Exclusive and Common Subsets of Zika Virus Polyprotein Mutants
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Abstract
Two subsets of Zika virus (ZIKV) polyprotein amino acid positions are identified. One subset (Exclusive) consists of mutating amino acid positions which were found only in polyproteins isolated from ZIKV of human origin. A second subset (Common) consists of mutating amino acid positions which were found both in ZIKV polyproteins of human origin and in ZIKV polyproteins of Aedes species mosquito origin. The dominance of the Exclusive subset in the polyprotein was found to range from the N-terminus structural proteins until non-structural protein NS3. Although no longer greater than the Common subset, elements of the Exclusive subset existed almost to the C-terminus of non-structural protein NS3. These results are considered in the context of reported epitopic and other biological characteristics of ZIKV.

Keywords: Zika virus; ZIKV; Polyprotein; Information entropy H; Shannon entropy; Exclusive subset; Common subset

Introduction
Zika virus (ZIKV) causes microcephaly, brain abnormalities and other neural diseases in gestating infants who are infected in utero [1, 2]. Because of the highly adverse effects of ZIKV on gestating infants, it is important to develop preventive vaccines and therapeutic agents.

Reported here is a bioinformatics analysis of ZIKV polyprotein that sorts mutations which occurred exclusively in ZIKV isolated from infected human hosts (Exclusive subset) and mutations which occurred in ZIKV isolated both from infected humans and from infected Aedes mosquito vectors (Common subset). It is proposed that the exclusive mutations may reflect immunological and other biological processes which occur exclusively in human hosts but not in the mosquito vector. These exclusive mutational objects thus may help guide development of human host-specific targets for vaccines and drugs.

Materials and Methods
Complete sets of full-length ZIKV polyprotein sequences isolated from humans (n=212) and from Aedes mosquitos (n=34) were downloaded from the NCBI Zika Virus Resource (http://www.ncbi.nlm.nih.gov/genome/viruses/variation/Zika) on 8 April 2017. Polyprotein domains were assigned by alignment with the default MR 766 reference sequence [3]. Polyprotein sequence management was facilitated with Jalview 2.9.0b2 [4].

Computations were performed using Anaconda 2.4.0 (64-bit) with Python 2.7.10, Numpy 1.10.1, Scipy 0.16.0, and Matplotlib 1.4.3. Information entropy (H) was computed by the equation of Shannon [5] and is expressed in bits. Amino acid positions with H>0.0 in the complete polyprotein dataset. The summed exclusive and common subset counts account for all 344 amino positions where H>0.0 in the complete polyprotein dataset. The mean and standard deviation for H in the Exclusive subset are 0.0745 ± 0.0804 bits. The mean and standard deviation for H in the Common subset are 0.1433 ± 0.1650 bits. In addition to containing fewer subset elements, the average H in the Common subset is significantly greater than the average H in the Exclusive subset (Mann-Whitney U=6525.5, p=7.9471 × 10-17).

Distributions of information entropy (H) in ZIKV polyproteins isolated from humans are shown in Figure 2. The elements of the Common subset predominated in the C-terminal half of the polyprotein amino acid chain (Figure 2a), in non-structural regions of the polyprotein (Figure 2d). H summation curves (Figure 2b) were approximated (+) by the following two indefinite integrals, where x represents the polyprotein amino acid position:

\[ \int H(\text{exclusive}) \, dx = x^4(2.3142 \times 10^{-12}) - x^3(1.2929 \times 10^{-9}) + x^2(1.4392 \times 10^{-6}) + x(5.5747 \times 10^{-3}) + 6.0842 \times 10^{01} \]

\[ \int H(\text{common}) \, dx = x^4(2.6759 \times 10^{-12}) - x^3(1.4469 \times 10^{-9}) + x^2(2.6463 \times 10^{-6}) + x(2.2839 \times 10^{-3}) + 3.8470 \times 10^{01} \]

In equations 1 and 2, the discrete summation curves are used.

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Figure 1: Definition of exclusive and of common subsets of amino acid mutants detected in ZIKAV polyproteins isolated from humans. (top) the positive H-value, objects in the Exclusive subset are represented vertically on the ordinate; (bottom) the positive H values in the Common subset are distributed on the graph surface area, indicating common occurrence of mutant amino acid positions in ZIKV polyproteins isolated from humans and from Aedes mosquitoes.

