Analyses of Entry Mechanisms of Novel Emerging Viruses Using Pseudotype VSV System

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Abstract: Emerging infectious diseases include newly identified diseases caused by previously unknown organisms or diseases found in new and expanding geographic areas. Viruses capable of causing clinical disease associated with fever and bleeding are referred to as viral hemorrhagic fevers (VHF). Arenaviruses and Bunyaviruses, both belonging to families classified as VHF are considered major etiologies of hemorrhagic fevers caused by emerging viruses; having significant clinical and public health impact. Because these viruses are categorized as Biosafety Level (BSL) 3 and 4 pathogens, restricting their use, biological studies including therapeutic drug and vaccine development have been impeded. Due to these restrictions and the difficulties in handling such live viruses, pseudotype viruses bearing envelope proteins of VHF viruses have been developed using vesicular stomatitis virus (VSV) as a surrogate system. Here, we report the successful developments of two pseudotype VSV systems; bearing the envelope proteins of Lujo virus and severe fever with thrombocytopenia syndrome (SFTS) virus, both recently identified viruses of the family Arenaviridae and Bunyaviridae, respectively. My presentation will summarize the characterization of the envelope proteins of Lujo virus including its cellular receptor use and cell entry mechanisms. In addition, I will also present a brief introduction of SFTS reported in Japan and the diagnostic studies in progress using these newly pseudotype VSV system.

Key words: Arenavirus, Bunyavirus, pseudotype, Cell fusion, entry, SFTSV, Lujo virus

Thank you, Dr. Morikawa. First of all, thank you very much for giving me this opportunity to present our recent data in this meeting. Today, I’d like to talk about the entry mechanisms of two novel emerging viruses using pseudotype VSV system.

Today’s presentation:
1. Characterization of Lujo virus GP and cell entry
2. Severe fever with thrombocytopenia syndrome (SFTS) in Japan

Today’s my presentation is two topics. The first part is the characterization of novel Arenavirus, Lujo virus, GP and its cell entry. The second part is the recent sensational infectious disease, severe fever with thrombocytopenia syndrome, SFTS, in Japan (Fig. 1).

Okay, let’s start the first part. The Arenaviral hemorrhagic fevers are caused by various Arenaviruses, which are distributed in South American and African continents as shown here [1, 2]. In this study, we focused on four significant arenaviruses shown in red letter, especially the Lujo virus which is recently isolated in Southern Africa [3] (Fig. 2).

Arenaviruses are categorized as New World arenaviruses and Old World arenaviruses [2]. Some of the arenaviruses defined host in nature, disease, case fatality, and entry receptors. However, the Lujo virus is still largely unknown [3, 4] (Fig. 3).

The Lujo virus is a novel arenavirus discovered in 2008 at Lusaka and Johannesburg. The first letter of the place name joined and designated Lujo. The Lujo virus shows high case fatality rate and is classified as a BSL-4 pathogen. Lujo virus is a highly novel genetic lineage apart from the Lassa virus or Junin virus. The Lujo virus is unknown for the entry mechanisms including receptors. So, the aim of this study is to clarify the characteristics of Lujo virus GP and to examine the entry mechanisms of Lujo virus by using pseudotype VSV (Fig. 4).

This is the schematic representation of the construction of the pseudotype VSVs [5]. The parental VSV encoding the luciferase gene instead of the VSV-G infect to the producing cells expressing arenavirus GPs. The viruses budded from these cells are arenavirus-GP pseudotype VSV.

The characteristics of the pseudotype virus is a useful tool examination of viral entry because of single infection,
and is safety because of using only envelope gene even if BSL-4 pathogen, and is easy construction just only single transfection of the plasmid and VSV infection, and is easy evaluation of the infectivity because of encoding the reporter gene such as GFP, luciferase, or secreted alkaline phosphatase [5] (Fig. 5).

