Protective antibodies against Eastern equine encephalitis virus bind to epitopes in domains A and B of the E2 glycoprotein

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Eastern equine encephalitis virus (EEEV) is a mosquito-transmitted alphavirus with a high case mortality rate in humans. EEEV is a biodefense concern because of its potential for aerosol spread and the lack of existing countermeasures. Here, we identify a panel of 18 neutralizing murine monoclonal antibodies (mAbs) against the EEEV E2 glycoprotein, several of which have ‘elite’ activity with 50 and 99% effective inhibitory concentrations (EC50 and EC99) of less than 10 and 100 ng ml−1, respectively. Alanine-scanning mutagenesis and neutralization escape mapping analysis revealed epitopes for these mAbs in domains A or B of the E2 glycoprotein. A majority of the neutralizing mAbs blocked infection at a post-attachment stage, with several inhibiting viral membrane fusion. Administration of one dose of anti-EEEV mAb protected mice from lethal subcutaneous or aerosol challenge. These experiments define the mechanistic basis for neutralization by protective anti-EEEV mAbs and suggest a path forward for treatment and vaccine design.

The EEEV is a mosquito-transmitted New World alphavirus in the Togaviridae family and is closely related to the Western (WEEV) and Venezuelan (VEEV) equine encephalitis viruses. Although relatively few human infections are reported annually, EEEV is one of the most severe mosquito-transmitted diseases with a 50–70% mortality rate and significant brain damage in most survivors1–6. Florida is now considered one of the major sources of EEEV epidemics in the USA, with transmission occurring throughout the year.

EEEV is an enveloped virus with a 11.5 kilobase single-stranded, positive-sense RNA genome that generates two RNA transcripts: a full-length genomic RNA; and a subgenomic RNA encoding the structural genes, C-E3-E2-6K-E1. After translation, the structural polyprotein C-E3-E2-6K-E1 is cleaved at the endoplasmic reticulum into the capsid protein and E3-E2-6K-E1. Additional protein processing in the endoplasmic reticulum and the Golgi apparatus results in transport of E2–E1 heterodimers to the plasma membrane7 where encapsidation of the genomic viral RNA occurs. The surface of the mature virion displays 80 spikes of trimers of E2–E1 heterodimers8. Structural studies of related alphaviruses have established an architecture with T = 4 icosahedral symmetry9–12. The E2 glycoprotein projects from the viral surface and consists of three domains: A, B and C13. Binding of EEEV E2 to poorly characterized host receptors is believed to initiate entry and endocytosis14. The acidic environment of the endosome induces conformational changes in the alphavirus E1 and E2 glycoproteins, which allow for the exposure of the fusion loop, insertion into the host membrane15 and nucleocapsid escape into the cytoplasm.

Few anti-EEEV mAbs have been described14–16 and only one has protective activity in mice17. These anti-EEEV mAbs have been mapped using peptides to three linear epitopes on E2: the N-terminal of domain A; the N- and C-terminal arches of domain B; and the C terminus of domain C14,15. In comparison, the epitopes of several murine and human mAbs against VEEV, WEEV or the more distantly related arthritogenic alphaviruses, for example, chikungunya virus (CHIKV), with therapeutic efficacy in vivo have been mapped18–21. These neutralizing mAbs predominantly recognize epitopes in domains A (residues 58–80) or B (residues 180–215) of the E2 glycoprotein, and inhibit infection at multiple steps including viral attachment, entry, fusion and egress18–21.

We isolated and purified a panel of murine mAbs against EEEV. Among these, 18 type-specific mAbs neutralized EEEV infection with 50% effective inhibitory concentration (EC50) values < 100 ng ml−1 and did not bind to WEEV or VEEV. Ten of these mAbs potently inhibited infection with EC50 values < 10 ng ml−1. In cell culture, most inhibited EEEV predominantly by blocking viral infection at a post-attachment step. We localized the epitopes of the majority of potently neutralizing mAbs to two solvent-exposed regions in domains A and B of the E2 glycoprotein. In vivo studies demonstrated that many of the neutralizing mAbs could protect mice against lethal subcutaneous or aerosol challenges by EEEV. Our results define the molecular basis for EEEV neutralization by protective mAbs and provide insight into the epitopes that could be

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targeted for immunotherapy and vaccine development against this highly lethal virus.

**Results**

**Generation of anti-EEEV mAbs.** We hypothesized that antibodies generated in the context of a live EEEV infection might have inhibitory activity. As EEEV is a biosafety level 3 select agent pathogen, performing B cell–myeloma cell fusions from infected animals presents technical challenges. To circumvent these issues, we engineered a chimeric biosafety level 2 pathogen that incorporates the non-structural genes and RNA replication control elements of a Sindbis virus (SINV, strain TR339) with the structural genes (C–E3–E2–6K–E1) of an EEEV isolate (strain FLN93-939) (Supplementary Fig. 1a)\(^2\). SINV-EEEV replicated efficiently in cell culture but did not cause disease in outbred and If3^-/- immunodeficient inbred mice (Supplementary Fig. 1b–e, and see text that follows).

