Evaluation of the Safety and Efficacy of a Live Attenuated Thermostable Rift Valley Fever Vaccine in Sheep, Goats and Cattle

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Abstract

Rift valley fever (RVF) is a highly significant vector-borne disease causing huge economic losses in livestock (ruminants and camels) and also human fatalities. The disease is endemic in most Sub-Saharan African countries, including West Africa, and has been present in the Middle East since 2010. Vaccination is considered to be the most effective way to prevent and control the expansion of the disease.

Currently available attenuated live vaccines for RVF have significant limitations in that they are either thermolabile (CL13 strain vaccine) or causes abortion and teratogenic effects (Smithburn strain vaccine). This study therefore set out to develop a safe and effective thermostable live attenuated RVF vaccine. The existing CL13 vaccine, which is a naturally attenuated strain, was made thermostable through three cycles of heating (56°C) and selection. The resulting candidate vaccine (CL13T) was stable at 4°C for 20 months and shows significantly improved levels of thermostability over the existing CL13 vaccine.

A pilot batch of the CL13T vaccine was produced and tested for safety and efficacy in cattle, sheep and goats. The vaccine was found to be safe, with no clinical signs or side effects observed in vaccinated animals, and there was no evidence for circulation of the virus in the blood of animals post-vaccination. On testing for efficacy in cattle, sheep and goats, through the detection of neutralizing antibodies post-vaccination, good levels of neutralizing antibodies were detected for a minimum of one year in sheep and goats, and neutralizing antibodies were detected for least 4 months in cattle.

This new thermostable vaccine could represent an efficient tool for the control of rift valley fever in endemic countries. The vaccine also has the potential to be used, along with an appropriate diagnostic test, to differentiate vaccinated from infected animals (DIVA).

Keywords: Rift valley fever; Thermostable; Clone 13T vaccine; Sheep; Goat; Cattle

Background

Rift valley fever virus (RVFV) belongs to the Phlebovirus genus of the Bunyaviridae family. It is a segmented, single-stranded RNA virus composed of three segments: the large (L), the medium (M), and the small (S) [1]. RVFV causes a viral zoonotic vector-borne disease known as rift valley fever (RVF). The disease is characterized by high levels of mortality of young animals and abortions in pregnant females. Affected species include sheep, goat, cattle and camel. Humans can also be infected by mosquito bites, and importantly by exposure to the blood and tissues from infected ruminants during slaughter, necropsy or while assisting aborting animals [2]. Naturally infected animals develop high viremia sufficient to infect arthropod vectors, even if the infection is clinically in apparent [3]. In contrast to human disease, RVF manifests as “abortion storms” in several adult livestock species and in infected newborns, mortality approaches 100% [4-6].

Initially confined to sub-Saharan Africa and Egypt [6], the virus emerged out of the African continent in 2000 to cause a major enzootic/epidemic in the Arabian Peninsula [7]. In 2006/07, in Kenya, Somalia and Tanzania, the disease caused more than 1000 human infections with 323 deaths. The virus then spread to Madagascar, Comoros and Mayotte islands in 2008 [8,9]. Recent studies have shown the presence of RVF infected camels in North Africa in 2011 [10], as well as outbreaks in Mauritania and Senegal [11].

Vaccines currently being used in RVF endemic countries such as South Africa and Kenya include the live-attenuated Smithburn strain vaccine and a formalin inactivated aluminum hydroxide adjuvanted vaccine, based on a low passage wild RVF strain [12]. Though effective, both of these vaccines have limitations preventing their potential use. The live attenuated vaccine (Smithburn strain) induces early and long-term immunity after a single injection [13] however it is not recommended for use in early-stage pregnant ewes [5,14]. The inactivated whole virus vaccine requires at least two initial administrations followed by annual revaccination [15,16].
The RVFV clone 13 (CL13) strain is a naturally attenuated strain of RVFV. It is a mutant strain carrying a large deletion in the non-structural protein coded by the S segment (NSs), which was isolated from a non-fatal human case of RVF in the Central African Republic [17]. Vaccination using the CL13 strain has been demonstrated to be safe and efficacious in sheep, goats and calves using a single vaccination [18,19]. However, a recent study revealed that this vaccine is unstable at temperatures above 22°C, raising the risk associated with using this vaccine in the tropical (hot) countries without strict maintenance of the cold chain during vaccine storage and delivery [20].

