

HHS Public Access

Author manuscript *Nature*. Author manuscript; available in PMC 2015 April 02.

Published in final edited form as:

Nature. 2014 October 2; 514(7520): 47-53. doi:10.1038/nature13777.

Reversion of advanced Ebola virus disease in nonhuman primates with ZMapp[™]

Xiangguo Qiu¹, Gary Wong^{1,2}, Jonathan Audet^{1,2}, Alexander Bello^{1,2}, Lisa Fernando¹, Judie B. Alimonti¹, Hugues Fausther-Bovendo^{1,2}, Haiyan Wei^{1,3}, Jenna Aviles¹, Ernie Hiatt⁴, Ashley Johnson⁴, Josh Morton⁴, Kelsi Swope⁴, Ognian Bohorov⁵, Natasha Bohorova⁵, Charles Goodman⁵, Do Kim⁵, Michael H. Pauly⁵, Jesus Velasco⁵, James Pettitt^{6,+}, Gene G. Olinger^{6,+}, Kevin Whaley⁵, Bianli Xu³, James E. Strong^{1,2,7}, Larry Zeitlin^{5,*}, and Gary P. Kobinger^{1,2,8,9,*}

¹National Laboratory for Zoonotic Diseases and Special Pathogens, Public Health Agency of Canada, Winnipeg, MB, Canada

²Department of Medical Microbiology, University of Manitoba, Winnipeg, MB, Canada

³Institute of Infectious Disease, Henan Centre for Disease Control, China

⁴Kentucky BioProcessing, Owensboro, KY, USA

⁵Mapp Biopharmaceutical Inc., San Diego, CA, USA

⁶United States Army Medical Research Institute of Infectious Diseases (USAMRIID), Frederick, MD, USA

⁷Department of Pediatrics and Child Health, University of Manitoba, Winnipeg, MB, Canada

⁸Department of Immunology, University of Manitoba, Winnipeg, MB, Canada

⁹Department of Pathology and Laboratory Medicine, University of Pennsylvania School of Medicine, Philadelphia, PA, USA

Abstract

Without an approved vaccine or treatment, Ebola outbreak management has been limited to palliative care and barrier methods to prevent transmission. These approaches, however, have yet

Financial disclosures

Users may view, print, copy, and download text and data-mine the content in such documents, for the purposes of academic research, subject always to the full Conditions of use:http://www.nature.com/authors/editorial_policies/license.html#terms

gary.kobinger@phac-aspc.gc.ca. Larry Zeitlin, Ph.D., Mapp Biopharmaceutical Inc., 6160 Lusk Blvd #C105, San Diego, CA 92121 USA, Phone: (443) 629-0104, larry.zeitlin@mappbio.com.

⁺current address: Integrated Research Facility, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Frederick, MD, USA

Author contributions

X.Q., G.K. and L.Z. designed the experiments. X.Q., G.W., J.A., A.B., L.F., J.B.A., H.F., H.W., J.A., J. P., G.G.O. and G.K. performed the experiments. X.Q., G.W., J.A., K.W., B.X., J. E. S., L.Z. and G.K. wrote the manuscript. E.H., A.J., J.M., K.S., O.B., N.B., C.G., D.K., M.H.P., J.V., K.W. and L.Z. contributed reagents for this study.

Her Majesty the Queen in right of Canada holds a patent on mAbs 2G4, and 4G7, PCT/CA2009/000070, "Monoclonal antibodies for Ebola and Marburg viruses." K.W. and L.Z. are the owners of Mapp Biopharmaceutical Inc. The authors declare no other competing interests.

to end the 2014 outbreak of Ebola after its prolonged presence in West Africa. Here we show that a combination of monoclonal antibodies (ZMappTM), optimized from two previous antibody cocktails, is able to rescue 100% of rhesus macaques when treatment is initiated up to 5 days post-challenge. High fever, viremia, and abnormalities in blood count and chemistry were evident in many animals before ZMappTM intervention. Advanced disease, as indicated by elevated liver enzymes, mucosal hemorrhages and generalized petechia could be reversed, leading to full recovery. ELISA and neutralizing antibody assays indicate that ZMappTM is cross-reactive with the Guinean variant of Ebola. ZMappTM currently exceeds all previous descriptions of efficacy with other therapeutics, and results warrant further development of this cocktail for clinical use.

Keywords

Monoclonal antibodies; Ebola; nonhuman primates; ZMapp[™]; post-exposure therapy

Introduction

Ebola virus (EBOV) infections cause severe illness in humans, and after an incubation period of 3 to 21 days, patients initially present with general flu-like symptoms before a rapid progression to advanced disease characterized by hemorrhage, multi-organ failure and a shock-like syndrome¹. In the spring of 2014, a new EBOV variant emerged in the West African country of Guinea², an area in which EBOV has not been previously reported. Despite a sustained international response from local and international authorities including the Ministry of Health (MOH), World Health Organization (WHO) and Médecins Sans Frontières (MSF) since March 2014, the outbreak has yet to be brought to an end after five months. As of 15th August 2014, there are 2127 total cases and 1145 deaths spanning Guinea, Sierra Leone, Liberia and Nigeria³. So far, this outbreak has set the record for the largest number of cases and fatalities, in addition to geographical spread⁴. Controlling an EBOV outbreak of this magnitude has proven to be a challenge and the outbreak is predicted to last for at least several more months⁵. In the absence of licensed vaccines and therapeutics against EBOV, there is little that can be done for infected patients outside of supportive care, which includes fluid replenishment, administration of antivirals, and management of secondary symptoms⁶⁷. With overburdened personnel, and strained local and international resources, experimental treatment options cannot be considered for compassionate use in an orderly fashion at the moment. However, moving promising strategies forward through the regulatory process of clinical development has never been more urgent.

Over the past decade, several experimental strategies have shown promise in treating EBOV-challenged nonhuman primates (NHPs) after infection. These include recombinant human activated protein C (rhAPC)⁸, recombinant nematode anticoagulant protein c2 (rNAPc2)⁹, small interfering RNA (siRNA)¹⁰, positively-charged phosphorodiamidate morpholino oligomers (PMO*plus*)¹¹, the vesicular stomatitis virus vaccine (VSV Δ G-EBOVGP)¹², as well as the monoclonal antibody (mAb) cocktails MB-003 (consisting of human or human-mouse chimeric mAbs c13C6, h13F6 and c6D8)¹³ and ZMAb (consisting of murine mAbs m1H3, m2G4 and m4G7)¹⁴. Of these, only the antibody-based candidates have demonstrated substantial benefits in NHPs when administered greater than 24 hours

Page 3

past EBOV exposure. Follow-up studies have shown that MB-003 is partially efficacious when administered therapeutically after the detection of two disease "triggers"¹⁵, and ZMAb combined with an adenovirus-based adjuvant provides full protection in rhesus macaques when given up to 72 hours after infection¹⁶.

The current objective is to develop a therapeutic superior to both MB-003 and ZMAb, which could be utilized for outbreak patients, primary health-care providers, as well as high-containment laboratory workers in the future. This study aims to first identify an optimized antibody combination derived from MB-003 and ZMAb components, before determining the therapeutic limit of this mAb cocktail in a subsequent experiment. In order to extend the antibody half-life in humans and to facilitate clinical acceptance, the individual murine antibodies in ZMAb were first chimerized with human constant regions (cZMAb). The cZMAb components were then produced in *Nicotiana benthamiana*¹⁷, using the large-scale, cGMP-compatible Rapid Antibody Manufacturing Platform (RAMP) and magnICON vectors that currently also manufactures the individual components of cocktail MB-003, before efficacy testing in animals.

Selecting for the best mAb combinations

Our efforts to down-select for an improved mAb cocktail comprising components of MB-003 and ZMAb began with the testing of individual MB-003 antibodies in guinea pigs and NHPs. In guinea pig studies, animals were given one dose of mAb c13C6, h13F6, or c6D8 individually (totaling 5 mg per animal) at 1 day post-infection (dpi) with 1000 × LD₅₀ of guinea pig-adapted EBOV, Mayinga variant (EBOV-M-GPA). Survival and weight loss were monitored over 28 days. Treatment with c13C6 or h13F6 yielded 17% survival (1 of 6 animals) with a mean time to death of 8.4 ± 1.7 and 10.2 ± 1.8 days, respectively. The average weight loss for c13C6 or h13F6-treated animals was 9% and 21% (Table 1). In nonhuman primates, animals were given three doses of mAb c13C6, h13F6, or c6D8, beginning at 24 hours after challenge with the Kikwit variant of EBOV (EBOV-K)¹⁸, and survival was monitored over 28 days. Only c13C6 treatment yielded any survivors, with 1 of 3 animals protected from EBOV challenge (Table 1), confirming in two separate animal models that c13C6 is the component that provides the highest level of protection in the MB-003 cocktail.

