Briefly, Madin-Darby bovine kidney (MDBK) cells were seeded in 96-well tissue culture plates (Corning) at a cell suspension of 3 × 10^5 cells/mL and incubated overnight at 37°C in a 5% CO_2_ incubator. All sample and control sera were heat-inactivated at 56°C for 30 min and diluted 1:10 in culture medium (DMEM supplemented with 100 IU/mL penicillin, 100 µg/mL streptomycin and 2.5% heat-inactivated fetal bovine serum; Life Technologies). Twofold serial dilutions (1:10 to 1:1280) were then prepared from the test sera. To all serum samples, an equal volume of EGFP-095-LSDV Neethling virus (7 × 10^2 PFU/mL) was added, and samples incubated at 37°C for 1 h in a 5% CO_2 incubator. In addition to the serum controls, a cell and virus only control were included. All sample and control sera were tested in duplicate.

To corresponding wells, 150 µL of the test and control serum samples were added to prepared MDBK cells and plates incubated at 37°C in a 5% CO_2 incubator for 4 days. Fluorescent foci (indicative of cytopathic effect) were determined using a fluorescent UV light microscope (Olympus CKX53). The neutralizing antibody titre for each sample was determined as the highest dilution at which no foci were identified, indicative of complete neutralization.

### 2.5.1 Statistical analysis

Considering the FVNT test as the gold standard, diagnostic sensitivity (Dse) and specificity (Dsp) of the ELISA was calculated.

McNemar’s chi-squared test for paired data was used to assess whether there was a statistically significant difference in the proportion positive between (i) FVNT and ELISA test results from SCVL in Mongolia (n = 45), (ii) FVNT and ELISA test results from Pirbright in the UK (n = 45) and (iii) ELISA test results from SCVL in Mongolia and Pirbright in the United Kingdom (n = 400).

Descriptive statistics were obtained for all sheep tested in both laboratories (n = 400). Frequency distribution of values were explored for time between vaccination and sampling, and %SP values for the ELISA test considering the results from each lab separately. For samples in which the exact date of vaccination was not available and a range of potential time was given (e.g. May 2016 or between 15 May and 15 June), the midpoint date within this range was considered. The extent to which sex, age, time (in days) since vaccination and province were
associated with seroconversion (positive/negative) was determined using univariable mixed effects binomial models including herd as random effect. Age was re-categorized as ≤1 year and >1 year to consider that sheep aged more than 1 year might have been vaccinated more than once. Time since vaccination was re-categorized in to three categories (<90 days, 91–180 days and >180 days) based on the expected dynamics of the immune response (Haegeman et al., 2020; Kali et al., 2019; Milovanovic et al., 2019). Collinearity was assessed between all predictor variables for which \( p < 0.2 \) in the univariate analysis and, when present, only one of the variables was kept in the model. Multivariable analysis was used to assess the relationship between the individual predictor variables and the outcome, accounting for the potential confounding effect of other variables. The %SP values were used to classify animals as positive/negative. The analysis was conducted using the results from each laboratory separately.

Statistical analysis was performed in R.3.3.2 (R Development Core Team 2017) using packages EpiR, car and lme4.

### 3 RESULTS

Blood samples were collected between 23 and 30 August 2016 from 400 sheep that had all been vaccinated with a live attenuated capripoxvirus vaccine. The 400 samples came from sheep herds belonging to 77 herders from seven provinces. Two thirds of the sheep (266; 66.5%) were female and 134 (33.5%) were male. Age ranged between 1 and 9 years (median 4 years) with similar age distribution between male and females. Time between vaccination and sampling ranged from 40 to 243 days (median 116 days). Day of vaccination was not recorded in four animals.

The 400 serum samples were tested in both the SCVL (Mongolia) and Pirbright (UK) using the ID Screen® Capripox Double Antigen Multi-species ELISA kit (IDvet) following the manufacturer’s instructions, and the results from the two laboratories compared. A higher number of samples were classified as positive when tested in SCVL \( (n = 188; 47\%) \) compared to Pirbright \( (n = 165; 41.3\%) \), this difference was statistically significant \( (p = 0.02) \) and the agreement was moderate (Table 1).