The objectives of this study were firstly to develop a thermostable CL13 vaccine, through the isolation of a derivative thermostable clone from the original CL13 strain, and secondly to evaluate the safety and efficacy of the new thermostable vaccine through the measurement of vaccine related RVFV, RVFV RNA and neutralizing antibodies in the blood post-vaccination in target species (sheep, goats and cattle).

Materials and Methods

Isolation and characterization of the thermostable RVFV CL13T candidate vaccine

The RVFV CL13 strain was propagated on Monkey African Green kidney (Vero) cells, heated to 56°C and resistant viral particles were selected by the limited dilution method. Serial dilutions of the virus were prepared and titrated in 96 well plates. Virus from the highest dilution showing cytopathic effect was considered a resistant clone and was selected for the next passage. Three cycles of heating and selection were carried out and the most stable clone, which was named Clone 13T (CL13T), was passaged on Vero cells and its kinetics of multiplication, infectious titre and thermostability was compared with the original CL13 strain. The RVFV was handled under BSL-3 laboratory conditions in biosafety class III gloved boxes.

To determine their replication kinetics, each of the selected clones were inoculated at a multiplicity of infection (MOI) of 0.01 on 4 Vero cell flasks grown and maintained in DMEM with 10% irradiated fetal calf serum. The inoculated flasks were incubated at 37°C in a 5% CO₂ atmosphere. Every 24 hours one of the four flasks was frozen at -80°C and titrated. Titres were calculated by the Reed-Muench method and expressed by TCID₅₀/ml.

Thermostability of the RVFV CL13T candidate vaccine

Viral suspensions of both vaccine strains (CL13T and CL13) were tested for stability by periodic titration after heating in water bath at 37°C, 45°C and 56°C as follows:

- At 37°C, with aliquots of virus titrated daily for 4 days.
- At 45°C, with aliquots of virus titrated every 30 min for 180 min.
- At 56°C, with aliquots of virus titrated every 10 min for 90 min.

Both strains were mixed V/V with a stabilizer (4% peptone, 8% sucrose and 2% glutamate), lyophilized and tested for stability after storage 20 months at 4°C and 7 days at 37°C.

Safety and efficacy testing of the CL13T candidate vaccine in animals

All animal experiments were carried out in accordance with guidelines for the care and handling of experimental animals. The animal experiments were approved by the ethics committee in charge of the control and supervision of experiments on animals and the experiments were conducted in high containment level 3 facilities. Prior to the experiment, all the animals were confirmed to be negative for antibodies to RVFV by ELISA and virus-neutralization (VN) as described in the OIE Manual [21]. After vaccination, rectal temperatures was recorded daily, and the animals were monitored daily for clinical signs including weakness, respiratory signs and general wellbeing. Antibody responses were analyzed weekly for 8 weeks, then monthly.

Six to nine month-old calves were randomly divided in 5 groups of 6 calves. Each group of 6 calves was vaccinated by the subcutaneous (SC) route using different doses of the CL13T vaccine, from 10⁴ to 10⁸ TCID₅₀. Vaccinated animals were observed and sampled for a period of 4 months. A group of 6 sheep and a group of 6 goats, all 3-6 months old, were vaccinated with 1ml of 10⁸ TCID₅₀ of the RVFVCL13 vaccine by the SC route. A further group of 2 sheep and 2 goats were used as unvaccinated controls. The 4 unvaccinated controls were housed in the same pen as the vaccinated animals.

Blood collected in EDTA tubes was tested for the presence of the vaccine virus genome by real-time RT-PCR [22] from day 0 to day 15 post vaccination. Blood samples were also passed onto Vero cells and the cultures were observed daily for cytopathic effect.

Statistical Analysis

The proportion of animals showing detectible antibody titres was compared between the vaccine groups using the Student t-test. A significance level of p=0.05 was used.

Results

Kinetics of CL13 and CL13T virus multiplication

The RVFV CL13T reached a titer of 10⁶.7 TCID₅₀/ml two days post inoculation (dpi) of Vero cells. The highest titre of CL13 virus (10⁷.0 TCID₅₀/ml) was obtained at 3 dpi of the Vero cells. The titres of both viruses then decreased progressively; at 4 dpi the CL13 titre was 10⁶.6 TCID₅₀/ml and the CL13 was 10⁷.7 TCID₅₀/ml (Figure 1).

