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Abstract: Zika virus (ZIKV), a mosquito-borne flavivirus, causes devastating congenital birth defects. We isolated a human monoclonal antibody (mAb), ZKA190, that potently cross-neutralizes multi-lineage ZIKV strains. ZKA190 is highly effective in vivo in preventing morbidity and mortality of ZIKV-infected mice. NMR and cryo-electron microscopy show its binding to an exposed epitope on DIII of the E protein. ZKA190 Fab binds all 180 E protein copies, altering the virus quaternary arrangement and surface curvature. However, ZIKV escape mutants emerged in vitro and in vivo in the presence of ZKA190, as well as of other neutralizing mAbs. To counter this problem, we developed a bispecific antibody (FIT-1) comprising ZKA190 and a second mAb specific for DII of E protein. In addition to retaining high in vitro and in vivo potencies, FIT-1 robustly prevented viral escape, warranting its development as a ZIKV immunotherapy.

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A human bi-specific antibody against Zika virus with high therapeutic potential

Jiaqi Wang^{1,2,11}, Marco Bardelli^{3,11}, Diego A. Espinosa⁴, Mattia Pedotti³, Thiam-Seng Ng^{1,2}, Siro Bianchi⁵, Luca Simonelli³, Elisa X.Y. Lim^{1,2}, Mathilde Foglierini³, Fabrizia Zatta⁵, Stefano Jaconi⁵, Martina Beltramello⁵, Elisabetta Cameroni⁵, Guntur Fibriansah^{1,2}, Jian Shi^{2,6}, Taylor Barca⁴, Isabel Pagani⁷, Alicia Rubio⁷, Vania Broccoli^{7,8}, Elisa Vicenzi⁷, Victoria Graham⁹, Steven Pullan⁹, Stuart Dowall⁹, Roger Hewson⁹, Simon Jurt¹⁰, Oliver Zerbe¹⁰, Karin Stettler⁵, Antonio Lanzavecchia³, Federica Sallusto³, Andrea Cavalli³, Eva Harris⁴, Shee-Mei Lok^{1,2,12}, Luca Varani^{3,12}, Davide Corti^{5,12,13}

¹Program in Emerging Infectious Diseases, Duke–National University of Singapore Medical School, Singapore 169857, Singapore

²Centre for BioImaging Sciences, National University of Singapore, Singapore 117557, Singapore

³Institute for Research in Biomedicine, Università della Svizzera italiana, Via Vincenzo Vela 6, 6500 Bellinzona, Switzerland

⁴Division of Infectious Diseases and Vaccinology, School of Public Health, University of California, Berkeley, 185 Li Ka Shing Center, 1951 Oxford Street, Berkeley, California, 94720-3370, United States of America

⁵Humabs BioMed SA, Via Mirasole 1, 6500 Bellinzona, Switzerland

⁶CryoEM unit, Department of Biological Sciences, National University of Singapore, Singapore 117557.

⁷Viral Pathogens and Biosafety Unit, San Raffaele Scientific Institute, Via Olgettina 58, 20132 Milan, Italy

⁸CNR-Institute of Neuroscience, Via Vanvitelli 32, 20129, Milan, Italy

⁹National Infection Service, Public Health England, Porton Down, Salisbury, Wiltshire, United Kingdom

¹⁰Department of Chemistry, University of Zurich, Zurich, Switzerland

¹¹These authors contributed equally

¹²These authors contributed equally

¹³Lead Contact

*Correspondence: sheemei.lok@duke-nus.edu.sg, luca.varani@irb.usi.ch, davide.corti@humabs.ch

SUMMARY

Zika virus (ZIKV), a mosquito-borne flavivirus, causes devastating congenital birth defects. We isolated a human monoclonal antibody (mAb), ZKA190, that potently cross-neutralizes multi-lineage ZIKV strains. ZKA190 is highly effective *in vivo* in preventing morbidity and mortality of ZIKV-infected mice. NMR and cryo-electron microscopy show its binding to an exposed epitope on DIII of the E protein. ZKA190 Fab binds all 180 E protein copies, altering the virus quaternary arrangement and surface curvature. However, ZIKV escape mutants emerged *in vitro* and *in vivo* in the presence of ZKA190, as well as of other neutralizing mAbs. To counter this problem, we developed a bispecific antibody (FIT-1) comprising ZKA190 and a second mAb specific for DII of E protein. In addition to retaining high *in vitro* and *in vivo* potencies, FIT-1 robustly prevented viral escape, warranting its development as a ZIKV immunotherapy.

INTRODUCTION

Effective therapeutics and vaccines against Zika virus (ZIKV) are urgently needed. Although ZIKV generally causes asymptomatic or mild disease, it can lead to neurological complications both in adulthood and during fetal development (Honein et al., 2016), likely due to infection of neural progenitor cells (Frontera and da Silva, 2016; Mlakar et al., 2016). Further, infection of the testes has been shown to cause infertility in mice (Govero et al., 2016).

Cryo-electron microscopy (cryoEM) of ZIKV revealed high structural similarity to other flaviviruses (Kostyuchenko et al., 2016; Sirohi et al., 2016). The smooth-surfaced mature ZIKV particle comprises 180 copies of E protein organized in icosahedral symmetry, each asymmetric unit containing three E proteins. Three E protein dimers lie parallel to each other building a raft, and thirty rafts form a herringbone pattern. The E protein consists of three domains – DI, DII and DIII (Barba-Spaeth et al., 2016; Kostyuchenko et al., 2016; Rey et al., 1995; Zhao et al., 2016). DIII contains the receptor-binding site and mediates virus attachment (Crill and Roehrig, 2001), whereas the fusion loop in DII facilitates virus-endosomal membrane fusion (Allison et al., 2001). After cell entry, the low endosomal pH triggers dramatic structural rearrangement of E protein dimers into trimers, leading to membrane fusion (Bressanelli et al., 2004; Modis et al., 2004; Stiasny et al., 1996). The post-fusion structure shows DIII rotated $\sim 70^\circ$ relative to DI, which requires movement of the DI-DIII linker (Bressanelli et al., 2004; Modis et al., 2004).

We recently isolated a panel of anti-ZIKV neutralizing human mAbs and identified a DIII-binding mAb, ZKA190, that showed strong neutralizing activity across ZIKV strains (Stettler et al., 2016). Here, we further characterized ZKA190 *in vitro* and *in vivo*. Using a combination of NMR, cryoEM and molecular simulations, we mapped the ZKA190 epitope to an exposed region consisting of the DI-DIII linker and the lateral ridge (LR) region of DIII. Different from other DIII antibodies, ZKA190 Fab binds to all 180 E proteins on the virus surface, altering the E protein quaternary interactions, causing separation of the rafts and flattening their curvature. *In vivo* studies of ZIKV infection in lethal mouse models demonstrated strong protection of ZKA190 from multiple

strains. To minimize the risks of viral escape and antibody-mediated disease enhancement, we engineered a bispecific antibody, which combines ZKA190 with the DII-binding mAb ZKA185 and also includes a mutation in the Fc region (LALA) abolishing binding to Fc γ receptor and complement (Hessell et al., 2017). FIT-1 retains the breadth and potencies of both parental antibodies and was not observed to generate escape mutants, encouraging FIT-1 development as a ZIKV therapeutic.

RESULTS

In vitro characterization of the DIII-specific mAb ZKA190

Recombinant ZKA190 synthesized in the IgG1 wild-type (wt) and IgG Fc-LALA format potently neutralized African, Asian and American ZIKV strains, with IC₅₀ ranging from 0.004 to 1.5 nM (**Figure 1A**; 0.6 to 8 ng/ml). Although both the ZKA190 IgG and its Fab bound to E protein with similar affinities (**Figure 1C**; K_D values of 0.3 and 0.2 nM, respectively), the Fab neutralized ZIKV infection less potently with a 300-fold higher IC₅₀ (**Figure 1B**). Since ZIKV has been shown to infect human neural progenitor cells (hNPC) (Garcez et al., 2016; Nowakowski et al., 2016; Onorati et al., 2016; Tang et al., 2016), we tested ZKA190 wt and LALA, and both fully abolished infection and replication of ZIKV in hNPCs (**Figure 1D**).

Antibody-dependent enhancement (ADE) of infection may contribute to disease severity in heterologous secondary DENV infections (Halstead, 2014), caused by the binding of non-neutralizing or sub-neutralizing concentrations of E- and prM-specific antibodies to mature, partially mature and fully immature viruses. ZKA190 supports ADE in K562 cells from 0.0001 to 1 nM, while ZKA190-LALA did not show any ADE activity (**Figure 1E**) and completely inhibited ADE induced by plasma antibodies from four ZIKV-immune donors (**Figure 1F**).

Anti-prM antibodies are elicited during the human immune response against flaviviruses, and they enhance *in vitro* infection of immature or partially immature virions (Dejnirattisai et al., 2010). ZKA190-LALA can fully block ADE of ZIKV infection induced by a flavivirus cross-

reactive anti-prM mAb, DV62, isolated from a dengue patient (Beltramello et al., 2010) (**Figure 1G**). Finally, we compared the ability of ZKA190 wt, LALA and Fab to cause or block ADE of ZIKV by human anti-DENV2 plasma or DV62 mAb (**Figure 1H**). Low concentrations of ZKA190 increased DV62-mediated ADE of ZIKV infection, consistent with its ability to promote the entry of both immature and mature virions, while at concentrations above 1.3 nM (i.e., 200 ng/ml), ZKA190 blocked ADE induced by both anti-DENV plasma and DV62. At concentrations above 1.6 nM, ZKA190 Fab was equally effective as ZKA190-LALA in reducing ADE induced by DV62, indicating that inhibition occurs at a post-attachment step, likely fusion.

ZKA190 binds to a conserved and highly accessible region of DIII

The ZKA190 epitope was determined by solution NMR spectroscopy (Bardelli et al., 2015; Simonelli et al., 2010; 2013). Differences in the NMR spectra of free and mAb-bound DIII identified DIII residues affected by ZKA190 binding, either directly or through allosteric effects (**Figure S1A**). The ZKA190 epitope consists of the LR of DIII (the BC, DE and FG loops), as well as part of the DI-DIII linker (**Figure 2A**). These residues are conserved among 217 known ZIKV strains, with the exception of V341I and E393D substitutions in the Uganda 1947 and the MR766 strains (**Figure S1D**), which ZKA190 was shown to neutralize (**Figure 1A**). The ZKA190 epitope is highly accessible on the ZIKV surface, except for the FG loop in the 5-fold vertex (**Figure 2B and S1C, molecule A**).

Computational docking followed by molecular dynamics simulation, guided and validated by both NMR- and cryoEM-derived epitope information as well as DIII mutagenesis, showed that ZKA190 binds through an interface characterized by shape and charge complementarity (**Figure 2B and S1E**). The computational model fits well to the cryoEM density map of ZIKV-ZKA190 Fab complex (**Figure 2C and 3D**). Docking suggests that there are no direct contacts between ZKA190 and the FG loop on DIII, thus changes in NMR signals likely derive from allosteric effects.

Consistently, mutations of FG loop residues, but not of other NMR-affected residues in the recombinant DIII, did not affect the binding affinity of ZKA190 (**Figure 2B and S2**).

CryoEM structure of ZKA190 complexed with ZIKV

CryoEM micrographs showed that ZKA190 binds to the virus at 4°C, 37°C and 40°C (**Figure S3A**). Structures of ZIKV complexed with ZKA190 Fab at 4°C and 37°C were determined to ~22 Å resolution and found to be similar (**Fig 2D, S3B and S3C**). At this resolution, the Fabs and E protein rafts can only be roughly fitted into the density; the footprint on DIII cannot be accurately evaluated using cryoEM results alone. In contrast to other mAbs against DIII LR of other flaviviruses, Fab ZKA190 bound to all 180 copies of E protein on the viral surface (**Figure 2D and S3**), achieving full occupancy.