The virus neutralization test (VNT) is considered the gold standard for assessing the level of protective immunity (neutralizing antibodies) to CPPV with high specificity (OIE, 2019). Therefore, 45 serum samples, representing a sample spread across time post-vaccination and ELISA test result, were tested for the presence of neutralizing antibodies using a FVNT. Out of the 45 samples selected to be tested by FVNT, 20 were deemed positive. Using the ELISA performed in SCVL (Mongolia), 22 of these 45 samples tested positive. Using the ELISA performed at Pirbright (UK), 18 tested positive (Tables 2 and 3). The range of values for those samples that were deemed positive are presented in Figure 2. There was a substantial agreement between the FVNT and ELISA tests regardless of the lab where the samples were tested (Kappa 0.82 and 0.73; Table 3). Considering FVNT as the gold standard, the Dse of the ELISA test was slightly better when used in Mongolia (0.90; 95% CI 0.66–0.98) than when used in the UK (0.85; 95% CI 0.61–0.96); while the Dsp was better in the UK (0.96; 95% CI 0.78–0.99) than in Mongolia (0.84; 95% CI 0.63–0.95). In other words, there were less false positives when the samples were tested in the United Kingdom and less false negatives when the samples were tested in Mongolia.

The 188 positive samples from the SCVL dataset are represented in Figure 3. Positive ELISA results were detected from day 40 to 243 post-vaccination. The ELISA dataset was used to identify factors which might influence seroconversion. An inverse correlation was found between the length of time since vaccination and serological status (Figure 3) with a statistically significant difference between short (up to 90 days) and long periods of time (>180 days) (Table 4; Supplementary material Tables S1 and S2). There was no significant difference between serological status (positive/negative) and sex or age (Table 4). Differences were found between serological status and province (Table 4); however, length of time since vaccination and province exhibited strong collinearity (Supplementary material Table S3) and therefore the univariate models were kept. The same patterns were found using ELISA results from both labs (Table 4).

### 4 DISCUSSION

This study examined the humoral immune response of sheep to vaccination with a live-attenuated capripoxvirus vaccine. Most importantly, the study evaluates sheep pox vaccination under field condition, therefore addressing a key gap in literature.

The inter-assay repeatability of the ELISA was examined by testing the same 400 samples using the same kit and protocol but in two different laboratories. The results from the two datasets differed slightly when compared, with a higher number of samples classified as positive when tested in SCVL compared to Pirbright. The differences between the ELISA results reported from the SCVL and Pirbright laboratories may be due to differences in equipment, users, environment (such as storage conditions of samples) or the quality control regimes in the two laboratories. Unfortunately, the limited volume of sera precluded the

### TABLE 1 Number of positive and negative results using ELISA in two labs (Mongolia and UK) (n = 400)

| ELISA UK | ELISA Mongolia | Number of negative samples | Number of positive samples | p value* | Kappa test |
|----------|----------------|------------------------------|---------------------------|----------|------------|
| Number of negatives samples | 178 | 57 | .02 | 0.54 |
| Number of positive samples | 34 | 131 |

*McNemar’s chi squared test for paired data.