To enhance the replication and immunogenicity of the attenuated SINV-EEEV in vivo, we inoculated If3^-/- C57BL/6 mice\(^2\). After infection and homologous boosting four weeks later, serum from If3^-/- mice had robust neutralizing activity against SINV-EEEV (end point titre >1:10,000). Splenocytes were collected from mice, fused to myeloma cells and 76 hybridomas producing anti-EEEV antibodies were isolated (Fig. 1a and Supplementary Table 1). Supernatant from 32 of the 76 hybridomas bound to EEEV virions purified from SINV-EEEV-infected cells and in a single end point dilution test, inhibited SINV-EEEV infection by 80% or more (Supplementary Table 1). These 32 mAbs were isolated (all of the immunoglobulin IgG2c or IgG3 subclass) and purified by protein A affinity chromatography for subsequent study.

We evaluated the purified mAbs for their ability to recognize the EEEV E2 glycoprotein. To do this, we purified recombinant EEEV E2 glycoprotein after expression in bacteria and oxidative refolding (Fig. 1b). Notably, 18 of 32 mAbs bound to the recombinant E2 glycoprotein in an enzyme-linked immunosorbent assay (ELISA) (Fig. 1c). We also tested a set of 24 mAbs for cross-reactivity with VEEV or WEEV (55 and 56% amino acid identity in the E2 glycoprotein required for mAb binding (Fig. 3a). Cells were transfected with plasmids encoding individual alanine (or serine for alanine residues) substitutions (360 residues) in the E2 gene in the context of a pE2-6K-E1 expression plasmid. We used critical residues as those with <25% binding to a given individual mAb that retained >70% binding to an anti-EEEV oligoclonal antibody control (Table 1, Supplementary Fig. 3 and Supplementary Table 2). We excluded from analysis mutations of cysteine residues and substitutions that globally altered E2 conformation, as defined by reduced binding of an oligoclonal antibody. A majority (13 of 16) of the neutralizing mAbs tested mapped to the ‘wing insertion’ of domain A (residues 52–82) or the distal region of domain B (β-strands A, B and E)\(^1\) of the E2 glycoprotein (Fig. 3a–c). The key loss-of-binding residues were highly conserved between the four (I, II, III and IV) EEEV subtypes (Fig. 3a). Although the domain B residues (I180, H181, S182, H213 and T215) required for mAb binding showed clear loss-of-binding phenotypes (Fig. 3d), some of the domain A residue changes (for example, D58, G59, D61 and M68) resulted in only partial loss-of-binding phenotypes (Fig. 3e). To extend these findings, we substituted selected residues in the A and B domains with bulkier and charged amino acids that might disrupt mAb interactions to a greater extent. We observed more profound loss-of-binding phenotypes when key domain B residues were substituted with arginine (Fig. 3f). Similarly, when the residues in domain A (D58, G59, D61, M68, K74 and L81) were mutated to arginine or glutamic acid, more pronounced loss of mAb binding was observed with EEEV-5, EEEV-58, EEEV-66, EEEV-82, EEEV-102 and EEEV-107 paradoxically enhanced plasma membrane fusion of the virus.

**Epitope mapping by alanine-scanning mutagenesis.** We used alanine-scanning mutagenesis coupled with HEK-293T cell-based expression and flow cytometry\(^1,8\) to identify residues in the E2 glycoprotein required for mAb binding (Fig. 3a). Cells were transfected with plasmids encoding individual alanine (or serine for alanine residues) substitutions (360 residues) in the E2 gene in the context of a pE2-6K-E1 expression plasmid. We used critical residues as those with <25% binding to a given individual mAb that retained >70% binding to an anti-EEEV oligoclonal antibody control (Table 1, Supplementary Fig. 3 and Supplementary Table 2). We excluded from analysis mutations of cysteine residues and substitutions that globally altered E2 conformation, as defined by reduced binding of an oligoclonal antibody. A majority (13 of 16) of the neutralizing mAbs tested mapped to the ‘wing insertion’ of domain A (residues 52–82) or the distal region of domain B (β-strands A, B and E)\(^1\) of the E2 glycoprotein (Fig. 3a–c). The key loss-of-binding residues were highly conserved between the four (I, II, III and IV) EEEV subtypes (Fig. 3a). Although the domain B residues (I180, H181, S182, H213 and T215) required for mAb binding showed clear loss-of-binding phenotypes (Fig. 3d), some of the domain A residue changes (for example, D58, G59, D61 and M68) resulted in only partial loss-of-binding phenotypes (Fig. 3e). To extend these findings, we substituted selected residues in the A and B domains with bulkier and charged amino acids that might disrupt mAb interactions to a greater extent. We observed more profound loss-of-binding phenotypes when key domain B residues were substituted with arginine (Fig. 3f). Similarly, when the residues in domain A (D58, G59, D61, M68, K74 and L81) were mutated to arginine or glutamic acid, more pronounced loss of mAb binding was observed with EEEV-5, EEEV-58, EEEV-66, EEEV-82, EEEV-102 and EEEV-107 paradoxically enhanced plasma membrane fusion of the virus.

