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Identifying allosteric small molecule binding sites of inactive NS2B-NS3 proteases of pathogenic *Flaviviridae*

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Abstract: Dengue, West Nile, Zika, Yellow fever, and Japanese encephalitis viruses persist as a significant global health threat. The development of new therapeutic strategies based on inhibiting essential viral enzymes or viral-host protein interactions is problematic due to the fast mutation rate and rapid emergence of drug resistance. This study focuses on the NS2B-NS3 protease as a promising target for antiviral drug development. A promising allosteric binding sites were identified in two conformational distinct inactive states and characterized for five flaviviruses and four Dengue virus subtypes. Their shapes, druggability, inter-viral similarity, sequence variation, and susceptibility to drug-resistant mutations have been studied. Two identified allosteric inactive state pockets appear to be feasible alternatives to a larger closed pocket near the active site, and they can be targeted with specific drug-like small molecule inhibitors. Virus-specific sequence and structure implications, and the feasibility of multi-viral inhibitors is discussed.

Keywords: ZIKA virus; protease inhibitors; mutation rates; allosteric druggable pockets; Dengue; Yellow Fever; Japanese encephalitis; NS2B; NS3

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1. Introduction

Finding small molecule therapeutics to treat emerging flaviviral diseases, dangerous human health, remains an unsolved problem. The most problematic mosquito-vectored *Flaviviridae* are the following: Zika virus (ZIKV), four subtypes of Dengue virus (DENV), West Nile virus (WNV), Japanese Encephalitis virus (JEV), and Yellow Fever virus (YFV) [1–3]. The existing supportive care and symptomatic treatments have limited efficacy, calling for the exploration of new viral targets essential for viral infectivity and growth, as well as identifying the relevant functional states for structure-based drug design [4]. These treatments should be able to reduce or prevent severe viscerotropic injuries caused by YFV, protect the human brain from the neurotropic injuries accompanying JEV, ZIKV, DENV, and WNV-related diseases [5], and prevent immune-mediated complications [6], cytokine storm, systemic vascular leakage, organ failure, or sepsis [7].

The genomes of the eight flaviviruses of interest consist of single-stranded **positive-sense RNA** coding for the polyprotein (PP) precursor [8,9]. The length and protein composition of the polyprotein **are** similar for the eight viruses [10–15]. The polyprotein consists of three structural proteins, Capsid (C), the precursor membrane glycoprotein (prM), and the envelope glycoprotein (E), and seven nonstructural proteins (NS1, NS2A, NS2B, NS3, NS4A, NS4B, and NS5) [16–20] (Figure 1). The structural proteins are assembled in the endoplasmic reticulum and processed in the Golgi, **where the viral particles mature into infectious virions**. Non-structural proteins form replication complexes in the endoplasmic reticulum to replicate viral RNA [21]. The polyprotein is cleaved by the viral protease NS2B-NS3-pro and human proteases. This cleavage is essential for viral maturation

and growth, thus making the NS2B-NS3 inhibition an attractive target for an antiviral agent [22–24].

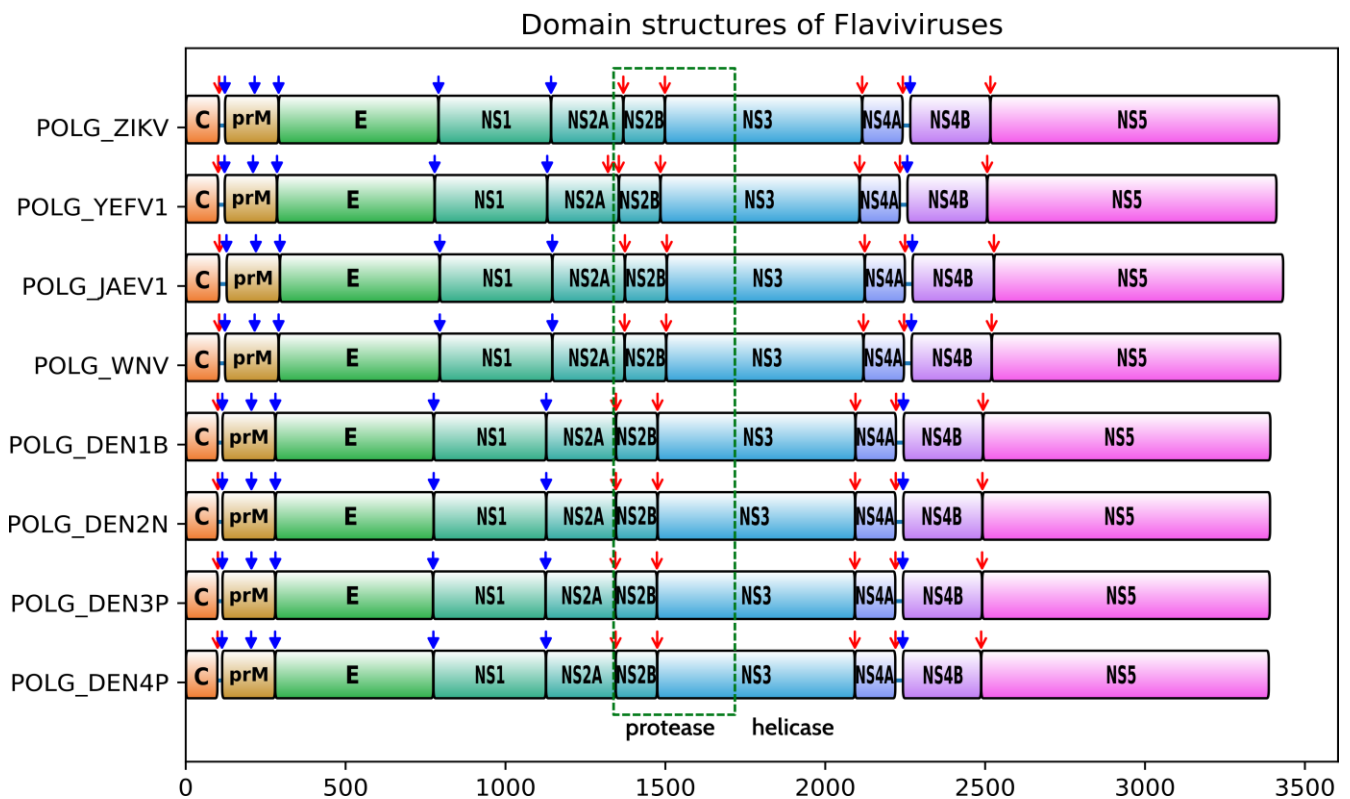


Figure 1. Domain organization of the polyprotein precursor encoded by different flaviviruses. Red arrows indicate the cleavage by the NS2B-NS3 protease and blue arrows indicate the cleavage by the host cell proteases. A green box shows the two domains of the protease.

The viral protease domain is designed in a particularly intricate manner. The active protease requires an NS2B fragment that wraps around the rest of the protease domain that includes the catalytic triad (His51, Asp75, and Ser135) [8,19]. NS3, a dual-function protein, consists of two domains: the N-terminal trypsin-like serine protease (NS3-pro) that binds to NS2B, and the C-terminal helicase (NS3hel). NS2B acts as a cofactor for NS3-pro, undergoing a conformational change necessary to activate the protease. The multiple sequence alignment of NS2B-NS3-pro-NS3hel revealed a high level of sequence conservation [8,17,19,25]. **However, it remains uncertain whether the inhibitor binding sites exhibit an even higher degree of conservation, warranting efforts to develop a single pan-flaviviral inhibitor.**

The inhibition of the active site **pockets** of NS2B-NS3 protease has been studied recently as a therapeutic strategy [17,21,26,27]. Highly potent macrocyclic inhibitors of the Zika protease active site with Ki under 5nM were identified [28]. However, the antiviral efficacy **in cell-based assays was low, only in the micromolar range, according to a follow-up study from the same laboratory** [29]. Therefore, a search for an alternative “inactive” conformational state and related allosteric pocket began. Our previous study identified compound R107 as a potential inhibitor of the “super-open” and inactive allosteric pocket of the NS2B-NS3-pro protease of ZIKV [24]. In the current study, we explored the feasibility of extending the initial finding for ZIKV to other infectious flaviviruses by analyzing over 13,600 sequences and 92 three-dimensional structures for the sequence variability of each virus in the vicinities of three identified pockets. The side chains lining the pockets of all proteases were studied for both intra-species variations, and inter-species differences for the five viruses and three additional Dengue types. The impact of that variation

on drug resistance, the “druggability” of the allosteric pockets, and the feasibility of small molecule pan-viral inhibitors is presented. 74
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2. Materials and Methods 76

2.1. Collection, superposition, and analysis of 3D structures of NS2B-NS3 proteases 77

The NS3-pro domain sequences extracted from the flavivirus sequences were searched against the Protein Data Bank, PDB [30], to identify close sequence matches with experimentally determined protein structures by NCBI Blast with a threshold of 90% sequence identity. The structures were split into individual conformationally different conformers of NS2B-NS3 and superimposed by NS3 to further classify them into the three conformational and functional states: closed active state, transient inactive state, and a fully opened inactive state (also known as “super-open”). The initial preparation has been performed as follows. 78
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For ZIKV, 59 structures were used for analysis. These structures contained NS2B and NS3-pro subunits. Most of those structures had NS2B chain as a separate polypeptide cleaved from the NS3 domain, while some contained one continuous NS2B-NS3 polypeptide chain. We assigned uniform residue numbers for residues of each domain. 86
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Three structures available for YFV were kept as is. For WNV, six structures were analyzed. One JEV protease structure, 4R8T [31], was used. 90
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DENV-1 was represented by 3L6P [32] and 3LKW [32] entries, each containing a single chain with both protease domains. For DENV-2, eleven protease structures were retained. For Dengue 3, two structures, 3U1I [33] and 3U1J [33], were identified. Finally, ten DENV-4 structures were used for the study. 92
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2.2. Sequence analysis 96