We then tested mAb c13C6 in combination with two of three mAbs from ZMAb in guinea pigs. The individual antibodies composing ZMAb were originally chosen for protection studies based on their *in vivo* protection of guinea pigs against EBOV-M-GPA¹⁹, and all three possible combinations were tested: ZMapp1 (c13C6+c2G4+c4G7), ZMapp2 (c13C6+c1H3+c2G4) and ZMapp3 (c13C6+c1H3+c4G7), and compared to the originator cocktails ZMAb and MB-003. Three days after challenge with $1000 \times LD_{50}$ of EBOV-M-GPA, the animals received a single combined dose of 5 mg of antibodies. This dosage is purposely given to elicit a suboptimal level of protection with the cZMAb and MB-003 cocktails, such that potential improvements with the optimized mAb combinations can be identified. Of the tested cocktails, ZMapp1 showed the best protection, with 4 of 6 survivors and less than 5% average weight loss (Table 1). ZMapp2 was next with 3 of 6 survivors and 8% average weight loss, and ZMapp3 protected 1 of 6 animals (Table 1). The level of

protection afforded by ZMapp3 was not a statistically significant increase over cZMAb (p = 0.224, log-rank test compared to ZMAb, $\chi^2 = 1.479$, df = 1), and showed the same survival rate along with a similar average weight loss (Table 1). As a result, only ZMapp1 and ZMapp2 were carried forward to NHP studies.

ZMapp1 or ZMapp2-treated NHPs

Rhesus macaques were used to determine whether administration of ZMapp1 or ZMapp2 was superior to ZMAb and MB-003 in terms of extending the treatment window. Due to mAb availability constraints, m4G7 was utilized in place of c4G7 for this NHP experiment. The experiment consisted of six NHPs per group receiving three doses of ZMapp1 (Group A) or ZMapp2 (Group B) at 50 mg/kg intravenously (IV) at 3-day intervals, beginning 3 days after a lethal intramuscular (IM) challenge with 4000 × TCID₅₀ (or 2512 PFU) of EBOV-K. Control animals were given phosphate-buffered saline (PBS) or mAb 4E10 (C1 and C2, respectively). Mock-treated animals succumbed to disease between 6–7 dpi with symptoms typical of EBOV (Figure 1a), characterized by high clinical scores but no fever (Figures 1b and 1c), in addition to viral titers up to ~10⁸ and ~10⁹ TCID₅₀ by the time of death (Figure 1d). Analysis of blood counts and serum biochemistry revealed leukocytopenia, thrombocytopenia, severe rash, decreased levels of GLU, as well as increased levels of ALP, ALT, BUN and CRE at end-stage EBOV disease (Figures 1e to 1o, Table 2).

All six Group A NHPs survived the challenge with mild signs of disease (Figure 1a, Table 2) (p = 0.0039, log-rank test, χ^2 = 8.333, df = 1, comparing to Group C), with the exception of A1 which showed an elevated clinical score (Figure 1b), increased levels of ALT, TBIL, and decreased PHOS (Figure 1, Table 2). However, this animal recovered after the third ZMapp1 dose and the clinical score dropped to zero by 15 dpi (Figure 1b). A fever was detected in all but one of the NHPs (A4) at 3 dpi, the start of the first ZMapp1 dose (Figure 1c). Viremia was also detected beginning at 3 dpi by TCID₅₀ in all but one animal from blood sampled just before the administration of the treatment (A3) (Figure 1d), and similar results were observed by RT-qPCR (Extended Data Table 1). The viremia decreased to undetectable levels by 21 dpi. EBOV shedding was not detected from oral, nasal and rectal swabs by RT-qPCR in any of the Group A animals (Extended Data Tables 2–4).

For Group B, 5 of 6 NHPs survived with B3 succumbing to disease at 9 dpi (Figure 1a) (p = 0.0039, log-rank test, χ^2 = 8.333, df = 1, comparing to Group C). Surviving animals showed only mild signs of disease (Table 2). The moribund animal showed increased clinical scores (Figure 1b), in addition to a drastic drop in body temperature shortly before death (Figure 1c). At the time of death, animal B3 had elevated levels of ALT, TBIL, BUN, CRE, in addition to decreased levels of GLU, suggesting multi-organ failure (Figure 1). All six Group B animals showed fever in addition to viremia at 3 dpi by TCID₅₀ and RT-qPCR (Figure 1D, Extended Data Table 1). It was interesting to note that in B3, the viremia reached ~10⁶ TCID₅₀ after 3 dpi (Figure 1d), suggesting that this NHP was particularly susceptible to EBOV infection. No escape mutants were detected with this animal. The administration of ZMapp2 at the reported concentrations was unable to effectively control viremia at this level. Virus shedding was also detected from the oral and rectal swabs by RT-

qPCR in the moribund NHP B3 (Extended Data Tables 2–4). Since ZMapp1 demonstrated superior protection to ZMapp2 in this survival study, ZMapp1 (now trademarked as ZMapp[™] by MappBio Pharmaceuticals) was carried forward to test the limits of protection conferred by this mAb cocktail in a subsequent investigation.

ZMapp[™]-treated NHPs

In this experiment, rhesus macaques were assigned into three treatment groups of six and a control group of three animals, with all treatment NHPs receiving three doses of ZMappTM (c13C6+c2G4+c4G7, 50 mg/kg per dose) spaced three days apart. After a lethal IM challenge with 1000 × TCID₅₀ (or 628 PFU) of EBOV-K¹⁸, we treated the animals with ZMappTM at 3, 6 and 9 dpi (Group D); 4, 7, and 10 dpi (Group E); or 5, 8 and 11 dpi (Group F). The control animals (Group G) were given mAb 4E10 as an IgG isotype control (n = 1) or PBS (n = 2) in place of ZMappTM starting at 4 dpi (Figure 2a). All animals treated with ZMappTM survived the infection, whereas the three control NHPs (G1, G2 and G3) succumbed to EBOV-K infection at 4, 8 and 8 dpi, respectively (p = 3.58E-5, log-rank test, $\chi^2 = 23.25$, df = 3, comparing all groups) (Figure 2b). At the time ZMappTM treatment was initiated, fever, leukocytosis, thrombocytopenia and viremia could be detected in the majority of the animals (Figure 2c-f, Table 3). All animals presented with detectable abnormalities in blood counts and serum biochemistry during the course of the experiment (Figure 2g-l, Table 3).

Based on clinical scores (Figure 2c, Extended Data Figure 1), the Group F animals did not appear to be as sick as animals E4 and E6, both of whom were near the clinical limit for IACUC mandated euthanasia at 5 and 7 dpi, respectively. Animal E4 had a flushed face and severe rash on more than 40% of its body surface between 5 to 8 dpi in addition to nasal hemorrhage at 7 dpi, whereas animal E6 had a flushed face and petechiae on its arms and legs between 7 to 9 dpi, in addition to jaundice between 10 to 14 dpi. This indicates that host genetic factors may play a role in the differential susceptibility of individual NHPs to EBOV-K infections. Fever, leukocytosis, thrombocytopenia, and a severe rash symptomatic of EBOV disease progression was detected in both E4 and E6 (Table 3). Increases in the level of liver enzymes ALT (10- to 30-fold increase), ALP (2- to 3-fold), and total bilirubin (TBIL, 3- to 11-fold) indicate significant liver damage (Figure 2g-l), a hallmark of filovirus infections. However, ZMappTM was successful in reversing observed disease symptoms and physiological abnormalities after 12 dpi, 2 days after the last ZMapp[™] administration (Table 3). Furthermore, ZMappTM treatment was able to lower the high virus loads observed in animals F2 and F5 (up to 10⁶ TCID₅₀/ml) to undetectable levels by 14 dpi (Figure 2f, Extended Data Figure 2).

ZMapp[™] cross-reacts with Guinea EBOV

While the results were very promising with EBOV-K infected NHPs, it was unknown whether therapy with ZMappTM would be similarly effective against the Guinean variant of EBOV (EBOV-G), the virus responsible for the West African outbreak. Direct comparison of published amino acid sequences between EBOV-G and EBOV-K showed that the epitopes targeted by ZMapp^{TM 2021} were not mutated between the two virus variants (Figure

3a), suggesting that the antibodies should retain their specificity for the viral glycoprotein. To confirm this, *in vitro* assays were carried out to compare the binding affinity of c13C6, c2G4 and c4G7 to sucrose-purified EBOV-G and EBOV-K. As measured by ELISA, the ZMappTM componentsshowed slightly better binding kinetics for EBOV-G than for EBOV-K (Figure 3b). Additionally, the neutralizing activity of individual mAbs was evaluated in the absence of complement for c2G4 and c4G7, and in the presence of complement for c13C6, as they have previously been shown to neutralize EBOV only under this condition¹³ (Figure 3c). The results supported the ELISA binding data, with comparable neutralizing activities between the two viruses.