**Epitope mapping by neutralization escape.** Alanine-scanning mutagenesis failed to map the epitopes of three inhibitory mAbs (EEEV-18, EEEV-82 and EEEV-102). As an alternative approach, we selected for neutralization escape mutants. We passaged SINV-EEEV in the presence of individual neutralizing mAbs until cytopathogenic effects were observed (3–4 passages), at which point the virus became resistant to neutralization. Remarkably, all three viral escape variants were reciprocally resistant to neutralization by the other mAbs in this group, suggesting they bound to an overlapping or shared epitope (Fig. 4a). To identify the escape muta-
Fig. 1 | Characterization of anti-EEEV mAbs. a, Supernatant from anti-EEEV hybridoma cells was screened for binding to a mixture of SINV-EEEV-infected and uninfected BHK-21 cells by flow cytometry. Shown are antibody staining from representative negative (1A2) and positive (13B10, subcloned as EEEV-10) hybridomas. Data are representative of two independent experiments. b, Recombinant EEEV E2 (residues 1-338) was refolded and purified by size exclusion chromatography (left panel), and analysed by SDSPolyacrylamide gel electrophoresis under non-reducing (NR) and reducing (R) conditions (right panel). Data are representative of two independent experiments. c, Purified anti-EEEV mAbs were tested for binding to recombinant EEEV E2 glycoprotein by ELISA. Data are the mean and s.d. of two independent experiments performed in duplicate. d, HEK-293T cells were transfected with EEEV, VEEV or WEEV pE2-6K-E1 structural genes and stained with EEEV mAbs, anti-VEEV mAb (3B4C-4), anti-WEEV mAb (WEEV-23) or an isotype control mAb (anti-HCV, H77.39). Data are from three independent experiments.

Discussion, we cloned and sequenced the viral RNA. Unexpectedly, all of the sequenced EEEV-18 escape variants (16 of 16 clones) contained a 6-amino acid repeat insertion (VGAQVKY197) in domain B (Fig. 4a and Supplementary Fig. 5). All EEEV-82 escape variant clones (13 of 13 clones) contained a G192R mutation in E2, whereas the EEEV-102 escape variant contained mutations in both domain A (M68T; 3 of 4 clones) and domain B (L227R; 4 of 4 clones) (Fig. 4b,c and Supplementary Fig. 5). The M68R and G192R mutations were introduced individually into the pE2-6K-E1 plasmid to confirm the loss-of-function phenotype. Mutations in M68R or G192R of the E2 gene resulted in abolished binding of EEEV-18, EEEV-82 and EEEV-102 to cells transfected with the pE2-6K-E1 expression plasmid (Fig. 4d). When the M68T, G192R and L227R mutations were introduced into the SINV-EEEV infectious complementary DNA (cDNA) clone, the resultant viruses showed diminished neutralization by EEEV-18, EEEV-82 and EEEV-102 (Fig. 4e). Finally, we tested whether the four neutralization escape variants were resistant to inhibition by the remaining potently neutralizing mAbs. Although all of the strongly neutralizing domain B mAbs (EEEV-3, EEEV-10, EEEV-22, EEEV-69 and EEEV-86) completely neutralized the escape variants with EC50 values similar to the parental virus, domain A (EEEV-5 and EEEV-66) and domain A/B (EEEV-18, EEEV-82 and EEEV-107) mAbs failed to neutralize the escape variants as efficiently (Supplementary Fig. 6).

MAb protection in mice. We assessed whether the mAbs could confer protection against EEEV infection in vivo (Fig. 5). We tested a subset of mAbs with differing neutralization potencies using a lethal challenge model in five-week-old CD-1 mice with a highly pathogenic EEEV (strain FL93-939) engineered to express nanoluciferase with little effect on virulence30. Mice received a single 100 µg (5 mg kg−1) dose of EEEV mAbs via the intraperitoneal route either before (−24 h) or after (+24 h) or after (+24 h) subcutaneous (106 plaque-forming units (PFU) of EEEV) or aerosol (50–100 median lethal dose, LD50) inoculation of EEEV. Mice treated with neutralizing anti-EEEV mAbs (EEEV-3, EEEV-22, EEEV-43, EEEV-58, EEEV-73, EEEV-82 and EEEV-86; EC50 values of 2.2–761 ng ml−1) before subcutaneous challenge had 80–100% survival rates, whereas administration of EEEV-26B, a poorly neutralizing mAb (EC50 > 12,500 ng ml−1) showed little protection (Fig. 5a). When mice were subjected to
a subcutaneous challenge and administered a single dose of mAb 24 h after infection (Fig. 5b) most neutralizing mAbs (EEEV-3, EEEV-18, EEEV-43, EEEV-58, EEEV-73 and EEEV-82) exhibited moderate-to-high levels of protection (40–100% survival rates), whereas EEEV-22, EEEV-86 and EEEV-26B exhibited less protection. Unexpectedly, the modestly neutralizing EEEV-43 mAb (EC_{50} of 761 ng ml^{-1}) still conferred protection (70% survival rate) when administered as post-exposure therapy in this model. Additions of mAb combinations targeting domain A (EEEV-18) and domain B (EEEV-3) and subcutaneous challenge resulted in 100% protection as prophylaxis and 75% protection as post-exposure therapy (Fig. 5a,b).

As EEEV is also highly pathogenic via the aerosol route, we examined the efficacy of the mAbs on an aerosol challenge with 50–100 LD_{50} of EEEV FL93-939. Among the mAbs tested, a majority (EEEV-3, EEEV-5, EEEV-18, EEEV-58 and EEEV-82) protected against death (70–100% survival) when administered as prophylaxis (Fig. 5c). Administration of a mAb combination (EEEV-3 + EEEV-18) as prophylaxis resulted in a 94% survival rate (Fig. 5c). In vivo imaging of mice treated with mAbs EEEV-3, EEEV-18, EEEV-82 and EEEV-86, but not the isotype control mAb, showed marked reductions in viral replication as judged by a decrease in light signal 4 days post-infection (Fig. 5e). However, in the most stringent model of protection, post-exposure therapy at 1 day after aerosol challenge, lower survival rates (10–20%) were observed with individual neutralizing mAbs EEEV-3, EEEV-5, EEEV-18, EEEV-22, EEEV-58, EEEV-69, EEEV-82 and EEEV-86 or a combination of neutralizing mAbs (EEEV-3 + EEEV-18) (Fig. 5d).