The deposited RNA sequences of all studied viruses have been downloaded from the NCBI Virus community portal that combined RefSeq, GenBank, and other NCBI repositories [34]. In total of 13,692 sequences were used for the pocket conservation analysis. 97
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2.2.1. Zika Virus NS2B-NS3 sequence set compilation and alignment 100

The ZIKV set contained 2,483 polyprotein amino acid sequences. They were preprocessed to extract only the NS2B and NS3-pro regions (residues 1,369-1,676 according to the InterPro entry PS51527). Nine hundred eighty protease sequences (82 unique sequences) fully representing the set have been aligned using the zero-gap global alignment algorithm ZEGA within ICM-PRO v3.9-3b [35]. The alignment was analyzed for position-dependent variability and attribution to three different pockets according to a 3D model of NS2B-NS3-pro. 101
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2.2.2. West Nile Virus NS2B-NS3 sequence set compilation and alignment 108

The WNV set contained 7,627 polyprotein amino acid sequences. They were preprocessed to extract only the NS2B and NS3-pro regions (residues 1,371-1,502 for NS2B and 1,502-1,679 for NS3-pro according to the InterPro [36] entry PS51527). Four hundred eighty-six protease sequences (67 unique sequences) fully representing the set have been aligned using the zero-gap global alignment algorithm ZEGA within ICM-PRO v3.9-3b [35]. The alignment was analyzed for position-dependent variability and attribution to three different pockets according to a 3D model of NS2B-NS3-pro. 109
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2.2.3. Yellow Fever Virus NS2B-NS3 sequence set compilation and alignment 116

The YFV set contained 1,967 polyprotein amino acid sequences. They were preprocessed to extract only the NS2B and NS3-pro regions (residues 1,371- 1,665 according to the InterPro [36] entry PS51527). Nine hundred ten protease sequences (119 unique sequences) fully representing the set have been aligned using the zero-gap global alignment algorithm ZEGA within ICM-PRO v3.9-3b [35]. The alignment was analyzed for position- 117
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dependent variability and attribution to three different pockets according to a 3D model of NS2B-NS3-pro. 122
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2.2.4. Japanese Encephalitis Virus NS2B-NS3 sequence set compilation and alignment 124

The JEV set contained 3,292 polyprotein amino acid sequences. They were preprocessed to extract only the NS2B and NS3-pro regions (residues 1,374-1,504 for NS2B and 1,505-1,682 for NS3-pro according to the InterPro [36] entry PS51527). Two hundred forty-six protease sequences (53 unique sequences) fully representing the set have been aligned using the zero-gap global alignment algorithm ZEGA within ICM-PRO v3.9-3b [35]. The alignment was analyzed for position-dependent variability and attribution to three different pockets according to a 3D model of NS2B-NS3-pro. 125
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2.2.5. Dengue -1, -2, -3, -4 Viruses NS2B-NS3 sequence set compilation and alignment 132

The DENVs set contained 15,387 for DENV-1, 13,326 for DENV-2, 7,126 for DENV-3 and 3,889 for DENV-4 polyprotein amino acid sequences. They were preprocessed to extract only the NS2B and NS3-pro regions (residues 1,346-1,475 for NS2B and 1,476-1,653 for NS3-pro according to the InterPro [36] entry PS51527). The fully representing sequence set have been aligned using the zero-gap global alignment algorithm ZEGA within ICM-PRO v3.9-3b [35]: 133
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DENV-1: 4,628 sequences (614 unique sequences) 139

DENV-2: 3,816 sequences (553 unique sequences) 140

DENV-3: 1,739 sequences (263 unique sequences) 141

DENV-4: 862 sequences (202 unique sequences) 142

The alignment was analyzed for position-dependent variability and attribution to three different pockets according to a 3D model of NS2B-NS3-pro. 143
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2.3. Analysis and visualization 145

The analysis, superposition, and visualization were done using ICM-Pro 3.9-3b molecular modeling software [35,37,38]. The domain plots of the polyprotein precursor were generated with Python and matplotlib [39] based on the data from InterPro and UniProt [36,40]. The assignment of the three functional and conformational states to each protease structure was done as follows. 146
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- The active *Closed* state is defined by the distance between C α -atoms of D75 of NS3-pro and G82 of NS2B being smaller than 10Å. All other models of the inactive state with the NS2B loop unwrapped are further sub-divided by conformation of the C-terminal end of NS3-pro according to the values of angle defined below. 151
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- The inactive *Transient* state is characterized by the C-terminal fragment hairpin displacement following the unwrapping of NS2B. The extension of the C-terminal end region for this state is defined by the C α atoms of G148, N152 and S158 (numbering from DENV-2) residues of NS3-pro being over 90°. 155
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- The inactive *Fully Opened* state is characterized by the hairpin unfolding and rearrangement of the NS3-pro C-terminal fragment with the angle defined above being less than 90°. 159
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3. Results 162

To analyze drug-targetable allosteric binding sites in the inactive states of the protease, we started by analyzing all experimentally determined structures of the proteases of five flaviviruses, introducing a uniform NS2B-NS3-pro numbering for each virus type and subtype, classifying them by the conformations of NS2B and the NS3-pro C-terminal fragment relative to the protease core, and assigning each protease one of three conformational states: closed, transient, and fully opened. Finally, new druggable pockets in the transient and fully opened state were predicted and sequence variability of immediate pocket surfaces was analyzed. A total of 92 models for five flaviviruses, three subtypes, in 163
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up to three conformational states have been collected and superimposed, and 13,692 sequences of the NS2B-NS3 proteases analyzed.

3.1. Three targetable conformational states of NS2B-NS3-pro for the five viruses

Ninety-two PDB entries for ZIKV, WNV, YFV, JEV, DENV-1, -2, -3, and -4, eight groups in total were analyzed. Each of the entries was processed to extract the best single representative of the NS2B-NS3-pro domain, superimposed to a common reference conformation of the core NS3-pro domain, and assigned one of three conformational states, one active state, and two inactive ones (Figure 2). The geometrical definitions of the conformational states defined by the NS2B and C-Hairpin of NS3-pro are described in Methods 2.3. Each of the eight viruses had PDB structures in at least one of the three states.

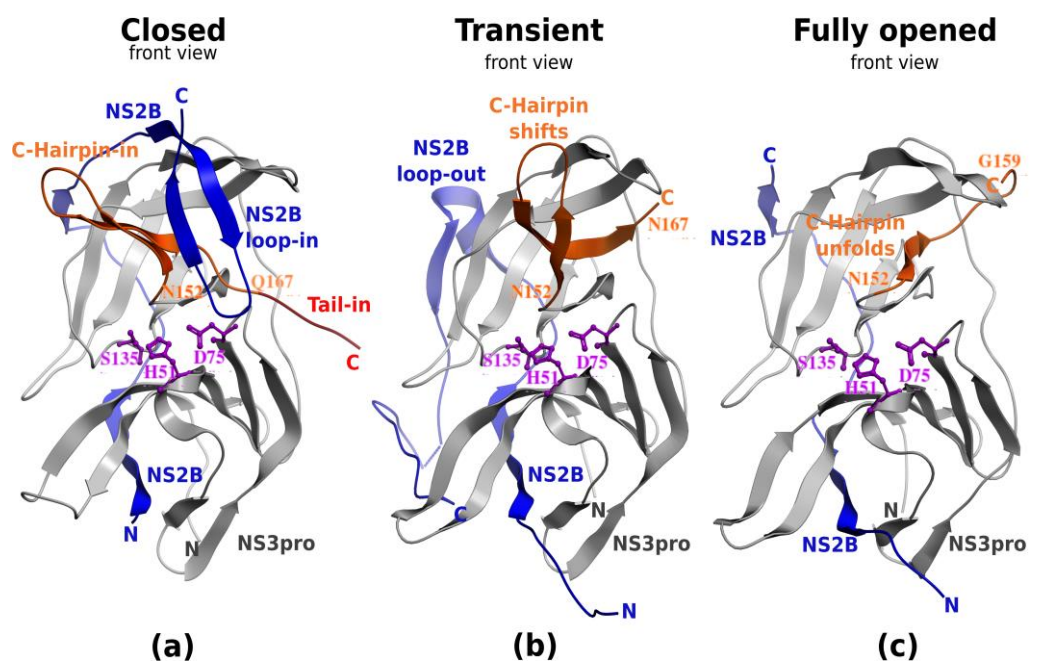


Figure 2. The mobile structural determinants of the three states in existing crystallographic structures of NS2B-NS3 protease domains are shown by blue and orange backbone ribbons. The Closed state (a) is exemplified by PDB ID 5YOF, the Transient state (b) by 2FOM, and the Fully opened (c) by 7M1V. The backbone ribbon is colored as follows: blue ribbon – mobile NS2B, gray ribbon – static NS3 core, orange ribbon – mobile NS3-pro-C-terminal hairpin. The catalytic triad is shown and labeled for reference.

Most structures (69 out of 92) represented the closed/active state, and most co-crystallized inhibitors were bound in the close vicinity of the active site (Table 1). Most closed-state structures were determined for ZIKV protease (53). The other three viruses and four Dengue subtypes had the following number of structures of the closed state: WNV (5), YFV (1), JEV (0), DENV-1 (0), DENV-2 (2), DENV-3 (2), and DENV-4 (6). We used those structures to predict allosteric binding sites of the closed state (see Section 3.2.2)

The inactive transient state was identified in 16 structures out of 92. All those structures were analyzed for alternative allosteric binding pockets that may be targeted for stabilizing the inactive state and/or interfering with the substrate binding. Those pockets, if identified, may represent promising sites for drug design.