Superposition of the structure of the ZKA190 Fab:E protein complex onto the uncomplexed ZIKV structure showed that Fabs bound to the E protein molecule around the 5-fold vertex clashed with each other (**Figure 2G**). On the contrary, cryoEM data showed equally strong densities of the variable region of the Fabs at all three individual E proteins in the asymmetric unit (**Figure 2F and S4**). To accommodate binding of all five Fabs without clashes, the E proteins have to separate from each other in the 5-fold vertex (**Figure 2G**). Comparison of the cryoEM structures of uncomplexed and ZKA190-bound ZIKV showed that in the latter, the E protein rafts move to a higher radius (**Figure 2E**) and are separated from each other (**Figure 2I**); the curvature of the rafts is also flattened (**Figure 2H**). In the uncomplexed ZIKV cryoEM structure, the E protein ectodomain interacts with the membrane-associated stem regions of E and M proteins on the virus, and this interaction might be disrupted by ZKA190 Fab binding, as suggested by the spatial gap observed between the E protein and the lipid bilayer (**Figure 2E**). The dissociation may increase the motion of the rafts, explaining the inability to obtain a higher resolution cryoEM map.

Mechanisms of ZKA190 neutralization

The ability of ZKA190 to efficiently neutralize the virus may involve inhibition of either cell attachment or membrane fusion. Another mechanism might involve virus inactivation through aggregation. Virus aggregation due to simultaneously engaging epitopes on different particles by ZKA190 was assessed by dynamic light scattering (DLS). Particles with size similar to single virus particle were detected at low mAb concentration, whereas large viral aggregates appeared at higher concentration (≥ 33 antibodies per virus) and correlated with neutralizing activity (**Figure 3A**). One caveat is that the contribution of aggregation to neutralization may be different *in vivo*, as the amount of virus could differ.

IgG ZKA190 may also engage neighboring E proteins on the viral surface. As shown by cryoEM, the distance between the Fabs' CH1 regions bound on the viral surface is compatible with that of a full IgG (**Figure 3B**). However, bivalent binding of a full IgG depends not only on the distance between two Fab molecules, but also on their relative orientation. Full atomistic molecular dynamics simulations of a homology model of ZKA190 IgG showed that a single ZKA190 molecule can bridge adjacent E proteins around the 5- and 3-fold vertices, but not the 2-fold vertex (**Figure 3C and 3D**). The Fab regions of the modelled ZKA190 IgG correlated well with the cryoEM structure (**Figure 3D**). This suggests that ZKA190 cross-linking of neighboring E proteins might prevent the E dimers from rearrangement into trimers that is required for membrane fusion. Both proposed mechanisms imply bivalent binding of ZKA190 IgG. Since ZKA190 Fab also neutralizes ZIKV, albeit less efficiently than the full IgG, additional mechanisms may be involved. ZKA190 might prevent the attachment of ZIKV to target cells or viral fusion to the endosomal membrane.

Neutralization at a post-attachment step is supported by data showing that ZKA190 inhibits *in vitro* ZIKV infection with similar potency regardless of whether the mAb was added to the virus pre- or post- attachment to cells (**Figure 3E**). This result is further supported by the qRT-PCR experiments that detect the amount of virus on the cell surface, which does not decrease at increasing antibody concentrations (**Figure 3F**), but rather increased compared to the virus-only

control, likely due to antibody-mediated virus aggregation (**Figure 3A**). Moreover, confocal microscopy analysis showed that ZIKV-complexed ZKA190 Fab can enter cells at neutralizing concentrations exceeding the IC₅₀ by 10,000-fold (**Figure 3G**).

***In vivo* characterization of the DIII-specific mAb ZKA190**

ZKA190 and ZKA190-LALA were tested in A129 mice challenged with a lethal dose of ZIKV strain MP1751 (African lineage). Prophylaxis with ZKA190 and ZKA190-LALA one day before virus challenge protected mice from mortality and morbidity at mAb concentrations of 5, 1 or 0.2 mg/kg (**Figure 4A-B**). ZKA190-LALA, and to a lesser extent ZKA190, delayed morbidity and mortality as compared to the control at 0.04 mg/kg. Viral titers in blood and organs were reduced significantly, even in the presence of serum antibody levels below 1 µg/ml (**Figure S5A-D**). To determine if low doses of ZKA190 could potentially enhance ZIKV infection, we administered 1 mg/kg, 0.04 mg/kg, 0.02 mg/kg and 0.01 mg/kg of ZKA190 to AG129 mice and challenged them with a recently isolated ZIKV strain from Nicaragua (Nica 2-16, American lineage). Mice receiving 1 mg/kg of ZKA190 or ZKA190-LALA were completely protected from ZIKV disease (**Fig 4C-D**). ZKA190 or ZKA190-LALA at 0.04 mg/kg and 0.02 mg/kg of partially protected mice from mortality and morbidity, and none of the lower antibody doses resulted in enhanced disease or infection. Blood viral titers on day 3 post-infection were reduced compared to control antibody-treated animals in a dose-dependent fashion (**Figure S5E**).

To evaluate therapeutic potential, we administered ZKA190 and ZKA190-LALA to A129 mice at different time-points following ZIKV infection. At a dose of 15 mg/kg, survival rates of 80%-100% were achieved, and morbidity was greatly reduced even when treatment was given four days post-infection (**Figure 4E-G**). ZKA190 and ZKA190-LALA treatment at all post-infection time-points resulted in significantly reduced viral titers with a clear trend for greater reduction with earlier treatment (**Figure S6A-S5C**). Of note, ZKA190-LALA showed a reduced antiviral activity

on day 5 as compared to ZKA190 when mAbs were given four days post-infection, a result that might relate to the impaired ability of the LALA mAb to facilitate rapid clearance of coated virions.

Of the 16 treated mice, one *in vivo* escape mutant (Monoclonal Antibody Resistant Mutant 1, MARM1) was isolated containing the amino acid substitution T335R in DIII, which was shown to abrogate ZKA190 binding (**Figure S2**). Viruses from the other treated mice did not contain any E mutations (**Figure S6D-S5E**).

***In vitro* selection of ZIKV escape mutants.**

To assess the ability of ZKA190 to select for resistant mutants (MARMs) *in vitro*, ZIKV (H/PF/2013) was passaged in the presence of sub-neutralizing concentrations of ZKA190. MARM2, carrying the DIII E370K mutation, was isolated after three rounds of selection. The mutation abolished neutralization by ZKA190, but not its binding to DIII (**Figure S2**). The mutations in the *in vivo* (T335R) and *in vitro* (E370K) MARMs are located on the BC and DE loops of DIII, respectively, and are consistent with the epitope identified by NMR.

Development of the bispecific FIT-1 antibody

Generation of viral escape mutants can render antibodies useless for treatment. This may be prevented by making a bispecific antibody combining two highly neutralizing mAbs, each specific for a distinct epitope. We selected ZKA185 and ZKA230, which potently neutralize African, Asian and American strains with IC₅₀ ranging from 0.02 to 0.62 nM (**Figure 5A**). ZKA185 binds with high affinity to recombinant ZIKV E protein and to Zika virus-like particles (VLP) but not to the isolated DIII (**Figure 5B**). ZKA230 bound to ZIKV VLPs, but not to recombinant E or DIII, suggesting that it recognizes a quaternary epitope displayed only on the viral surface (**Figure 5B**). ZKA185 IgG and Fab were shown to bind to E and VLP antigens with similar high affinity by ELISA. Conversely, ZKA230 Fab bound to VLPs with markedly reduced affinity compared to its IgG.

We isolated MARMs of ZKA185 (MARM3) and ZKA230 (MARM4) by passaging virus in the presence of sub-neutralizing mAb concentrations. MARM3 contained substitutions at K84E and D67H on DII (**Figure 5D**). MARM4 showed a mixture of different amino acid substitutions at position 84 (from K to G, E or R). ZKA190 neutralized ZKA185 and ZKA230 MARMs as well as the parental virus (**Figure 5C**). ZKA185 neutralized both ZKA190 and ZKA230 MARMs. In contrast, ZKA230 neutralized only ZKA190 MARM2.

To gain insight into the development of MARMs capable of escaping from the pressure of multiple mAbs, we serially passaged ZKA190 MARM2 (E370K) in the presence of ZKA185 or ZK230 and found that double MARMs emerged after 3 to 4 passages. ZKA230 introduced the K84E mutation while ZKA185 selected for the D76G mutation. These findings indicate that ZIKV can escape multiple mAbs targeting distinct sites when the selection is performed in a stepwise fashion.

MAb ZKA185 was selected to combine with ZKA190 for the development of a bispecific antibody since it potently cross-neutralizes ZIKV strains and does not compete with ZKA190. We produced a bispecific antibody in the tetravalent symmetric format Fabs-in-tandem-Ig (FIT-Ig) (Gong et al., 2017). ZKA190 was placed in the outer and ZKA185 in the inner Fab position (**Figure 5E**), using the backbone IgG1 antibody in LALA format (here designated FIT-1). FIT-1 bound DIII, E and VLPs in ELISA (**Figure 5F**) and retained high neutralizing potency against ZIKV strains, with IC_{50} values largely similar to those of the parental mAbs (**Figure 5G**). FIT-1 bound to E protein with higher affinity than either of the parental mAbs (K_D values: ZKA185 1.8 nM, ZKA190 9.3 nM and FIT-1 $K_D < 1$ pM due to slower dissociation rate, presumably through avidity effects) and effectively neutralized all the ZKA190, ZKA185 and ZKA230 MARMs (**Figure 5H**). In addition, the neutralizing activity of the Fab fragment of FIT-1 (comprising one ZKA190 and one ZKA185) was reduced only by about 6-fold (**Figure 5I**, right panel), whereas those of the individual ZKA190 and ZKA185 Fabs were reduced by about 100-fold compared to their full-

length IgG1 (**Figure 1B** and **Figure 5I**). Taken together, these findings suggest the ZKA190 and ZKA185 moieties in FIT-1 are both active.

No MARMs were isolated despite 8 rounds of serial passages in the presence of FIT-1. In contrast, MARMs appeared after 3 to 4 passages with individual mAbs. Simultaneous mutations in both DIII and DII are thus unlikely to occur, suggesting that FIT-1 is a safer therapeutic. Confocal microscopy studies showed that FIT-1, like ZKA190, inhibits virus infection at a post-attachment step, likely fusion (**Figure 5J**). Finally, FIT-1 and its Fab blocked ADE of human anti-DENV2 plasma or DV62 at concentrations above 0.1 and 10 nM, respectively (**Figure 5K**).

We evaluated the therapeutic potential of FIT-1 by administering three different doses (15, 5 and 1 mg/kg) at three different time-points to mice following ZIKV infection. At a dose of 15 mg/kg, survival rates were 100% without signs of morbidity even when treatment was given three days post-infection (**Figure 6**); viral titers were abrogated and no escape mutants were detected, indicating high *in vivo* efficacy. Administration of 5 mg/kg resulted in survival rates of 70-100%; again, no escape mutants were detected on day 5 after infection. The lowest dose tested (1 mg/kg) protected when administered on day 1 or 2, but not day 3, after infection.

DISCUSSION

The highly potent ZKA190 mAb binds to the lateral ridge in DIII of the E protein in a unique orientation compared to other flavivirus-neutralizing LR-binding mAbs (Edeling et al., 2014; Nybakken et al., 2005; Zhao et al., 2016) (**Figure 7A**), resulting in full occupancy of all 180 E proteins. Unlike ZKA190, the cryoEM structures of E16:WNV (Kaufmann et al., 2006) and ZV23:ZIKV (Wang et al., 2016) showed that their Fabs are unable to bind to DIII around the 5-fold vertex. Superposition of the E106:DIII (Edeling et al., 2014), ZV23:DIII and ZV67:DIII (Zhao et al., 2016) structures onto an uncomplexed ZIKV cryoEM structure showed clashes with the neighboring E proteins around the 5-fold vertex (**Figure 7B**).