Discussion

The West African outbreak of 2014 has highlighted the troubling absence of available vaccine or therapeutic options to save thousands of lives and stop the spread of EBOV. The lack of a clinically acceptable treatment offer limited incentive for people who suspect they might be infected to report themselves for medical help. Several previous studies have showed that antibodies are crucial for host survival from EBOV²²²³²⁴. Prior NHP studies have also demonstrated the ZMAb cocktail could protect 100% or 50% of animals when dosing was initiated 1 or 2 dpi, while the MB-003 cocktail protected 67% of animals with the same dosing regimen. Before the success with mAb-based therapies, other candidate therapeutics had only demonstrated efficacy when given within 60 minutes of EBOV exposure.

Our results with $ZMapp^{TM}$, a cocktail comprising of individual mAbs selected from MB-003 and ZMAb, demonstrate for the first time the successful protection of NHPs from EBOV disease when intervention was initiated as late as 5 dpi. In the preceding ZMapp1/ZMapp2 experiment, 11 of 12 treated animals had detectable fever (with the exception of A4), and live virus could be detected in the blood of 11 of 12 animals (with the exception of A3) by 3 dpi. Therefore, for the majority of these animals, treatment was therapeutic (as opposed to post-exposure prophylaxis), initiated after two detectable triggers of disease. ZMapp2 was able to protect 5 of 6 animals when administered at 3 dpi. For reasons currently unknown, the lone non-survivor (B3) experienced a viremia of 10^6 TCID₅₀ at 3 dpi, which is 100-fold greater than all other NHPs and approximately 10-fold higher than what ZMAb has been reported to suppress in a previous study¹⁶. This indicates enhanced EBOV replication in this animal, possibly due to host factors. It was important to note that despite the high levels of live circulating virus detected in B3, ZMapp2 administration was still able to prolong the life of this animal to 9 dpi, and suggests that in cases of high viremia such as this, the dosage of mAbs should be increased.

The highlight of these experimental results is undoubtedly $ZMapp^{TM}$, which was able to reverse severe EBOV disease as indicated by the elevated liver enzymes, mucosal hemorrhages and rash in animals E4 and E6. The high viremia (up to $10^6 \text{ TCID}_{50}/\text{ml}$ of blood in some animals at the time of intervention) could also be effectively controlled without the presence of escape mutants, leading to full recovery of all treated NHPs by 28 dpi. In the absence of direct evidence demonstrating $ZMapp^{TM}$ efficacy against lethal EBOV-G infection in NHPs, results from ELISA and neutralizing antibody assays show that

binding specificity is not abrogated between EBOV-K and EBOV-G, and therefore the levels of protection should not be affected. The compassionate use of ZMappTM in two infected American healthcare workers with apparently positive results pertaining to survival and reversion of EBOV disease²⁵, supports this assertion. Rhesus macaques have approximately 55–80 ml of blood per kg of body weight²⁶; at a dose of 50 mg/kg of antibodies, the estimated starting concentration is approximately 625–909 µg/ml of blood (total; ~200–300 µg/ml for each antibody). Therefore, the low EC₅₀ values for EBOV-G (0.004 – 0.02 µg/ml) bode well for treating EBOV-G infections with ZMappTM.

Since the host antibody response is known to correlate with and is required for protection from EBOV infections²³²⁴, mAb-based treatments are likely to form the centerpiece of any future therapeutic strategies for fighting EBOV outbreaks. However, whether ZMappTM-treated survivors can be susceptible to re-infection is unknown. In a previous study of murine ZMAb-treated, EBOV-challenged NHP survivors, a re-challenge of these animals with the same virus at 10 and 13 weeks after initial challenge yielded 6 of 6 survivors and 4 of 6 survivors, respectively²⁷. While specific CD4⁺ and CD8⁺ T-cell responses could be detected in all animals, the circulating levels of glycoprotein (GP)-specific IgG were shown to be 10-fold lower in non-survivors compared to survivors, suggesting that antibody levels may be indicative of protective immunity²⁷. Sustained immunity with experimental EBOV vaccines in NHPs remain unknown, however in a recent study, a decrease in GP-specific IgG levels due to old age or a suboptimal reaction to the VSV Δ G/EBOVGP vaccine in rodents also appear to be indicative of non-survival²⁸.

ZMapp[™] consists of a cocktail of highly purified mAbs; which constitutes a less controversial alternative than whole blood transfusions from convalescent survivors, as was performed during the 1995 EBOV outbreak in Kikwit²⁹. The safety of mAb therapy is welldocumented, with generally low rates of adverse reactions, the capacity to confer rapid and specific immunity in all populations, including the young, the elderly and the immunocompromised, and if necessary, the ability to provide higher-than-natural levels of immunity compared to vaccinations³⁰. The evidence presented here suggests that ZMapp[™] currently offers the best option of the experimental therapeutics currently in development for treating EBOV-infected patients. We hope that initial safety testing in humans will be undertaken soon, preferably within the next few months, in order to enable the compassionate use of ZMapp[™] as soon as possible.

Materials and Methods

Ethics statement

The guinea pig experiment, in addition to the second and third NHP study (ZMapp1, ZMapp2 and ZMappTM) were performed at the National Microbiology Laboratory (NML) as described on Animal use document (AUD) #H-13-003, and has been approved by the Animal Care Committee (ACC) at the Canadian Science Center for Human and Animal Health (CSCHAH), in accordance with the guidelines outlined by the Canadian Council on Animal Care (CCAC). The first study with MB-003 in NHPs was performed at United States Army Medical Research Institute of Infectious Diseases (USAMRIID) under an Institutional Animal Care and Use Committee (IACUC) approved protocol in compliance

with the Animal Welfare Act, Public Health Service Policy, and other federal statutes and regulations relating to animals and experiments involving animals. The facility where this research was conducted in accredited by The Association for Assessment and Accreditation of Laboratory Animal Care International and adheres to principles stated in the 8th edition of the *Guide for the Care and Use of Laboratory Animals*, National Research Council 2011.

mAb production

The large-scale production of mAb cocktails cZMAb, MB-003, ZMapp1, ZMapp2 and ZMapp[™] in addition to control mAb 4E10 (anti-HIV) from *N. benthamiana* under GMP conditions was done by Kentucky BioProcessing (Owensboro, KY) as described previously¹³¹⁵³¹. The large-scale production of m4G7 was performed by the Biotechnology Research Institute (Montreal, QC) using a previously described protocol¹⁶.

Viruses

The challenge virus used in NHPs was Ebola virus *H.sapiens*-tc/COD/1995/Kikwit-9510621 (EBOV-K) (order *Mononegavirales*, family *Filoviridae*, species *Zaire ebolavirus*; GenBank accession #AY354458)¹⁸. Passage three from the original stock was used for the studies at the NML and passage four was used for the study performed at USAMRIID (the NHP study with the individual MB-003 mAbs). Sequencing of 112 clones from the passage three stock virus revealed that the population ratio of 7U:8U in the EBOV GP editing site was 80:20; sequencing for the passage four stock virus was not performed, and therefore the ratio of 7U:8U in the editing site was unknown. The virus used in guinea pig studies was guinea pig-adapted EBOV, Ebola virus VECTOR/*C.porcellus*-lab/COD/1976/Mayinga-GPA (EBOV-M-GPA) (order *Mononegavirales*, family *Filoviridae*, species *Zaire ebolavirus*; Genbank accession number AF272001.1)³². The Guinean variant used in IgG ELISA and neutralizing antibody assays was Ebola virus *H.sapiens*-tc/GIN/2014/Gueckedou-C05 (EBOV-G) (order *Mononegavirales*, family *Filoviridae*, species *Zaire ebolavirus*; GenBank accession #KJ660348.1)².

Animals

Outbred 6–8 week old female Hartley strain guinea pigs (Charles River) were used for these studies. Animals were infected IP with $1000 \times LD_{50}$ of EBOV-M-GPA. The animals were then treated with one dose of ZMAb, MB-003, ZMapp1, ZMapp2, c13C6, h13F6 or c6D8 totaling 5 mg per guinea pig, and monitored every day for 28 days for survival, weight and clinical symptoms. This study was not blinded, and no animals were excluded from the analysis.

For the MB-003 study performed at USAMRIID, thirteen rhesus macaques (*Macaca mulatta*) were obtained from the USAMRIID primate holding facility, ranging from 5.1 to 10 kg. This study was not blinded, and no animals were excluded from the analysis. Animals were given standard monkey chow, primate treats, fruits, and vegetables for the duration of the study. All animals were challenged IM with a target dose of 1000 PFU. Treatment with either monoclonal antibody, MB-003 cocktail, or PBS was administered on 1, 4, and 7 dpi via saphenous intravenous infusion. Animals were monitored at least once daily for changes

in health, diet, behavior, and appearance. Animals were sampled for chemical analysis, complete bloods counts and viremia on 0, 3, 5, 7, 10, 14, 21, and 28 dpi.