Seven structures exhibited a fully opened inactive state with NS2B fully unwrapped, and the C-terminal hairpin unfolded and extended. Interestingly, this C-terminal extension creates a new cavity that may be targeted by a small molecule. The fully opened state conformational was found in six ZIKV PDBs and one JEV structure. The predicted allosteric pockets in inactive conformations overlapped with the pockets identified in the transient states but were different from them. Given the variable nature of the inactive states

and the related allosteric pockets, these additional conformations and pockets presented alternative inactive-state pockets for ligand screening.

To deduce “druggability” of three different types of allosteric pockets, we analyzed the known co-crystallized ligands. Table 1 summarizes the distribution of the conformational states of the proteases for the five viruses and Dengue subtypes with and without the co-crystallized ligands.

Closed conformation/Active State.

Most of the closed conformations with the active site pocket (ASP) were solved with a peptide-like inhibitor. A smaller number of closed structures were crystallized with small molecule ligands only for the Zika virus; however, all the ligands were notably weak binders. The affinities span a micromolar range of 1.5 mM to 200 mM [41–44] (PDB: 5YOD, 5H4I, 7V VX, 6L4Z, 6L50), e.g. 5YOD [41] ligand has IC₅₀ of 1.5 mM, the 5H4I [42] ligand has IC₅₀ of 14.08 mM, 7V VX [43] of 48.7 mM, and 6L4Z [44] and 6L50 [44] were co-crystallized with ligands of IC₅₀ of 200 mM. These values indicate the limitations associated with targeting the active site pocket of the Zika protease, and likely for the other viruses given the ASP pocket similarities.

Structural reasons making the ASP difficult for targeting with small molecules include its charge and shape properties, namely three negatively charged residues, two aspartic acids in NS3-pro, and one glutamic acid in NS2B for YFV, DENV-1, -2, and -4 [41]. In addition, the pocket is relatively flat. While the research aimed at the identification of both covalent and non-covalent inhibitors with promising therapeutic potential continues [45–48], we focused on identifying a different location for a small molecule inhibitor.

Open conformation/Inactive State.

The two structures in a fully opened conformation were crystallized in an apo inactive state, and with a weak (IC₅₀ over 1 μM) [49] small-molecule inhibitor (PDB: 7M1V), both for the Zika protease [25]. This allosteric pocket, distant from the active site, was recently targeted with a structure-based docking screen and experimental testing and a weak allosteric inhibitor was identified [24]. However, this promising binding site can be targeted for more potent and selective inhibitors.

Each of the DENV-1, -2, -4 subtypes of the Dengue virus protease, had at least one transient or fully opened state structure determined experimentally (Table 1). Notably, three DENV-2 Transient state structures, 6MO0, 6MO1, 6MO2 [50] were cocrystallized with IC₅₀ from 200 to 860 nM small-molecule inhibitors [50] useful for both the pocket definition and attempts to improve the inhibition efficacy. The distinction between the two inactive states for these three structures was challenging because the C-terminal hairpin residues were missing; we used the visible stem strands of the loop for the assignment to the Transient state. These inhibitors helped to define an allosteric pocket for all Dengue viruses. This ligand-defined pocket is different from the ligand-defined ZIKV pocket [24,51] even though the two pockets overlapped partially.

The inactive state proteases were crystallized for other flaviviruses, but the structural information is insufficient and completely missing for YFV. Only an apo-structure in the inactive transient conformation (2GGV) [19] was determined for the West Nile virus protease, and one fully opened inactive structure of JEV has been solved (4R8T) [31].

Table 1. Comparative analysis of the existing crystallographic structures of NS2B-NS3 in relation to their conformational states. The sorting of the crystallographic structures of NS2B-NS3 into three conformational states was conducted based on criteria described in section 2.3.

Flaviviruses	Closed		Transient		Fully opened	
Abbreviation	<i>with a peptide like inhibitor</i>	<i>with a ligand</i>	<i>without a ligand</i>	<i>with small-molecule inhibitor</i>	<i>without a ligand</i>	<i>with small-molecule inhibitor</i>
ZIKV	5LC0, 5GJ4, 5H6V, 5YOF, 5ZMQ, 5ZMS, 5ZOB, 6JPW, 6KK2, 6KK3, 6KK4, 6KK5, 6KK6, 6KPQ, 6Y3B, 7DOC, 7O2M, 7O55, 7OBV, 7OC2, 7PFQ, 7PFY, 7PFZ, 7PG1, 7PGC, 7VLG, 7VLH, 7VLI, 7VXY, 7ZLC, 7ZLD, 7ZMI, 7ZNO, 7ZPD, 7ZQ1, 7ZQF, 7ZTM, 7ZUM, 7ZV4, 7ZVV, 7ZW5, 7ZWK, 7ZYS, 8A15, 8AQA, 8AQB, 8AQK	5H4I, 5YOD, 6L4Z, 6L50, 7VXX	5GPI		7M1V	5GXJ, 5T1V, 5TFN, 5TFO, 6UM3
WNV	2FP7, 2IJO *, 2YOL, 3E90, 5IDK				2GGV	
YFV			6URV			
JEV						4R8T
DENV-1					3L6P, 3LKW	
DENV-2	2M9P, 2M9Q			6MO0, 6MO1, 6MO2	2FOM, 4M9F, 4M9I, 4M9K, 4M9M, 4M9T	
DENV-3	3U1I, 3U1J *					
DENV-4	5YVU *, 5YW1 *		5YVV, 5YVW, 5YVY, 7VMV		2VBC, 2WHX, 2WZQ, 5YVJ	

* Crystallographic structure of NS2B-NS3 with bovine pancreatic trypsin inhibitor.

In summary, there is sufficient structural information for targeting allosteric pockets in one of the two inactive states with small molecules as a strategy for developing novel small-molecule inhibitors against all five flaviviruses studied. For proteases with multiple structures in the inactive state, we analyzed the structural variation of each protease in order to be able to evaluate more and less conformationally defined surfaces of the

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protease, and to select the most suitable locations and representatives for further docking screens.

3.2. Conformational variations among flaviviral NS2B-NS3 proteases across the three states

For comparative analysis, representative structures for each of the five viruses and Dengue 1,2,3,4 subtypes were selected with a preference for better resolution and fewer residues missing from the model, or zero occupancy for each of the three states. The following models were prepared and converted to full atom models: the closed state: ZIKV (5YOF) [41], WNV (2FP7) [52], YFV (6URV) [53], DENV-2 (2M9P) [54], DENV-3 (3U1I) [33], DENV-4 (5YW1) [55]; the transient state: WNV (2GGV) [19], DENV-1 (3L6P) [32], DENV-2 (2FOM) [52], DENV-4 (2VBC) [56]; and fully opened state: ZIKV (7M1V) [25], JEV (4R8T) [31]. The closed active state in this analysis was included for comparison with the two inactive states.

The active state conformations of ZIKV, WNV, YFV, DENV-3, and -4, reveal that the closed conformation is structurally conserved not only inside each of the Dengue subtypes, but also across the five species (Figure 3a). The catalytic triad is aligned perfectly between the viruses.

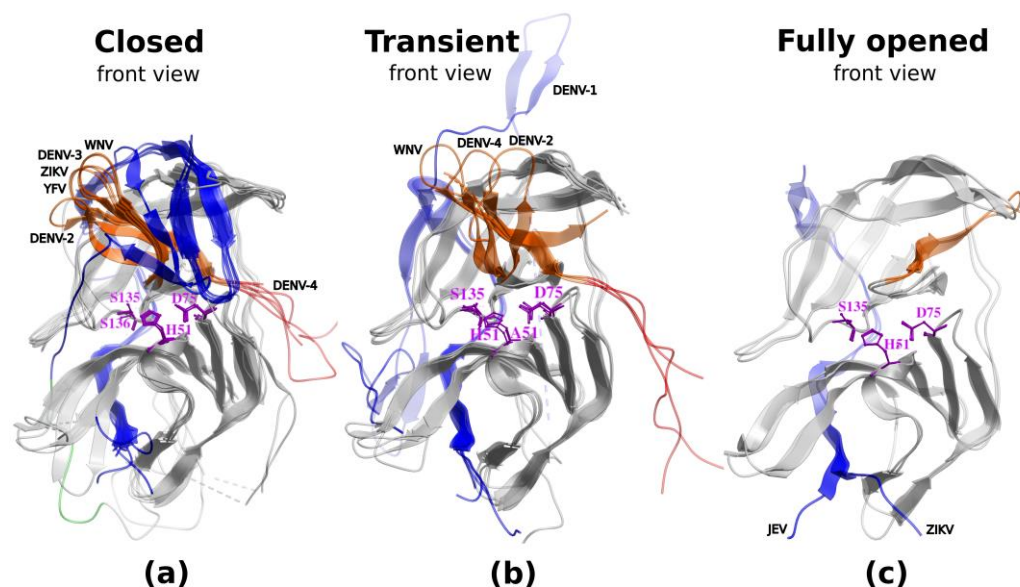


Figure 3. Structural variations within each of the Closed (a), Transient (b), and Fully opened (c) states of the flaviviral NS2B-NS3-pro domains for all five flaviviruses and four Dengue subtypes under study. Some differences in residue numbers (YFV S136 instead of S135 for other viruses) are due to sequence length variations in both NS2B and NS3-pro subunits (a). The NS2B loop in one of the closed-state DENV-2 constructs was conformationally shifted due to an inserted linker in a crystallized construct (see a green backbone fragment in (a)).