Figure 2: Distribution of information entropy (H) in ZIKV polyprotein subsets. (a) H distributions in exclusive (red) and in common (black) subsets of ZIKV polyprotein amino acid positions; (b) summation curves are shown for the exclusive subset of amino acid positions (red) and for the common subset of amino acid positions (black); (c) differences (blue) between the exclusive H subset summation and the common H subset summation; (d) a reference diagram of the sequential organization protein components of the polyprotein. In graphs (b) and (c), the solid lines denote observed values while closed circles denote values fit according to equation 1 (Exclusive subset) and equation 2 (Common subset). In figure (d), the ZIKV precursor structural protein components of the polyprotein are, with amino acid positions in parentheses: C (capsid protein, 6-122), prM (precursor membrane protein, 128-290) and E (envelope protein, 292-794); the non-structural precursor protein components are NS1 (797-1148), NS2A (1158-1307), NS2B (1377-1502), NS3 serine protease, helicase (1521-2124), NS4A (2125-2242), protein 2K (2243-2265), NS4B (2270-2512), FtsJ-like methyl transferase (2575-2746) and NS5 ribonuclease (2774-3412).

Figure 3: Predicted H distributions in the exclusive and common subsets of ZIKV polyprotein mutating amino acid positions. H values were predicted from first derivatives of equations 1 and 2 by treating those polynomials as indefinite integrals.
to generate approximations as continuous, indefinite integrals. Difference curves (Δsum, Figure 2c) for the observed summation and approximated polynomials were obtained by subtraction: Δsum = equation (1) - equation (2). The amino acid position with maximum value in the directly computed curve position 2068, which is within the NS3 non-structural protein domain of the polyprotein (Figure 2d). The amino acid position with maximum value in the Δsum curve computed from the polynomial approximation is position 1609, which also is within the NS3 non-structural protein domain of the polyprotein. There were 69 amino acid position in the human Exclusive subset, from position 2068 to the C-terminal, with an H sum of 5.4103 bits, which is 33.17% of the Exclusive subset observed total.

Differentiation of the indefinite integrals in Equations 1 and 2 yield the following 2 equations, which were used to produce the H distribution curves in Figure 3:

\[ \text{H(exclusive)} = x^2(9.2569 \times 10^{-11}) - x^3(3.8788 \times 10^{-10}) + x(2.8785 \times 10^{-10}) + 0.0056 \]

\[ \text{H(common)} = x^2(1.0704 \times 10^{-12}) - x^3(4.3407 \times 10^{-10}) + x(5.2926 \times 10^{-10}) + 0.0023 \]

The polyprotein position at which H(exclusive)=H(common) in the continuous model of equations 3 and 4 is between amino acid positions 1608 and 1609 which, as noted above, is within the NS3 domain.

The data in Figures 2 and 3 suggest that dominance of the human Exclusive subset of mutations extends beyond the structural protein components of the polyprotein. Precursors of the structural proteins comprise amino acids 1-794 of the polyprotein. The greater mutability of the elements of the human Exclusive subset, in comparison with the Common subset, probably represents functions of processes occurring in the human host but not in the mosquito vector. Such processes may include immunological escape [7], antibody dependent enhancement [8] and other biochemical and biophysical factors present in the host but not in the mosquito vector. For example, viral interactions with interferons are important determinants of outcome of ZIKV infection [9]. There are other, specific immunological human host factors that affect the Zika virus. Neutralizing antibodies against the prM precursor membrane protein (prM) and envelope E protein have been reported [10-12]. The prM and E proteins initially exist as precursors within the structural domain of the polyprotein. Antibody and T-cell responses against the NS1, NS2, NS3, NS4B and NS5 non-structural polyproteins have been reported [10,11,13,14].

The distribution of the elements of the human Exclusive subset is in agreement with the physiological and immunological Zika virus-host interaction processes discussed above. The activity of the Exclusive subset is detectable well into the non-structural domain of the polyprotein. A more detailed elucidation of the mutational pattern in the human Exclusive subset may increase our understanding of Zika virus biology, thereby facilitating the development of anti-Zika strategies, preventives and therapeutics.

References