This slide shows their incorporation and glycosylation of flag-tagged arenavirus GP into the virions. All of the arenavirus GPs were efficiently incorporated into the virions and showed high-mannose type of glycosylation because of being sensitive to the endoglycosidase-H treatment. In contrast, VSV-G showed complex-type of glycosylation because of resistance to the endoglycosidase-H treatment (Fig. 6).

Next, we examined the infectivities of arenavirus pseudotype in various mammalian cells. All of the arenavirus pseudotypes can infect many types of cells, but Lujo virus pseudotype is not susceptible to the mouse-derived cell lines, NIH3T3 and normal murine liver cells, NMuLi, and human T-cell derived Molt-4 cells. The Lassa virus pseudotype is also not susceptible to the Jurkat cells as reported previously [6] (Fig. 7).

Next to determine whether the arenavirus pseudotypes show the pH-dependent entry, the bafilomycin, ammonium chloride or chloroquine, which are the inhibitors of endosomal acidification, were utilized [7]. As you can see, all of the arenavirus pseudotype infections were inhibited by these inhibitors in dose-dependent manner. The control virus, murine leukemia virus pseudotype, which causes direct fusion at the plasma membrane, has no influence (Fig. 8).

I’d like to remind you again that New World arenaviruses use the transferrin receptor 1 as a receptor for the entry, and Old World arenaviruses use alpha-dystroglycan (α-DG) as a receptor [2]. So, next we examined whether the Lujo virus uses these receptors (Fig. 9).

At first, we examined the inhibition of arenavirus pseudotype infection by anti-human TIF1 monoclonal antibody. As you can see, the Junin virus and Chapare virus pseudotype infections were inhibited by the anti-human TIF1 antibody. In contrast, the Lujo virus as well as Lassa virus pseudotype infections exhibited no inhibition (Fig. 10).

Furthermore, we confirmed involvement of TIF1 by using the CHO cells expressing human TIF1. As you can see, the Junin virus and Chapare virus pseudotype infec-
tions were increased in the CHO cells expressing human TfR1 compared with the parenteral CHO cells. In contrast, Lujo virus as well as Lassa virus pseudotype infection was no significant increase in these cells. These results indicate that Lujo virus pseudotype infection is independent of TfR1-mediated entry (Fig. 11).

Next, we examined the involvement of α-DG. Many Old World arenaviruses such as Lassa virus can use O-mannosylated α-DG as a receptor for the entry [2, 8]. So, to determine whether Lujo virus can use it, we utilized the Raji or Jurkat cells expressing LARGE. The LARGE is one of the glycosylation enzymes and causes hyperglycosylation of α-DG.

As you can see, the Lassa virus pseudotype infection is increased in both Raji and Jurkat cells expressing LARGE. In contrast, the Lujo virus and other pseudotype virus infection was no significant increase (Fig. 12).
Fig. 7. Inhibition of AREpv infection by H⁺-ATPase inhibitors

| Virus       | Host in Nature | Disease                                      | Case Fatality | Receptor |
|-------------|----------------|----------------------------------------------|---------------|----------|
| New World arenavirus (NWA) |                |                                              |               |          |
| Junin (JUNV) | Calomys musculinus | Argentine hemorrhagic fever (AHF) | 15-30%        | TR1      |
| Machupo (MACV) | Calomys-caudatus | Argentine hemorrhagic fever (AHF) | 15%           | TR1      |
| Guanarito (GTOV) | Zygadonotomys brevicaudata | Venezuelan hemorrhagic fever (VHF) | 25%          | TR1      |
| Sabia (SABV) | Unknown         | Brachyhemorrhagic fever                      | 1/3          | TR1      |
| Chupana (CHIPV) | Unknown | Not yet named                               | 1/5           | hTIR1    |
| Old World arenavirus (OWA) |                |                                              |               |          |
| Lassa (LASV) | Montomyidae species | Lassa fever                                  | 10-15%        | α-DG     |
| lymphocytic choriomeningitis virus (LCMV) | Mastomys natalensis | Lymphocytic choriomeningitis | <1%           | α-DG     |
| Lujo (LUJV) | Unknown         | Not yet named                               | 4/5           |          |