Our previous structure of uncomplexed ZIKV revealed that the E protein shell is more compact than that of DENV (Kostyuchenko et al., 2016). Binding of ZKA190 disrupts the tight interactions between E proteins in the 5-fold vertices at both 4°C and 37°C, thus allowing full occupancy. ZKA190 may either select a transient viral conformation where the 5-fold vertex is “open” (conformational selection) or cause surface structural changes (allosteric effects). The latter appears more likely since binding occurs even at 4°C, where the thermal motions required for E protein conformational changes are limited. NMR studies and DIII mutants indicated that binding of ZKA190 causes an allosteric effect on the DIII FG loop, which points towards the center of the 5-fold vertex. ZKA190 binding to one DIII near the 5-fold vertex may alter the FG loop conformation, opening the vertex and exposing additional ZKA190 binding sites. Alternatively, the antibody could first bind to the 3- and/or 2-fold vertices, causing allosteric effects throughout the viral surface resulting in opening of the 5-fold vertex, to which other antibodies could then bind; however, experimental evidence for this is lacking.

Multiple mechanisms may contribute to ZKA190 activity. We showed here that high concentrations of ZKA190 induced aggregation of virus particles by crosslinking different particles with the two Fab arms. This correlated with increased neutralization activity, suggesting a role for aggregation in viral neutralization. Molecular dynamics showed that the two Fab arms of ZKA190 can bind within a viral particle, possibly preventing the dramatic rearrangement of the E proteins on the virus surface required for fusion.

RT-PCR and confocal microscopy data indicate that the virus remains attached to cells in the presence of ZKA190 and is able to enter cells in the presence of neutralizing concentrations of ZKA190. Furthermore, similar neutralization profiles of ZKA190 (and FIT-1) in the post- and pre-attachment assays, as well as inhibition by the ZKA190 Fab fragment of DENV plasma and anti-prM antibody-mediated ADE, support the notion that ZKA190 inhibits a post-attachment step, likely fusion. The epitope of ZKA190 encompasses the LR of DIII and the DI-DIII linker region. In a normal infection, the low-pH endosomal environment provokes the E proteins to rearrange from a

dimeric structure to a trimeric post-fusion structure. Binding of ZKA190 to the DI-DIII linker and DIII may hinder the movement of DIII, preventing rearrangement and thus membrane fusion, as previously suggested in structural studies of E16:WNV (Kaufmann et al., 2006; Zhang et al., 2015).

Although antibody effector functions can enhance the efficacy of anti-flavivirus antibodies (Chung et al., 2006), comparison of ZKA190 and ZKA190-LALA *in vivo* efficacy indicates that Fc-dependent mechanisms, such as ADCC or CDC, likely do not play a major role in prophylaxis or therapy. Despite clear evidence of *in vitro* ADE by ZKA190, and in contrast to a recent report (Bardina et al., 2017), our *in vivo* results indicate that ZKA190-mediated disease enhancement does not occur even when the mAb is administered at doses expected to provide only partial neutralization.

Numerous flavivirus MARMs are readily isolated in *in vitro* systems (de Alwis et al., 2012; Zou et al., 2012), similarly to what we observed with ZKA190, ZKA185 and ZKA230 mAbs. However, only a few studies described isolation of MARMs *in vivo* (Lai et al., 2007). Here, we isolated a ZKA190 MARM from one animal treated therapeutically, although no MARMs were detected in animals treated prophylactically. To reduce the risk of emergence of escape mutants, we produced a bispecific antibody coupling ZKA190 to the anti-DII ZKA185 mAb. The FIT-1 bispecific antibody is highly potent *in vitro* and *in vivo* and did not generate escape mutants even after 8 passages.

Flavivirus particles in *in vitro* preparations represent different maturation states (mature, partially immature, and fully immature) (Lok, 2016). Only partially and fully immature virions have prM on their surfaces, and anti-prM antibodies can confer infectivity to the otherwise non-infectious immature virions (Rodenhuis-Zybert et al., 2010). Since anti-prM antibodies are elicited in anti-flavivirus immune responses (Dejnirattisai et al., 2010), they can contribute significantly to pathogenesis (Dejnirattisai et al., 2010). We showed here that ZKA190 and FIT-1 can block *in vitro* ADE mediated by the cross-reactive prM mAb DV62, indicating that ZKA190 can effectively block prM antibody-mediated infection by at least partially immature ZIKV particles.

Our current studies *in vitro* and in lethal adult mouse models showed that the FIT-1 bispecific antibody is a good candidate for development of antibody-based therapy for ZIKV infection. Further studies in pregnant rhesus macaques (Dudley et al., 2016; Nguyen et al., 2017) would enable testing FIT-1 efficacy in blocking fetal infection. The *in vivo* protective activity of FIT-1 indicates that alternative delivery mechanisms, such as DNA or RNA vectors, or half-life-extending modification of the Fc, might be achievable to make it cost-effective for the prevention of congenital ZIKV disease.

ACCESSION NUMBERS

Coordinates of Fab ZKA190:ZIKV complex incubated at 37°C was deposited in the Protein Data Bank under accession code 5Y0A and the NMR structure of DIII under PDB code 5OMZ and chemical shift used for structural calculation and mapping under BMRB code 34167. The cryoEM maps of Fab ZKA190:ZIKV complexes incubated at 4°C and 37°C were deposited in the Electron Microscopy Database under accession number EMD-6794 and EMD-6793, respectively.

Star Methods

Detailed methods are provided in the online version of this paper and include the following:

- Key resources table
- Contact for reagent and resource sharing
- Experimental model and subject details
- Methods details
- Statistical analysis

AUTHOR CONTRIBUTIONS

FZ and **SJ** produced and purified antibodies. **MF** produced figures and carried out bioinformatic analysis. **SB** performed *in vitro* cellular neutralization, ADE assays and MARMs selection. **DE**, **TB**, **VG**, **SP**, **SD** and **RH** designed and carried out *in vivo* studies. **JW**, **TN**, **EL**, **GF**, **JS** designed and performed structural cryoEM research. **MB**, **MP**, **LS** designed and performed NMR structural research, docking, molecular dynamics, mutagenesis, binding and immunofluorescence studies; **AC** designed and performed molecular dynamics simulations; **SJu** and **OZ** performed NMR studies. **IP**, **AR**, **VB** and **EV** designed and performed *in vitro* studies on hNPCs infection models. **KS** carried out binding studies, *in vitro* cellular neutralization, MARMs isolation and sequencing, analyzed the data and coordinated *in vitro* characterization activities. **EC** and **MB** provided supervision and contributed analyzing data. **FS** and **AL** edited the paper and provided supervision. **JW**, **EH**, **SL**, **LV** and **DC** designed research, analyzed data, and wrote the paper.

CONFLICT OF INTEREST

The authors declare conflict of interest. **AL** is the scientific founder of Humabs BioMed SA. **AL** holds shares in Humabs BioMed. **KS**, **FZ**, **SJ**, **EC**, **MB** and **DC** are employees of Humabs Biomed.

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Figure Legends

Figure 1. Neutralization and enhancement of ZIKV infection by ZKA190 mAb.

(A) ZKA190 was tested for neutralization of four strains of ZIKV, as determined by the percentage of infected Vero cells in the presence of increasing amounts of mAbs. Data are representative of at least two independent experiments. (B) Neutralization of ZIKV H/PF/2013 strain by ZKA190 IgG and Fab determined as in (A). (C) Binding of ZKA190 IgG and Fab to recombinant ZIKV E and DIII proteins as assessed by SPR. (D) Neutralization of ZIKV strain PRVABC59 infection of hNPCs by ZKA190, ZKA190-LALA and a control mAb as determined by plaque assay on Vero cells (left panel) and indirect immunofluorescence of infected hNPCs using fluorophore-labelled anti-E antibody (right panel). (E) ADE of ZIKV infection of non-permissive K562 cells by ZKA190 and ZKA190-LALA. (F) ADE induced in K562 cells when ZIKV is pre-incubated with serial dilutions of plasma or serum from different ZIKV-positive patients (left panel). When ZKA190 LALA is added to the ZIKV-serum complexes, ADE is inhibited (right panel). (G) ADE induced in K562 cells when ZIKV is pre-incubated with serial dilutions of a prM cross-reactive mAb (DV62) derived from a DENV-immune donor. ZKA190-LALA inhibits ADE of ZIKV when complexed with prM-reactive mAb DV62. (H) Effect on ADE induced by peak enhancing dilution of a DENV2 plasma sample (left panel) or anti-prM DV62 mAb (right panel) by serial dilutions of indicated mAbs.

Figure 2. ZKA190 Epitope Identified By NMR, CryoEM and Docking.

(A) Cartoon representation of the 12 lowest-energy NMR structures of ZIKV DIII, with residues affected by ZKA190 binding in red. Flexibility in the N-terminus of the construct is apparent. (B) Model of the ZKA190:DIII complex derived by computational docking and molecular simulation validated by cryoEM and NMR results. The NMR-identified epitope on DIII (grey) is indicated in red. The ZKA190 heavy and light chains are colored in dark and light green, respectively. DIII

residues that affect or do not affect antibody binding when mutated are shown as orange and blue sticks, respectively. (C) NMR-identified ZKA190 epitope (red) is accessible on the virus surface (white). (D-I) The $\sim 22\text{\AA}$ resolution cryoEM map of the ZIKV:ZKA190 Fab complex at 37°C. (D) Surface of the cryoEM map. One icosahedral asymmetric unit is indicated by a white triangle. Light and dark blue correspond to the E protein shell and the Fab densities, respectively. (E) A quarter of the central section of the uncomplexed (left) and complexed (right) cryoEM maps showing that the E protein layer in the latter moved outwards with the highest protrusion at the 5-fold vertex. (F) CryoEM structure of ZIKV:ZKA190 Fab complex. The DIIs of all E proteins are bound by Fabs (green). E proteins in an asymmetric unit are labelled A, B and C, and the neighboring asymmetric unit within a raft A', B' and C'. DI, DII and DIII are colored in red, yellow and blue, respectively. (G) E proteins at the 5-fold vertices move apart upon Fab ZKA190 binding. (Left) Superposition of the Fab ZKA190:E complex onto the uncomplexed ZIKV shows clashes between the Fabs. (Right) In contrast, no clashes are observed in the cryoEM complex structure. E proteins are in grey and Fabs are colored. (H) Superposition of the E protein raft in the free (grey) and ZKA190-complexed (colors) ZIKV; the curvature flattens in the complex structure. E proteins of one asymmetric unit (A, B and C) are in red, yellow and blue; those from another unit are in pink. (I) The rafts separate from each other in the complex (right) compared to the uncomplexed ZIKV structure (left). The inter-raft interface is indicated with a green dotted line, and the gap between rafts is indicated with an arrow. See also Figure S1, S2, S3 and S4.