The calculation of the backbone deviations of the active site vicinity of the Closed state including the NS2B and C-terminal fragment shows the conformational conservation of the active site vicinity between the viruses and subtypes (Table 2). The Transient state characterized by the NS2B unwrapping and NS3-hairpin preservation, shows the hairpin, forming a complete loop directed upwards. Furthermore, its absence in several flavivirus crystallographic structures [6MO0-2, 2WHX [57], 2WZQ [57], 5YVJ [58]] confirms this flexibility. The NS2B loop rotates and extends backward rather than closing the upper part of the active site. This flexibility creates ample space for the NS3-loop to adopt varying conformations across different flaviviruses, as observed in crystallographic structures from WNV, DENV-1, -2, and -4. In the Transient conformation of DENV-1, the NS2B exhibits a distinct conformation different from other flaviviruses and is directed upwards due to binding to crystallographic neighbors (PDB IDs: 3L6P, 3LKW).

Although there are fewer examples of inactive complexes for Transient and Fully Opened conformations, those conformations are sufficiently conserved to explore alternative mechanisms of inhibition of flavivirus proteases by stabilizing the inactive state. The most promising state for stronger small molecule binders is the Transient state despite some variability of the C-terminal fragment of NS3-pro.

3.3. Identification of druggable pockets in two inactive states of the NS2B-NS3 protease

To identify possible druggable pockets, we performed pocket prediction using ICM-PocketFinder [59]. ICM pocket finder relies on generating cumulative grid-potential based maps that evaluate van der Waals interactions between the receptor and a virtual ligand and contouring the maps to identify contiguous density regions. Detailed characteristics of the active site, and further evaluation of their druggability is performed with the DLID measure [60]. Three pocket locations, denoted as allosteric pockets 1, 2, and 3 (AP1, AP2, AP3) were further analyzed for druggability and sequence conservation between the viruses and Dengue subtypes. The intraviral sequence conservation around those pockets for each of the 8 groups was analyzed in Section 3.4.

3.3.1. Pocket locations, composition

The three allosteric pockets were identified by the following procedure. First, the ICM pocket finder tool generated the likely pocket shapes, then we analyzed the overlap of those shapes with the co-crystallized ligands, analyzed for a consistent location in different virtual protease structures, and translated into a particular surface in terms of amino acid positions (Figure 4). The pockets are shown below, and the intra-viral conservation is shown by alternative amino acid codes in Figure 4.

The AP1 pocket within NS3-pro (Figure 4) was the pocket we (R.A.) screened to identify an allosteric inhibitor [24]. AP1 is formed by 18 amino acids, combining hydrophobic and polar features. Nine of those amino acids are conserved between three viruses with X-ray structures, and the other nine amino acids differ in ZIKV and JEV, thus changing the shape of AP1. Hydrophobic residues, such as L76 (R in JEV), W83, and V147, create a patch that can be targeted to increase the binding affinity of a tentative inhibitor. The polar residues, including K74, and E86 in Dengue 2 are different in ZIKV and JEV. The AP2 pocket (Figure 4) overlaps with AP1 and is only visible for ZIKV co-crystallized with a binder (PDB: 7M1V). Eleven out of twelve amino acids forming the pocket are common with AP1, but the shape is somewhat different, which may be essential in a structure-based docking screen.

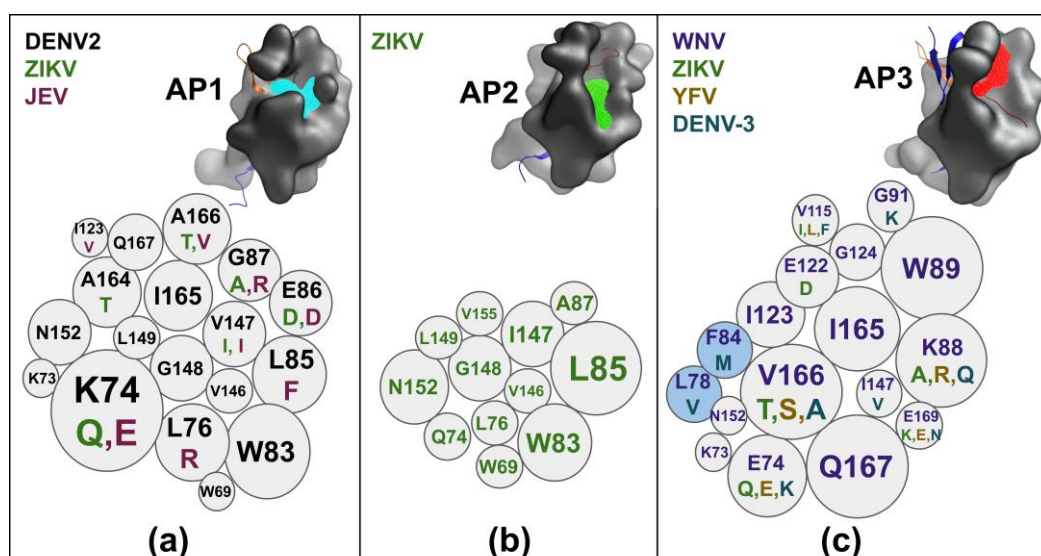


Figure 4. Allosteric Pockets (AP1, AP2, AP3) on flavivirus NS2B-NS3-pro proteins (gray surfaces) highlight conserved and variable residues. AP1 (a): Residues across DENV2 (black), ZIKV (green), and JEV (magenta) include highly variable sites like K74, E86, and L85. AP2 (b): Conserved residues in ZIKV (green) include L85, with surrounding residues contributing to structural integrity. AP3 (c): Variability across WNV (purple), ZIKV (green), YFV (dark yellow), and DENV3 (dark cyan) is shown, with key residues like W89, V166, and Q167. Bubble sizes reflect residue contribution to the pocket surface, with colors denoting specific viruses.

The AP3 pocket has not been characterized previously, but it is clearly visible in structure analysis and pocket prediction in at least four viruses, ZIKV, WNV, Dengue 3, YFV. The amino acid composition of AP3 is different and overlaps with AP1 for only seven amino acids out of 17. Several hydrophobic residues, including W89, I165, I123, are well conserved and can be exploited for pan-flaviviral inhibitors. A large-scale screen for AP3 binders is in progress.

3.3.2. Backbone conformational conservation of the pockets between flaviviruses

In addition to the sequence variation around the active site pocket, AP1, AP2, we further studied the conformational variation of the backbone surrounding the pockets to evaluate the prospect of identifying pan-flaviviral protease inhibitors.

For each pair of protease representatives of each virus and four Dengue protease states we calculated the root-mean-square-deviation (RMSD) measure of the pocket vicinity (Table 2). The NS2B-NS3 protease of DENV-2 exhibits the largest deviations among the proteases studied, potentially shedding light on the challenges of effectively targeting DENV-2 through binders.

Table 2. Root mean square deviation (RMSD, in Å units) calculations of the backbone atoms between residues forming the ASP in the closed/active state of ZIKV, WNV, YFV, DENV-2, DENV-3, DENV-4 flaviviruses, the AP1 in the transient state (WNV, DENV-1, DENV-2, DENV-4), and the AP2 in the fully opened state (ZIKV, JEV).

ASP, CLOSED STATE						
	ZIKV	WNV	YFV	DENV-2	DENV-3	DENV-4
DENV-4	0.9	0.7	0.7	3.7	0.5	0.0
DENV-3	0.8	0.6	0.7	3.97	0.0	
DENV-2	3.8	3.99	3.9	0.0		
YFV	0.5	0.6	0.0			
WNV	0.4	0.0				
ZIKV	0.0					
AP1, TRANSIENT STATE						
	WNV	DENV-1	DENV-2	DENV-4		
DENV-4	0.7	0.7	2.9	0.0		
DENV-2	2.97	2.9	0.0			
DENV-1	0.7	0.0				
WNV	0.0					
AP2, FULLY OPENED STATE						
	ZIKV	JEV				
JEV	0.7	0.0				
ZIKV	0.0					

In contrast, the pockets of ZIKV, WNV, and YFV proteases show smaller structural deviations. Some variations may be attributed to a construct and crystal conditions of a particular model, other variations suggest a separate structure-based screen for the deviated conformations of the pocket.

The predicted AP3 pocket overlaps with AP1 substantially with the exception of the distal end of the AP3 pocket proximal to the NS3-pro C-terminal hairpin. Therefore, the conformational conservation of AP1 and AP3 backbone is similar.

The X-ray structures of NS3-protease may also be analyzed for average B-factors of residues lining the AP1 and AP2 pockets. For each of the PDB entries studied, we derived an average B-factor and separately calculated the mean B-factor of residues covering the pockets. The ratios of the pocket residue B-factors to the protein-wide mean B-factor value indicated the relative mobility of the binding site residues in each pocket and in each virus. Let us describe a few structures chosen as best pocket representatives. The average *relative* B-factors of residues in the AP1 pocket were as low as 0.80 for DENV2 (6MO0, ligand-bound state) which means that the pocket is more structurally conserved compared to an average B-factor for all protein residues. The same relative AP1 3D-conservation number for DENV2 (2FOM, apo-state) was 1.28 when compared to the average protein atom B-factor. This number is higher than 1. or 0.9 because the 2FOM structure is in an unliganded apo-state. That results in less ordered residues in the pocket. The average *relative* B-factors of residues in the AP2 pocket were as low as 0.88 for ZIKV (7M1V, ligand-bound state) which means that the pocket is more structurally conserved compared to an average B-factor for all protein residues. Overall, the pocket residues in the apo-state are comparable, or more structurally conserved than the surface residues, and only slightly less ordered than the buried amino acids.