## References

- Haegeman et al., 2020
- Kali et al., 2019
- Milovanovic et al., 2019
- OIE, 2019
| Days post-vaccination | ELISA Mongolia lab (%S/P) | ELISA Mongolia lab | ELISA UK lab (%S/P) | ELISA UK lab | FVNT Titre | FVNT |
|-----------------------|---------------------------|-------------------|---------------------|--------------|-----------|------|
| 40                    | 41.93 Positive            | Positive          | 83                  | Positive     | 80        | Positive |
| 40                    | 0                         | Negative          | 2.09 Negative       | 0            | 80        | Negative |
| 40                    | 3.01 Negative             | Negative          | 0                   | Negative     | 0         | Negative |
| 40                    | 5.32 Negative             | Negative          | 2.58 Negative       | 0            | 0         | Negative |
| 74                    | 45.7 Positive             | Positive          | 0.48 Negative       | 0            | 0         | Negative |
| 75                    | 17.2 Negative             | Negative          | 0                   | Negative     | 0         | Negative |
| 76                    | 1.61 Negative             | Negative          | 0                   | Negative     | 0         | Negative |
| 76                    | 9.07 Negative             | Negative          | 1.25 Negative       | 0            | Negative |
| 76                    | 29.23 Negative            | Negative          | 0                   | Negative     | 0         | Negative |
| 77                    | 232.86 Positive           | Positive          | 210.9 Positive      | 480          | Positive  |
| 77                    | 160.18 Positive           | Positive          | 32.81 Positive      | 120          | Positive  |
| 80                    | 7.81 Negative             | Negative          | 0.55 Negative       | 0            | 0         | Negative |
| 80                    | 117.84 Positive           | Positive          | 325.5 Positive      | 320          | Positive  |
| 80                    | 156.96 Positive           | Positive          | 49.47 Positive      | 80           | Positive  |
| 81                    | 83.47 Positive            | Positive          | 82.25 Positive      | 120          | Positive  |
| 92                    | 26.77 Negative            | Negative          | 1.42 Negative       | 0            | 0         | Negative |
| 92                    | 72.25 Positive            | Positive          | 143.3 Positive      | 30           | Positive  |
| 92                    | 62.25 Positive            | Positive          | 0.32 Negative       | 0            | 0         | Negative |
| 102                   | 1.91 Negative             | Negative          | 3.7 Negative        | 0            | 0         | Negative |
| 107                   | 46.07 Positive            | Positive          | 0                   | Negative     | 0         | Negative |
| 110                   | 105.24 Positive           | Positive          | 54.56 Positive      | 30           | Positive  |
| 114                   | 3.63 Negative             | Negative          | 0.26 Negative       | 0            | 0         | Negative |
| 114                   | 24.4 Negative             | Negative          | 0                   | Negative     | 0         | Negative |
| 119                   | 33.47 Positive            | Positive          | 0                   | Negative     | 10        | Positive |
| 121                   | 14.76 Negative            | Negative          | 22.32 Negative      | 20           | Positive  |
| 122                   | 0.11 Negative             | Negative          | 139.5 Positive      | 60           | Positive  |
| 122                   | 34.83 Positive            | Positive          | 181.6 Positive      | 160          | Positive  |
| 122                   | 173.3 Positive            | Positive          | 400.1 Positive      | 1280         | Positive  |
| 130                   | 88.57 Positive            | Positive          | 130.7 Positive      | 60           | Positive  |
| 131                   | 0                         | Negative          | 3.76 Negative       | 0            | Negative  |
| 131                   | 0                         | Negative          | 0                   | Negative     | 0         | Negative |
| 131                   | 158.88 Positive           | Positive          | 252.2 Positive      | 30           | Positive  |
| 132                   | 0                         | Negative          | 6.6 Negative        | 0            | Negative  |
| 133                   | 50.27 Positive            | Positive          | 101.2 Positive      | 160          | Positive  |
| 135                   | 40.82 Positive            | Positive          | 46.3 Positive       | 0            | 0         | Negative |
| 135                   | 0                         | Negative          | 0                   | Negative     | 0         | Negative |
| 193                   | 43.25 Positive            | Positive          | 3.49 Negative       | 20           | Positive  |
| 229                   | 132.41 Positive           | Positive          | 169.2 Positive      | 80           | Positive  |
| 229                   | 132.19 Positive           | Positive          | 185.3 Positive      | 60           | Positive  |
| 229                   | 0                         | Negative          | 2.07 Negative       | 0            | 0         | Negative |
| 229                   | 0                         | Negative          | 0.03 Negative       | 0            | 0         | Negative |
| 234                   | 0                         | Negative          | 0.19 Negative       | 0            | 0         | Negative |
| 234                   | 0                         | Negative          | 1.53 Negative       | 0            | 0         | Negative |
| 236                   | 121.73 Positive           | Positive          | 255 Positive        | 640          | Positive  |
| 243                   | 0                         | Negative          | 0.3 Negative        | 0            | 0         | Negative |

