Phylogenetically based establishment of a dengue virus panel, representing all available genotypes, as a tool in dengue drug discovery

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Abstract

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Dengue fever is the most widespread of the human arbovirus diseases, with approximately one third of the world's population at risk of infection. Dengue viruses are members of the genus Flavivirus (family Flaviviridae) and, antigenically, they separate as four closely related serotypes (1-4) that share 60 to 75 % amino acid homology. This genetic diversity complicates the process of antiviral drug discovery. Thus, currently no approved denguespecific therapeutic treatments are available. With the aim of providing an efficient tool for dengue virus drug discovery, a collection of nineteen dengue viruses, representing the genotypic diversity within the four serotypes, was developed. After phylogenetic analysis of the full-length genomes, we selected relevant strains from the EVAg collection at Aix-Marseille University and completed the virus collection, using a reverse genetic system based on the infectious sub-genomic amplicons technique. Finally, we evaluated this dengue virus collection against three published dengue inhibitory compounds. NITD008, which targets the highly conserved active site of the viral NS5 polymerase enzyme, exhibited similar antiviral potencies against each of the different dengue genotypes in the panel. Compounds targeting less conserved protein subdomains, such as the capsid inhibitor ST-148, or SDM25N, a ∂ opioid receptor antagonist which indirectly targets NS4B, exhibited larger differences in potency against the various genotypes of dengue viruses. These results illustrate the importance of a phylogenetically based dengue virus reference panel for dengue antiviral research. The collection developed in this study, which includes such representative dengue viruses, has been made available to the scientific community through the European Virus Archive to evaluate novel DENV antiviral candidates.

Introduction

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35 Dengue virus (DENV) is a major threat to human health, with approximately one third of the world's population at risk of being infected. DENV is the causative agent of dengue fever, as 36 well as the more severe dengue haemorrhagic fever (DHF)¹ and dengue shock syndrome 37 (DSS). It belongs to the genus Flavivirus (Flaviviridae family), which comprises other 38 clinically important human pathogens, such as yellow fever virus, West Nile virus and the 39 40 recently emerging Zika virus². DENV is an arthropod borne virus transmitted through the bite of infected mosquitoes from the genus Aedes (Stegomyia). Epidemiological transmission of 41 42 DENV is confined to urban and peri-urban cycles for which Aedes aegypti and Ae albopictus mosquitoes, respectively, are the primary transmission vectors³. Dengue is a positive-sense 43 44 single stranded RNA virus with a 10.7 kb genome encoding a single polyprotein which is post-translationally processed into three structural proteins, viz., capsid (C), pre-membrane 45 (prM), envelope (E) and seven non-structural proteins (NS1, NS2A, NS2B, NS3, NS4A, 46 NS4B and NS5)⁴. Four antigenically closely related serotypes of DENV (1-4) which share 60 47 to 75 % amino acid homology, have been identified⁵. Within this serotype demarcation, the 48 DENV are also grouped into genotypes, with varying terminology between authors^{6,7} 49 (hereunder we refer to the grouping proposed by Weaver and Vasilakis⁷). 50

- 51 Hence, many of the DENV diagnostic tools do not readily distinguish between DENV serotypes. Moreover, co-circulation of different serotypes during DENV epidemics⁸ increases 52 53 the complexity of virus identification. Added to these factors, antibody dependent 54 enhancement of the disease i.e., when patients contract a heterotypic secondary DENV is a potential additional complication for effective treatment of patients. 55 56 Consequently, scientists are faced with the challenge of developing Directly Active Antivirals 57 (DAA) that can inhibit the entire spectrum of genetically diverse serotypes and/or genotypes of DENV. However, despite the tireless efforts to provide an antiviral therapy 10-13, there are 58 59 still no approved drugs on the market to treat dengue infections. At present, the treatments available are merely supportive¹⁴. 60
- 61 A major barrier to evaluating the activity spectrum of potential DENV-inhibitory molecules 62 arises from the non-availability of a well-defined panel of viruses that specifically represents 63 the genetic variability of all characterised DENV isolates. With the aim of providing a tool for 64 DENV research, with which to assess the antiviral activity of potential inhibitory molecules, 65 we have developed a collection of DENV with sequences that include representative genotypes from within the four DENV serotypes (figure 1). Wherever possible, we selected 66 clinical strains with a limited number of passages in cell culture. Strains were selected from 67 either the European Virus Archive (EVA) collection¹⁵, the French National Reference Centre 68 for arboviruses (CNR), or the World Reference Center for Emerging Viruses and Arboviruses 69 70 (WRCEVA). Viruses that could not be obtained but for which full length genome sequences 71 were available, were re-created using the versatile infectious sub-genomic amplicons (ISA)
- 72 reverse genetics technology^{16,17}.