Figure 3. ZKA190 aggregates virus at high antibody:virus molar ratios and inhibits a post-attachment stage of virus infection

(A) Dynamic light scattering experiments measuring the hydrodynamic size of the virus complexes shows ZKA190 IgG aggregates ZIKV at IgG:virion molar ratio of $\geq 33:1$. The neutralization activity of antibody correlates with the formation of virus-antibody aggregates. The neutralization is

expressed as the \log_{10} -transformed fold-reduction of the virus titer compared to the virus-only control. **(B)** ZKA190 Fab (green) is shown on the viral surface (grey). The close proximity ($<86\text{\AA}$) between CH1 domains of two Fabs (red lines) in the cryoEM ZIKV:ZKA190 Fab structure indicates possible bivalent binding of IgG on the same virus particle. However, due to the orientation of the Fabs, only the sites (cyan dotted circles) around the 5-fold and the 3-fold vertices can accommodate bivalent binding of a full IgG. **(C)** Molecular Dynamics (MD)-based atomic model of the full ZKA190 IgG (green) bridging across two neighboring E proteins on the viral surface (grey). For clarity, only one IgG binding is shown. Bivalent binding to other sites within these vertices is possible. **(D)** Superposition of the MD-based full IgG model (green cartoon) on the cryoEM map (blue mesh) near the 3-fold vertex shows good fitting of the modelled Fabs to the cryoEM density. **(E)** Neutralization test of ZIKV when ZKA190 is added pre- or post-attachment of virus to cells that were incubated at 4°C . Results show no difference in the neutralization profile. **(F)** RT-PCR detecting the amount of virus remaining on cells when ZKA190 was added pre- and post- virus attachment to cells. When increasing concentrations of mAb were mixed with virus before attaching to cells (“pre-attach.”), there is a gradual increase in the amount of virus on cell surface, likely due to aggregation of virus. Addition of the mAb to virus that had already bound to cells (“post-attach.”) showed that the mAb is unable to strip the virus off the cell surface. Therefore, the mAb can also inhibit a post-attachment stage of the infection cycle. **(G)** Confocal microscopy experiments. ZIKV incubated with a concentration exceeding 10,000-fold the IC_{50} value of either ZKA190 Fab or full IgG were added to Vero cells. The ZIKV:antibody complex is detected inside the cells (green) and co-localizes with endosomes (red, yellow overlay). Endosomes and acidic organelles are marked by LysoTracker red; Alexa-488-conjugated ZKA190 is in green. Nuclei are stained with DAPI (blue). Data in (A) and (F) are represented as mean \pm SD of at least two independent experiments. Data in (E) are represented as mean \pm SEM of three independent experiments performed in triplicate.

Figure 4. Prophylactic and therapeutic efficacy of ZKA190.

(A to D) ZKA190 is strongly protective against ZIKV infection when administered prophylactically to mice (A129 in (A) and AG129 in (C, D)) challenged with a lethal dose of ZIKV strain MP1745 (A, B) or Nica 2-16 (C, D). Experiments used N=4-8 mice per group. **(A, C)** Kaplan-Meier survival curves are shown. Significance was determined using the Mantel-Cox log-rank test. Panel A: ZKA190 at 5, 1 and 0.2 mg/kg versus Ctr mAb, $P = 0.0031$; ZKA190 at 0.04 mg/kg versus Ctr mAb, $P = 0.0116$; ZKA190-LALA at 5, 1, 0.2 and 0.04 mg/kg versus Ctr mAb, $P = 0.0031$. Panels C: ZKA190 and ZKA190 LALA at 1 mg/kg versus Ctr mAb, $P = 0.0091$; ZKA190 at 0.02 mg/kg versus Ctr mAb, $P = 0.0035$; ZKA190-LALA at 0.04 mg/kg versus Ctr mAb, $P = 0.0142$. **(B, D)** Morbidity score of mice monitored over a 14-15 day period (two different scoring methods were used; see (Dowall et al., 2016) for panel B and (Orozco et al., 2012) for panel D). Control mAb is MPE8 mAb (A, specific for RSV F protein (Corti et al., 2013)) and a human IgG polyclonal isotype control (C). **(E to G)** ZKA190 or ZKA190-LALA were administered at 15 mg/kg at different time-points after ZIKV infection to A129 mice challenged with a lethal dose of ZIKV strain MP1745. **(E)** A Kaplan-Meier survival curve is shown. Experiments used N=5 mice per group. Significance was determined by using the Mantel-Cox log-rank test. ZKA190 and ZKA190-LALA administered on day 1, 2, 3 or 4 versus Ctr., $P = 0.0016$. **(F)** Morbidity score of mice monitored over a 14-day period according to (Dowall et al., 2016). **(G)** Mice were monitored over a 14-day period for body weight loss. Control mAb is MPE8 specific for RSV F protein (Corti et al., 2013). See also Figures S5 and S6.

Figure 5. Engineering of ZKA190 into the bispecific FIT-1 mAb.

(A) ZKA185 and ZKA230 mAbs were tested for neutralization of four strains of ZIKV, as determined by the percentage of infected Vero cells in the presence of increasing amounts of mAbs.

Data are representative of at least two independent experiments. **(B)** Binding of ZKA185 and ZKA230 IgG and Fab to recombinant ZIKV VLP, E and DIII antigens as assessed by ELISA. **(C)** ZKA190, ZKA185 and ZKA230 were tested for neutralization of H/PF/2013 (wt) and MARMs 1-4. **(D)** Surface representation of two E protein dimers bound by ZKA190 (green); the ZKA190 NMR derived epitope is in red; positions mutated in MARMs are indicated in yellow (E370), blue (T335), orange (D67) and magenta (K84). **(E)** Model of FIT-1. The natural linkers between inner and outer Fabs allow flexible movement of Fabs in the FIT-1 antibody. The variable regions of ZKA185 and ZKA190 are highlighted in blue and green, respectively. **(F)** Binding of FIT-1 IgG and Fab to recombinant ZIKV VLP, E and DIII antigens as assessed by ELISA. **(G)** ZKA190, ZKA185 and FIT-1 mAbs were tested for neutralization of four strains (IC_{50} values, **G**) and four MARMs (**H**) of ZIKV. **(I)** Neutralization of ZIKV H/PF/2013 strain by ZKA185, ZKA230 and FIT-1 IgG and Fab determined as in **(A)**. **(J)** Confocal microscopy experiments as shown in Figure 3G. **(K)** Effect on ADE induced by peak enhancing dilution of anti-prM DV62 mAb or DENV2 plasma by serial dilutions of FIT-1 IgG and Fab.

Figure 6. Therapeutic efficacy of FIT-1.

FIT-1 is strongly effective against ZIKV infection when administered therapeutically at different time-points to mice (A129) challenged with a lethal dose of ZIKV strain MP17451. Experiments used N=5-6 mice per group. **(A)** Kaplan-Meier survival curves are shown. Significance was determined by using the Mantel-Cox log-rank test. FIT-1 at 15, 5 and 1 mg/kg given either on day 1 or 2 versus Ctr mAb, $P = 0.0012$; ZKA190 at 15 and 5 mg/kg given on day 3 versus Ctr mAb, $P = 0.0012$; ZKA190 at 1 mg/kg given on day 3 versus Ctr mAb, $P = 0.0170$. **(B)** Morbidity score of mice monitored over a 21-day period (Dowall et al., 2016). **(C)** Viral load was measured as PFUs on day 5 in blood of all animals. **(D)** Mice were monitored over a 21-day period for body weight loss. Control mAb in panel A is MPE8 mAb (specific for RSV F protein (Corti et al., 2013)). **(E)** Viral load was measured as genomic copies by qPCR on day 5 in blood of all animals and in blood

and indicated tissues when animals were culled at the end of the study or when the humane endpoints were met. Significance was determined compared to control antibody treatment by nonparametric unpaired Mann-Whitney U test. * $p < 0.05$; ** $p < 0.01$.

Figure 7. Comparison of ZKA190 to other DIII lateral ridge antibodies.

(A) Superposition of the structures of the Fabs E16 (orange, PDB 1ZTX), E106 (cyan, PDB 4L5F), ZV23 (blue, PDB 5GZR) and ZK67 (purple, PDB 5KVG) onto DIII of ZKA190-E protein cryoEM structure. (B) Fv regions of ZKA190, E106, ZV23 and ZV67 (colors as indicated) over the ZIKV surface, obtained by superimposing their structural complex with DIII to E protein on the viral surface (white surface, the 5-fold vertex is shown). All mAbs, but not ZKA190, clashes with adjacent E proteins when superimposed onto molecule A near the 5-fold vertex.

LIST OF SUPPLEMENTAL FIGURES

Figure S1. Identification of ZKA190 epitope and analysis of its conservation in ZIKV strains.

Figure S2. Binding of wt or mutated DIII to ZKA190 IgG.

Figure S3. CryoEM reconstruction of ZIKV:Fab ZKA190 complex.

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Figure S5. Prophylactic efficacy of the anti-ZIKV DIII-specific mAb ZKA190 against ZIKV strains MP1741 and Nica 2-16.

Figure S6. Therapeutic efficacy of the anti-ZIKV DIII-specific mAb ZKA190.

SUPPLEMENTAL FIGURE LEGENDS:

Figure S1. Identification of ZKA190 epitope and analysis of its conservation in ZIKV strains.

(A) Overlay of [15N,1H]-HSQC spectra of 15N-labeled ZIKV DIII in the absence (black) or presence (red) of unlabelled ZKA190 Fab. Differences identify DIII residues affected by mAb binding. (B) NMR epitope mapping of ZKA190 Fab in complex with ZIKV DIII. The chemical shift perturbation (CSP, y-axis) is plotted against the DIII residue number. Residues affected by antibody binding are in red. (C) Residues in the FG loop identified by NMR epitope mapping are partially hidden in E protein mol A but largely exposed in mols B and C. DIII of E protein is coloured in blue. Residues identified by NMR epitope mapping are coloured in magenta, except those in the FG loop, which are coloured in green. Adjacent E proteins are shown as grey surface. (D) Level of amino acid residue conservation in ZKA190 epitope as calculated by the analysis of sequences from 217 ZIKV strains found in ZIKV Resources (NCBI) databases as of November 24th 2016. (E) Open-book representation showing charge complementarity between the epitope and paratope of the docking result. Boundaries of the epitope and paratope are circled in green. The borders between heavy and light chains of Fab and its corresponding footprint on DIII are shown as yellow dashed lines. Related to Figure 2.

Figure S2. Binding of wt or mutated DIII to ZKA190 IgG. (A-E) SPR data and binding kinetics are shown. (F) DIII mutants that affect (red highlights) or do not affect binding are shown as indicated in the figure. Related to Figure 2.

Figure S3. CryoEM reconstruction of ZIKV:Fab ZKA190 complex. (A) ZKA190 Fab binds to ZIKV at 4°C, 37°C and 40°C. The uncomplexed ZIKV particles mostly have smooth surfaces at all temperatures. In contrast, in the ZKA190-complexed samples, most particles appear spiky, indicating Fab binding. The scale bar represents 50 nm. (B) The cryoEM maps of ZIKV:ZKA190 Fab complex at 4°C and 37°C, displayed at multiple contour sigma levels. The Fab densities at all positions are similarly reduced at higher contour levels, suggesting equal occupancies. The densities of virus surface and Fab molecules are colored in cyan and blue, respectively. The white triangle highlights one asymmetric unit. Vertices are indicated. (C) Fourier shell correlation (FSC) of the ZIKV:ZKA190 Fab complexes at 4°C and 37°C were plotted against spatial frequency. Related to Figure 2.

Figure S4. Interpretation of the cryoEM map and validation of the docking model in the cryoEM density map.

(A) Interpretation of the cryoEM ZIKV:ZKA190 map by fitting the crystal structure of E protein dimers (PDB 5JHM) and homology model of the Fab into the map. Left, overall view of the fit. Middle, density map displayed at a higher contour level showing the E protein raft is well fitted. Right, density map displayed at a lower contour level showing that the Fab density above molecule C has the best shape corresponding to Fab, compared to those above molecules A and B. The map was first interpreted by displaying the map at high contour so that the shape of the E protein dimers could be easily observed for fitting of the E protein raft, then by using

lower contours, Fab ZKA190 was fitted into the density above E protein molecule C. The Fab-E protein (molecule C) complex was subsequently superimposed onto molecules A and B. The positions of the Fab above E protein molecules A and B were slightly adjusted to optimize the fit. Fab molecules are shown as green ribbons. The three individual E proteins in one asymmetric unit are labelled as A, B and C while those in the neighboring asymmetric unit within a raft, A', B' and C'. The cryoEM map is shown as a transparent grey surface. The black triangle represents one asymmetric unit. **(B)** The docking model of the DIII complexed with the variable region of the Fab (Fv) fits well into the cryoEM density map. The docking model was first superimposed onto E protein dimers (A-C' and B-B') by using their DIIs. Each E protein dimer:Fv ZKA190 complex was then fitted into the cryoEM density as a rigid body. Left, the overall fit. Middle, E proteins raft shown with the density map displayed at a high contour level. Right, Fv molecules fit well to the density map. E proteins within one raft are colored in purple whereas Fv molecules are colored in green. Related to Figure 2.