![Figure 1: Kinetics of RVFV CL13T and RVFV CL13 growth on Vero cells.](image-url)
Thermostability of CL13 and CL13T candidate vaccine viruses

No loss in infectivity was observed after incubating the liquid form of the CL13T candidate vaccine virus for 4 days at 37°C, while at the same temperature and incubation time the CL13 vaccine virus lost all its infectivity (Figure 2A).

The freeze-dried CL13T vaccine, stored at 4°C for a 20 month period, showed a drop of 1 log TCID$_{50}$ in infectious titre (7.8 to 6.8/ml), while the freeze-dried CL13 vaccine, stored at 4°C for a 12 month period, decreased in titre from 5.8 to less than 4.0 (data not shown). When stored for 7 days at 37°C, the titre of the freeze-dried CL13T vaccine decreased from 7.8 to 6.2, whereas during the same time period the CL13 vaccine lost all its infectivity (Figure 2B). Additionally, the CL13T vaccine virus remained stable for more than 3 hours at 45°C, during which time the CL13 lost over a log of infectivity (Figure 2D). The CL13T vaccine virus remained stable for 90 min at 56°C, whereas during the same time period the CL13 vaccine virus lost nearly 3 logs of infectivity (Figure 2C).

Not enough sensitivity to differentiate between CL13 and CL13T, the obtained value for the melting point is 167.5°C for RVF CL13T and 166.8°C for RVF CL13.

Safety testing of CL13T candidate vaccine

Normal body temperatures were recorded in all the sheep, goats and cattle post-vaccination with different doses (10$^4$ to 10$^8$ TCID$_{50}$) of the CL13T candidate vaccine virus (data not shown). No abnormal behavior, clinical signs or local reactions at the injection sites were observed in any of the vaccinated animals.

EDTA blood, collected from day 0 to day 15 post-vaccination, was tested for the presence of RVFV by virus isolation on Vero cells and for RVFV RNA by real-time RT-PCR. No live virus or viral RNA was detected in any of the post-vaccination samples, indicating that the vaccine virus was not circulating in the vaccinated animals.

Serological responses in animals vaccinated with the CL13T candidate vaccine virus

All cattle that were vaccinated with different doses of the CL13T candidate vaccine seroconverted as shown in Figure 3. Only two of the six cattle vaccinated with 10$^4$ TCID$_{50}$ of the CL13T candidate vaccine tested positive by VN, with both of the positive cattle having a titre of 1.5 at 42 and 56 days post vaccination (dpv) and one of the cattle having a titre of 1.02 at 86 dpv. Five out of the six cattle vaccinated with 10$^5$ TCID$_{50}$ of the CL13T candidate vaccine tested positive with VN titres between 1.02 to 1.26 by 28 dpv.

A good neutralizing antibody response was recorded in the majority of vaccinated animals from 7 dpv following the administration of a single dose of 10$^6$, 10$^7$ and 10$^8$ TCID$_{50}$ of the CL13T candidate vaccine (Figure 3). Neutralizing antibody were detected in the vaccinated cattle up to 116 dpv in all groups, except for the group of cattle that were administrated a dose of 10$^4$ TCID$_{50}$.
No significant difference in the proportion of animals with detectible antibodies was found between the groups that were vaccinated with $10^5$ TCID$_{50}$ and $10^6$ TCID$_{50}$ ($p=0.147$) and the groups vaccinated with $10^5$ TCID$_{50}$ and $10^7$ TCID$_{50}$ ($p=0.065$). Statistically significant differences ($p<0.05$) were however observed when comparing the proportion of animals with detectible antibodies between the group vaccinated with $10^5$ TCID$_{50}$ and the groups vaccinated with $10^4$ TCID$_{50}$ and $10^8$ TCID$_{50}$.

High titres of antibodies, measured by both ELISA and VN, were recorded in all the vaccinated sheep and goats from 7 dpv (Figure 4). The kinetics of the antibody responses were similar in sheep and goats and antibodies persisted in both species for up to a year post-vaccination (Figure 4). All the unvaccinated control animals remained negative throughout the study, indicating that the vaccine virus was not being spread between the vaccinated and the unvaccinated animals. No statistically significant differences in serum neutralizing antibody titres were observed between sheep and goat groups ($p=0.320$).