Fig. 8. Arenaviruses known to be human pathogens

Fig. 9. Inhibition of AREpv infection by anti-human TIR1 antibody

Fig. 10. Infectivities of AREpv in CHO cells expressing hTIR1

Fig. 11. Infectivities of AREpv in Raji or Jurkat cells expressing O-mannosylated α-DG
Furthermore, we confirmed involvement of DG by using the DG-knockout ES cells. The cells were transfected with control or DG expressing plasmid and infected with pseudotype viruses [9]. As you can see, only Lassa virus pseudotype infection was increased in DG-expressing cells. These results indicate that Lujo virus pseudotype infection is independent of both the DG and TfR1 (Fig. 13).

Many viral envelope proteins have a potential to change the conformation and cell fusion activities in low pH exposure. To examine cell fusion activities of arenavirus GPs, the GP expressing cells were treated with the indicated pH buffers. As you can see, the syncytium formations were observed in Lassa, Junin, Chapare virus GP, and VSV-G expressing cells under low pH conditions. In contrast, surprisingly, no syncytium formation was observed in Lujo virus GP expressing cells even at pH4.0 buffer treatment (Fig. 14).

Similarly, by using two different plasmids encoding, the luciferase gene under the control of the T7 promoter and T7 polymerase, the cell fusion activities were evaluated by the luciferase reporter assay [10]. As you can see, the cell fusion activities were observed in Lassa, Junin, Chapare virus GP, and VSV-G expressing cells, but not Lujo virus GP expressing cells under low pH conditions (Fig. 15).

We also examined the effects of low pH exposure on arenavirus pseudotype infection. The viruses were treated with the indicated pH buffer before infection. After neutralization with the culture medium, the infectivities were evaluated.

As you can see, all of the arenavirus pseudotypes including the Lujo virus pseudotype infections were abolished after low pH buffer treatment. In contrast, VSV pseudotype can infect to the cells even at the low pH buffer treatment, because VSVG is reversible to change the pH-triggered conformation. Together with the cell fusion experiments, the Lujo virus GP is changed conformation at low pH, but some kind of molecules, which do not express at the plasma membrane, maybe needed for the membrane fusion (Fig. 16).
To further examine the entry pathways of arenavirus pseudotype, we utilized chlorpromazine [7, 11]. This chemical is broadly known not only inhibitor for the clathrin-mediated endocytosis, but also inducer for lipidosis. As you can see, all of the arenavirus pseudotype infections were inhibited by the treatment with chlorpromazine in a dose dependent manner, especially the Lujo virus and Lassa virus pseudotype infections drastically reduced (Fig. 17).

Recently, it is reported that Ebola virus infection is involved in the cholesterol transporter protein, Niemann-Pick C1, NPC1, and acid sphingomyelinase, which is the hydrolase enzyme involved in the sphingolipid metabolism [12–14] (Fig. 18).

So, to examine the role of lipid metabolism in arenavirus infection, we utilized the other lipidosis-inducing drugs, imipramine, desipramine, amitriptyline. These drugs are known to the anti-depressant drugs, and cause lipid accumulation in vitro. As you can see, the Lujo virus pseudotype as well as the Ebola virus pseudotype infections were inhibited by these drugs (Fig. 19).

Furthermore, we utilized the cholesterol transport inhibitor, U18666A, to examine the effects of arenavirus pseudotype infection. This inhibitor has a similar activity to inhibit the function of NPC1, so the Ebola virus entry was abolished. As you can see, the Lujo virus pseudotype infection as well as Ebola virus pseudotype infection was
inhibited by this inhibitor in a dose-dependent manner. Now, we are investigating the involvement of NPC1 with Lujo virus pseudotype entry (Fig. 20).