Figure S5. Prophylactic efficacy of the anti-ZIKV DIII-specific mAb ZKA190 against ZIKV strains MP1741 and Nica 2-16. **(A-D)** Results related to Figure 4A-B. **(A)** Shown is the viremia measured as PFU/ml on day 5 in blood of all animals. **(B)** Viral load was measured as genomic copies/ml by qPCR on day 5 in blood of all animals and in blood and indicated tissues when animals were culled at the end of the study or when the humane end-points were met. **(C)** Mice were monitored over a 14-day period for body weight loss. **(D)** Serum concentrations of human IgG in day 5 blood samples. **(E)** Results related to Figure 4E-G. Shown is the viremia measured as genomic copies/ml by qRT-PCR on days 3 and 6 in blood of all animals. Significance was determined compared to control antibody treatment by nonparametric unpaired Mann-Whitney U test. * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$. Related to Figure 4.

Figure S6. Therapeutic efficacy of the anti-ZIKV DIII-specific mAb ZKA190. **(A)** Viral load was measured as PFUs on day 5 in blood of all animals. **(B)** Viral load was measured as genomic copies by qPCR on day 5 in blood of all animals and in blood and indicated tissues when animals were culled at the end of the study or when the humane end-points were met. Significance was determined compared to control antibody treatment by nonparametric unpaired Mann-Whitney U test. * $p < 0.05$; ** $p < 0.01$. **(C)** Serum levels of human IgG in day 5 blood samples. **(D)** Sequencing of the DIII region from ZIKV *ex vivo* isolates from ZKA190-treated animals. Full description of sequenced samples. **(E)** Amino acid sequence alignment of DIII from the 18 selected samples (derived from 16 animals) as compared to the parental sequence of the inoculum virus (MP1751 strain). Related to Figure 4.

STAR METHODS

CONTACT FOR REAGENTS AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Davide Corti (davide.corti@humabs.ch)

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Reagents

Cell lines were obtained from American Tissue Culture Collection. Vero and K562 cells were cultured in DMEM or RPMI 1640 medium (Invitrogen) supplemented with 10% FCS (Hyclone), 1% penicillin, 1% streptomycin and L-glutamine (complete DMEM or RPMI; all from Invitrogen). The ZIKV H/PF/2013, MR766 and MRS_OPY_Martinique_PaRi_2015 (MRS_OPY) strains used in this work were obtained from European-Virus-Archive. The PV10552 strain was isolated in 2016 from the semen of an international traveler in Italy and was kindly provided by Fausto Baldanti from the Policlinico San Matteo, Italy (accession number KY003154). The Puerto Rican 2013 PRVABC59 strain was obtained from the CDC (GenBank Accession #KU501215). The virus had been passaged three times in Vero cells prior to our acquisition. Vero cells were maintained in Eagle Minimal Essential Medium (EMEM) supplemented with 10% fetal bovine serum and 1% penicillin-streptomycin. The virus was passaged two more times in Vero cells to establish a viral stock. Viral stocks were titered by limiting dilution plaque assay on Vero cells. Infections with ZIKV were performed at 37°C for 4 hours (h) before cells were overlaid with methylcellulose. After 6 days, plaques were visualized by crystal violet staining. The murine cross-reactive mAb used to detect ZIKV and DENV infection was produced from the 4G2 hybridoma purchased from ATCC and purified using protein G affinity chromatography (Thermo Scientific). The antigens used in ELISA were: recombinant E protein from ZIKV (Meridian LifeScience, R01635), recombinant DIII synthesized using the sequence from ZIKV Brazil 2015 strain (SPH2015). The DIII of ZIKV was expressed in E. coli Rosetta and Rosetta pLysS cells respectively, with a pET21 vector (Novagen) either in LB or M9 minimal media with aptly labelled ingredients as required for NMR (¹⁵N NH₄Cl, ¹³C Glucose, 70% D₂O). The cells were induced at OD₆₀₀ = 0.6 with 0.5 mM of IPTG and harvested after 4 hours post-induction. After sonication and centrifugation, the pellet was washed and centrifuged two times in sodium phosphate buffer pH 7.2, 150 mM NaCl, 1 M urea and 1% Triton-X100. The pellet was resuspended in phosphate buffer pH 7.2 and 8 M urea (buffer A). Following addition of 0.2% of PEI and centrifugation to remove DNA contamination, ammonium sulphate to 60% was added to the supernatant to remove traces of PEI. After centrifugation, the pellet was resuspended in buffer A, and denaturated protein was diluted in refolding buffer (20 mM phosphate buffer pH 7.2, 150 mM NaCl, 200 mM arginine) in order to reach a final protein concentration of 0.2 mg/mL. DIIs of DENV and ZIKV were purified on a Superdex-75 size exclusion column (GE) in PBS buffer. The elution and dynamic light scattering profiles of DIII were consistent with monomeric species.

METHOD DETAILS

Production of recombinant IgG and LALA variant

Briefly, the VH and VL sequences were cloned into human Ig γ 1, Ig κ and Ig λ expression vectors (kindly provided by Michel Nussenzweig, Rockefeller University, New York, NY, USA), essentially as described (47). Recombinant mAbs were produced by transient transfection of EXPI293 cells (Invitrogen), purified by Protein A chromatography (GE Healthcare) and desalted against PBS.

Production of recombinant FIT bispecific antibodies.

The three genes encoding for FIT were codon optimized and synthesized by Genscript and cloned as follow: i) the VL of the outer Fab, followed by the full constant region (lambda or kappa), is fused with the VH of the inner and cloned into the Ig γ 1 expression vector (modified to encode for the LALA mutation). The resulting polypeptide 1 is formed by VL and CL of the outer Fab, VH of the inner Fab fused to IgG1 CH1-hinge-CH2-CH3 domains; ii) the VH gene of the outer Fab (encoding for polypeptide 2 formed by VH and CH1 of the outer Fab) was cloned into the Fab expression vector (Ig γ 1 expression vector in which a stop codon is introduced after the codon encoding for the CH1 cysteine residue 220); iii) the VL gene of the inner Fab is cloned into the Ig κ or Ig λ expression vectors (encoding for polypeptide 3 formed by VL and CL of the inner Fab). Recombinant FIT-Ig mAbs were produced by transient transfection of EXPI293 cells (Invitrogen) using a molar ratio of 1:3:3 of the three constructs described above (as described in PCT/US2014/072336), purified by Protein A chromatography (GE Healthcare) and desalted against PBS. The proteins were analyzed by SDS-PAGE in both reduced and non-reduced conditions and their concentrations determined by BCA (Pierce, Rockford, IL). In non-reduced conditions, FIT-Ig migrated as a major single band of approximately 250 KDa. In reducing conditions, each of the FIT-Ig proteins yielded two bands, one higher MW band is polypeptide 1 of approximately 75 KDa, and one lower MW band corresponds to both polypeptide 2 and 3 overlapped at approximately 25 KDa. To further study the physical properties of FIT-Ig in solution, size exclusion chromatography (SEC) was used to analyze each protein. Purified FIT-Ig, in PBS, was applied on a Superdex 200 Increase 5/150 GL. All proteins were determined using UV detection at 280 nm and 214 nm. FIT-Ig proteins exhibited a single major peak, demonstrating physical homogeneity as monomeric proteins.

Infection of human neural progenitor cells (hNPCs)

Adult male fibroblasts obtained from the Movement Disorders Bio-Bank (Neurogenetics Unit of the Neurological Institute 'Carlo Besta', Milan) were reprogrammed using the CytoTune-iPS 2.0 Sendai kit (Life Technologies). hiPSCs were maintained in feeder-free conditions in mTeSR1 (Stem Cell Technologies). To generate embryoid bodies (EBs), dissociated hiPSCs were plated into low-adhesion plates in mTeSR1 supplemented with N2 (0.5x) (ThermoFisher Scientific), human Noggin (0.5 mg/ml, R&D System), SB431542 (5 μ M, Sigma), Y27632 (10 μ M, Miltenyi Biotec) and penicillin/streptomycin (1%, Sigma) (Marchetto et al., 2010). To obtain rosettes, EBs were plated after 10 days onto matrigel-coated plates (1:100, matrigel growth factor reduced, Corning) in DMEM/F12 (Sigma) with N2 (1:100), non-essential amino acids (1%, ThermoFisher Scientific) and penicillin/streptomycin. After 10 days, cells were passaged with Accutase (Sigma) and seeded onto matrigel coated-flasks in NPC media containing DMEM/F12, N2 (0.25%), B27 (0.5%, ThermoFisher Scientific), penicillin/streptomycin and FGF2 (20 ng/ml, ThermoFisher Scientific).

hNPCs (3×10^4) were plated on coverslips in 24-well plates 3 days prior to infection with PRVABC59 strain. Virus stock was incubated with the mAbs 1h prior to addition to hNPCs to obtain an MOI of 0.5. After 4h of virus adsorption, culture supernatant was removed and fresh medium containing the mAbs was re-added. Supernatant was collected 96h post-infection to measure virus titers by plaque assay on Vero cells. Cells were fixed in 4% paraformaldehyde (PFA, Sigma) solution in phosphate-buffered saline (PBS, Euroclone) for 30 min for indirect immunofluorescence. Fixed cells were permeabilized for 30 minutes (min) in blocking solution, containing 0.2% Triton X-100 (Sigma) and 10% donkey serum (Sigma), and incubated overnight at 4°C with the primary antibodies in blocking solution. The following antibody was used for detection: anti-envelope (1:200, Millipore, MAB10216). Then, cells were washed with PBS and incubated for 1h with Hoechst and anti-mouse Alexa Fluor-488 secondary antibodies (1:1,000 in blocking solution, ThermoFisher Scientific). After PBS washes, cells were washed again and mounted.

Virus neutralization and antibody-dependent enhancement

Neutralization of ZIKV infection by mAbs was measured using a micro-neutralization flow cytometry-based assay. Different dilutions of mAbs were mixed with ZIKV (MOI of 0.35) for 1 hour at 37°C and added to 5000 Vero cells/well in 96-well flat-bottom plates. After four days for ZIKV, the cells were fixed with 2% formaldehyde, permeabilized in PBS containing 1% fetal calf serum (Hyclone) and 0.5% saponin, and stained with the mouse mAb 4G2. The cells were incubated with a goat anti-mouse IgG conjugated to Alexa Fluor488 (Jackson Immuno- Research, 115485164) and analyzed by flow cytometry. The neutralization titer (50% inhibitory concentration [IC₅₀]) is expressed as the antibody concentration that reduced the infection by 50% compared to virus-only control wells. ADE was measured by a flow cytometry-based assay using K562 cells. mAbs and ZIKV (MOI 0.175) were mixed for 1 hour at 37°C and added to 5000 K562 cells/well. After four days, cells were fixed, permeabilized, and stained with mAb m4G2. The number of infected cells was determined by flow cytometry, as described above.