Discussion

EEEV is a highly pathogenic, encephalitic alphavirus that lacks approved vaccines or therapies. We generated a panel of 76 mAbs that bound to EEEV-infected cells, including 18 strongly neutralizing mAbs. Ten of the 18 mAbs exhibited potent neutralizing activity with EC_{50} values of <10 ng ml^{-1}. Mapping studies show that these strongly neutralizing mAbs principally recognized epitopes in domains A and/or B of the E2 glycoprotein. Mechanism of action studies revealed that most of the inhibitory mAbs blocked
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for non-neutralizing antibodies against other arthritogenic alphaviruses, to residues within the A and B domains on CHIKV E2 glycoprotein. Using a combination of alanine-scanning and targeted mutagenesis of E2 and neutralization escape selection, we mapped the epitopes for neutralizing anti-EEEV mAbs to residues within these domains. Regions in the E2 domains A and B have been implicated as epitopes for neutralizing mAbs against other alphaviruses including VEEV, CHIKV, SINV and Ross River virus. Our most potently neutralizing mAbs (EEEV-5, EEEV-8, EEEV-82, EEEV-102 and EEEV-107) recognize an epitope in the ‘wing region’ (residues 51–81) on E2, a receptor for several arthritogenic alphaviruses, including Semliki Forest virus and CHIKV. Whereas others have immunized mice with recombinant EEEV E2 glycoprotein or inactivated EEEV to obtain mAbs, we speculate that we obtained a large number of neutralizing mAbs because mice were immunized with a replicating virus that displayed EEEV structural proteins in their native form. At present, it remains unclear why we obtained only type-specific neutralizing mAbs.

Neutralizing antibodies against alphaviruses inhibit infection at several stages in the viral replication cycle including attachment, entry, fusion or egress. Our most inhibitory neutralizing mAbs to E2 domains A and/or B did not block viral attachment to cells; instead, they inhibited infection at a post-attachment stage. Plasma membrane fusion assays showed that several of these mAbs block pH-dependent fusion with membranes. Among the mAbs tested that inhibited infection at a post-attachment step, generally, those recognizing epitopes in domain B (EEEV-5, EEEV-10, EEEV-22, EEEV-66 and EEEV-86) showed less potency when antibody was added after the virus attached to the cells. A previous study with domain B mAbs against CHIKV suggested that bivalent engagement of the virion was necessary for potent neutralization. It is possible that the anti-EEEV mAbs may also require bivalent engagement for complete neutralization; this mode of recognition may be technically difficult to achieve once the virion has attached to cells because some epitopes are unavailable for binding. One of the neutralizing mAbs, EEEV-69, paradoxically increased virus attachment to Vero cells; unexpectedly, increased plasma membrane fusion was observed with EEEV-66, EEEV-82, EEEV-102 and EEEV-107. These results are analogous to prior reports with anti-VEEV and anti-SINV mAbs, both of which increased attachment by stabilizing the interaction between the virus and cells. The increase in fusion could be due to antibody-induced exposure of cryptic epitopes that facilitates virus binding to the plasma membrane, a mechanism previously reported with a flavivirus. This phenomenon may not impact the neutralizing activity of these mAbs if neutralization occurs at a stage in the entry pathway before fusion or plasma membrane fusion is not equivalent to endosomal fusion.

Some reports have speculated that domains A and B on the E2 glycoprotein contain a site of receptor engagement for multiple alphaviruses. A recent study mapped the binding site of Mrx8a, a receptor for several arthritogenic alphaviruses, to residues within the A and B domains on CHIKV E2 glycoprotein. Using a combination of alanine-scanning and targeted mutagenesis of E2 and neutralization escape selection, we mapped the epitopes for neutralizing anti-EEEV mAbs to residues within these domains. Regions in the E2 domains A and B have been implicated as epitopes for neutralizing mAbs against other alphaviruses including VEEV, CHIKV, SINV and Ross River virus. Our most potently neutralizing mAbs (EEEV-5, EEEV-8, EEEV-66, EEEV-82, EEEV-102 and EEEV-107) recognize an epitope in the ‘wing region’ (residues 51–81) on E2, a solvent-exposed site at the distal tip of the A domain. The neutralizing mAbs that mapped to domain B preferentially bound to two epitopes at residues 180–182 (EEEV-3, EEEE-10, EEEE-21, EEEE-22 and EEEE-86) or residues 213–215 (EEEV-4, EEEE-19, EEEE-21, EEEE-60 and EEEE-69). Cryo-electron microscopy (cryo-EM) studies with two neutralizing anti-VEEV mAbs (F5 and B4C-4) showed binding to sites proximal to and within the wing region of domain A (residues 73–120) or to residues 177–223 in domain B, respectively. These mAbs are thought to neutralize VEEV infection by preventing the structural rearrangements required for fusion.