Bold values indicate positive findings.
TABLE 3 Number of positive and negative results using FVNT and ELISA in two labs (Mongolia and UK) (n = 45)

|                      | ELISA Mongolia   | ELISA lab UK     |
|----------------------|------------------|-------------------|
|                      | Number of        | Number of         |
|                      | negatives samples| positive samples  | p value*   | Kappa test |
| FVNT                 |                  |                   |           |            |
| Number of negatives  | 21               | 4                 | .68       | 0.73       |
| Number of positives  | 2                | 18                |           |            |
|                      |                  |                   |           |            |

*McNemar’s chi squared test for paired data.

FIGURE 2 Frequency distributions of values among samples, belonging to the 45-sample subset, that were deemed positive by each test (a); and values among those samples with contradictory results between the two labs considering all samples tested (b). Red vertical lines show cut-off value used to classify samples as positive/negative (ELISA S/P30% and FVNT 1:10).

The FVNT was used as a gold standard to determine the Dse and Dsp of the ELISA. The two sets of ELISA results were in substantive agreement with the FNVT across the 45 sera tested. The values calculated using data from SCVL and Pirbright were Dsp 84% and 96%, and Dse 90% and 85%, respectively. These are similar to parameters previously reported for ELISA conducted in cattle with LSD (Dsp 87% and Dse 91%) (Milovanovic et al., 2019) but lower than the Dsp data reported by IDvet (Dsp 99.7%) (Haegeman et al., 2019). This difference in Dsp estimate may be attributed to the origin of the samples used (field versus experimental), the level of antibodies present in the sample and the sample quality (e.g. haemolysis) (Bowden et al., 2009; Haegeman et al., 2020).
Occasional disagreements between the ELISA and FVNT were not unexpected as they are fundamentally different tests that detect different subsets of antibodies. The ELISA detects antibodies against immunogenic CPPV antigens, whereas the FVNT detects antibodies with the ability to neutralize CPPV. Sporadic disagreement between CPPV serological tests, particularly at early times post-infection or post-vaccination, has been reported previously (Haegeman et al., 2019; Milovanovic et al., 2019). This should be a factor to consider when designing the timing of a CPPV ELISA-based post-vaccination testing programme.

ELISAs, Immunoperoxidase Monolayer Assays, virus/serum neutralization tests and immunofluorescent antibody tests have all been published for detecting humoral response to CPPV and recently comprehensively reviewed (Haegeman et al., 2019). The ELISA is the easiest and cheapest of the four techniques and the most suitable for inter-laboratory standardization. The IDvet ELISA has been validated in a number of peer-reviewed publications, showing good concordance with the VNT (Haegeman et al., 2020; Milena et al., 2019; Milovanovic et al., 2019; Moller et al., 2019). Most of these studies use sera from cattle vaccinated or infected with LSDV. This study reports the use of the IDvet ELISA with CPPV vaccine field sera, demonstrating that it provided a high throughput means of assessing the overall CPPV antibody status of a large cohort of animals, therefore broadening the potential utility of the test.

Examination of factors which might influence seroconversion in the sheep identified no significant difference between serological status (positive/negative) and sex or age but did find an inverse correlation between serological status and the length of time since vaccination. Differences were also found between serological status and province, with the data suggesting this is due to timing of vaccination in different provinces. There were no important different geographic, environmental, or cultural features or husbandry practices identified between the provinces, which are all in the eastern and central region of Mongolia.