Size measurement of ZKA190 IgG – ZIKV complex by dynamic light scattering

ZIKV was purified as described above and diluted in NTE buffer (corresponding to ~0.2 mg/ml E protein). Immune complexes were formed by mixing ZIKV with serial dilutions of ZKA190 IgG in IgG/virion molar ratios ranging from 900:1 to 1:1 (concentration of virion was determined using SDS-PAGE gel by comparing the virus E proteins with BSA standards). After one hour incubation at 4°C, the diameter of the immune complexes were measured using dynamic light scattering (DLS) and the infectivity was determined using standard plaque assay in BHK-21 cells. Neutralization is represented as the fold reduction of the remaining virus infectivity in each condition compared to the virus only control (\log_{10} (virus titer (PFU/ml) in the absence of mAb) / (virus titer (PFU/ml) in the presence of mAb). For the size measurement, 15 μ l of the sample was loaded into a quartz cuvette and measured at 25°C by the Zetasizer Nano S machine (Malvern). Data were analyzed using Zetasizer Nano software version 6.01. The deconvolution of the measured correlation curve was done using a non-negative least-squares algorithm. Each sample is measured 10 times and each data point is a repeat of two independent experiments. Size of the immune complex is shown as intensity weighted mean hydrodynamic size of the ensemble collection of particles in diameter. Our purified virus alone sample has a polydispersity index of 0.2, indicating that the sample is monodisperse and acceptable for DLS size measurement. Aggregation of immune complex has very

broad size distribution, resulting in a polydispersity index higher than 0.6. Therefore, the measured size is not very accurate for the aggregations. However, DLS was only used to probe the formation of aggregation in this experiment and the exact size of the aggregation is less important.

Pre- and postattachment neutralization assays.

The pre- and post-attachment neutralization assays were conducted as previously described (Fibriansah et al., 2015). For the pre-attachment neutralization assay, BHK-21 cells were grown in 24-well plates. Different dilutions of ZKA190 IgG were incubated with 100 plaque forming units (PFU) of ZIKV for 1 hour at 4°C and subsequently added to the cells and further incubated for 1 hour at 4°C. For the post-attachment neutralization assays, the cell monolayer was pre-chilled at 4°C for 15 mins before 100 PFU of ZIKV was added. The cells were co-incubated with the virus for 1 hour at 4°C and then washed twice with ice-cold serum-free medium to get rid of the unbound virus. Subsequently, different dilutions of ZKA190 IgG were added to the cells and further incubated for 1 hour at 4°C. The cells were finally washed twice with ice-cold serum-free medium and overlaid with 1% aquacide in RPMI medium supplemented with 2% fetal bovine serum. The cell monolayer was fixed and stained after 5-days incubation at 37°C in order to visualize the plaque formed by un-neutralized ZIKV.

RT-PCR to quantitate virus on cell surface

The number of ZIKV remaining on the surface of BHK-21 cells after ZKA190 treatment was measured by quantitative RT-PCR as previously described (Fibriansah et al., 2015). The experiment procedure is similar to the pre- and post-attachment neutralization assay described above but the quantification of the amount of virus particles attached on the cell surface were conducted immediately after the final wash of the cells. In pre-attachment condition, 10⁵ PFU of ZIKV was incubated with neutralizing concentrations of ZKA190 IgG for 1 hour at 4°C and then added to the cells pre-chilled at 4°C. After one hour of incubation at 4°C, the cells were washed twice and total RNA was purified using RNeasy mini kit (Qiagen). In post-attachment condition, the cells were first incubated with the same amount of ZIKV at 4°C for 1 hour and then washed twice to get rid of the unbound virus. Neutralizing concentrations of ZKA190 IgG were then added to the cell and incubated at 4°C for 1 hour. Finally, the cells were washed twice and total RNA was purified. Complementary DNA was synthesized using qScript cDNA SuperMix (Quantabio). Real-time quantitative PCR (qPCR) was performed using iQTM SYBR Green supermix kit (Bio-Rad) in a CFX 96 Real-Time System (Bio-Rad). Glyceraldehyde-3-phosphate dehydrogenase was used as the housekeeping gene to normalize samples. The analysis of relative levels of ZIKV RNA in different samples was performed by comparative 2- $\Delta\Delta$ CT method (Livak and Schmittgen, 2001). Gene specific primers (ZIKV-NS1-Forward: 5'-TGGAGTTCAACTGACGGTCG-3'; ZIKV-NS1-Reverse: 5'-TACCCCGAACCCATGATCCT-3'; *Gapdh*-Forward: 5'-GGCAAGTTCAAAGGCACAGTC-3'; *Gapdh*-Reverse: 5'-CACCAGCATCACCCCATTT-3') were used for the real-time qPCR experiment.

Effect of LALA antibodies on ADE

ZIKV (MOI 0.175) was mixed with plasma from primary ZIKV-infected donors for 30 minutes at 37°C. ZKA190-LALA was added at 50 µg/ml, mixed with 5000 K562 cells/well and incubated for three days. Cells were then stained with 4G2 and analyzed by flow cytometry, as described above.

ELISA

A standard ELISA was used to determine binding of mAbs to ZIKV DIII or E proteins. ELISA plates were coated overnight at 4°C with 3 µg/ml of antigens. Plates were blocked with a 1% w/v solution of Bovine Serum Albumin (BSA; Sigma) in PBS, incubated with serial dilutions of mAbs for 1.5 hours at room temperature. After washing, antibody binding was revealed using goat anti-human IgG coupled to alkaline phosphatase (Jackson ImmunoResearch). Plates were then washed, substrate (p-NPP, Sigma) was added and plates were read at 405 nm. Serum levels of human IgG in blood samples of infected mice was determined using a sandwich ELISA specific for human IgG antibodies. ELISA plates were coated overnight at 4°C 5 µg/ml of goat anti-human IgG Fc-specific, mouse-adsorbed, antibodies (Southern Biotech). A standard based on the spiked-in mAb used in the challenge study diluted in mouse serum was used to interpolate the concentration of human mAbs in mouse sera samples. Plates were blocked with a 1% w/v solution of Bovine Serum Albumin (BSA; Sigma) in PBS, incubated with serial dilutions of sera samples for 1.5 hours at room temperature. After washing, human IgG levels were revealed using goat anti-human IgG Fc-specific (mouse-adsorbed) antibodies coupled to alkaline phosphatase (Southern Biotech). Plates were then washed, substrate (p-NPP, Sigma) was added and plates were read at 405 nm.

Surface Plasmon Resonance

Binding properties of the complexes between mAbs IgG or Fab and DIII (wild-type and mutants) or E protein were determined at 25°C with a ProteOn XPR-36 instrument (Bio-Rad). The antibodies were immobilized on the surface of a GLM sensor chip through standard amine coupling. Serial dilution of DIII and E-protein in the nanomolar range were injected at a flow rate of 100 µl/min (contact time 4 minutes); dissociation was followed for 10 minutes. Analyte responses were corrected for responses from buffer-only injection both on a channel with antibody immobilized and on a channel with no antibody immobilized. Curve fitting and data analysis were performed with Bio-Rad ProteOn Manager software (version 3.1.0.6).

NMR spectroscopy

Spectra were recorded on a Bruker Avance 700 MHz NMR spectrometer at 300 K. For assignments of backbone resonances standard triple resonance experiments (HNCO, HN(CA)CO, HN(CO)CACB, HNCACB) were used, while sidechains were annotated using HCCH-TOCSY and HBHA(CO)NH experiments. All NMR experiments were processed using Topspin 2.1 (Bruker Biospin) and analysed with CARRA. NOESY cross peaks were automatically assigned using the CYANA “noeassign” macro based on the manually assigned chemical shifts. Upper-distance restraints used for the structure calculations in CYANA using the standard simulated annealing protocol were derived from 70 ms ¹⁵N- and ¹³C-resolved NOESY spectra.

Backbone dynamics of ZIKV DIII were derived from ¹⁵N relaxation measurements recorded on 600 and 700 MHz spectrometers. Proton-detected versions of the CPMG (R2), inversion-recovery (R1) and ¹⁵N{¹H}-steady-state NOE were utilized.

Delay settings for the T2 series were in the range of 0 to 0.25 sec and for the T1 series between 0.02 to 2 sec. The ¹⁵N{¹H}-NOE experiment used a relaxation delay of 5 s. The R1 and R2 relaxation rates were derived from least-squares fits of corresponding exponential functions to the measured data using home-written scripts. The relaxation data were analyzed in a model-free approach using the software package DYNAMICS. The program ROTDIF was used to calculate the overall

correlation time from the relaxation data (8.5 ns). NMR epitope mapping was performed as previously described (Bardelli et al., 2015; Simonelli et al., 2010; 2013). Briefly, overlay of ^{15}N -HSQC spectra of labelled DIII free or bound to ZKA190 Fab allowed identification of DIII residues whose NMR signal changed upon complex formation, indicating that they were affected by ZKA190 binding. Changes were identified by manual inspection and by the Chemical Shift Perturbation (CSP), $\text{CSP} = ((\Delta\delta_{\text{H}})^2 + (\Delta\delta_{\text{N}}/10)^2)^{1/2}$. NMR samples were typically 800 μM of [^{15}N , ^{13}C]-labeled DIII in 20 mM sodium phosphate, 50 mM NaCl, pH 6.0. Perdeuterated (nominally 70%) ^2H , ^{15}N DIII samples were used for NMR epitope mapping with a DIII:ZKA190 Fab ratio of 1:1.1; DIII concentration was typically 0.4 mM.

Antibody modelling, computational docking and Molecular Dynamics simulations

The ZKA190, ZKA185 and ZKA230 variable fragments were modelled according to the canonical structure method with the program RosettaAntibody (Sircar et al., 2009) as previously described (Pedotti et al., 2011). Ten models, mainly differing in the H3 loop conformation, were generated and independently used for docking. For ZKA190, either the NMR calculated or X-ray derived structure of DIII (PDB 5JHM) was used for docking. Docking was performed using RosettaDock 2.3 as previously described (Simonelli et al., 2010). In summary, each antibody model was independently docked to DIII. Amongst the thousands of computationally generated complexes, those in better agreement with NMR and cryoEM experimental data were selected and further refined by computational docking. Models were selected to maximize fit to the cryoEM density map and presence of NMR-defined epitope residues at the computational interface in an iterative process. For ZKA185 and ZKA230, four adjacent E protein monomers extract from the CryoEM structure of ZIKV were used for docking. The best solutions according to the algorithm were evaluated in terms of their agreement with the experimental data (MARMs) After refinement, selected models of all the three antibodies were subjected to a 100ns molecular dynamics (MD) simulation to adjust the local geometry and verify that the structure was energetically stable. MD was performed with the AMBER package proteins were centered in a cubic box filled with TIP3P water model and 0.15M Na+Cl- ions using the ff14SB force field. After energy minimization and temperature and pressure equilibration of 100ps, 100ns MD simulations were performed. The trajectory files generated were analyzed after removal of periodic boundary conditions.

Modelling of ZKA190 full IgG bound to the viral surface

A homology model of ZKA190 full IgG was generated using the x-ray structure of a full IgG1 (PDB 1HZH) as template. Analysis of the cryo-EM, NMR and docking data showed that adjacent ZKA190 Fab binding sites on the viral surface are likely compatible with distances and orientation of the two arms of a full IgG; a pair on the 3-fold ($\sim 60 \text{ \AA}$) and one on the 5-fold vertex ($\sim 56 \text{ \AA}$) (**Figure 3B**). Models of the full IgG were aligned with the Fabs on these binding sites and restrained molecular dynamics simulations were used to generate atomistic models of the full antibody bridging across two E proteins on the viral surface. A 1ns simulation with restraints on the distance between the two Fabs of the full IgG was first performed to allow movement of the two arms into an appropriate orientation. Another 1 ns simulation of 16641 C α -C α distance restraints moved the arms in their final position. Finally, 10'000 steps of energy minimizations were performed. The final result shows good agreement to both cryo-EM and NMR data.

Confocal Microscopy

Vero cells were plated at 7,500 cells/well on 12 mm-diameter coverslips in 24-well plates and incubated overnight. Cells were infected with ZIKV H/PF/2013 (MOI of 100) in the presence or absence of neutralizing concentrations of Alexa-488 conjugated mAbs (0.7 μ M) at 37°C for 3 h, washed with PBS, and fixed with 2% paraformaldehyde in PBS for 30 min at room temperature. Acidified endosome were identified with LysoTracker red (Invitrogen) by adding the dye (50 nM) to the cells for the last 30 min of the incubation prior to fixation. Fixation was followed by extensive washes in PBS and 50 mM glycine and finally the coverslips were prepared for microscopy analysis using Vectashield mounting medium for fluorescence with DAPI (Vector Laboratories). Samples were analyzed by confocal microscopy using a Leica TCS SP5 microscope with a 63 \times /1.4 N.A. objective. Image analysis and processing was performed with FIJI software.