Through neutralization escape selection, we also mapped neutralizing mAbs (EEEV-18, EEEE-58 and EEEE-102) to residues spanning domains A and B (residues 68, 192–197 and 227). We note that the corresponding M68 residue on the CHIKV p62-E1 structure is located beneath the β-strand i6 (residues 74–79) and is not solvent-exposed. Residue M68 is tightly packed against residue L81, a key binding residue for mAbs EEEE-58, EEEE-66, EEEE-82, EEEE-102 and EEEE-107. We hypothesize that the mutation of either residue (M68 or L81) perturbs the conformal display of the domain A ‘wing region’ epitope. Mutation of the

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Table 1 | Profiles of strongly neutralizing antibodies against EEEV

| Antibody | Isotypea | E2 domain | E2 alanine/arginine residues which reduced mAb binding | EC50 (ng ml−1) | EC90 (ng ml−1) | EC99 (ng ml−1) |
|----------|----------|------------|--------------------------------------------------|----------------|----------------|----------------|
| EEEV-3   | IgG2c    | B          | I180, H181, S182                                  | 5.6            | 53.2           | 619.5          |
| EEEV-5   | IgG2c    | A          | K74                                              | 31.8           | 126.1          | 566.3          |
| EEEV-10  | IgG2c    | B          | I180, H181, S182                                  | 3.4            | 33.5           | 411.6          |
| EEEV-18  | IgG3     | A/Bb       | M68, G192, A193, Q194, V195, K196, Y197          | 7.7            | 23.2           | 78.1           |
| EEEV-22  | IgG2c    | B          | I180, H181, S182                                  | 6.3            | 42.7           | 341.9          |
| EEEV-58  | IgG2c    | A/B        | K56, T57, D58, G59, D61, M68, K74, S75, L81, G192 | 4.3            | 66.3           | 1302           |
| EEEV-66  | IgG2c    | A          | D58, L81                                         | 1.9            | 19.6           | 244.8          |
| EEEV-69  | IgG3     | B          | H213, T215                                       | 9.3            | 17.7           | 35.7           |
| EEEV-82  | IgG3     | A/B        | M68, L81, G192                                   | 6.8            | 17.2           | 47.2           |
| EEEV-86  | IgG2c    | B          | I180, H181, S182                                  | 2.2            | 12.5           | 82.1           |
| EEEV-102 | IgG3     | A/B        | M68, L81, G192, L227                              | 4.3            | 20.3           | 110.7          |
| EEEV-107 | IgG2c    | A/B        | T57, D58, M68, Q73, S75, L81, G192                | 11.4           | 96.2           | 985.3          |

aThe Ig isotype was determined by ELISA. bA/B indicates domains A and B. ND, not done.
Neutralizing mAbs map to domain A or B on the E2 glycoprotein. a. Epitope residues of anti-EEEV mAbs identified by alanine-scanning mutagenesis and viral escape are indicated on the EEEV subtype I (strain FL93-939, GenBank accession no. EF151502), subtype II (strain BR56-BeAn512, GenBank accession no. AF159559), subtype III (strain PE-0.0155, GenBank accession no. DQ241304) and subtype IV (BR85-436087, GenBank accession no. AF159561) E2 glycoprotein sequences. Anti-EEEV mAbs mapped to domain A or A/B are depicted as circles and mAbs mapped to domain B are depicted as squares. b. Key domain B residues necessary for mAb engagement are highlighted in purple on the CHIKV p62-E1 monomer (PDB 3N41) and trimer (PDB 5ANY). c. The E1 glycoprotein is in grey, the E2 glycoprotein is in cyan and the E1 fusion loop is in orange. d. The binding data of key domain A residues identified from alanine-scanning mutagenesis are shown for potently neutralizing mAbs. e. The binding data of key domain A residues identified from alanine-scanning mutagenesis are shown for potently neutralizing mAbs. f. The binding data of key domain B identified from arginine or glutamic acid mutagenesis are shown for potently neutralizing mAbs. g. The binding data of key domain A residues identified from arginine or glutamic acid mutagenesis are shown for potently neutralizing mAbs. Residues were identified as critical if <25% mAb binding was observed and >70% binding was retained by the oligoclonal EEEV mAb control. Data are the mean and s.d. from two independent experiments.
neutralization escape variants. Although we speculate that the binding site of EEEV-66 may be similar to or overlap that of mAbs EEEV-18, EEEV-82 and EEEV-102, higher resolution structural studies (for example, X-ray crystallography or cryo-EM) will be required to determine the precise antibody footprints.

The composite AB domain epitope, which bridges the two domains, is analogous to the site recognized by the neutralizing anti-CHIKV mAb (CHK-265), which binds and cross-links these domains on adjacent spikes on the virion surface. The cross-linking of two E2 subunits by CHK-265 restricts domain B from undergoing conformational changes and prevents the exposure of the fusion loop located underneath in the E1 subunit. A similar mechanism may occur with the strongly neutralizing EEEV mAbs EEEV-18, EEEV-82, EEEV-102 and EEEV-107.
Several of our highly neutralizing mAbs showed substantial protective efficacy when mice were challenged with EEEV by a subcutaneous or aerosol route. In the lethal subcutaneous challenge models, mAb protection correlated most consistently with potent neutralization activity and binding to residues spanning domains A and B of the E2 glycoprotein (EEEV-18, EEEV-58 and EEEV-82). One strongly neutralizing domain B mAb (EEEV-3) also protected efficiently in these models. Most of these mAbs (EEEV-3, EEEV-18 and EEEV-58) neutralized infection at a post-attachment stage and efficiently blocked viral plasma membrane fusion. Unexpectedly, EEEV-43, a weakly neutralizing mAb (EC_{50} of 761 ng ml^−1), and EEEV-73 (EC_{50} of 49.7 ng ml^−1), a moderately neutralizing mAb, both protected when administered as prophylaxis or therapy. Analogously, a non-neutralizing anti-EEEV mAb protected against subcutaneous EEEV challenge in mice when administered 1 day before infection. Although further studies are warranted, we...
speculate that FC effector functions may contribute to the in vivo efficacy of weakly to moderately neutralizing protective mAbs. Alternatively, the neutralization assays with Vero cells may not fully reflect the inhibitory activity against cell targets in vivo.