3.4. Sequential variations around pockets and the risk of drug resistance

Sequence conservation of amino acids for each of the five flaviviruses and additional Dengue subtypes can be derived from sequence data deposited to the NCBI Virus database. We were particularly interested in relative conservation of the amino acids proximal to the three main targetable pockets. This variation needs to be taken into consideration to avoid sequence dependent resistance to inhibitors. We analyzed the sequence conservation and diversity of the polyprotein precursor within and across five viruses and three additional Dengue subtypes (Figure 5) for the pocket vicinities, ASP, AP1 and AP2.

We retrieved a large number of amino acid sequences from the NCBI Virus database: 980 for ZIKV, 486 for WNV, 910 for YFV, 226 for JEV, 4628 for DENV-1, 3819 for DENV-2, 1780 for DENV-3, and 863 for DENV-4. The sequences were processed to extract the NS2B-NS3-pro fragments and further filtered to identify unique sequences. The next step was to extract amino acids in positions proximal to each of the three pockets studied.

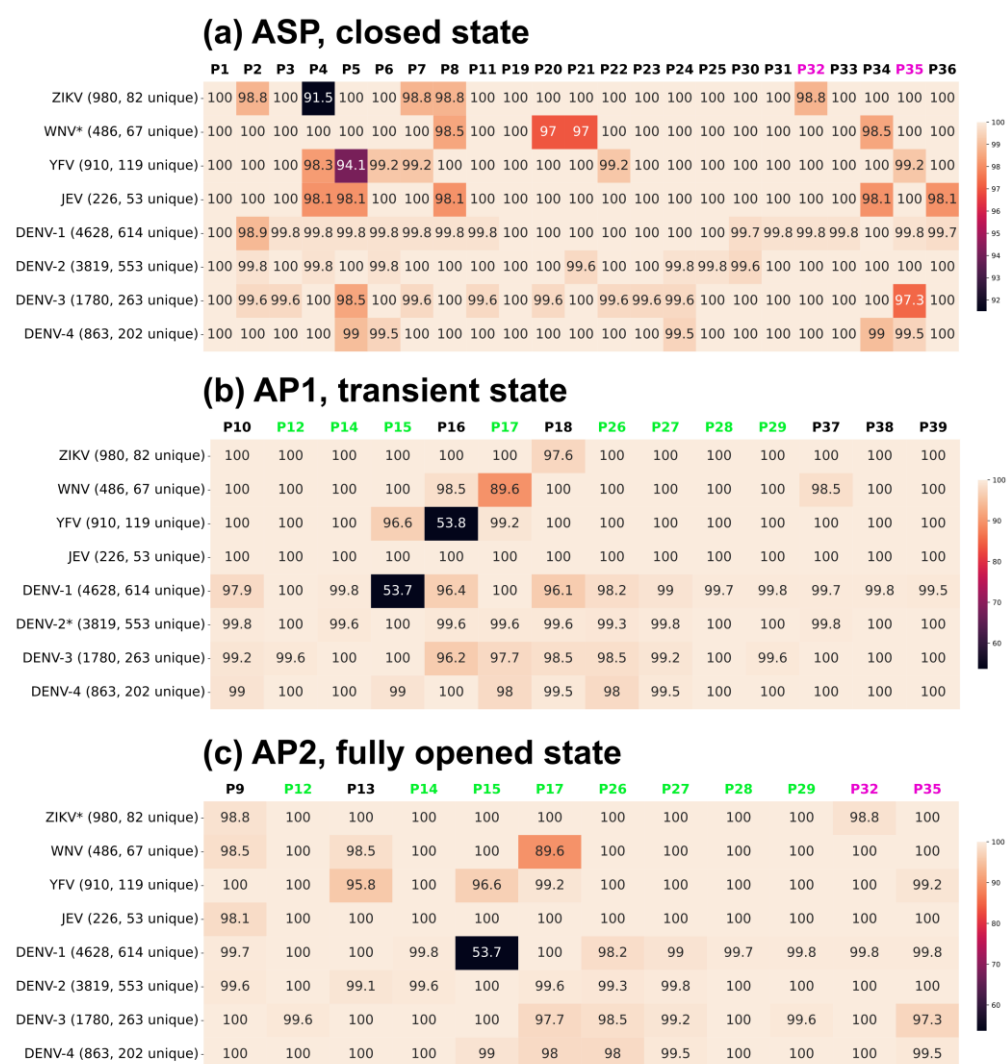


Figure 5. Sequence conservation (%) analysis of (a) Active site pocket (closed state), (b) Allosteric pocket 1 (transient), (c) Allosteric pocket 2 (fully opened state) within various flaviviruses. The letter "P" denotes the amino acid position within the pocket. The asterisk indicates the flaviviruses used to define the amino acids in the three pockets (closed – 5IDK (WNV), transient – 6MO2 (DENV-2), fully opened – 7M1V (ZIKV)).

For each of the conformational states, amino acids forming the pockets ASP, AP1, and AP2 were refined to subsets neighboring the co-crystallized ligands. A comparison of the amino acids forming ASP in the closed state across flaviviruses reveals that WNV's pocket is the most conserved, with variation observed at only four positions. In contrast, DENV-1 has the most variable pocket. Despite the significant conservation of ASP in ZIKV, the largest mutation rate is observed at position ASP-P4, making this site prone to mutations. Similarly, a low conservation percentage was found in YFV at position ASP-P5. These sites should be considered and, if necessary, avoided during inhibitor design.

In the transient state, the most conserved AP1 is observed in JEV, with 100% conservation for all residue positions, while the least conserved one is found in DENV-1 and -3. The conserved surface of the AP1 pocket in JEV makes it a good target.

AP2 was identified in the fully opened conformation. The most conserved AP2 was found for JEV, with mutability at only one position, while DENV-1-AP2 exhibits the most overall variability. In all three states, DENV-1-AP2 exhibited the highest variability, likely due to the availability of a larger number of amino acid sequences. In the transient state, the most variable sites include YFV at position 16, where D is present in 53.8% of sequences and E in 46.2%, and DENV-1 at position 15, with F in 53.7% and L in 46.3%.

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High variability is also observed at position P17 in WNV (89.6%) and at position P16 in YFV (53.8%). Those positions also need to be considered and avoided during drug development. Despite JEV and DENV-4 having the highest percentage of unique sequences, indicating their lower amino acid sequence conservation within the species, these viruses exhibit relative stability and sequence conservation in all three pockets, especially for JEV (Figure 5).

We also compared sequences and subsets around the pockets of the most frequent amino acids for each of the viruses to evaluate a potential for multi-viral or even pan-viral inhibitors. Figure 6 shows pocket surroundings for ZIKV, WNV, YFV, JEV, and DENV-1 through DENV-4. The number of positions with identical amino acids in different viruses is only 47.83% around ASP, 35.71% for AP1, and 41.67% for AP2 (Figure 6).

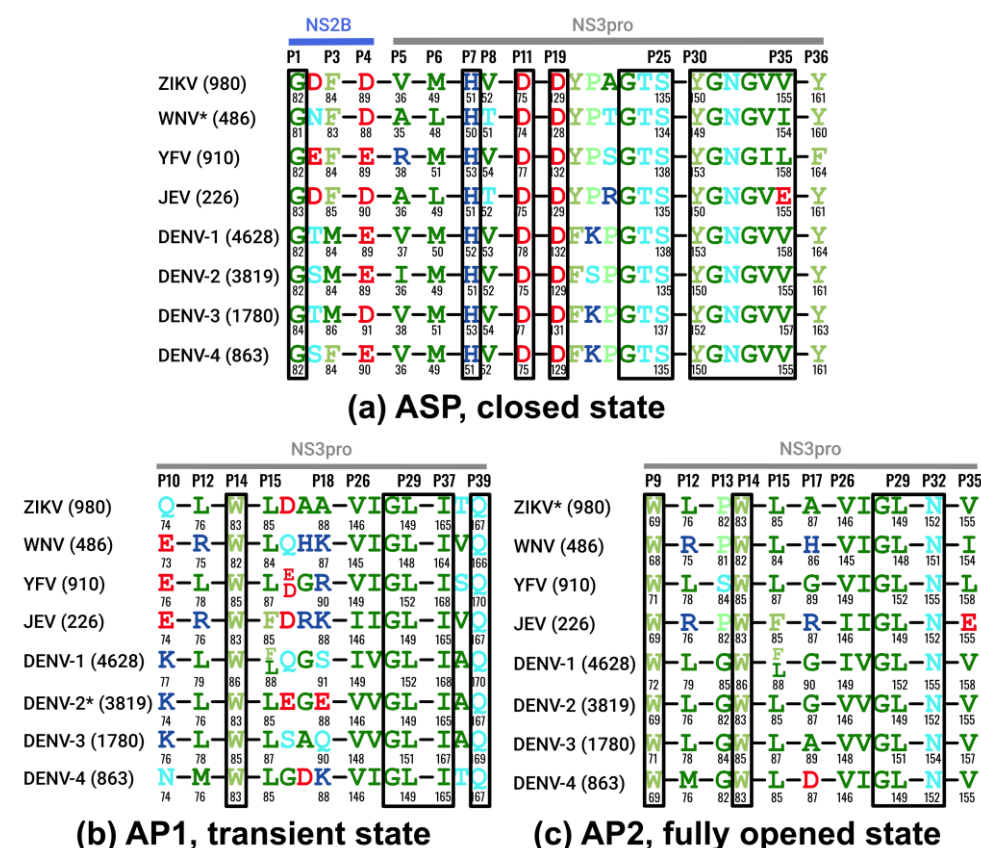


Figure 6. Amino acid sequence conservation analysis of (a) Active site pocket (Closed state), (b) Allosteric pocket 1 (Transient), (c) Allosteric pocket 2 (Fully opened state) among various flaviviruses. The letter "P" denotes the amino acid position within the pocket. The asterisk indicates the flaviviruses used to define the amino acids in the three pockets (Closed – 5IDK (WNV), Transient – 6MO2 (DENV-2), Fully opened – 7M1V (ZIKV)). To simplify the analysis, we focused on these specific flaviviruses as representatives for the determination of amino acid residues within the pockets. Amino acids identical among all flaviviruses are outlined in black rectangles. Each flavivirus has its own local numbering for NS2B and NS3.