This is the conclusion to the first part: The Lujo virus GP contains high-mannose type of glycosylation. The Lujo virus pseudotype is not susceptible to some cell lines. The entry is pH dependent. The Lujo virus GP does not have cell-to-cell fusion activities. The Lujo virus pseudotype utilizes neither α-DG nor TIR1 as receptors. The lipidosis in endosomes inhibits Lujo virus pseudotype entry (Fig. 21).

In summary, the Lujo virus entered the target cells via unknown receptors but not TIR1 and α-DG. The viruses were endocytosed at low pH, but some kind of intracellular receptor molecules such as NPC1 for Ebola virus may be needed in endosomal membrane fusion. And, this interaction may be needed to exclude the cholesterol or phospholipid accumulation. Lujo virus may have a unique entry mechanism and further precise studies will be needed (Fig. 22).

Okay, second part of my presentation is SFTS in Japan. This is the paper of the New England Journal of Medicine published in 2011 [15]. The SFTS emerged in 2009–2010 at six provinces in China. The novel Bunyavirus designated SFTSV, was isolated from the blood samples of SFTS patients (Fig. 23).

In Japan, last autumn, the first diagnosed SFTS patient was found in Yamaguchi Prefecture [16]. Until last

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**Fig. 19.** Inhibition of AREpv infection by cholesterol transport inhibitor

**Fig. 20.** Conclusion (part I)

**Fig. 21.** Schematic model of arenavirus (Lujo virus) cell entry
Fever with Thrombocytopenia Associated with a Novel Bunyavirus in China

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BACKGROUND

Heightened surveillance of acute febrile illness in China since 2009 has led to the identification of a severe fever with thrombocytopenia syndrome (SFTS) with an unknown cause. Infections with Angiostrongylus cantonensis have been suggested as a cause, but the pathogen has not been detected in most patients on laboratory testing.

METHODS

We obtained blood samples from patients with the case definition of SFTS in six provinces in China. The blood samples were used to isolate the causal pathogen by inoculation of cell cultures and for detection of viral RNA by polymerase-chain-reaction assay. The pathogen was characterized by electron microscopy and nucleotide sequencing. We used enzyme-linked immunosorbent assay, indirect immunofluorescence assay, and neutralization testing to analyze the level of virus-specific antibody in patients’ serum samples.

RESULTS

We isolated a novel virus, designated SFTS bunyavirus, from patients who presented with fever, thrombocytopenia, leukopenia, and multisystem dysfunction. RNA sequence analysis revealed that the virus was a newly identified member of the genus Phlebovirus in the Bunyaviridae family. Electron-microscopic examination revealed virions with the morphologic characteristics of a bunyavirus. The presence of the virus was confirmed in 47 patients with SFTS from six provinces by detection of viral RNA, specific antibodies in the virus in blood, or both. Serologic assays showed a virus-specific immune response in all 15 pairs of serum samples collected from patients during the acute and convalescent phases of the illness.

Fig. 22. Published paper of SFTS in China

| Survived | 25 |
| Succumbed | 18 |
| Total | 43 |
| (As of Sep. 13, 2013) |

Fig. 23. Age and area incidence of SFTS in Japan

month, including the retrospective studies, total 43 people were infected with the SFTSV and 18 people died. Up to the present, we do not know the exact reason, but the SFTS occurs almost in western part of Japan including the Nagasaki Prefecture (Fig. 24).

This slide summarizes the characteristics of SFTSV.
The SFTSV belongs to the Family Bunyaviridae and the Genus Phlebovirus. The Rift Valley fever virus is the same genus. This virus has three segmented negative-stranded RNA; L, M, and S shown here. This virus is visualized with a diameter of 80 to 100 nanometer as shown here and detected from the ixodid tick such as Haemaphysalis longicornis, the Japanese name is Futatogechimadani. The fatality rate is around 10% to 30%. The major clinical symptoms of SFTS are fever and thrombocytopenia which means lower platelet count and leucopenia which means low white blood cell count. In the laboratory, the virus is easily isolated from Vero cells as shown here. But this virus is classified as a BSL-3 pathogen. So, to further operate easily, we tried to construct pseudotype virus possessing the SFTSV GP (Fig. 25).