Discussion

Live attenuated vaccines against RVF have been shown to be more efficient for disease control compared to inactivated vaccines that require an initial course of two vaccinations, followed by an annual booster vaccination [15,16]. The live-attenuated vaccines for RVF that are currently available either have safety problems, as with the Smithburn strain [23] or stability issues, as with the thermolabile CL13 vaccine [20].

In this study, we report the isolation and characterization of a new thermostable RVF CL13T candidate vaccine, derived from the CL13 vaccine strain through a heating and cloning method. This ‘heating and cloning’ technique has been used in the past to develop thermostable vaccines that have been successfully used in the field; examples include the thermostable Rinderpest and Sheep pox vaccines [24,25].

The kinetics of virus replication on Vero cells, as well as the thermostability at 4°C, 37°C, 45°C and 56°C has been compared between the CL13T candidate vaccine strain and the original CL13 vaccine virus. When the CL13 vaccine virus was propagated in cell culture, the maximum viral titre occurred after 2 or 3 dpi, depending on the MOI [26]. A similar result was obtained in this current study, with the titre of the CL13 virus peaking at 3 dpi and the titre of the CL13T virus peaking at 2 dpi. Interestingly, the CL13T reached a titre of $10^{8.5}$ TCID$_{50}$/ml, whereas the CL13 reached a maximum titre of $10^{7.5}$ TCID$_{50}$/ml. This indicates that the cloning process also increased the fecundity of the selected virus (CL13T), producing a virus that replicated more rapidly, reaching higher peak viral titres on Vero cells. These properties would have obvious advantages for vaccine production.

As regards thermostability, the CL13T vaccine candidate strain was more stable for longer time periods than the CL13 strain at normal storage conditions (4°C). The CL13T strain was also more resistant to heating than the CL13 strain. A previous study also confirmed the lack of thermostability of the CL13 vaccine strain [20].
This study indicates that the CL13T vaccine strain is safe for use in cattle and small ruminants. No side effects, clinical signs or rises in body temperature were observed in the vaccinated animals, even when very high doses of the vaccine were administered. Additionally, RVFV or viral RNA was not detected in the blood of the vaccinated animals by virus isolation or real-time RT-PCR, showing that the vaccine virus was not circulating post-vaccination in the blood of vaccinated animals. The CL13 vaccine has previously been demonstrated to be safe to use in pregnant animals, so it is highly likely that the CL13T vaccine virus, which is derived from the CL13 vaccine virus and contains the same deleted NSs segment as CL13, will also be safe for use in pregnant animals [18].

The optimum protective dose of the CL13T candidate vaccine has been established in cattle, sheep and goats. In cattle the effect of the vaccine dose is clear; earlier and higher neutralizing antibody titres were observed in cattle as the dose of virus in the vaccine was increased. A dose of 10⁵ TCID₅₀ resulted in a good serological response in the vaccinated cattle, while in small ruminants (sheep and goats) a lower dose of 10³ TCID₅₀ conferred a good and long-lasting serological response. These findings are in accordance with reported results for the Smithburn and CL13 strains [18,19], however the neutralizing titres observed post-vaccinated with the CL13T vaccine were more pronounced and longer lasting than that seen with the Smithburn and CL13 strains, This serological response may be enhanced by the use the specific adjuvant such [27].

In conclusion, this study reports the development and characterization of a thermostable live attenuated RVF vaccine. The new CL13T candidate vaccine, which was derived from the deleted CL13 vaccine, is stable at 4°C for 20 months and shows improved levels of thermostability over the existing CL13 vaccine. The CL13 vaccine was safe, with no clinical signs or side effects observed in vaccinated animals, and there was no evidence for the circulation of the virus in the blood of animals post-vaccination. On testing for efficacy in cattle, sheep and goats, through the detection of neutralizing antibodies post-vaccination, good levels of neutralizing antibodies were detected for a minimum of one year in sheep and goats, and neutralizing antibodies were detected for at least 4 months in cattle. This new thermostable vaccine could represent an efficient tool for the control of rift valley fever in endemic countries and also has the potential to be used, along with an appropriate diagnostic test to differentiating infected from vaccinated animals (DIVA).

References