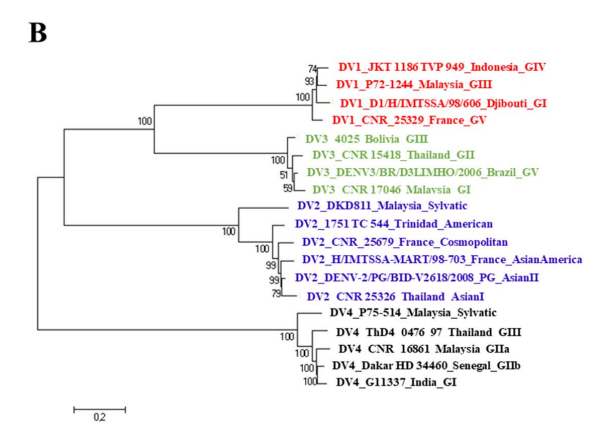


Figure 1: Global serotype-representative DENV collection A: Pipeline of the workflow employed for the virus collection. B: Maximum likelihood phylogenetic tree (GTR+G+I model with 500 bootstraps), based on the complete nucleic acid sequences of the virus collection. Strain information: DENV-1: Genotype I: (Djibouti, 1998, AF298808); Genotype III (Malaysia,1972, EF457905.1); Genotype IV (Indonesia, 1977, EUO74031); Genotype V (France, 2014, MF004384); DENV-2: Genotype Asian-America (France Martinique, 1998, AF208496); Genotype American (Trinidad, 1953, EU073981.1); Genotype Cosmopolitan (France, 2014, MF004385); Genotype Asian 1 (Thailand, 2014, MH888331); Genotype Asian 2 (Papua New Guinea, 2008, FJ906959.1); Genotype Sylvatic (Malaysia, 2008, FJ467493.1); DENV-3: Genotype I (Malaysia, 2012, MF004386); Genotype II (Thailand, 2012 MH888332); Genotype III (Bolivia,2011, MH888333); Genotype V (Brazil,2006, JN697379.1); DENV-4: Genotype IIb (Senegal, 1981, MF004387,); Genotype IIa (Malaysia, 2013, MH888334); Genotype III (Thailand, 1997, AY618988.1); Genotype Sylvatic (Malaysia, 1975, JF262779.1); Genotype I (INDIA, 1961, JF262783.1). Complete information relevant to the strains of the collection are more fully detailed in the supplemental material.