Hence, developing drugs that target all flaviviruses simultaneously is a challenging task. For the ASP pocket it would mean avoiding positions P3 to P6, and P20 to P22. However, it is quite possible to develop a drug that targets one of the three pockets in more than one flavivirus. A drug could be developed to bind to the ASP in both ZIKV and YFV. It is also feasible to find a compound that would inhibit the closed state of five flaviviruses at once: WNV and the four DENV serotypes.

Targeting AP1 in the transient state with a drug effective across multiple flaviviruses presents a significant challenge. However, DENV-1 and DENV-3 would be good

candidates for a single AP1-binding inhibitor targeting two subtypes. Inhibiting allosteric pocket 2 in the fully opened state across multiple flaviviruses appear to be attractive for simultaneous inhibition of DENV-1, -2, and -3 because all positions around AP2 are identical or similar (G-P17-A) between the three Dengue subtypes. DENV-4 AP2 pocket residues include two additional mutations, with P17:G-to-D being a more significant change.

In summary, the AP1 is the most promising pocket for targeting, even though it may be difficult to find a pan-flaviviral NS3-pro inhibitor with same efficacy.

4. Discussion

The global impact of flaviviruses remains substantial, affecting hundreds of millions of people each year. This underscores the urgent need for effective antiviral therapies. Developing selective small-molecule compounds capable of targeting and inhibiting flaviviruses is a critical priority. Among potential therapeutic targets, the NS2B-NS3 protease complex stands out due to its pivotal role in viral replication, making it a highly promising focus for anti flaviviral drug development. Unfortunately, no approved anti flaviviral drugs are currently available, prompting ongoing research efforts to identify selective and potent inhibitors for NS2B-NS3-pro proteases of the flaviviruses [18,24,46,47,61].

Here we studied NS2B-NS3-pro protease-activity-specific allosteric binding sites in ZIKV, WNV, YFV, JEV, DENV-1, -2, -3, -4 proteases, in three distinct conformational states closed, transient, fully opened identified previously [25,44,53,56,62–64]. Our main objective was to identify and characterize the binding sites alternative to the pocket near the closed active site pocket. The main reason for that search was that the active site inhibitors identified so far were not sufficiently potent [41,42,65,66]. Among several obstacles, three negatively charged amino acids around ASP (Figure 6) create a total negative charge [41] at the pocket, and the pocket shape is relatively open. Nevertheless, research is still ongoing to search for covalent (S135) or non-covalent inhibitors [45–48].

On the other hand, the open state allosteric pockets AP1 [24] and AP2 pockets provide a better opportunity, as Figure 6 shows, their surface is not charged, and the shape is more enclosed. **We followed the definitions of the allosteric pocket as formulated in [67] in a sense that the allosteric pockets were sufficiently close to the active site and the substrate binding site, but distinct from it. In addition, we had preliminary evidence that some of the allosteric site ligands identified so far inhibited the protease activity by either stabilizing the inactive state or directly interfering with the substrate binding.** Some inhibitors against those open state pockets have been found: AP1 [24,26] and AP2 [51]. Both allosteric pockets are primarily hydrophobic, with AP2 being even more hydrophobic than AP1. For the AP1 in the Transient state, there are crystallographic structures with small molecule compounds [50] for DENV-2. However, not every structure of the transient state was suitable for structure-based docking screens due to the missing fragments around the AP2 pocket [68]. Overall, the AP1 pocket models in the transient state can be found in the PDB for three flaviviruses (DENV2, ZIKV, JEV) already and can be modelled for the remaining two viruses and subtypes. AP2 is a less investigated pocket [25] because it is visible only in one crystal structure in ZIKA in a fully open state.

Finally, the AP3 is the only pocket that is allosteric, yet it is present in both the closed active state, and in the transient state. The AP3 pocket is the least studied one yet is promising in terms of its composition and shape in comparison with the active site pocket.

The sequence conservation within each species (intraspecies analysis) revealed that two allosteric pockets of the inactive state for all viruses are well conserved for targeting (Figure 5), with JEV AP1 surface being the most conserved (100%) among the flaviviruses. The backbone conformation around the AP1 and AP2 pockets was also conserved sufficiently between different viral proteases, with one notable exception of the AP1 pocket of Dengue 2 structure (RMSD to other subtypes and viruses close to 3Å) in which a C-terminal fragment of NS3-pro moves upward. With this caveat, the AP1 pocket appears to be a prime target for structure-based screening for potential inhibitors, **against large compound databases or peptides and peptidomimetics.**

While targeting multiple flaviviruses simultaneously is a challenging task, it is not impossible if inhibitor interactions are focused on the conserved pocket patches only.

5. Conclusions

Targeting the NS2B-NS3 protease of flaviviruses remains a promising antiviral strategy due to its critical role in viral replication. However, targeting the active site of the closed state of the protease proved to be challenging. Our study explored allosteric binding sites, focusing on two distinct conformational states of the inactive protease—transient, and fully opened, and compared them with the closed (active) state. The two allosteric pockets in the inactive states emerged as "druggable" due to their hydrophobic nature and enclosed shape, with a transient state pocket showing notable sequence conservation across flaviviruses. Our analysis of both intra-viral and inter-viral variations of pocket sequences and 3D conformations revealed that virus-specific allosteric inhibitors can achieve sufficient efficacy by structure-guided improvements. Continued exploration of these sites and structure-guided screening of those pockets against small molecule libraries or peptides/peptidomimetics, holds promise for advancing anti-flaviviral drug discovery.

Author Contributions: S.G. and H.G. performed the data analysis; R.A. designed the study; S.G., H.G., and R.A. drafted the manuscript and reviewed the final version. All authors have read and agreed to the published version of the manuscript. All authors have read and agreed to the published version of the manuscript.

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References

- Pierson, T.C.; Diamond, M.S. The Continued Threat of Emerging Flaviviruses. *Nat Microbiol* **2020**, *5*, 796–812, doi:10.1038/s41564-020-0714-0.
- Qian, X.; Qi, Z. Mosquito-Borne Flaviviruses and Current Therapeutic Advances. *Viruses* **2022**, *14*, 1226, doi:10.3390/v14061226.
- CDC About Viral Hemorrhagic Fevers Available online: <https://www.cdc.gov/viral-hemorrhagic-fevers/about/index.html> (accessed on 17 November 2024).
- Wang, Q.-Y.; Shi, P.-Y. Flavivirus Entry Inhibitors. *ACS Infect. Dis.* **2015**, *1*, 428–434, doi:10.1021/acsinfecdis.5b00066.
- Slon Campos, J.L.; Mongkolsapaya, J.; Screaton, G.R. The Immune Response against Flaviviruses. *Nat Immunol* **2018**, *19*, 1189–1198, doi:10.1038/s41590-018-0210-3.
- Narayan, R.; Tripathi, S. Intrinsic ADE: The Dark Side of Antibody Dependent Enhancement During Dengue Infection. *Front. Cell. Infect. Microbiol.* **2020**, *10*, 580096, doi:10.3389/fcimb.2020.580096.