Sequence analysis from ex vivo ZIKV samples

Envelop protein (E) sequences were obtained from National Center for Biotechnology Information (NCBI) Zika Virus Variation Resource (Brister et al., 2014) as of November 24th, 2016. After aligning the protein sequences with Clustal Ω (Sievers et al., 2011) and purging the sequences that did not cover the EDIII domain, 217 sequences were obtained. An amino acid distribution analysis was performed for each residue of the EDIII protein sequences. The related 217 nucleotide sequences of the E protein were retrieved and aligned using MAFFT (Kato and Standley, 2013). The identical nucleotide sequences were collapsed into unique sequences (138 in total) and a phylogenetic tree was derived by maximum-likelihood method (bootstrapped 1,000 times) using MEGA 7 software (Kumar et al., 2016).

MARMs selection

Two-thousands TCID₅₀ of H/PF/2013 ZIKV in 500 μ l were incubated with 250 μ l containing varying concentrations of mAb (8 different concentrations, starting with a final concentration of 200 μ g/ml and performing serial 1:4 dilutions). The mixture was incubated for 45 minutes at 37°C, followed by the addition of 250 μ l of a Vero cells suspension (3.2×10^6 cells) and an incubation in a 24 well plate for three-four days at 37°C to allow virus propagation to occur. After each step of selection, 500 μ l of supernatants from three conditions were selected: the lowest concentration of mAb at which full protection of the monolayer was observed, one concentration at which we observed a partial CPE effect on the cell monolayer and one concentration at which 100% of the cell monolayer was destroyed by the ZIKV CPE. The tube was spun down for 5 minutes at 1000 x g, aliquoted and stored at -80°C. Half of the volume was again pre-mixed with varying concentrations of mAb to repeat the selection and propagation process. The remaining supernatant was used for micro-neutralization assays and subsequent sequencing of the virus.

Sequencing of MARMs virus

To identify the escape mutations of the selected MARMs virus, a genomic RNA extraction was done followed by a one-step-PCR to amplify and sequence the ZIKV E protein amplicon. Cell supernatant (140 μ l) from the MARMs selection was used for RNA extraction with the QIAamp Viral RNA mini kit (Qiagen). cDNA synthesis and PCR amplification were performed together using the SuperScript III One-Step RT-PCR with Platinum Taq (Invitrogen). For one reaction 25 μ l reaction mix, 8 μ l sterile water, 2 μ M of each primer, 1 μ l RNase out (Life Technologies), 2 μ l

Superscript III RT/Platinum TaqMix and 12 µl RNA were used giving a final reaction volume of 50 µl. For the E protein N-terminal part, the primer pair Zika-E-F1 5'-TGCAAACGCGGTCGCAAACCTGGTTG-3' and ZIKV-E-R1 5'-CGTGCCAAGGTAATGGAATGTCGTG-3' and for the C-terminal part the primer pair ZIKV-E-f1530 5'-AGCCTAGGACTTGATTGTGAACCGA-3' and ZIKV-E-R2769 5'-TTACAGATCCCACAACGACCGTCAG-3' were used. The cycling conditions were 54°C for 40 minutes, 94°C for 2 min followed by 45 cycles of 94°C for 45 seconds, 50°C for 45 seconds and 68°C for 1.5 minutes with a final elongation step at 68°C for 5 minutes and a final cooling step at 4°C. The PCR products were analyzed and extracted from a 1.5 % agarose gel and further purified with the GFX PCR DNA and Gel Band Purification kit (GE Healthcare). For the sequencing reaction 8 µl of purified PCR product was mixed with 2 µM primer in a final volume of 10 µl and sent for sequencing (Microsynth). E protein N-terminal products were sequenced with ZIKV-E-F2 5'-ACTTGGTCATGATACTGCTGATTGC-3' and ZIKV-E-R2 5'-TCGGTTCACAATCAAGTCCTAGGCT-3', C-terminal PCR products with ZIKV-E-f2058 5'-GCTAACCCCGTAATCACTGAAAGCA-3' and ZIKV-E-r2248 5'-AAGACTGCCATTCTCTTGGCACCTC-3'. Sequences were assembled and analyzed using CLC Main Workbench software (CLC Bio, version 5).

ZIKV infection of A129 mice

ZIKV infection studies were conducted in the laboratories of Public Health England (PHE), UK. Female A129 mice (IFN-alpha/beta receptor -/-) aged 5-8 weeks were supplied from a UK approved breeding colony (B&K Universal Ltd, UK). All procedures with animals were undertaken according to the United Kingdom Animals (Scientific Procedures) Act 1986. These studies were approved by the ethical review process of Public Health England, Porton Down, UK and the Home Office, UK via an Establishment Licence (PEL PCD 70/1707) and project Licence (30/2906). PHE has Animal Welfare Assurance that has been reviewed and approved by the Office of Laboratory Animal Welfare (OLAW) – ref #A5560-01. Humane endpoints were defined as 20% weight loss, neurological signs or immobility. When any of these endpoints were reached, animals were culled to prevent unnecessary suffering.

Assessment of prophylactic and therapeutic activity of ZKA190 and FIT-1 in A129 mice

Mice were administered mAbs (ZKA190, ZKA190-LALA and MPE8) diluted in PBS at different doses via the intraperitoneal (i.p.) route in a volume of 500 µl. MAbs were administered either 1 day before or 1, 2, 3 or 4 days after virus challenge. Animals were challenged subcutaneously with 10² pfu ZIKV (strain MP1751) and followed for 14 days. Weights and temperatures were monitored daily and clinical observations were recorded at least twice per day. On day 5 post-challenge, 50 µl of blood was collected from each animal into a RNAprotect tube (Qiagen,UK) and frozen at -80°C. At the end of the study (14 days post-challenge) or when animals met human endpoints, necropsies were undertaken, and blood and sections of brain, spleen, liver, kidney and ovary were collected for virological analysis.

Assessment of prophylactic activity of ZKA190 in AG129 mice

To test whether ZKA190 was capable of inducing antibody-dependent enhancement (ADE) of ZIKV infection *in vivo*, 4- to 6-week-old AG129 mice lacking the interferon α/β and γ receptors

(van den Broek et al., 1995) were used to test low doses of mAb prior to challenge. Animals were bred at the University of California, Berkeley, and all experimental procedures involving the use of AG129 mice were pre-approved and performed according to the guidelines of the Institutional Animal Care and Use Committee of University of California, Berkeley. Mice were injected intraperitoneally (i.p.) with antibodies or phosphate buffered saline (PBS) in a total volume of 100 μ l and then infected 20-24 hours later with 10^3 focus-forming units (FFU) of ZIKV strain Nica 2-16 (Tabata et al., 2016) in a 50- μ l volume by subcutaneous (s.c.) foot pad injection. Mice were monitored over a 15-day period post-infection; body weight was recorded daily, and morbidity was scored using a standardized 5-point system (Orozco et al., 2012). Mice were euthanized if they lost more than 20% of their initial weight and/or if they progressed to stage 5 on the morbidity scale (moribund, consistent with partial paralysis and neurological symptoms).

Viral load quantification from AG129 ZIKV infected mice

Blood samples from ZIKV-infected mice or controls were collected from the facial (submandibular) vein of anesthetized mice on days 3 and 6 post-infection. Blood specimens were allowed to clot at room temperature, and serum was separated by centrifugation. ZIKV RNA was extracted from serum samples using the QIAamp Viral RNA Mini Kit (Qiagen), and RNA levels were determined by TaqMan one-step quantitative reverse transcriptase PCR (qRT-PCR) using a standard curve with 10-fold dilutions of ZIKV Nica 2-16 RNA. Primer sets used for this assay: ZIKV_1086_Fwd, 5'-CCGCTGCCCAACACAAG-3', ZIKV_1162c_Rev, 5'- CCACTAACGTTCTTTTGCAGACAT-3'; Probe ZIKV_1107-FAM_UG&Nica 5'- AGCCTACCTTGACAAGCAATCAGACACTCAA-3' (Lanciotti et al., 2008).

Viral load quantification from A129 ZIKV infected mice

Tissue samples from A129 mice were weighed and homogenized into PBS using ceramic beads and an automated homogenizer (Precellys, UK) using six 5 second cycles of 6500 rpm with a 30 second gap. Two hundred μ l of tissue homogenate or blood solution was transferred into 600 μ L RLT buffer (Qiagen, UK) for RNA extraction using the RNeasy Mini extraction kit (Qiagen, UK); samples were passed through a QIAshredder (Qiagen, UK) as an initial step. A ZIKV specific real-time RT-PCR assay was utilized for the detection of viral RNA from subject animals. The primer and probe sequences were adopted from a published method (52) with in-house optimization and validation performed to provide optimal mastermix and cycling conditions. Real-time RT-PCR was performed using the SuperScript III Platinum One-step qRT-PCR kit (Life Technologies, UK). The final mastermix (15 μ l) was comprised of 10 μ l of 2x Reaction Mix, 1.2 μ l of PCR-grade water, 0.2 μ l of 50 mM MgSO₄, 1 μ l of each primer (ZIKV 1086 and ZIKV 1162c both at 18 μ M working concentration), 0.8 μ l of probe (ZIKV 1107-FAM at 25 μ M working concentration) and 0.8 μ l of SSIII enzyme mix. Five μ l of template RNA was added to the mastermix, yielding a final reaction volume of 20 μ l. The cycling conditions used were 50°C for 10 minutes, 95°C for 2 minutes, followed by 45 cycles of 95°C for 10 seconds and 60°C for 40 seconds, plus a final cooling step of 40°C for 30 seconds. Quantification analysis using fluorescence was performed at the end of each 60°C step. Reactions were run and analyzed on the 7500 Fast platform (Life Technologies, UK) using 7500 software version 2.0.6. Quantification of viral load in samples was performed using a dilution series of quantified RNA oligonucleotide (Integrated DNA Technologies). The oligonucleotide comprised the 77 bases of ZIKV RNA targeted by the assay, based on GenBank accession AY632535.2 and was synthesized to a scale of 250 nmole with HPLC purification.

EDIII coding region sequencing from biological samples

For each sample, an attempt was made to reverse-transcribe and amplify the whole genome using the v2 amplicon sequencing protocol kindly supplied by J. Quick and N. Loman of the Zibra project (Faria et al., 2016) (www.zibraproject.org/data/amplicon_sequencing_protocol_v2.pdf). The two amplicon pools produced for each sample were quantified using the dsDNA HS assay (Life Technologies) and combined, before 1 ug of DNA was used as input to the EXP-NBD002 barcoding kit with barcodes NB01-NB09. Samples 1-9 and 10-18 were pooled separately and two sequencing libraries produced using the SQK-NSK007 kit. Each library was sequenced using a FLO-MIN105 Flow cell on a MinION Mk1B device and base calling was performed using the Metrichor workflow “2D Basecalling plus Barcoding for FLO-MIN105 250bps” (Oxford Nanopore Technologies). Between 3989 and 12801 2D reads were generated per sample. FastQ data from 2D reads was extracted using poretools (Loman and Quinlan, 2014) and mapped using BWA-MEM (Li and Durbin, 2009) to the sequence of the original MP1741 stock used in this study (Genbank KY288905). Coverage across the genome varied due to differences in amplification efficiency between amplicons, but mean coverage of the EDIII coding region was greater than 60-fold (range 64-fold to 410-fold) for all samples other than 14 and 16, which were not included for further analysis. The `align_trim.py` script from the Zibraproject.org pipeline (<https://github.com/zibraproject/zika-pipeline>, branch master, commit ID a4364fa8901b06d6d8bd058c19acd2c5e780179e) was used with an MP1741 adapted BED file of primer binding sites to trim primer sequences from the mapped reads. Samtools (Li et al., 2009) was used to convert the trimmed SAM file to a sorted and indexed BAM which was used as input along with FAST5 files for nanopolish v0.5 variants module (Loman et al., 2015) (Quick et al., 2016) (<https://github.com/jts/nanopolish>, branch master, commit ID 6406f78c01d05f968bc841c03e4ccb4ea208677a) to identify SNPs that met minimum criteria of read depth 20, support fraction 0.75 and minimum quality value of 200.