At first, we examined the localization of SFTSV-GP in Vero cells infected with the SFTSV or transfected with the GP expressing plasmid. As you can see, both GPs did not express at the plasma membrane but mainly localized in the ER or Golgi apparatus (Fig. 26).

Generally, VSV were budded from the plasma membrane, so the cell surface expressed GPs were efficiently incorporated into the virions, and these pseudotype viruses showed highly infectious [17]. In contrast, the cytosolic expressed GP such as the SFTSV-GP was not efficiently incorporated into the virions. But rarely a small portion of the VSV maybe budded from the ER or Golgi, and these VSVs incorporated the GP into the virions and showed moderately infectious (Fig. 27).

This slide shows the infectivities of SFTSV pseudotype in various mammalian cells. The Rift Valley fever virus and 100-diluted VSV pseudotypes were used as the control for the comparison. The SFTSV pseudotype virus can infect many types of cells, but are not susceptible to the lymphocyte cell lines (Fig. 28).

As shown in first part of my presentation, SFTSV pseudotype infection was also inhibited by the bafilomycin and ammonium chloride. That means this virus infection is required for the endosomal acidification (Fig. 29).
Finally, we examined neutralization tests using the pseudotype viruses. The SFTSV pseudotype is specifically neutralized by the serial dilution of patient serum. These results indicate that the neutralizing antibodies are present in the serum of convalescent patient and pseudotype virus is a useful tool for the neutralization test (Fig. 30).

This is the conclusion to the second part. The SFTS is endemic, especially to the western part of Japan. The pseudotype virus exhibits a high infectivity to various mammalian cells. The pseudotype virus is not susceptible to lymphocyte cell lines. The pseudotype entry is pH-dependent. The pseudotype virus is neutralized by SFTS patient serum.

SFTSV pseudotype virus is a useful tool not only to use diagnostic studies or drug screening but also to examine entry mechanisms of SFTSV.

Finally, thanks to all the members of Department of Virology I and Department of Veterinary Science, National Institute of Infectious Diseases, and the DG knockout ES cells were kindly provided by Drs. Kawaoka and Campbell.

Thank you very much for your attention.

Questions and Discussion

Male Participant Thank you for your nice presentation. About SFTS, SFTS-G is processed in the cells and do you confirm the GN and GC incorporated to the virus particle, both GN, GC is incorporated to the virus particle?
Jiro Yasuda

Other questions? I’ve got one question. While using the bafilomycin or some other treatment, the Lujo virus entry is associated with maybe NPC-1 or some other things. But at the same time, the Lassa virus pseudotype infection is also inhibited by the similar inhibitors. So, in this experiment, do you think the Lassa also utilized NPC or some other factors?

Hideki Tani

Yeah, that’s a good question. The Lassa virus pseudotype also partially inhibited this NPC inhibitor. But we don’t have the data to compare the Lassa virus GP and Lujo virus GP. We need to compare these GP in the future.

Shigeru Morikawa

That’s the same situation of Lujo virus infection. Raji cells are not susceptible to the SFTSV infection in our studies as well as reported in the previous studies, but DC or some macrophages cells infected with SFTSV, so that maybe different to the cell line and the primary cells. We need to study about it.

Hideki Tani

I have one question on the SFTS. Because in human patient, the main target of the cells in vivo is maybe lymphocyte, because in the lymph node, there are many cells infected, but not in the other epithelial cells like hepatocytes or some others. But in vitro basing this system or basing the SFTS virus itself, the virus can infect a variety of cells, but except for the lymphocyte. What do you think about this discrepancy?

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Shigeru Morikawa

Okay, anyway, maybe I think you can find a receptor for SFTS using this system. Thank you very much, Dr. Tani.

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