For the ZMapp1 and ZMapp2 study, fourteen male and female rhesus macaques (*Macaca mulatta*), ranging from 4.1 to 9.6 kg (4–8 years old) were purchased from Primgen (USA). This study was not blinded, and no animals were excluded from the analysis. Animals were assigned groups based on gender and weight. Animals were fed standard monkey chow, fruits, vegetables, and treats. Husbandry enrichment consisted of visual stimulation and commercial toys. All animals were challenged IM with a high dose of EBOV [backtiter: $4000 \times \text{TCID}_{50}$ or 2512 PFU] at 0 dpi. Administration of the first treatment dose was initiated at 3 dpi, with identical doses at 6 and 9 dpi. Animals were scored daily for signs of disease, in addition to changes in food and water consumption. On designated treatment days in addition to 14, 21, and 27 dpi, the rectal temperature and clinical score were measured, and the following were sampled: blood for serum biochemistry and cell counts and viremia. This study was not blinded, and no animals were excluded from the analysis.

For the ZMappTM study, twenty-one male rhesus macaques, ranging from 2.5 to 3.5 kg (2 years-old) were purchased from Primgen (USA). This study was not blinded, and no animals were excluded from the analysis. Animals were assigned groups based on gender and weight. Animals were fed standard monkey chow, fruits, vegetables, and treats. Husbandry enrichment consisted of visual stimulation and commercial toys. All animals were challenged IM with EBOV [backtiter: $1000 \times \text{TCID}_{50}$ or 628 PFU] at 0 dpi. Administration of the first treatment dose was initiated at 3, 4 or 5 dpi, with two additional identical doses spaced three days apart. Animals were scored daily for signs of disease, in addition to 14, 21, and 28 dpi, the rectal temperature and clinical score were measured, and the following were sampled: blood for serum biochemistry and cell counts and viremia.

Blood counts and blood biochemistry

Complete blood counts were performed with the VetScan HM5 (Abaxis Veterinary Diagnostics). The following parameters were shown in the figures: levels of white blood cells (WBC), lymphocytes (LYM), percentage of lymphocytes (LYM%), levels of platelets (PLT), neutrophils (NEU) and percentage of neutrophils (NEU%). Blood biochemistry was performed with the VetScan VS2 (Abaxis Veterinary Diagnostics). The following parameters were shown in the figures: levels of alkaline phosphatase (ALP), alanine aminotransferase (ALT), blood urea nitrogen (BUN), creatinine (CRE), and total bilirubin (TBIL).

Enzyme-linked immunosorbent assays (ELISAs)

IgG ELISA with c13C6, c2G4 or c1H3 was performed as described previously¹⁶ using gamma-irradiated EBOV-G and EBOV-K virions purified on a 20% sucrose cushion as the capture antigen in the ELISA. Each mAb was assayed in triplicate.

Neutralizing antibody assays

Two-fold dilutions of c13C6, c2G4 or c1H3 ranging from 0.0156 to 2 μ g/ml were first incubated with 100 PFU of EBOV-G at room temperature for 1 hour with or without complement, transferred to Vero E6 cells and incubated at 37°C for 1 hour, and then replaced with DMEM supplemented with 2% fetal bovine serum and scored for the presence of cytopathic effect (CPE) at 14 dpi. The lowest concentrations of mAbs demonstrating the absence of CPE were averaged and reported.

EBOV titration by TCID₅₀ and RT-qPCR

Titration of live EBOV was determined by adding 10-fold serial dilutions of whole blood to VeroE6 cells, with three replicates per dilution. The plates were scored for cytopathic effect at 14 dpi, and titers were calculated with the Reed and Muench method³³. Results were shown as median tissue culture infectious dose (TCID₅₀).

For titers measured by RT-qPCR, total RNA was extracted from whole blood with the QIAmp Viral RNA Mini Kit (Qiagen). EBOV was detected with the LightCycler 480 RNA Master Hydrolysis Probes (Roche) kit, with the RNA polymerase (nucleotides 16472 to 16538, AF086833) as the target gene. The reaction conditions were as follows: 63°C for 3 min, 95°C for 30 s, and cycling of 95°C for 15 s, 60°C for 30 s for 45 cycles on the ABI StepOnePlus. The lower detection limit for this assay is 86 genome equivalents/ml. The sequences of primers used were as follows: EBOVLF2 (CAGCCAGCAATTTCTTCCAT), EBOVLR2 (TTTCGGTTGCTGTTTCTGTG), and EBOVLP2FAM (FAM-ATCATTGGCGTACTGGAGGAGCAG-BHQ1).

Sequence alignment

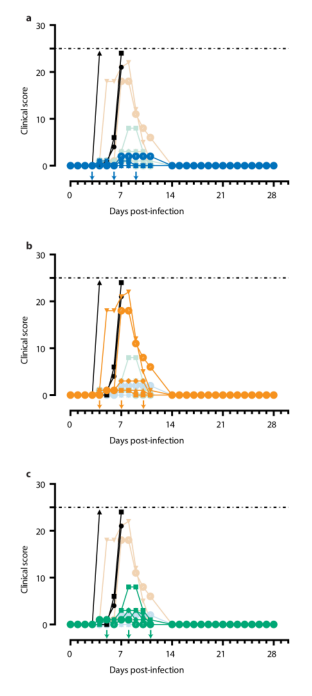
Protein sequences for EBOV-K and EBOV-G surface glycoproteins were obtained from GenBank, accession numbers AGB56794.1 and AHX24667.1 respectively. The sequences were aligned using DNASTAR Lasergene 10 MEGAlign using the Clustal W algorithm.

Statistical analysis

For the guinea pig and nonhuman primate studies, each treatment group consisted of six animals. Assuming a significance threshold of 0.05, a sample size of six per group will give >80% power to detect a difference in survival proportions between the treatment (83% survival or higher) and the control group using a one-tailed Fisher's exact test.

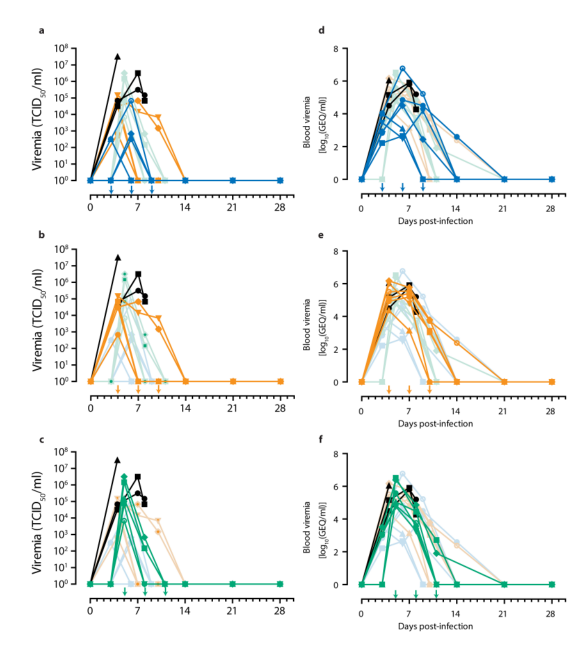
Survival was compared using the log-rank test in GraphPad PRISM 5, differences in survival were considered significant when the p-value was less than 0.05. Antibody binding to EBOV-G and EBOV-K was compared by fitting the data to a 4-parameter logistic regression using GraphPad PRISM 5. The EC_{50} were considered different if the 95% Confidence Intervals excluded each other. For all statistical analyses, the data conformed to the assumptions of the test used.

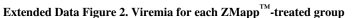
Extended Data



Extended Data Figure 1. Clinical scores for each ZMappTM-treated group

Arrows indicate treatment days. Dashed line represents humane endpoint threshold. Faded symbols/lines are the other two treatment groups, for comparison. Control group (Group G) is shown in black on all three panels. (a) Clinical score of Group D (blue); (b) Clinical score of Group E (orange); (c) Clinical score of Group F (green).





Arrows indicate treatment days. Faded symbols/lines are the other two treatment groups, for comparison. Control group (Group G) is shown in black on all three panels. (a) $TCID_{50}$ of Group D (blue); (b) $TCID_{50}$ of Group E (orange); (c) $TCID_{50}$ of Group F (green). (d) Viremia by RT-qPCR of Group D (blue); (e) Viremia by RT-qPCR of Group E (orange); (f) Viremia by RT-qPCR of Group F (green).

Extended Data Table 2

Oral swab viremia measured by RT-qPCR for the ZMapp1- and ZMapp2-treated NHPs.