Virus sample preparation for cryoEM studies

Aedes albopitus C6/36 cells (ATCC) were propagated in RPMI 1640 media supplemented with 10% fetal bovine serum at 29°C. The cells were inoculated with ZIKV strain H/PF/2013 (Baronti et al., 2014) at a MOI of 0.5 and incubated at 29°C with 2% of fetal bovine serum. The virus containing media were harvested 4 days post-infection and clarified from cell debris by centrifugation. The supernatant was then precipitated by 8% of polyethylene glycol 8000 in NTE buffer (12mM Tris, 120 mM NaCl, 1 mM EDTA, pH 8.0) overnight at 4°C and pelleted by centrifugation at 14,000 g for 1 h. The resulting pellet was then resuspended in NTE buffer and further purified through a 24% (w/v) sucrose cushion followed by a 10-30% (w/v) potassium tartrate density gradient. The virus band in the gradient was then extracted and buffer exchanged into NTE buffer and concentrated using a concentrator with 100-kDa molecular weight cut-off filter. The concentration of the virus was estimated by comparing the viral E protein band with bovine serum albumin at known concentrations as standard in a Coomassie blue-stained SDS-PAGE.

CryoEM sample preparation

For preparation of the antibody-virus complex for cryoEM imaging, an excessive amount of recombinant Fab fragment of ZKA190 was added to the purified ZIKV to achieve a molar ratio of three Fab molecules for two E proteins. The complexes were incubated at 4, 37 and 40°C for 30 min,

respectively. To prepare the cryoEM grids, 2.3 μl of sample was applied to a lacey copper grid with a thin carbon film, blotted for 1 s with an FEI Vitrobot in 100% humidity and then plunged into liquid ethane. The frozen grids were kept in liquid nitrogen before imaging.

CryoEM data collection and image processing

CryoEM images of the frozen ZKA190 Fab:ZIKV complexes were collected on a Titan Krios (FEI) microscope equipped with 300 kV field emission gun. Leginon was used for automated data collection (Carragher et al., 2000). The magnification was 81,871, giving a pixel size of 1.71 Å. The images were recorded in single image mode on Falcon II direct electron detector (FEI) with a total dose of 20 $\text{e}^{-}\text{Å}^{-2}$. The images were taken at underfocus range between 1.0 and 3.5 μm . A total of 902 and 738 micrographs were collected for ZKA190 Fab:ZIKV complexes at 4°C and 37°C, respectively. The astigmatic defocus parameters were estimated using ctffind3 (Mindell and Grigorieff, 2003b) and used in orientation search and 3D reconstruction in Relion (Scheres, 2012). Particles were picked using Relion template-based autopicking program, visually examined to discard nonviral particles, and subsequently subjected to Relion 2D classification to produce 2D class averages. Classes containing junks and broken particles were excluded from further processing. In total, 10,641 and 6,728 particles in the ZKA190Fab:ZIKV complex samples that were incubated at 4°C and 37°C were selected for further processing. The 3D orientation search and reconstruction was done with icosahedral symmetry using Relion auto-refine procedure, where two models are refined independently for two random halves of the data to prevent over-fitting (Scheres, 2012). The mature ZIKV map (EMDB: EMD-8139) was filtered to 60 Å and used as the starting model. The 3D refinement produced structures with resolutions of 22 Å for both complexes at Fourier shell correlation (FSC) cut-off of 0.5.

Model fitting

As the E protein raft was intact (**Figure S4A**), the cryoEM map was first interpreted by fitting in the E protein dimers using the “fit-in-map” in Chimera (Pettersen et al., 2004). We tried fitting the E protein densities of both ZKA190 Fab:ZIKV complex cryoEM maps at 4°C and 37°C by using either the uncomplexed ZIKV (PDB 5IZ7) or the crystal structure of the ZIKV E protein dimers (PDB 5JHM). The crystal structure of the ZIKV E protein dimer fits better to the raft density than that of the cryoEM uncomplexed ZIKV. The difference between the cryoEM ZIKV E protein dimer structure and the crystal structure of ZIKV E ectodomain is in the DI-DII hinge angle. In the uncomplexed ZIKV cryoEM structure, the DI-DII hinge is fixed at a certain angle because its ectodomain interacts with the membrane associated stem regions of E and M proteins on the virus. However, in the ZKA190 Fab:ZIKV structure, the E protein raft is lifted up, as observed by the separation of the E protein shell density from the viral lipid bilayer membrane (**Figure 2E**). This may explain why the crystal structure of recombinant E protein fits better to the cryoEM density map of the complex. We then fitted the variable and constant domains of the Fab ZKA190 homology model into their corresponding densities above the DIII of E protein molecule C around the 3-fold vertex, as this density shows the best shape for the Fab (**Figure S4A**). The fitted molecule C and ZKA190 Fab were then locked and this E protein-ZKA190 Fab complex was superimposed as a rigid body to E molecules A and B, respectively. These Fab molecules correlated well with their corresponding densities. Slight adjustment of the positions of the Fab above molecules A and B was done to further optimize the fit (**Figure S4A**). No crashes between the symmetry related molecules were detected.

Structure analysis

Electrostatic potentials of protein surfaces were calculated using Coulombic Surface Coloring in Chimera (Pettersen et al., 2004).

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical analysis

Mortality data were expressed using Kaplan-Meier survival curves, and log-rank (Mantel-Cox) tests were used to determine statistical significance using Prism 7.0 software (GraphPad).

DATA AND SOFTWARE AVAILABILITY

Coordinates of Fab ZKA190:ZIKV complex incubated at 37°C was deposited in the Protein Data Bank under accession code 5Y0A and the NMR structure of DIII under PDB code 5OMZ and chemical shift used for structural calculation and mapping under BMRB code 34167. The cryoEM maps of Fab ZKA190:ZIKV complexes incubated at 4°C and 37°C were deposited in the Electron Microscopy Database under accession number EMD-6794 and EMD-6793, respectively.

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
ZKA190	(Stettler et al., 2016); this study	N/A
ZKA190-LALA	This study	N/A
ZKA185	(Stettler et al., 2016); this study	N/A
ZKA230	(Stettler et al., 2016); this study	N/A
FIT-1	This study	N/A
4G2	Absolute Antibody Ltd	Cat# Ab00230-2.0-BT
Bacterial and Virus Strains		
ZIKV H/PF/2013	European-Virus-Archive	Cat# 001v-EVA1545
ZIKV MR766	European-Virus-Archive	Cat# 001v-EVA143
ZIKVMRS_OPY_Martinique_PaRi_2015	European-Virus-Archive	Cat# 001v-EVAg1589
ZIKV PV10552	Baldanti F., IRCCS, Pavia (IT) (Acc. Number KY003154)	
ZIKV PRVABC59	CDC	
ZIKV MP1751	(Stettler et al., 2016); this study	
ZIKV Nica 2-16	(Tabata et al., 2016)	
Biological Samples		
Chemicals, Peptides, and Recombinant Proteins		
recombinant E protein	Meridian LifeScience	Cat# R01635
ZIKV DIII from ZIKV Brazil 2015 strain (SPH2015)	(Stettler et al., 2016); this study	N/A
Critical Commercial Assays		
RNeasy mini kit	Qiagen	Cat# 74106
qScript cDNA SuperMix	Quantabio	Cat# 95048-100
iQ™ SYBR® Green Supermix	Bio-Rad	Cat# #1708880
Deposited Data		
Fab ZKA190:ZIKV complex incubated at 37°C	This paper	PDB: 5Y0A
CryoEM maps of Fab ZKA190:ZIKV complexes incubated at 4°C	This paper	EMD-6794
CryoEM maps of Fab ZKA190:ZIKV complexes incubated at 4°C	This paper	EMD-6793
NMR structure of DIII under PDB code 5OMZ	This paper	5OMZ

Chemical shift used for structural calculation and mapping	This paper	BMRB code 34167
Experimental Models: Cell Lines		
EXPI293	Invitrogen	
Adult male fibroblasts	Movement Disorders Bio-Bank	
<i>Aedes albopitus</i> C6/36	ATCC	
Vero cells	ATCC	
K562 cells		
BHK-21	ATCC	
Experimental Models: Organisms/Strains		
A129 mice, (IFN-alpha/beta receptor -/-)		
AG129 mice	(van den Broek et al., 1995)	
Oligonucleotides		
ZIKV_1086_Fwd, 5'-CCGCTGCCCAACACAAG-3'	This study	
ZIKV_1162c_Rev, 5'-CCACTAACGTTCTTTTGCAGACAT-3'	This study	
Probe ZIKV_1107-FAM_UG&Nica 5'-AGCCTACCTTGACAAGCAATCAGACACTCAA-3'	(Lanciotti et al., 2008)	
Zika-E-F1 5'-TGCAAACGCGGTTCGCAAACCTGGTTG-3'	This study	
ZIKV-E-R1 5'-CGTGCCAAGGTAATGGAATGTCGTG-3'	This study	
ZIKV-E-f1530 5'-AGCCTAGGACTTGATTGTGAACCGA-3'	This study	
ZIKV-E-R2769 5'-TTACAGATCCCACAACGACCGTCAG-3'	This study	
ZIKV-E-F2 5'-ACTTGGTCATGATACTGCTGATTGC-3'	This study	
ZIKV-E-R2 5'-TCGGTTCACAATCAAGTCCTAGGCT-3'	This study	
ZIKV-E-f2058 5'-GCTAACCCCGTAATCACTGAAAGCA-3'	This study	
ZIKV-E-r2248 5'-AAGACTGCCATTCTCTTGGCACCTC-3'	This study	
ZIKV-NS1-Forward: 5'-TGGAGTTCAACTGACGGTCG-3'	(Xu et al., 2016)	
ZIKV-NS1-Reverse: 5'-TACCCCGAACCCATGATCCT-3'	(Xu et al., 2016)	
<i>Gapdh</i> -Forward: 5'-GGCAAGTTCAAAGGCACAGTC-3'	(Xu et al., 2016)	
<i>Gapdh</i> -Reverse: 5'-CACCAGCATCACCCATTT-3'	(Xu et al., 2016)	
Recombinant DNA		
Software and Algorithms		
Pymol Molecular Graphics Systems	Schrodinger, LLC	
Prism 7.0	GraphPad Software	
CLC Main Work Bench	CLC bio	

UCSF Chimera (v1.11.2)	(Pettersen et al., 2004)	https://www.cgl.ucsf.edu/chimera/
Relion	(Scheres et al., 2012)	http://www2.mrc-lmb.cam.ac.uk/relion/index.php/Main_Page
Ctffind3	(Mindell et al., 2003)	http://grigoriefflab.janelia.org/ctf
Leginon	(Carragher et al., 2000)	http://emg.nysbc.org/redmine/projects/leginon/wiki/Leginon_Homepage
Rosetta	(Gray et al., 2003)	
Amber	(Case et al., 2017)	http://ambermd.org/doc12/Amber17.pdf
CARA	(Keller et al., 2004)	http://cara.nmr-software.org/downloads/3-85600-112-3.pdf
Cyana	(Guntert et al., 2015)	
Zetasizer Nano software version 6.01	Malvern Instruments	
Other		