A temporal response similar to that reported previously (Abdelwahab et al., 2016; Boumart et al., 2016; Kali et al., 2019; Milovanovic et al., 2019) was seen in our study, with a correlation found between length of time since vaccination and serological status as measured by commercial ELISA. The ELISA detected seroconversion between 40 and 76 days post-vaccination albeit with borderline positive %SP values at these time points. Animals categorized into medium time period post-vaccination (between 90 and 180 days) were more likely to be positive and animals categorized into the group with long time period (>180 days) were less likely to be positive. Antibodies against CPPV were still detected in this study in four samples at 236 days post-vaccination. A limitation of this study is that sheep were only tested at one point in time. In the future, longitudinal studies with fewer animals tested but at frequent and uniform intervals post-vaccination, particularly at early timepoints, would be beneficial and should be implemented in future post-vaccination monitoring programmes.

Correlating antibody levels with protection against viral challenge requires further study. Antibodies alone are known to provide protection against poxviral disease including SP (Kempe et al., 1961; Law et al., 2005; Kitching, 1986), and levels of poxvirus antibodies, measured either by ELISA or neutralization assay, are often used as a correlate of protective immunity in people and animals. However, the level of antibodies that confers protection against poxviruses is unknown. One study found people with pre-existing neutralizing titres <1:32 against vaccinia virus were more susceptible to smallpox infection than those with antibody titers ≥1:32 (Mack et al., 1972). A neutralization index (calculated as the log titre difference between the titre of the virus in the negative serum and in the test serum) of ≥1.5 is considered positive for LSDV, SPPV and GTPV (OIE, 2019) although there is no data to link this index with protection from challenge. Importantly, a low antibody response by ELISA or neutralization tests post-vaccination may not necessarily mean absence of protection as (i) there may be sufficient memory B cells present to provide a rapid anamnestic antibody response and therefore protection post challenge (Crotty et al., 2003), (ii) it is likely only a low amount of neutralizing antibodies is required for protection (Boshra et al., 2015; Hammarlund et al., 2003) and (iii) cell-mediated immunity or non-neutralizing antibodies may provide protection. The levels of antibodies detected at later time points in this study (>180 days) were lower than earlier time points. While this indi-

![FIGURE 3 Median %SP value using results from SCVL (black line), first and third quartile (grey dashed lines) and days post-vaccination. Red horizontal line represent %S/P cut-off used to classify samples as positive/negative. Blue vertical dashed lines are cut-off used to classified days post-vaccination in categories (<90 days, 91–180 days and > 180 days) based on the expected dynamics of the immune response](image-url)
### Table 4: Distribution of factors considered for sheep pox serological status after vaccination following univariable analysis. All models include herder as random effect.