DAYS	A1	A2	A3	A4	A5	A6	B1	B2	B3	B4	B5	B6	C1	C2
0	UD	UD	UD	UD	UD	UD								
3	UD	UD	UD	UD	UD	UD								
6	UD	UD	UD	UD	UD	UD								
7													5.05E+03	
9		UD	4.81E+04	UD	UD	UD								
14	UD		UD	UD	UD									
21	UD		UD	UD	UD									
27	UD		UD	UD	UD									

Footnote: UD = Undetectable.

Extended Data Table 3

Nasal swab viremia measured by RT-qPCR for the ZMapp1- and ZMapp2-treated NHPs.

	_													
DAYS	A1	A2	A3	A4	A5	A6	B1	B2	B3	B4	B5	B6	C1	C2
0	UD	UD	UD	UD	UD									
3	UD	UD	UD	UD	UD									
6	UD	UD	UD	UD	3.75E+02									
7													1.98E+04	2.16E+03
9		UD	UD	UD										
14	UD		UD	UD	UD									
21	UD		UD	UD	UD									
27	UD		UD	UD	UD									

Footnote: UD = Undetectable.

Extended Data Table 4

Rectal swab viremia measured by RT-qPCR for the ZMapp1- and ZMapp2-treated NHPs.

DAYS	A1	A2	A3	A4	A5	A6	B1	B2	B3	B4	B5	B6	C1	C2
0	UD	UD	UD	UD	UD	UD								
3	UD	UD	UD	UD	UD	UD								
6	UD	UD	UD	UD	4.16E+02	8.17E+03								
7													4.38E+04	
9		UD	3.90E+02	UD	UD	UD								
14	UD		UD	UD	UD									
21	UD		UD	UD	UD									
27	UD		UD	UD	UD									

Footnote: UD = Undetectable.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

The authors thank Kevin Tierney, Allen Grolla, Shane Jones, Jessica Dong, and Dr. Darwyn Kobasa for their excellent technical assistance, Dr. Victor Klimyuk and Dr. Yuri Gleba for access to the magnICON expression system, and Dr. Herta Steinkellner for access to transgenic *N. benthamiana*. This work was supported by the Defense Threat Reduction Agency (DTRA contract HDTRA1-13-C-0018), the National Institutes of Health (U19AI109762), the Public Health Agency of Canada (PHAC), and a Canadian Safety and Security Program (CSSP) grant to G.P.K. and X.Q. G.W. is the recipient of a Doctoral Research Award from the Canadian Institute for Health Research (CIHR).

References

- Bausch DG, Sprecher AG, Jeffs B, Boumandouki P. Treatment of Marburg and Ebola hemorrhagic fevers: a strategy for testing new drugs and vaccines under outbreak conditions. Antiviral Res. 2008; 78:150–161.10.1016/j.antiviral.2008.01.152 [PubMed: 18336927]
- Baize S, et al. Emergence of Zaire Ebola Virus Disease in Guinea Preliminary Report. N Engl J Med. 201410.1056/NEJMoa1404505
- 3. WHO.int. WHO Ebola virus disease (EVD). 2014. <http://www.who.int/csr/don/archive/disease/ ebola/en/>
- 4. CDC.gov. Chronology of Ebola Hemorrhagic Fever Outbreaks. 2014. <<u>http://www.cdc.gov/vhf/ebola/resources/outbreak-table.html</u>>
- 5. Reliefweb.int. W. African Ebola epidemic 'likely to last months': UN. 2014. http://reliefweb.int/report/guinea/w-african-ebola-epidemic-likely-last-months-un
- Clark DV, Jahrling PB, Lawler JV. Clinical management of filovirus-infected patients. Viruses. 2012; 4:1668–1686.10.3390/v4091668 [PubMed: 23170178]
- Guimard Y, et al. Organization of patient care during the Ebola hemorrhagic fever epidemic in Kikwit, Democratic Republic of the Congo, 1995. J Infect Dis. 1999; 179(Suppl 1):S268– 273.10.1086/514315 [PubMed: 9988194]
- Hensley LE, et al. Recombinant human activated protein C for the postexposure treatment of Ebola hemorrhagic fever. J Infect Dis. 2007; 196(Suppl 2):S390–399.10.1086/520598 [PubMed: 17940975]
- Geisbert TW, et al. Treatment of Ebola virus infection with a recombinant inhibitor of factor VIIa/ tissue factor: a study in rhesus monkeys. Lancet. 2003; 362:1953–1958.10.1016/ S0140-6736(03)15012-X [PubMed: 14683653]
- Geisbert TW, et al. Postexposure protection of non-human primates against a lethal Ebola virus challenge with RNA interference: a proof-of-concept study. Lancet. 2010; 375:1896– 1905.10.1016/S0140-6736(10)60357-1 [PubMed: 20511019]
- Warren TK, et al. Advanced antisense therapies for postexposure protection against lethal filovirus infections. Nat Med. 2010; 16:991–994.10.1038/nm.2202 [PubMed: 20729866]
- Feldmann H, et al. Effective post-exposure treatment of Ebola infection. PLoS Pathog. 2007; 3:e2.10.1371/journal.ppat.0030002 [PubMed: 17238284]
- Olinger GG Jr, et al. Delayed treatment of Ebola virus infection with plant-derived monoclonal antibodies provides protection in rhesus macaques. Proc Natl Acad Sci U S A. 2012; 109:18030– 18035.10.1073/pnas.1213709109 [PubMed: 23071322]
- 14. Qiu X, et al. Successful treatment of ebola virus-infected cynomolgus macaques with monoclonal antibodies. Sci Transl Med. 2012; 4:138ra181.10.1126/scitranslmed.3003876
- Pettitt J, et al. Therapeutic intervention of Ebola virus infection in rhesus macaques with the MB-003 monoclonal antibody cocktail. Sci Transl Med. 2013; 5:199ra113.10.1126/scitranslmed. 3006608
- Qiu X, et al. mAbs and Ad-Vectored IFN-alpha therapy rescue ebola-infected nonhuman primates when administered after the detection of viremia and symptoms. Sci Transl Med. 2013; 5:207ra143.10.1126/scitranslmed.3006605
- Giritch A, et al. Rapid high-yield expression of full-size IgG antibodies in plants coinfected with noncompeting viral vectors. Proc Natl Acad Sci U S A. 2006; 103:14701–14706.10.1073/pnas. 0606631103 [PubMed: 16973752]

- Jahrling PB, et al. Evaluation of immune globulin and recombinant interferon-alpha2b for treatment of experimental Ebola virus infections. J Infect Dis. 1999; 179(Suppl 1):S224– 234.10.1086/514310 [PubMed: 9988188]
- Qiu X, et al. Ebola GP-specific monoclonal antibodies protect mice and guinea pigs from lethal Ebola virus infection. PLoS Negl Trop Dis. 2012; 6:e1575.10.1371/journal.pntd.0001575 [PubMed: 22448295]
- 20. Wilson JA, et al. Epitopes involved in antibody-mediated protection from Ebola virus. Science. 2000; 287:1664–1666. [PubMed: 10698744]
- 21. Qiu X, et al. Characterization of Zaire ebolavirus glycoprotein-specific monoclonal antibodies. Clin Immunol. 2011; 141:218–227.10.1016/j.clim.2011.08.008 [PubMed: 21925951]
- 22. Dye JM, et al. Postexposure antibody prophylaxis protects nonhuman primates from filovirus disease. Proc Natl Acad Sci U S A. 2012; 109:5034–5039.10.1073/pnas.1200409109 [PubMed: 22411795]
- 23. Wong G, et al. Immune parameters correlate with protection against ebola virus infection in rodents and nonhuman primates. Sci Transl Med. 2012; 4:158ra146.10.1126/scitranslmed.3004582
- 24. Marzi A, et al. Antibodies are necessary for rVSV/ZEBOV-GP-mediated protection against lethal Ebola virus challenge in nonhuman primates. Proc Natl Acad Sci U S A. 2013; 110:1893– 1898.10.1073/pnas.1209591110 [PubMed: 23319647]
- 25. Promedmail.org. EBOLA VIRUS DISEASE WEST AFRICA (117): WHO, NIGERIA, LIBERIA, DRUG, MORE. 2014. http://www.promedmail.org/direct.php?id=2666073>
- 26. NC3RS.org. Practical blood sample volumes for laboratory animals, domestic species and nonhuman primates. 2014. http://www.nc3rs.org.uk/bloodsamplingmicrosite/page.asp?id=426
- 27. Qiu X, et al. Sustained protection against Ebola virus infection following treatment of infected nonhuman primates with ZMAb. Sci Rep. 2013; 3:3365.10.1038/srep03365 [PubMed: 24284388]
- 28. Wong G, et al. Immunization with Vesicular Stomatitis Virus vaccine expressing the Ebola glycoprotein provides sustained long-term protection in rodents. Vaccine. 2014 in press.
- Mupapa K, et al. International Scientific and Technical Committee. Treatment of Ebola hemorrhagic fever with blood transfusions from convalescent patients. J Infect Dis. 1999; 179(Suppl 1):S18–23.10.1086/514298 [PubMed: 9988160]
- UPMChealthsecurity.org. Next-Generation Monoclonal Antibodies: Challenges and Opportunities. UPMC Center for Biosecurity; 2013.
- Zeitlin L, et al. Enhanced potency of a fucose-free monoclonal antibody being developed as an Ebola virus immunoprotectant. Proc Natl Acad Sci U S A. 2011; 108:20690–20694.10.1073/pnas. 1108360108 [PubMed: 22143789]
- Connolly BM, et al. Pathogenesis of experimental Ebola virus infection in guinea pigs. J Infect Dis. 1999; 179(Suppl 1):S203–217.10.1086/514305 [PubMed: 9988186]
- 33. Reed LJ, Muench H. A simple method of estimating fifty per cent endpoints. American journal of hygiene. 1938; 27:493–497.