The post-exposure mAb therapy trials in the context of aerosol challenge of mice showed limited efficacy. After aerosol challenge, encephalitic alphavirus-infected mice rapidly enter the brain from the olfactory neuroepithelium via olfactory nerves whereas antibody entry is limited by the blood–brain barrier. The combination of high levels of virus and limiting amounts of a single mAb in the brain may result in rapid neutralization escape. Indeed, the use of a single neutralizing anti-CHIKV mAb promoted escape variants in vivo. However, since combination therapy with highly neutralizing domain A- and domain B-reactive antibodies failed to improve clinical outcome after aerosol challenge, virus entry into the brain may represent a point after which mAb therapy has limited efficacy against EEEV in mice.

Currently, there are no approved vaccines against EEEV. Vaccine efforts against HIV, hepatitis C virus (HCV) and influenza virus focus on eliciting antibodies to protective epitopes on viral envelope proteins through “reverse vaccinology.” Our study identifies specific epitopes on the E2 glycoprotein that can be engaged by potently neutralizing EEEV mAbs. Studies are planned to apply this information to the next generation of vaccine design against EEEV and other encephalitic alphaviruses.

Methods
Animal ethics statement. All animal procedures were carried out in accordance with Association for Assessment and Accreditation of Laboratory Animal Care-approved institutional guidelines for animal care and use and approved by the Institutional Animal Care and Use Committees at the University of Pittsburgh and Washington University School of Medicine. Animals were used in accordance with the American Council on Science and Health, which has been approved by the US National Institutes of Health Guide for the Care and Use of Laboratory Animals. All efforts were made to minimize suffering.

Cell lines and plasmids. Vero, HEK 293T and BHK-21 cells were obtained from the American Type Culture Collection and propagated in DMEM supplemented with 5% (Vero and BHK-21) or 10% (HEK 293T) foetal bovine serum (FBS; Omega Scientific), 100 U ml⁻¹ penicillin, 100 µg ml⁻¹ streptomycin and 10 mM HEPES. All cell lines were tested and judged free of Mycoplasma contamination using a commercial kit. The plasmids pKR780-2-E2, pKR780-2-VEEVE and pKR780-2-VEEVE are comprised of the codon-optimized pE2-6K-E1 genes of EEEV FL93-939, VEEV and WEEV, respectively, under the control of a chicken β-actin promoter, which have been cloned into the pCAGGS expression vector (Addgene).

Expression of WT or mutant structural proteins. E2 receptor-encoding plasmids were transfected in HEK-293T cells using Lipofectamine 3000 reagent (Invitrogen). Cells were harvested 48 h after transfection. Recombinant E2 glycoprotein (5 µg ml⁻¹) was immobilized onto Maxisorp ELISA plates (Thermo Fisher Scientific) overnight in sodium bicarbonate buffer, pH 9.3. Plates were washed three times with PBS, 0.05% Tween 20 and blocked with 5% BSA/PBS for 1 h at 37 °C. Anti-EEEV mAbs were diluted in 2% BSA in PBS and incubated for 1 h at room temperature. After serial washing, horseradish peroxidase-conjugated goat anti-mouse IgG (1:2,000 dilution; Thermo Fisher Scientific) was added and incubated for 1 h at room temperature. After washing, plates were developed with Dako 3,3',5,5'-tetramethylbenzidine substrate (Agilent); the reaction was stopped with 2N H₂SO₄ and absorbance was read at 450 nm with a TriStar Microplate Reader (Berthold Technologies). For virus capture ELISA, ultracentrifuged SINV-EEEV virions were immobilized directly on Maxisorp ELISA plates for 1 h at room temperature. Virus ELISAs were performed similarly, but Tween 20 detergent was omitted from the wash buffer.

Expression of WT or mutant structural proteins. Alkaline-scanning mutagenesis was performed on EEEV E2 residues 1–180 with alanine residues mutated to serine. EEEV E2 alanine mutants that exhibited a partial loss of binding phenotype (residues 56–62, 64, 68, 73–79, 81, 192, 180–182, 212–213 and 215) were substituted with arginine residues. For residues with positive charges (K56 and K74), a glutamic acid substitution was made. Plasmids containing the codon-optimized EEEV, VEEV or WEEV pE2-6K-E1 structural proteins or EEEV E2 alanine mutants were transfected in HEK-293T cells using Lipofectamine 3000 (Thermo Fisher Scientific). Sixteen hours post-transfection, cells were washed with PBS and fixed with the F03/Transfection Factor Staining Buffer Set (Thermo Fisher Scientific). Cells were washed twice with PBS followed by another wash with permeabilization buffer (Thermo Fisher Scientific). Cells were stained with anti-EEEV mAbs 3B4C-4 and anti-VEEV mAb (VEEV-23; S.K.A. and M.S.D., unpublished results) were used as positive controls. After two washes with permeabilization buffer, antibodies were detected with Alexa Fluor 647 conjugated goat anti-mouse IgG (1:2,000 dilution; Thermo Fisher Scientific). After two washes, cells were resuspended in 100 µl of permeabilization buffer and stained with anti-chorionic villi cells (MC1Mx; Biogenezza) to rule out contamination. As previously published criteria, alanine mutants with >25% reactivity compared to WT that exhibited >70% reactivity to a polyclonal anti-EEEV mAb cocktail were deemed as key binding residues.