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In order to select representative genotypes, we collected dengue full-length genome sequences from the NCBI database and complemented this database with those of our, still unpublished, "in house" and CNR strains. We performed phylogenetic reconstructions with the maximum likelihood method to assign all available genome sequences to a genotype in a serotype (supplementary material Fig 1, 2, 3 and 4). Within each genotype, we focused on strains that were not subjected to extensive cell passage and were either available as biological isolates in virus collections or as full-length sequences in GenBank. Six dengue genotypes were available only as complete genome sequences in the NCBI database without any biological strain counterparts in referenced collections (DENV-1 genotype III, DENV-2 genotype sylvatic and Asian II, DENV-3 genotype V and DENV-4 genotype III and sylvatic). Two genotypes were not available at all because of incomplete genome sequence (DENV-1 genotype II and DENV-3 genotype IV). To obtain the biological viruses from the completely sequenced strains, we designed reverse genetics systems based on the ISA technique^{16–18} and generated synthetic overlapping DNA fragments that covered each of the entire genome, bordered by a CMV promoter on the 5' end and a Ribozyme and poly-adenylation signal on the 3' end. The overlapping fragments were co-transfected into a mix of human and hamster embryonic kidney cell lines (HEK 293 and BHK-21 purchased from the American Cell Culture Collection). This enabled us to recover the missing biological strains to complete the collection. The initial viral stocks were amplified in Vero E6 cells and fully sequenced. All the DENV strains used, have been made available through the EVAg collection (https://www.european-virus-archive.com/).

109 Various specific dengue inhibitors that target several viral proteins involved in different replication steps, have been discovered. ST-148, an inhibitor targeting the capsid structural 110 111 protein, has been reported to inhibit all DENV serotypes in cell culture, although with varying efficiency. This inhibitor also appears promising in the AG-129 mouse model when infected 112 with a strain of DENV-2¹⁹. NITD008, an adenosine analogue inhibitor that targets the RNA-113 114 dependent RNA polymerase activity, was shown to be inhibitory against all dengue serotypes as well as other flaviviruses, including West Nile virus, yellow fever virus and tick-borne 115 Powassan virus¹³. SDM25N, a ∂ opioid receptor antagonist, has been reported to target the 116 NS4B protein, probably indirectly through a cellular factor. Thus far, it has only been shown 117 to be active against a DENV-2 strain¹². Based on the different mechanisms of action of the 3 118 119 compounds, their respective target and its associated sequence variability across the different 120 genotypes, we hypothesize that the antiviral activity of the compounds might differ between 121 all of the genotypes of DENV. Therefore, the antiviral activity of these three compounds was 122 assessed using a single common protocol based on a viral RNA yield reduction assay²⁰. The 123 assay did not depend on the cytopathogenic potential of the strain, thus allowing for the 124 inclusion of any dengue strain in the panel tested. The compounds were assayed from 10 to 125 0.004µM, with 3-fold step-dilution in triplicate. The amount of viral RNA in the supernatant medium, sampled at intervals during the growth cycle, was quantified by qRT-PCR to 126 127 determine the 50% maximal effective concentration (EC₅₀) (Table 1).

Table 1 Dengue virus collection-susceptibility to three antiviral compounds assessed by yield reduction assay. Anti-capsid ST-148, nucleoside analogue NITD008 and ∂ opioid receptor antagonist

SDM25N were independently tested twice, with 3 replicates per experiment, against the dengue collection from $10\mu M$ to $0.005\mu M$. AA: Asian American, A: American, C: cosmopolitan.

			ST-148	NITD008	SDM25N
	Virus	Genotype	EC50 (µM)	EC50 (µM)	EC50 (µM)
Dengue 1	D1/H/IMTSSA/98/606 Djibouti	I	>10	0,9± 0,1	>10
	JKT 1186 TVP 949 Indonesia	IV	>10	0,3 ±0,03	5,5 ±3,67
	CNR_25329 France	\mathbf{V}	>10	2,7± 4	7,4± 0,04
	P72-1244 Malaysia	III	3± 0,5	0,9± 0,2	>10
Dengue 2	H/IMTSSA-MART/98-703 France	AA	0,8 ±0,5	0,9± 0,3	2,9± 0,95
	_1751 TC 544 Trinidad	A	1 ±0,7	0,3± 0,06	2,9± 0,01
	CNR_25679 France	C	1,1 ±0,3	0,2± 0,07	1,9± 0,03
	CNR 25326 Thailand	Asian I	0,1 ±0,03	0,9± 0,2	7,7± 0,04
	DENV-2/PG/BID-V2618/2008 Papua New Guinea	Asian II	0,2± 0,16	0,3± 0,5	4,1± 0,02
	DKD811 Malaysia	Sylvatic	0,4 ±0,18	0,4± 0,1	>10
Dengue 3	DENV3/BR/D3LIMHO/2006 Brazil	${f v}$	>10	1± 0,09	>10
	4025 Bolivia	Ш	>10	1± 0,05	>10
	CNR 17046 Malaysia	I	>10	2,8± 0,3	>10
	CNR 15418 Thailand	II	>10	1,2± 0,3	>10
Dengue 4	G11337 India	I	>10	1,2± 0,03	>10
	Dakar HD 34460 Senegal	IIb	>10	0,9± 0,3	>10
	CNR_16861 Malaysia	IIa	>10	0,4± 0,01	>10
	ThD4_0476_97 Thailand	Ш	0,3 ±0,08	0,2 ±0,08	>10
	P75-514 Malaysia	Sylvatic	>10	1± 0,05	>10