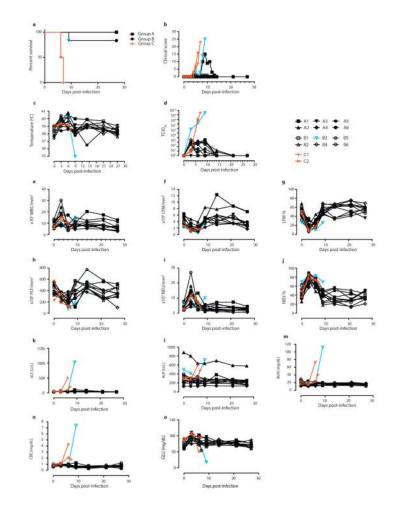


Figure 1. Post-exposure protection of EBOV-infected nonhuman primates with ZMapp1 and ZMapp2 $% \mathcal{A} = \mathcal{A} = \mathcal{A} + \mathcal{A$

Rhesus macaques were challenged with EBOV-K, and 50 mg/kg of ZMapp1 (Group A) or ZMapp2 (Group B) were administered on days 3, 6, and 9 (n=6 per treatment group, n=2 for controls). Non-specific IgG mAb or PBS was administered as a control (Group C). (a) Kaplan-Meier survival curves. (b) Clinical score. (c) Rectal temperature. (d) EBOV viremia by TCID₅₀. Blood parameters: (e) white blood cell count; (f) lymphocyte count; (g) lymphocyte percentage; (h) platelet count; (i) neutrophil count; (j) neutrophil percentage; (k) ALT; (l) ALP; (m) BUN; (n) CRE; (o) GLU.

Qiu et al.

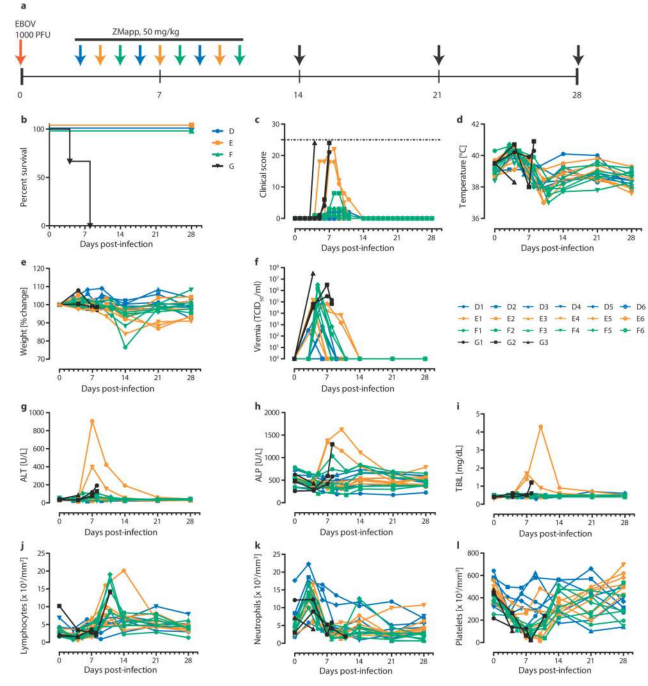


Figure 2. Post-exposure protection of EBOV-infected nonhuman primates with ZMapp $_{^{\rm TM}}$

(a to f) Rhesus macaques (n=6 per ZMappTM treatment group, n=3 for controls) were challenged with EBOV-K, and 50 mg/kg of ZMappTM were administered beginning at 3 (Group A), 4 (Group B) or 5 (Group C) days after challenge. Non-specific IgG mAb or PBS was administered as a control (Group D). (a) Timeline of infection, treatment and sample days. (b) Kaplan-Meier survival curves. (c) Clinical scores; the dashed line indicates the minimum score requiring mandatory euthanasia. (d) Rectal temperature. (e) Percentage

body weight change. (f) EBOV viremia by $TCID_{50}$. (g to l) Selected clinical parameters of Group A to D animals. (g) ALT. (h) ALP. (i) TBIL. Counts for (j) Lymphocytes, (k) Neutrophils and (l) Platelets over the course of the experiment.

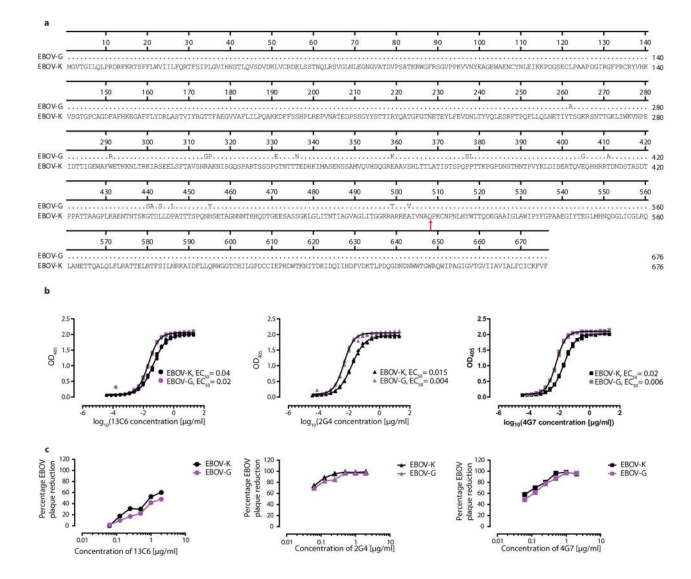


Figure 3. Amino acid alignment of the Kikwit and Guinea variants of EBOV, and *in vitro* antibody assays of mAbs c13C6, c2G4 and c4G7 with EBOV-G or EBOV-K virions (a) Sequence alignment of the EBOV glycoprotein from the Kikwit (EBOV-K) and Guinea (EBOV-G) variants, with the binding epitopes of ZMappTM pointed with an arrow, and (b) an ELISA (for each antibody, the EC₅₀ are different (p < 0.05, regression analysis) between the two antigens) as well as (c) a neutralizing antibody assay showing the activity of the individual mAbs composing ZMappTM against EBOV-K (black) and EBOV-G (purple), and the samples were run in triplicate.

⊳
Ē
Ľ,
9
_
\leq
Manu
Manu
Manuscr

Author Manuscript

Efficacy of individual and combined monoclonal antibody treatments in guinea pigs and nonhuman primates.

Treatment groups, time of treatment	Dose (mg)	Mean time to death (days)	# Survivors / # Total	% Survival	% Weight loss	p-value, con cZMAb	p-value, compared with cZMAb MB-003
Guinea pigs						,	,
PBS, 3 dpi	N/A	7.3 ± 0.5	0/4	0	%6	ı	I
cZMAb, 3 dpi	5	11.6 ± 1.8	1/6	17	7%	ı	ı
MB-003, 3 dpi	5	8.2 ± 1.5	9/0	0	40%	ı	ı
ZMapp1, 3 dpi	5	9.0 ± 0.0	4/6	67	<5%	0.190	0.0147
ZMapp2, 3 dpi	5	8.3 ± 0.6	3/6	50	8%	0.634	0.0692
ZMapp3, 3 dpi	5	8.6 ± 1.1	1/6	17	9%6	0.224	0.411
c13C6, 1 dpi	5	8.4 ± 1.7	1/6	17	%6	ı	ı
h13F6, 1 dpi	5	10.2 ± 1.8	1/6	17	21%	ı	ı
c6D8, 1 dpi	5	10.5 ± 2.2	0/6	0	38%	1	ı
Treatment groups, time of treatment	Dose (mg/kg)) Mean time to death (days)) # Survivors / # Total	l 🛛 % Survival	_		
Nonhuman primates							
PBS, 1 dpi	N/A	8.4 ± 1.9	0/1	0			
MB-003, 1 dpi	50	14.0 ± 2.8	1/3	33			
c13C6, 1 dpi	50	9.0 ± 1.4	1/3	33			
h13F6, 1 dpi	50	9.0 ± 2.0	0/3	0			
c6D8, 1 dpi	50	9.7 ± 0.6	0/3	0			

٦

ript	
Author	
Manuscrip	

Table 2

Clinical findings of EBOV-infected NHPs from 1 to 27 dpi.