Generation of virus escape mutants. To generate neutralization escape mutants, SINV-EEEV (1.2 × 10⁶ FFU) were incubated with 1 µg ml⁻¹ of EEEV mAbs for 1 h at 37 °C. The virus–mAb complexes were added to Vero cells. Cell populations were transferred to 24-well plates and cultured for 4 days. Then, the wells were fixed in 3% formaldehyde and permeabilized with 0.1% Triton X-100 and 0.5% Tween-20. After blocking with 1% bovine serum albumin, 1% horse serum and 0.02% NaN₃, the wells were incubated with a mixture of all EEEV mAbs and then washed with PBS. After four washes, the wells were incubated with horseradish peroxidase-conjugated goat anti-mouse antibodies (1:2,000 dilution; Thermo Fisher Scientific). After four washes, the wells were incubated with 3,3′,5,5′-tetramethylbenzidine (Thermo Fisher Scientific) followed by the addition of 2N H₂SO₄ to stop the reaction. After four washes, the absorbance was read at 450 nm with a TriStar Microplate Reader (Berthold Technologies).
The amplified structural genes were sequenced using four primer sets. Escape mutations were introduced into pK8R78-2-EEV containing the codon-optimized pE2 6K-E1 genes of EEEV FL93-939, expressed in HEK-293T cells, stained with anti-EEEV mAbs and analysed by flow cytometry as described earlier.

Mapping of mutations onto the CHIKV p62-E1 crystal structure. Figures were prepared using the atomic coordinates of the CHIKV p62-E1 monomer (PDB 3N41) and trimer (PDB 3ANY) using the PyMOL software (PyMOL Molecular Graphics System, version 1.7.4; Schrödinger).

Attachment inhibition assays. Vero cells were seeded at 3 × 10^4 cells per well 24 h before being assayed. Anti-EEEV mAbs, heparin (Sigma-Aldrich) and BSA (Sigma-Aldrich) were diluted to specified concentrations and incubated for 1 h at 37 °C with SINV-EEEV at a multiplicity of infection of 0.01. The virus–mAb complex was then chilled to 4 °C and added to pre-chilled Vero cells for 1 h at 4 °C. After six washes with chilled PBS, RNA was extracted using an RNeasy Mini Kit (QIAGEN). EEEV RNA levels were determined using a TaqMan RNA-to-Ct, i-Step Kit (Thermo Fisher Scientific) and an E2-specific primer/probe set. EEEV RNA levels were normalized against glyceraldehyde 3-phosphate dehydrogenase and the relative fold change was compared to cells treated with an isotype control mAb.

Pre/post-attachment and post-attachment neutralization assays. Pre/post-attachment neutralization assays were performed by first incubating diluted anti-EEEV mAbs with 10^6 PFU of SINV-EEEV for 1 h at 37 °C. The virus–mAb complex was then added to Vero cells for 1.5 h at 37 °C. Cells were washed with 1% (w/v) methylcellulose in MEM supplemented with 2% FBS. Post-attachment neutralization assays were performed by first incubating Vero cells with 10^6 PFU of SINV-EEEV for 1 h at 4 °C. Cells were washed extensively with cold DMEM to remove unbound virus. Diluted anti-EEEV mAbs were added to virus-adsorbed cells and incubated for 1 h at 4 °C. After a 15 min incubation at 37 °C to allow virus internalization, cells were overlaid with methylcellulose as previously described. Pre/post-attachment and post-attachment neutralization assays were processed similarly to the single end point neutralization assay described earlier.

Fusion inhibition assays. FFWO assays were performed by first allowing viral adsorption to BHK-21 cells (multiplicity of infection, 25) for 1 h at 4 °C. Unbound virus was removed by washing with chilled PBS. Diluted mAbs (50 μg/ml) were added to virus-adsorbed cells for 30 min at 4 °C. Cells were washed with chilled PBS. FFWO was induced by pulsing with fusion medium (Roswell Park Memorial Institute 1640, 10 mM HEPES, 0.2% BSA and 30 mM succinic acid, pH 5.5) for 2 min at 37 °C. A non-fusion control was included using control media (Roswell Park Memorial Institute 1640, 10 mM HEPES, 0.2% BSA, pH 7.6). After the 37 °C pulse, cells were washed twice with chilled PBS and incubated in DMEM supplemented with 5% FBS, 10 mM HEPES, 100 U/ml penicillin, 100 μg/ml streptomycin and 20 mM NaCl to prevent infection via endocytosis. Infection was allowed to proceed for 5 h and cells were detached and fixed with the Fix/3% Transcription Factor Staining Buffer Set (Thermo Fisher Scientific). Cells were stained with human mAb EEEV-53 (L.E.W. and J.E.C., unpublished results) at 1 μg/ml in permeabilization buffer and incubated for 1 h at 4 °C. After two washes with permeabilization buffer, viral antigen was detected with Alexa Fluor 647 conjugated goat anti-human IgG (1:2,000 dilution; Thermo Fisher Scientific). After two washes with permeabilization buffer, cells were resuspended in 100 μl and analysed on a MACSQuant Analyzer.