The DENV strains of the collection showed similar sensitivity towards the nucleoside analogue inhibitor NITD008 with EC₅₀'s ranging from $0.2\mu M$ to $2.8\mu M$, which is in accordance with previously published results²¹.

The capsid inhibitor ST-148 inhibited all DENV-2 genotypes with EC₅₀'s ranging from 0.25 to 1.1 μ M. However, only one genotype for DENV-1 (DENV-1 GIII at 0.5 μ M), and for DENV-4 (DENV-4 GIII at 0.3 μ M), were inhibited by this compound. Finally, no activity was observed against DENV-3 genotypes, with EC₅₀'s > 10 μ M. Although Byrd and co-workers¹⁹ found that the DENV-2 serotype was the most sensitive serotype to this capsid inhibitor and showed some variability in the inhibition against other serotypes, they did not fully evaluate

- the variation in susceptibility to other serotypes sufficiently comprehensively to draw
- 144 conclusions.
- 145 SDM25N showed moderate efficacy, EC₅₀'s ranging from 1.7 7.7 μM against a large
- proportion of the DENV-2 genotype strains, and half of the DENV-1 genotypes. However, no
- activity was observed against none of the DENV-3 and 4 genotypes with EC₅₀'s above 10
- 148 µM. This result suggests that the binding affinity of NS4B to the hypothetical cellular factor
- targeted by SDM25N varies greatly among various DENV genotypes and/or that this cellular
- factor might be dispensable for efficient replication of some DENV genotypes.
- Overall, the results demonstrate that compounds targeting highly conserved sites, exemplified
- by nucleoside analogue inhibitor NITD008 (targeting the active site of the polymerase), had a
- broader pan-serotypic activity, with similar EC₅₀'s regardless of the DENV genotype. In
- 154 contrast, compounds targeting less conserved proteins or protein subdomains, either directly
- 155 (e.g. the capsid) or indirectly through an interaction with a host factor of the cell (e.g.
- SDM25N), exhibited larger differences in activity towards the various genotypes of DENV.
- 157 Importantly, these data illustrate the fact that a sound in cellulo evaluation of anti-dengue
- 158 candidate molecules requires the use of a complete reference virus panel that enables
- estimates of the antiviral activity against each of the identified DENV genotypes to be
- 160 obtained. Modern reverse genetics techniques have enabled us to develop such a
- representative collection, and it has been made available to the scientific community through
- the European Virus Archive collection (EVA). We believe that the availability of this new
- tool will enable the independent assessment of pan-serotypic activity of anti-dengue
- 164 candidates in the future, fulfilling a critical requirement for a successful dengue antiviral
- small molecule.

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- 176 <u>virusarchive.com</u>).

Contributions

- OG, MVL, GQ and XDL generated the idea of the panel. FT, XDL and GQ conceived the
- experiments. XDL proposed the study design, FT, CB, and GO performed the experiments.
- 181 FT and GQ analyzed the results. FT and GQ wrote the paper. FT, CB, GQ, OG, MVL and
- 182 XDL reviewed and edited the paper.

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