7. Pan, Y.; Cai, W.; Cheng, A.; Wang, M.; Yin, Z.; Jia, R. Flaviviruses: Innate Immunity, Inflammasome Activation, Inflammatory Cell Death, and Cytokines. *Front. Immunol.* **2022**, *13*, 829433, doi:10.3389/fimmu.2022.829433. 537
538
8. Brecher, M.; Zhang, J.; Li, H. The Flavivirus Protease as a Target for Drug Discovery. *Viol. Sin.* **2013**, *28*, 326–336, doi:10.1007/s12250-013-3390-x. 539
540
9. Li, Z.; Zhang, J.; Li, H. Flavivirus NS2B/NS3 Protease: Structure, Function, and Inhibition. In *Viral Proteases and Their Inhibitors*; Elsevier, 2017; pp. 163–188 ISBN 978-0-12-809712-0. 541
542
10. Chernov, A.V.; Shiryayev, S.A.; Aleshin, A.E.; Ratnikov, B.I.; Smith, J.W.; Liddington, R.C.; Strongin, A.Y. The Two-Component NS2B-NS3 Proteinase Represses DNA Unwinding Activity of the West Nile Virus NS3 Helicase. *Journal of Biological Chemistry* **2008**, *283*, 17270–17278, doi:10.1074/jbc.M801719200. 543
544
545
11. Shi, Y.; Gao, G.F. Structural Biology of the Zika Virus. *Trends in Biochemical Sciences* **2017**, *42*, 443–456, doi:10.1016/j.tibs.2017.02.009. 546
547
12. Patkar, C.G.; Jones, C.T.; Chang, Y.; Warriar, R.; Kuhn, R.J. Functional Requirements of the Yellow Fever Virus Capsid Protein. *J Virol* **2007**, *81*, 6471–6481, doi:10.1128/JVI.02120-06. 548
549
13. Dethoff, E.A.; Boerneke, M.A.; Gokhale, N.S.; Muhire, B.M.; Martin, D.P.; Sacco, M.T.; McFadden, M.J.; Weinstein, J.B.; Messer, W.B.; Horner, S.M.; et al. Pervasive Tertiary Structure in the Dengue Virus RNA Genome. *Proc. Natl. Acad. Sci. U.S.A.* **2018**, *115*, 11513–11518, doi:10.1073/pnas.1716689115. 550
551
552
14. Murugesan, A.; Manoharan, M. Dengue Virus. In *Emerging and Reemerging Viral Pathogens*; Elsevier, 2020; pp. 281–359 ISBN 978-0-12-819400-3. 553
554
15. Solomon, T. Recent Advances in Japanese Encephalitis. *J Neurovirol* **2003**, *9*, 274–283, doi:10.1080/13550280390194037. 555
16. Samrat, S.K.; Xu, J.; Li, Z.; Zhou, J.; Li, H. Antiviral Agents against Flavivirus Protease: Prospect and Future Direction. *Pathogens* **2022**, *11*, 293, doi:10.3390/pathogens11030293. 556
557
17. Wahaab, A.; Mustafa, B.E.; Hameed, M.; Stevenson, N.J.; Anwar, M.N.; Liu, K.; Wei, J.; Qiu, Y.; Ma, Z. Potential Role of Flavivirus NS2B-NS3 Proteases in Viral Pathogenesis and Anti-Flavivirus Drug Discovery Employing Animal Cells and Models: A Review. *Viruses* **2021**, *14*, 44, doi:10.3390/v14010044. 558
559
560
18. Voss, S.; Nitsche, C. Inhibitors of the Zika Virus Protease NS2B-NS3. *Bioorganic & Medicinal Chemistry Letters* **2020**, *30*, 126965, doi:10.1016/j.bmcl.2020.126965. 561
562
19. Aleshin, A.E.; Shiryayev, S.A.; Strongin, A.Y.; Liddington, R.C. Structural Evidence for Regulation and Specificity of Flaviviral Proteases and Evolution of the *Flaviviridae* Fold. *Protein Science* **2007**, *16*, 795–806, doi:10.1110/ps.072753207. 563
564
565
20. Pastorino, B.; Nougairède, A.; Wurtz, N.; Gould, E.; De Lamballerie, X. Role of Host Cell Factors in Flavivirus Infection: Implications for Pathogenesis and Development of Antiviral Drugs. *Antiviral Research* **2010**, *87*, 281–294, doi:10.1016/j.antiviral.2010.04.014. 566
567
568
21. Fishburn, A.T.; Pham, O.H.; Kenaston, M.W.; Beesabathuni, N.S.; Shah, P.S. Let's Get Physical: Flavivirus-Host Protein-Protein Interactions in Replication and Pathogenesis. *Front Microbiol* **2022**, *13*, 847588, doi:10.3389/fmicb.2022.847588. 569
570
571
22. Starvaggi, J.; Previti, S.; Zappalà, M.; Ettari, R. The Inhibition of NS2B/NS3 Protease: A New Therapeutic Opportunity to Treat Dengue and Zika Virus Infection. *IJMS* **2024**, *25*, 4376, doi:10.3390/ijms25084376. 572
573
23. Da Silva-Júnior, E.F.; De Araújo-Júnior, J.X. Peptide Derivatives as Inhibitors of NS2B-NS3 Protease from Dengue, West Nile, and Zika Flaviviruses. *Bioorganic & Medicinal Chemistry* **2019**, *27*, 3963–3978, doi:10.1016/j.bmc.2019.07.038. 574
575
576

24. Meewan, I.; Shiryaev, S.A.; Kattoula, J.; Huang, C.-T.; Lin, V.; Chuang, C.-H.; Terskikh, A.V.; Abagyan, R. Allosteric Inhibitors of Zika Virus NS2B-NS3 Protease Targeting Protease in “Super-Open” Conformation. *Viruses* **2023**, *15*, 1106, doi:10.3390/v15051106. 577-579
25. Shiryaev, S.A.; Cieplak, P.; Cheltsov, A.; Liddington, R.C.; Terskikh, A.V. Dual Function of Zika Virus NS2B-NS3 Protease. *PLoS Pathog* **2023**, *19*, e1011795, doi:10.1371/journal.ppat.1011795. 580-581
26. Knyazhanskaya, E.; Morais, M.C.; Choi, K.H. Flavivirus Enzymes and Their Inhibitors. In *The Enzymes*; Elsevier, 2021; Vol. 49, pp. 265–303 ISBN 978-0-12-823468-6. 582-583
27. João, E.E.; Lopes, J.R.; Guedes, B.F.R.; da Silva Sanches, P.R.; Chin, C.M.; Dos Santos, J.L.; Scarim, C.B. Advances in Drug Discovery of Flavivirus NS2B-NS3pro Serine Protease Inhibitors for the Treatment of Dengue, Zika, and West Nile Viruses. *Bioorg Chem* **2024**, *153*, 107914, doi:10.1016/j.bioorg.2024.107914. 584-586
28. Braun, N.J.; Quek, J.P.; Huber, S.; Kouretova, J.; Rogge, D.; Lang-Henkel, H.; Cheong, E.Z.K.; Chew, B.L.A.; Heine, A.; Luo, D.; et al. Structure-Based Macrocyclization of Substrate Analogue NS2B-NS3 Protease Inhibitors of Zika, West Nile and Dengue Viruses. *ChemMedChem* **2020**, *15*, 1439–1452, doi:10.1002/cmdc.202000237. 587-589
29. Huber, S.; Braun, N.J.; Schmacke, L.C.; Murra, R.; Bender, D.; Hildt, E.; Heine, A.; Steinmetzer, T. Synthesis and Structural Characterization of New Macrocyclic Inhibitors of the Zika Virus NS2B–NS3 Protease. *Archiv der Pharmazie* **2024**, *357*, 2400250, doi:10.1002/ardp.202400250. 590-592
30. Burley, S.K.; Berman, H.M.; Kleywegt, G.J.; Markley, J.L.; Nakamura, H.; Velankar, S. Protein Data Bank (PDB): The Single Global Macromolecular Structure Archive. *Methods Mol Biol* **2017**, *1607*, 627–641, doi:10.1007/978-1-4939-7000-1_26. 593-595
31. Weinert, T.; Olieric, V.; Waltersperger, S.; Panepucci, E.; Chen, L.; Zhang, H.; Zhou, D.; Rose, J.; Ebihara, A.; Kuramitsu, S.; et al. Fast Native-SAD Phasing for Routine Macromolecular Structure Determination. *Nat Methods* **2015**, *12*, 131–133, doi:10.1038/nmeth.3211. 596-598
32. Chandramouli, S.; Joseph, J.S.; Daudenarde, S.; Gatchalian, J.; Cornillez-Ty, C.; Kuhn, P. Serotype-Specific Structural Differences in the Protease-Cofactor Complexes of the Dengue Virus Family. *J Virol* **2010**, *84*, 3059–3067, doi:10.1128/JVI.02044-09. 599-601
33. Noble, C.G.; Seh, C.C.; Chao, A.T.; Shi, P.Y. Ligand-Bound Structures of the Dengue Virus Protease Reveal the Active Conformation. *J Virol* **2012**, *86*, 438–446, doi:10.1128/JVI.06225-11. 602-603
34. Brister, J.R.; Ako-adjei, D.; Bao, Y.; Blinkova, O. NCBI Viral Genomes Resource. *Nucleic Acids Research* **2015**, *43*, D571–D577, doi:10.1093/nar/gku1207. 604-605
35. Abagyan, R.; Totrov, M.; Kuznetsov, D. ICM?A New Method for Protein Modeling and Design: Applications to Docking and Structure Prediction from the Distorted Native Conformation. *J. Comput. Chem.* **1994**, *15*, 488–506, doi:10.1002/jcc.540150503. 606-608
36. Finn, R.D.; Attwood, T.K.; Babbitt, P.C.; Bateman, A.; Bork, P.; Bridge, A.J.; Chang, H.-Y.; Dosztányi, Z.; El-Gebali, S.; Fraser, M.; et al. InterPro in 2017—beyond Protein Family and Domain Annotations. *Nucleic Acids Res* **2017**, *45*, D190–D199, doi:10.1093/nar/gkw1107. 609-611
37. Neves, M.A.C.; Totrov, M.; Abagyan, R. Docking and Scoring with ICM: The Benchmarking Results and Strategies for Improvement. *J Comput Aided Mol Des* **2012**, *26*, 675–686, doi:10.1007/s10822-012-9547-0. 612-613
38. Molsoft L.L.C. Available online: <https://www.molsoft.com/> (accessed on 17 November 2024). 614
39. Droettboom, M.; Caswell, T.A.; Hunter, J.; Firing, E.; Nielsen, J.H.; Varoquaux, N.; Root, B.; Elson, P.; Dale, D.; Jae-Joon Lee; et al. Matplotlib/Matplotlib V2.0.1. **2017**, doi:10.5281/zenodo.570311. 615-616
40. The UniProt Consortium UniProt: A Worldwide Hub of Protein Knowledge. *Nucleic Acids Research* **2019**, *47*, D506–D515, doi:10.1093/nar/gky1049. 617-618