				Clinical findings	ST		
Animal ID	Treatment group	Body temperature	Rash	White blood cells	Platelets	Biochemistry	Outcome
A1	50 mg/kg c13C6+c2G4+m4G7, 3 dpi	Fever (6, 9, 14 dpi)			Thrombocytopenia (6, 9 dpi)	ALT↑ (9, 14 dpi), TBIL↑ (9 dpi), PHOS↓ (6 dpi)	Survived
A2	50 mg/kg c13C6+c2G4+m4G7, 3 dpi	Fever (3 dpi)		Leukocytosis (3 dpi)		CRE↓ (14 dpi)	Survived
A3	50 mg/kg c13C6+c2G4+m4G7, 3 dpi	Fever (3 dpi)		Leukocytosis (3 dpi)	Thrombocytopenia (6 dpi)		Survived
A4	50 mg/kg c13C6+c2G4+m4G7, 3 dpi			Leukocytopenia (9 dpi)	Thrombocytopenia (3, 6, 14, 21, 27 dpi)		Survived
AS	50 mg/kg c13C6+c2G4+m4G7, 3 dpi	Fever (3, 6, 9 dpi)		Leukocytopenia (9 dpi)	Thrombocytopenia (3, 21 dpi)		Survived
A6	50 mg/kg c13C6+c2G4+m4G7, 3 dpi	Fever (3 dpi)					Survived
B1	50 mg/kg ZMapp2, 3 dpi	Fever (3, 14, 21 dpi)		Leukocytopenia (6, 14, 21, 27 dpi)	Thrombocytopenia (6 dpi)		Survived
B2	50 mg/kg ZMapp2, 3 dpi	Fever (3, 6 dpi)			Thrombocytopenia (6, 9 dpi)		Survived
B3	50 mg/kg ZMapp2, 3 dpi	Fever (3, 6 dpi), Hypothermia (9 dpi)	Severe rash (9 dpi)		Thrombocytopenia (6, 9 dpi)	ALTTT (9 dpi), TBIL (1 (9 dpi), BUNTT (9 dpi), CRETT (9 dpi), GLUUU (9 dpi)	Died, 9 dpi
B4	50 mg/kg ZMapp2, 3 dpi	Fever (3, 6 dpi)		Leukocytopenia (6 dpi)	Thrombocytopenia (6, 27 dpi)		Survived
B5	50 mg/kg ZMapp2, 3 dpi	Fever (3, 6, 14, 21 dpi)		Leukocytosis (3 dpi)	Thrombocytopenia (3, 6 dpi)		Survived
B6	50 mg/kg ZMapp2, 3 dpi	Fever (3 dpi)		Leukocytosis (3 dpi), Leukocytopenia (6, 9, 14, 21, 27 dpi)	Thrombocytopenia (6 dpi)	PHOS¢ (3 dpi), CRE↓ (27 dpi)	Survived
CI	PBS, 3 dpi		Moderate rash (6 dpi), Severe rash (7 dpi)	Leukocytosis (3 dpi)	Thrombocytopenia (6, 7 dpi)	ALB↓ (7 dpi), ALT↑ (7 dpi), BUN↑ (7 dpi)	Died, 7 dpi
C2	Control mAb, 3 dpi		Severe rash (6 dpi)	Leukocytopenia (6, 7 dpi)	Thrombocytopenia (6, 7 dpi)	ALP↑ (3 dpi), ALT↑↑↑ (6 dpi), BUN↑ (6 dpi), CRE↑↑↑ (6 dpi),	Died, 6 dpi

Author Manuscript

numbers of WBCs and platelets, respectively. Leukocytosis and thrombocytosis were defined as a twofold or greater increase in numbers of WBCs and platelets over baseline, where WBC count >11.000. petechiae covering 10 to 40% of the skin, and severe rash as areas of petechiae and/or ecchymosis covering >40% of the skin. Leukocytopenia and thrombocytopenia were defined as a >30% decrease in Hypothermia was defined as below 35°C. Fever was defined as >1.0°C higher than baseline. Mild rash was defined as focal areas of petechiae covering <10% of the skin, moderate rash as areas of 1, two- to threefold increase; 11, four- to fivefold increase; 11, two- to threefold decrease; 11, four- to fivefold decrease; albumin; AMY, amylase; TBIL, total bilirubin; BUN, blood urea nitrogen; PHOS, phosphate; CRE, creatinine; GLU, glucose; GLOB, globulin.

_
b
S
0
-
_ <u>`</u> .
_ <u></u>
_

Author Manuscript

Qiu et al.

က	
Ð	
P	
Та	

Clinical findings of EBOV-infected NHPs from 1 to 28 dpi.

				Clinical	Clinical findings		
Animal ID	I reatment group	Body temperature	Rash	White blood cells	Platelets	Biochemistry	Outcome
DI	50 mg/kg ZMapp ^{ni,} 3 dpi	Fever (3, 6, 14, 21 dpi)		Leukocytosis (3, 6, 21 dpi)	Thrombocytopenia (3, 6, 9, 14, 21 dpi)	ALB↓ (14, 21 dpi), ALP↓ (9, 14, 21, 28 dpi), AMY↓ (9 dpi), GLOB↑ (21, 28 dpi)	Survived
D2	$50 \text{ mg/kg ZMapp}^{TM}$, 3 dpi			Leukocytopenia (21, 28 dpi)	Thrombocytopenia (28 dpi)	(iqb (9 dpi)	Survived
D3	$50 \text{ mg/kg ZMapp}^{\text{TM}}$, 3 dpi	Fever (3 dpi)		Leukocytosis (3, 14 dpi)	Thrombocytopenia (3, 21, 28 dpi)	ALT↓ (6 dpi)	Survived
D4	$50 \text{ mg/kg ZMapp}^{TM}$, 3 dpi			Leukocytopenia (14 dpi)	Thrombocytopenia (14, 21 dpi)	ALT↓ (9 dpi), CRE↑ (14 dpi)	Survived
D5	50 mg/kg ZMapp [™] , 3 dpi	Fever (3 dpi)		Leukocytopenia (21, 28 dpi)	Thrombocytopenia (6, 9 dpi)	ALB↓ (9 dpi), BUN↓ (3, 6, 14, 21, 28 dpi)	Survived
D6	$50 \text{ mg/kg ZMapp}^{TM}$, 3 dpi				Thrombocytopenia (6 dpi)		Survived
EI	50 mg/kg ZMapp TM , 4 dpi				Thrombocytopenia (4, 7, 21 dpi)	AMY↓↓ (4, 21 dpi), AMY↓ (7, 10, 14 dpi), CRE↓ (21, 28 dpi)	Survived
E2	50 mg/kg ZMapp [™] , 4 dpi	Fever (4 dpi)		Leukocytosis (4, 10 dpi)	Thrombocytopenia (4, 7, 10, 21 dpi)	$ALT \downarrow \downarrow (4 dpi), GLU^{\uparrow} (4 dpi)$	Survived
E3	$50 \text{ mg/kg ZMapp}^{TM}$, 4 dpi	Fever (4 dpi)		Leukocytosis (4, 10 dpi)	Thrombocytopenia (7, 10, 14 dpi)	CRE↓ (14 dpi)	Survived
E4	50 mg/kg ZMapp ^{n,} , 4 dpi		Severe rash (5, 6, 7, 8 dpi), Mild rash (9 dpi)	Leukocytosis (10, 14, 21, 28 dpi)	Thrombocytopenia (4, 7, 10, 14 dpi)	$\begin{array}{c} ALP\uparrow \left(7, 10, 14 \text{ dpi}\right), ALT \\ \uparrow\uparrow\uparrow \left(7 \text{ dpi}\right), ALT \uparrow\uparrow \left(10 \text{ dpi}\right), \\ AMY\downarrow \left(4, 7, 10 \text{ dpi}\right), \\ TBIL\uparrow\uparrow\uparrow \left(7 \text{ dpi}\right), TBIL\uparrow\uparrow \left(10, 114 \text{ dpi}\right), \\ 14 \text{ dpi}\right), PHOS\downarrow \left(7, 10 \text{ dpi}\right), K \\ \downarrow\downarrow \left(4 \text{ dpi}\right) \end{array}$	Survived
ES	50 mg/kg ZMapp TM , 4 dpi	Fever (7 dpi)		Leukocytosis (4 dpi)	Thrombocytopenia (4, 7, 10, 14 dpi)	ALT† (7 dpi), AMY↓ (4, 7 dpi), PHOS↓ (10 dpi)	Survived
E6	50 mg/kg ZMapp ^{nx} , 4 dpi	Fever (4 dpi)	Mild rash (7, 8, 9 dpi)	Leukocytosis (4, 10, 14 dpi)	Thrombocytopenia (4, 7, 10, 14 dpi)	ALPf (7, 10 dpi), ALT $\uparrow\uparrow\uparrow\uparrow$ (7, 10 dpi), ALT $\uparrow\uparrow\uparrow\uparrow$ (7, 10, 10, 14 dpi), AMY4 (7, 10 dpi), TBIL $\uparrow\uparrow\uparrow$ (10 dpi), TBIL $\uparrow\uparrow\uparrow$ (10 dpi), (10 dpi), TBIL $\uparrow\uparrow\uparrow$ (10 dpi), CBIL $\uparrow\uparrow\uparrow$ (10 dpi), 0 dpi), GLOB† (21 dpi)	Survived
F1	50 mg/kg ZMapp [™] , 5 dpi			Leukocytosis (11 dpi)	Thrombocytopenia (3, 5, 8, 11 dpi)	AMY↓ (5 dpi), PHOS↓ (11 dpi), CRE↓ (28 dpi)	Survived
F2	50 mg/kg ZMapp ^{n,} , 5 dpi	Fever (3, 5 dpi)	Mild rash (8 dpi)	Leukocytosis (3, 5, 11 dpi)	Thrombocytopenia (3, 5, 8, 11, 14, 21 dpi)	PHOS↓ (11 dpi), CRE↓↓ (11 dpi)	Survived
F3	50 mg/kg ZMapp TM , 5 dpi			Leukocytopenia (8 dpi), Leukocytosis (3 dpi)	Thrombocytopenia (5, 8, 11, 21 dpi	ALT† (8 dpi), CRE↓↓ (14 dpi)	Survived