| Variable          | Mongolia laboratory |                  | UK laboratory |                  |
|-------------------|---------------------|------------------|---------------|------------------|
|                   | Number of negative (%) | Number of positive (%) | Estimate | Standard error | p value | Number of negative (%) | Number of positive (%) | Estimate | Standard error | p value |
| **Sex**           |                     |                  |               |                  |         |                      |                  |         |                  |         |
| Female            | 142 (67.0)          | 124 (66.0)       | Ref.          |                  |         | 155 (66.0)            | 111 (67.3)       | Ref.    |                  |         |
| Male              | 70 (33.0)           | 64 (34.0)        | −0.15         | 0.30             | .61     | 80 (34.0)             | 54 (32.7)        | −0.09   | 0.33             | .78     |
| **Age**           |                     |                  |               |                  |         |                      |                  |         |                  |         |
| Up to 1 year      | 4 (1.9)             | 7 (3.7)          | Ref.          |                  |         | 10 (5.3)              | 1 (0.6)          | *       | *                | *       |
| >1 year           | 208 (98.1)          | 181 (96.3)       | −0.31         | 0.93             | .74     | 225 (95.7)            | 164 (99.4)       |         |                  |         |
| **Days since vaccination** |                |                  |               |                  |         |                      |                  |         |                  |         |
| Up to 90 days     | 43 (20.7)           | 52 (27.8)        | Ref.          |                  |         | 58 (25.1)             | 37 (22.5)        | Ref.    |                  |         |
| 91–180 days       | 123 (59.1)          | 122 (65.2)       | −0.52         | 0.53             | .33     | 127 (55.0)            | 118 (72.0)       | 0.35    | 0.67             | .60     |
| >180 days         | 42 (20.2)           | 13 (6.9)         | −2.23         | 0.80             | .006    | 46 (19.9)             | 9 (5.5)          | −2.16   | 1.03             | .04     |
| **Province**      |                     |                  |               |                  |         |                      |                  |         |                  |         |
| Dornod            | 30 (14.2)           | 49 (26.1)        | Ref.          |                  |         | 50 (21.3)             | 29 (17.6)        | Ref.    | 0.87             |         |
| Dornogovi         | 52 (24.5)           | 9 (4.8)          | −3.00         | 0.73             | <.001   | 44 (18.7)             | 17 (10.3)        | −0.86   | 0.92             | .32     |
| Dundgovi          | 36 (17.0)           | 18 (9.6)         | −1.72         | 0.69             | .01     | 33 (14.0)             | 21 (12.7)        | −0.34   | 1.15             | .71     |
| Govisumber        | 17 (8.0)            | 10 (5.3)         | −1.49         | 0.84             | .08     | 18 (7.6)              | 9 (5.4)          | −0.50   | 1.15             | .66     |
| Khentii           | 48 (22.6)           | 39 (20.7)        | −1.16         | 0.59             | .04     | 38 (16.2)             | 49 (29.7)        | 1.06    | 0.78             | .18     |
| Sukbaatar         | 27 (12.7)           | 46 (24.5)        | −0.18         | 0.60             | 0.77    | 50 (21.3)             | 23 (17.0)        | −0.47   | 0.82             | .56     |
| Tuv               | 2 (0.9)             | 17 (9.0)         | 1.86          | 1.10             | .09     | 2 (0.8)               | 17 (10.3)        | 3.58    | 1.40             | .01     |

*Model does not converge.
Bold values indicate positive findings.
cates that >180 days would not be the ideal time to carry out a post-vaccination monitoring survey, we would caution against extrapolating this information to estimating protection against challenge. Further studies to understand the protective immune response (humoral and cell-mediated) to SPPV are required.

5 | CONCLUSIONS

Our results show that the use of a commercial CPPV ELISA kit provides a robust and reliable assay for post-vaccination surveillance on a regional or national level for SP in low resource settings. Our work builds on previous studies investigating the humoral immune response to CPPV vaccination and addresses particularly the limited number of studies assessing SP vaccination under field conditions. Our results have indicated that the timing of a post-vaccination SP testing survey is an important factor to consider when planning post-vaccination monitoring.

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None of the funding bodies played a role in the design of the study, collection, analysis or interpretation of the data, or in writing the manuscript.

ETHICAL APPROVAL

The animal samples used in this study were archived sera that had been collected in 2016 as part of a national Mongolian surveillance programme. Ethical approval for the national surveillance programme was granted by Mongolian authorities at the State Central Veterinary Laboratory and the Mongolian General Authority for Veterinary Services.

AUTHOR’S CONTRIBUTIONS

P.M.B, G.L., G.U., and O.M. contributed to the conception of the study; P.M.B, G.L., G.T., G.U., B.V., B.S., B.D. and P.F. designed the study; G.T. generated the fluorescently-labelled LSDV strain; P.F. developed and carried out the FVNTs; G.U., Bu.K., O.M. and J.F. carried out the ELISAs; G.L., P.F., P.M.B and G.U. analysed and interpreted the data; P.F., G.L., and P.M.B drafted the manuscript. All authors have approved the submitted version of the manuscript.

AVAILABLE OF DATA AND MATERIALS

The dataset supporting the conclusions of this article are included within this article, its additional files, and available in a public repository.

ORCID

Petra Fay 🏷️ https://orcid.org/0000-0001-5808-4284
Georgina Limon 🏷️ https://orcid.org/0000-0001-6230-2475
John Flannery 🏷️ https://orcid.org/0000-0002-5012-2829
Philippa M. Beard 🏷️ https://orcid.org/0000-0002-2285-810X

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**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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