41. Li, Y.; Zhang, Z.; Phoo, W.W.; Loh, Y.R.; Li, R.; Yang, H.Y.; Jansson, A.E.; Hill, J.; Keller, T.H.; Nacro, K.; et al. Structural Insights into the Inhibition of Zika Virus NS2B-NS3 Protease by a Small-Molecule Inhibitor. *Structure* **2018**, *26*, 555–564.e3, doi:10.1016/j.str.2018.02.005. 619–621
42. Zhang, Z.; Li, Y.; Loh, Y.R.; Phoo, W.W.; Hung, A.W.; Kang, C.; Luo, D. Crystal Structure of Unlinked NS2B-NS3 Protease from Zika Virus. *Science* **2016**, *354*, 1597–1600, doi:10.1126/science.aai9309. 622–623
43. Peng, M.; Li, C.-Y.; Chen, X.-L.; Williams, B.T.; Li, K.; Gao, Y.-N.; Wang, P.; Wang, N.; Gao, C.; Zhang, S.; et al. Insights into Methionine S-Methylation in Diverse Organisms. *Nat Commun* **2022**, *13*, 2947, doi:10.1038/s41467-022-30491-5. 624–626
44. Quek, J.P.; Liu, S.; Zhang, Z.; Li, Y.; Ng, E.Y.; Loh, Y.R.; Hung, A.W.; Luo, D.; Kang, C. Identification and Structural Characterization of Small Molecule Fragments Targeting Zika Virus NS2B-NS3 Protease. *Antiviral Research* **2020**, *175*, 104707, doi:10.1016/j.antiviral.2020.104707. 627–629
45. Lin, X.; Cheng, J.; Wu, Y.; Zhang, Y.; Jiang, H.; Wang, J.; Wang, X.; Cheng, M. Identification and In Silico Binding Study of a Highly Potent DENV NS2B-NS3 Covalent Inhibitor. *ACS Med. Chem. Lett.* **2022**, *13*, 599–607, doi:10.1021/acsmchemlett.1c00653. 630–632
46. Shin, H.J.; Kim, M.-H.; Lee, J.-Y.; Hwang, I.; Yoon, G.Y.; Kim, H.S.; Kwon, Y.-C.; Ahn, D.-G.; Kim, K.-D.; Kim, B.-T.; et al. Structure-Based Virtual Screening: Identification of a Novel NS2B-NS3 Protease Inhibitor with Potent Antiviral Activity against Zika and Dengue Viruses. *Microorganisms* **2021**, *9*, 545, doi:10.3390/microorganisms9030545. 633–635
47. Norshidah, H.; Leow, C.H.; Ezleen, K.E.; Wahab, H.A.; Vignesh, R.; Rasul, A.; Lai, N.S. Assessing the Potential of NS2B/NS3 Protease Inhibitors Biomarker in Curbing Dengue Virus Infections: In Silico vs. In Vitro Approach. *Front. Cell. Infect. Microbiol.* **2023**, *13*, 1061937, doi:10.3389/fcimb.2023.1061937. 636–638
48. Tang, H.; Liu, Y.; Ren, R.; Liu, Y.; He, Y.; Qi, Z.; Peng, H.; Zhao, P. Identification of Clinical Candidates against West Nile Virus by Activity Screening in Vitro and Effect Evaluation in Vivo. *Journal of Medical Virology* **2022**, *94*, 4918–4925, doi:10.1002/jmv.27891. 639–641
49. Shiryaev, S.A.; Cieplak, P.; Cheltsov, A.; Liddington, R.C.; Terskikh, A.V. DUAL FUNCTION OF ZIKA VIRUS NS2B-NS3 PROTEASE 2021. 642–643
50. Yao, Y.; Huo, T.; Lin, Y.-L.; Nie, S.; Wu, F.; Hua, Y.; Wu, J.; Kneubehl, A.R.; Vogt, M.B.; Rico-Hesse, R.; et al. Discovery, X-Ray Crystallography and Antiviral Activity of Allosteric Inhibitors of Flavivirus NS2B-NS3 Protease. *J. Am. Chem. Soc.* **2019**, *141*, 6832–6836, doi:10.1021/jacs.9b02505. 644–646
51. Bank, R.P.D. RCSB PDB - 7M1V: Structure of Zika Virus NS2b-NS3 Protease Mutant Binding the Compound NSC86314 in the Super-Open Conformation Available online: <https://www.rcsb.org/structure/7M1V> (accessed on 17 November 2024). 647–649
52. Erbel, P.; Schiering, N.; D'Arcy, A.; Renatus, M.; Kroemer, M.; Lim, S.P.; Yin, Z.; Keller, T.H.; Vasudevan, S.G.; Hommel, U. Structural Basis for the Activation of Flaviviral NS3 Proteases from Dengue and West Nile Virus. *Nat Struct Mol Biol* **2006**, *13*, 372–373, doi:10.1038/nsmb1073. 650–652
53. Noske, G.D.; Gawriljuk, V.O.; Fernandes, R.S.; Furtado, N.D.; Bonaldo, M.C.; Oliva, G.; Godoy, A.S. Structural Characterization and Polymorphism Analysis of the NS2B-NS3 Protease from the 2017 Brazilian Circulating Strain of Yellow Fever Virus. *Biochimica et Biophysica Acta (BBA) - General Subjects* **2020**, *1864*, 129521, doi:10.1016/j.bbagen.2020.129521. 653–656
54. Bank, R.P.D. RCSB PDB - 2M9P: NMR Structure of an Inhibitor Bound Dengue NS3 Protease Available online: <https://www.rcsb.org/structure/2M9P> (accessed on 17 November 2024). 657–658

55. Bank, R.P.D. RCSB PDB - 5YW1: Crystal Structure of Full Length NS3 Protein (eD4NS2BNS3) in Complex with Bovine Pancreatic Trypsin Inhibitor Available online: <https://www.rcsb.org/structure/5YW1> (accessed on 24 November 2024).
56. Luo, D.; Xu, T.; Hunke, C.; Grüber, G.; Vasudevan, S.G.; Lescar, J. Crystal Structure of the NS3 Protease-Helicase from Dengue Virus. *J Virol* **2008**, *82*, 173–183, doi:10.1128/JVI.01788-07.
57. Luo, D.; Wei, N.; Doan, D.N.; Paradkar, P.N.; Chong, Y.; Davidson, A.D.; Kotaka, M.; Lescar, J.; Vasudevan, S.G. Flexibility between the Protease and Helicase Domains of the Dengue Virus NS3 Protein Conferred by the Linker Region and Its Functional Implications. *Journal of Biological Chemistry* **2010**, *285*, 18817–18827, doi:10.1074/jbc.M109.090936.
58. Bank, R.P.D. RCSB PDB - 5YVJ: Crystal Structure of Full Length NS2B47-NS3 (gD4NS2BNS3) from Dengue Virus 4 in Open Conformation Available online: <https://www.rcsb.org/structure/5YVJ> (accessed on 17 December 2024).
59. An, J.; Totrov, M.; Abagyan, R. Pocketome via Comprehensive Identification and Classification of Ligand Binding Envelopes. *Mol Cell Proteomics* **2005**, *4*, 752–761, doi:10.1074/mcp.M400159-MCP200.
60. Sheridan, R.P.; Maiorov, V.N.; Holloway, M.K.; Cornell, W.D.; Gao, Y.-D. Drug-like Density: A Method of Quantifying the “Bindability” of a Protein Target Based on a Very Large Set of Pockets and Drug-like Ligands from the Protein Data Bank. *J. Chem. Inf. Model.* **2010**, *50*, 2029–2040, doi:10.1021/ci100312t.
61. Nunes, D.A.D.F.; Santos, F.R.D.S.; Da Fonseca, S.T.D.; De Lima, W.G.; Nizer, W.S.D.C.; Ferreira, J.M.S.; De Magalhães, J.C. NS2B-NS3 Protease Inhibitors as Promising Compounds in the Development of Antivirals against Zika Virus: A Systematic Review. *Journal of Medical Virology* **2022**, *94*, 442–453, doi:10.1002/jmv.27386.
62. Lei, J.; Hansen, G.; Nitsche, C.; Klein, C.D.; Zhang, L.; Hilgenfeld, R. Crystal Structure of Zika Virus NS2B-NS3 Protease in Complex with a Boronate Inhibitor. *Science* **2016**, *353*, 503–505, doi:10.1126/science.aag2419.
63. Su, X.-C.; Ozawa, K.; Qi, R.; Vasudevan, S.G.; Lim, S.P.; Otting, G. NMR Analysis of the Dynamic Exchange of the NS2B Cofactor between Open and Closed Conformations of the West Nile Virus NS2B-NS3 Protease. *PLoS Negl Trop Dis* **2009**, *3*, e561, doi:10.1371/journal.pntd.0000561.
64. Li, Q.; Kang, C. Insights into Structures and Dynamics of Flavivirus Proteases from NMR Studies. *IJMS* **2020**, *21*, 2527, doi:10.3390/ijms21072527.
65. Quek, J.-P.; Ser, Z.; Chew, B.L.A.; Li, X.; Wang, L.; Sobota, R.M.; Luo, D.; Phoo, W.W. Dynamic Interactions of Post Cleaved NS2B Cofactor and NS3 Protease Identified by Integrative Structural Approaches. *Viruses* **2022**, *14*, 1440, doi:10.3390/v14071440.
66. Xiong, Y.; Cheng, F.; Zhang, J.; Su, H.; Hu, H.; Zou, Y.; Li, M.; Xu, Y. Structure-Based Design of a Novel Inhibitor of the ZIKA Virus NS2B/NS3 Protease. *Bioorganic Chemistry* **2022**, *128*, 106109, doi:10.1016/j.bioorg.2022.106109.
67. Stank, A.; Kokh, D.B.; Fuller, J.C.; Wade, R.C. Protein Binding Pocket Dynamics. *Acc Chem Res* **2016**, *49*, 809–815, doi:10.1021/acs.accounts.5b00516.
68. Behnam, M.A.M.; Klein, C.D.P. Conformational Selection in the Flaviviral NS2B-NS3 Protease. *Biochimie* **2020**, *174*, 117–125, doi:10.1016/j.biochi.2020.04.014.

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