Anti-dengue drug: viral polyprotein, a potential target

Dear editor

We would like to add our views regarding the article “Identification of covalent active site inhibitors of dengue virus protease” by Koh-Stenta et al.¹ The article suggests the development of a possible drug to combat dengue virus. The drug will inhibit the ability of the virus to replicate by inhibiting the protease enzyme of the virus.

The genome of dengue virus is a single-stranded, positive-sense RNA that encodes for a polyprotein. The polyprotein is translated into three structural components, namely capsid, envelope, and membrane, and seven nonstructural proteins, namely NS1, NS2A, NS2B, NS3, NS4A, NS4B, and NS5. NS3 is a serine protease that has a critical role alongside host cell proteases in the protein breakdown of viral polyprotein that leads to replication.² The foundation of Koh-Stenta et al’s study was fundamentally based on the similarity of the amino acid sequence of proteases, especially at the active sites, of the dengue virus and the West Nile virus.³ As extensive knowledge was available on the West Nile virus and peptide-bound X-ray crystal structure confirmed many protein–ligand interactions similar to the two viruses,⁴ the knowledge could be applied on dengue virus.

The results of Koh-Stenta et al’s research are highly encouraging and show a possibility for the development of an anti-dengue drug in the near future. Another recent study by Dutch investigators suggested that a protein manufactured by the dengue virus NS4B would make a plausible target for antiviral drug.⁵ Their data suggested that a metabolite of acetaminophen (a common pain killer) AM404 was able to inhibit dengue virus replication. In our opinion, this is a ground-breaking discovery, and the study not only brings us one step closer to the development of an antiviral drug against dengue virus, but also enables the determination of when the replication of the virus is inhibited. The latter was achieved by using a derivative of dengue virus that expressed luciferase, a molecule that produces bioluminescence, during replication.

Hence, by correlating the two studies, we conclude that two proteins, ie, NS4B and NS2B, are potential targets for the development of an anti-dengue drug. Moreover, other areas should be explored, of the viral genome, to target other potential sites for drug development. It could be possible to manufacture drugs that stop the virus from entering the cell (entry inhibitors) or drugs that inhibit the strand of RNA (5‘ capping) inhibiting viral replication.⁶

Disclosure

The authors report no conflicts of interest in this communication.
References