\geq
Ę
5
9
2
≦ a
Man
Manu
Manusc
Manus

	E			Clinica	Clinical findings		
Animai IU	r reaument group	Body temperature	Rash	White blood cells	Platelets	Biochemistry	Outcome
F4	$50 \text{ mg/kg ZMapp}^{\text{TM}}$, 5 dpi	Fever (3, 5 dpi)		Leukocytopenia (8 dpi)	Thrombocytopenia (5, 8, 11, 28 dpi)	PHOS↓ (8 dpi)	Survived
FS	$50 \text{ mg/kg ZMapp}^{\text{TM}}$, 5 dpi	Fever (3 dpi)		Leukocytosis (3, 11, 14 dpi)	Thrombocytopenia (5, 8, 11 dpi)	PHOS↓ (5, 8 dpi), CRE↓ (8, 11, 21, 28 dpi)	Survived
F6	$50 \text{ mg/kg ZMapp}^{\text{TM}}$, 5 dpi	Fever (3 dpi)		Leukocytopenia (8, 21, 28 dpi)	Thrombocytopenia (8, 11, 21 dpi)	PHOS↓ (5, 8, 11 dpi), GLU↑ (5 dpi)	Survived
61	PBS, 4 dpi		Severe rash (4 dpi)	Leukocytopenia (4 dpi)	Thrombocytopenia (4 dpi)	AMY↓ (4 dpi)	Died, 4 dpi
G2	Control mAb, 4 dpi		Severe rash (8 dpi)	Leukocytopenia (7, 8 dpi)	Thrombocytopenia (4, 7, 8 dpi)	ALP \uparrow (8 dpi), ALT \uparrow (7 dpi), ALT $\uparrow\uparrow\uparrow$ (8 dpi), CRE \uparrow (8 dpi), dpi)	Died, 8 dpi
G3	PBS, 4 dpi	Fever (4, 8 dpi)	Severe rash (8 dpi)	Leukocytopenia (7, 8 dpi)	Thrombocytopenia (4, 7, 8 dpi)	ALP↑ (8 dpi), ALT↑ (7, 8 dpi), AMY↓ (7 dpi), AMY ↓↓ (8 dpi), TBIL↑ (8 dpi), PHOS↓ (7 dpi)	Died, 8 dpi

Hypothermia was defined as below 35°C. Fever was defined as >1.0°C higher than baseline. Mild rash was defined as focal areas of petechiae covering <10% of the skin, moderate rash was defined as areas of petechiae covering 10 to 40% of the skin, and severe rash was defined as areas of petechiae and/or ecchymosis covering >40% of the skin. Leukocytopenia and thrombocytopenia were defined as a >30% decrease in the numbers of WBCs and platelets, respectively. Leukocytosis and thrombocytosis were defined as a twofold or greater increase in numbers of WBCs and platelets above baseline, where WBC fivefold decrease. ALB, albumin; ALP, alkaline phosphatase; ALT, alanine aminotransferase; AMY, amylase; TBIL, total bilirubin; BUN, blood urea nitrogen; PHOS, phosphate; CRE, creatinine; GLU, counts are greater than 11.0. \uparrow , two- to threefold increase; $\uparrow\uparrow\uparrow$, four- to fivefold increase; $\uparrow\uparrow\uparrow$, greater than fivefold increase; $\downarrow\downarrow$, two- to threefold decrease; $\downarrow\downarrow$, four- to fivefold decrease; $\downarrow\downarrow\downarrow$, greater than glucose; K⁺, potassium; GLOB, globulin.

Author
Manuscript

Extended Data Table 1

Blood viremia measured by RT-qPCR for the ZMapp1- and ZMapp2-treated NHPs.

DAY	A1	A2	A3	A4	A5	9 6	B1	B2	B3	B4	B5	B6	C1	C3
0	ΠD	ΩŊ	Ð	GD	ΠD	ΩŊ	ΩŊ	ΩŊ	đŋ	Ð	ΠD	ΠŊ	Ð	ΩŊ
3	UD	3.98E+02 UD	Ð	Ð	9.99E+02	9.99E+02 1.27E+03		8.05E+03 1.65E+04	9.36E+03 9.77E+03	9.77E+03	9.27E+02	9.48E+02	Ð	4.34E+02
9	3.10E+03	3.10E+03 4.49E+02 UD	Ð	8.34E+02	5.81E+03	2.09E+03	Ð	1.22E+04	1.22E+04 1.04E+05 4.26E+03 3.14E+02	4.26E+03	3.14E+02	4.49E+03	5.57E+06	2.05E+07
7													5.50E+05	
6		UD	Ð	Ð	Π	5.24E+02	Ð	1.74E+05	5.03E+05	5.03E+05 1.87E+03	5.16E+02	ΠD		
14	14 3.62E+03	ΩŊ	Ð	Ð	Π	ΠD	Ð	Ð		Ð	ΠŊ	Π		
21	UD	ΠD	Ð	Ð	ΩŊ	ΠD	Ð	Ð		CD	ΠŊ	Π		
27	UD	UD	Ð	Ð	ΠD	ΠD	Ð	Ð		Ð	ΠD	ΠD		
Footnote	Footnote: UD = Undetectable.	stectable.												

∇
-
Autho
=
-
<u></u>
0
<
цц,
7
<u> </u>
S
0
Manuscri
0
Ť.

Extended Data Table 5

Blood viremia measured by RT-qPCR for the ZMapp $^{\mbox{\tiny TM}}\xspace$ -treated NHPs.

DAYS	A1	A2	A3	A4	A5	9V	B1	B2	B3	B4	B5	B6	CI	C3	C3	C4	C5	C6	DI	D2	D3
0		-	-	-	-	-	-	-	-	-	-	1	-	-	-	-	-	-		1	-
3	676.08	165.96	10233	2884	812.83	10965							1047.1			1122	3235.9 1148.2	1148.2			
4							85114	128825	23442	1E+06	1E+06	323594							31623	144544	1E+06
5													380189	3E+06	109648	58884	2E+06	69183			
9	70795	446.68	1230.3	316.23	32359	6E+06															
7							154882	257040	1380.4	63096	588844	363078							645654	812831	
8													3715.4	28184	29512	1862.1	72444	5888.4	158489	18621	
6	31623	-	-	15136	275.42	165959															
10							1	1071.5	-	1318.3	6166	5248.1									
Π													1	524.81	-	1	81.283	-			
14	398.11		-	-	-	-1	-	-		-	-	239.88	1	1	-	1	1	-			
21	1	-	-	1	-	1	-	-	-	-	-	1	-	1	-	-	-	-			
28	1	-	1	1	-	1	-	1	-	-	-	1	1	1	-	1	1	-			