Mouse protection studies. Five-week-old female CD-1 mice (Charles River) were administered 100 μl of anti-EEEV mAb or isotype control mAb via an intraperitoneal route 24 h pre- or post-challenge. For combined antibody testing, two washes with permeabilization buffer, cells were resuspended in 100 μl and analysed on a MACSQuant Analyzer.

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Acknowledgements
This work was supported by the Defense Threat Reduction Agency (grant no. HDTRA1-15-1-0013 to M.S.D. and W.B.K. and grant no. HDTRA1-13-1-0034 to J.E.C) and National Institutes of Health grant no. R01 AI095436 to W.B.K.

Author contributions
A.S.K., S.K.A., A.Z., M.K.S., D.S.R., D.W.T., W.B.K. and M.S.D. designed the experiments. A.S.K., S.K.A., C.L.G., A.Z., D.W.T., C.S. and K.B. performed the experiments. A.S.K., S.K.A., C.L.G., D.H.F., A.Z., D.W.T. and K.B. performed the data analysis. L.E.N., J.E.C. and D.H.F. contributed key reagents. D.S.R. and D.H.F. contributed the methodology. A.S.K. and M.S.D. wrote the initial draft of the manuscript, with the other authors providing comments and edits to the final version.

Competing interests
M.S.D. is a consultant for InBios International and is on the Scientific Advisory Board of Moderna. J.E.C. has served as a consultant for Takeda Vaccines, Sanofi Pasteur, Pfizer and Novavax, is on the Scientific Advisory Boards of CompuVax, GigaGen, Meissa Vaccines, PaxVax, and is the Founder of IDBiologics.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41564-018-0286-4.

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The authors declare that all data supporting the findings of this study are available within the paper and its Supplementary information. The Supplemental Tables provide data on the newly-generated mAbs, and mutagenesis (alanine and arginine) mapping of the mAb binding sites on EEEV E2 protein.
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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | We used a power calculation (80% power, 0.05 type I error) to see an 3 to 5-fold effect in vivo (depending on data distribution), which was an n=10. |
| Data exclusions | No data were excluded |
| Replication | All cell culture experiments were repeated multiple independent times. In vivo experiments were performed with independent repeat experiments. |
| Randomization | Not randomized. There was no need to randomized animals for this study. However, the animals were purchased commercially, age- and sex-matched. |
| Blinding | Not blinded. Although the study was not blinded, key experiments were repeated independently by multiple members of the laboratory |

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Materials & experimental systems

| Materials & experimental systems | n/a |
|---------------------------------|-----|
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| Antibodies | ☑ |
| Eukaryotic cell lines | ☑ |
| Palaeontology | ☐ |
| Animals and other organisms | ☐ |
| Human research participants | ☐ |

Methods

| Methods | n/a |
|---------|-----|
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Antibodies used | Anti-EEEV antibodies were validated by Western blotting, binding to recombinant protein, and binding to transfected or infected cells. These EEEV antibodies were generated in this paper |

Validation | Describe the validation of each primary antibody for the species and application, noting any validation statements on the manufacturer’s website, relevant citations, antibody profiles in online databases, or data provided in the manuscript. |

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Authentication | All cells were purchased from ATCC |
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Methodology

Sample preparation

Infection was allowed to proceed for 5 h and cells were detached and fixed with Foxp3/Transcription Factor Staining Buffer Set (Thermo Fisher). Cells were stained with human mAb EEEV-53 (L.E.W. and J.E.C, unpublished results) at 1 μg/ml in permeabilization buffer and incubated for 1 h at 4°C. After two washes with permeabilization buffer, viral antigen was detected with Alexa Fluor 647 conjugated goat anti-human IgG (1:2000, Thermo Fisher). After two washes with permeabilization buffer, cells were resuspended in 100 μl and analyzed on a MACSQuant Analyzer (Miltenyi Biotec).

Instrument

MACSQuant Analyzer (Miltenyi Biotec).

Software

FloJo

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Describe the abundance of the relevant cell populations within post-sort fractions, providing details on the purity of the samples and how it was determined.

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Describe the template used for normalization/transformation, specifying subject space or group standardized space (e.g. original Talairach, MNI305, ICBM152) OR indicate that the data were not normalized.

Noise and artifact removal

Describe your procedure(s) for artifact and structured noise removal, specifying motion parameters, tissue signals and
### Noise and artifact removal
(physiological signals, heart rate, respiration).

### Volume censoring
Define your software and/or method and criteria for volume censoring, and state the extent of such censoring.

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## Statistical modeling & inference

### Model type and settings
Specify type (mass univariate, multivariate, RSA, predictive, etc.) and describe essential details of the model at the first and second levels (e.g., fixed, random or mixed effects; drift or auto-correlation).

### Effect(s) tested
Define precise effect in terms of the task or stimulus conditions instead of psychological concepts and indicate whether ANOVA or factorial designs were used.

### Specify type of analysis:
- [ ] Whole brain
- [ ] ROI-based
- [ ] Both

### Statistic type for inference
Specify voxel-wise or cluster-wise and report all relevant parameters for cluster-wise methods.

(See Eklund et al. 2016)

### Correction
Describe the type of correction and how it is obtained for multiple comparisons (e.g., FWE, FDR, permutation or Monte Carlo).

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## Models & analysis

| n/a | Involved in the study |
|-----|-----------------------|
| x   | Functional and/or effective connectivity |
| x   | Graph analysis |
| x   | Multivariate modeling or predictive analysis |