RESEARCH ARTICLE

Nucleobases and corresponding nucleosides display potent antiviral activities against dengue virus possibly through viral lethal mutagenesis

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Abstract

Dengue virus affects millions of people worldwide each year. To date, there is no drug for the treatment of dengue-associated disease. Nucleosides are effective antivirals and work by inhibiting the accurate replication of the viral genome. Nucleobases offer a cheaper alternative to nucleosides for broad antiviral applications. Metabolic activation of nucleobases involves condensation with 5-phosphoribosyl-1-pyrophosphate to give the corresponding nucleoside-5'-monophosphate. This could provide an alternative to phosphorylation of a nucleoside, a step that is often rate limiting and inefficient in activation of nucleosides. We evaluated more than 30 nucleobases and corresponding nucleosides for their antiviral activity against dengue virus. Five nucleobases and two nucleosides were found to induce potent antiviral effects not previously described. Our studies further revealed that nucleobases were usually more active with a better tissue culture therapeutic index than their corresponding nucleosides. The development of viral lethal mutagenesis, an antiviral approach that takes into account the quasispecies behavior of RNA viruses, represents an exciting prospect not yet studied in the context of dengue replication. Passage of the virus in the presence of the nucleobase 3α (T-1105) and corresponding nucleoside 3β (T-1106), favipiravir derivatives, induced an increase in apparent mutations, indicating lethal mutagenesis as a possible antiviral mechanism. A more concerted and widespread screening of nucleobase libraries is a very promising approach to identify dengue virus inhibitors including those that may act as viral mutagens.

Author summary

Dengue virus is a world-wide public health menace estimated to infect hundreds of millions of people per year. Vaccines to prevent dengue virus infection have had limited success due in part to the requirement to elicit effective immune responses against the four dengue serotypes. There is an urgent unmet need for anti-dengue virus therapeutics. Nucleosides are effective antiviral small molecules which usually work by inhibiting the accurate
replication of the viral genome. Typically, nucleosides must be converted within the cell to their triphosphate form to inhibit virus replication, thus inefficient phosphorylation often leads to suboptimal activity. We screened a small library of nucleobases that require an activation pathway different from nucleosides to achieve the same active form. We identified some known and previously undescribed dengue virus nucleobase inhibitors and their corresponding nucleosides. Our investigation of the mechanism of action of one nucleobase and its corresponding nucleoside found evidence for enhanced mutagenesis of the dengue virus genome in the presence of the compounds in cell culture. A wide screening of nucleobases libraries is a promising strategy to discover dengue virus inhibitors including potential viral mutagens.

Introduction

Dengue virus (DENV) is a worldwide health threat, with hundreds of millions of people infected yearly in more than 100 countries [1]. There are four known DENV serotypes and a first infection with one serotype followed by a second infection with another serotype may result in severe disease [2, 3]. For these and other issues, vaccines designed for a pan-serotype protection, including the commercial dengue vaccine approved and used in a few countries, have yielded mixed results [4, 5]. Safety and partial efficacy concerns in addition to cost, storage and delivery issues may hinder implementation of vaccines in many countries.

There are currently no approved drugs to treat DENV infection. Thus far, classical antiviral approaches (e.g. NS5 polymerase inhibitors, entry inhibitors, protease inhibitors, etc.) have yet to provide treatments for DENV infection and therefore the investigation of new antiviral strategies is warranted [6–8]. One such strategy to explore is lethal mutagenesis [9]. The idea of viral lethal mutagenesis is to exploit the natural tendency of RNA viruses to mutate in order to favor the accumulation of deleterious mutations in the newly formed viruses, eventually leading to viral extinction (for review see [10]). DENV and other RNA viruses display a high mutation rate (10^{-4} to 10^{-6} mutations per bp per generation) [11, 12] as an evolutionary characteristic allowing these viruses to escape host immune defense mechanisms and adapt rapidly to new stress conditions [13, 14]. An error-prone viral polymerase combined with a high replication rate are considered to be the main sources of mutations. It is this critical source of viral adaptability (e.g. the virus high mutation rate) that makes RNA viruses a target of choice for antiviral lethal mutagenesis strategies [15–17]. RNA viruses maintain a delicate balance between their need to adapt and their need to preserve a level of genetic integrity put at risk by deleterious mutations [18]. Modifying this fragile equilibrium by increasing the viral mutation rate with mutagens has been proposed as an antiviral strategy [15]. The well-known antiviral nucleoside drug ribavirin induces lethal mutagenesis for different viruses [19–22].

The discovery of new nucleosides as antiviral mutagens has been impaired by several hurdles including the toxicity of the potential drugs as well as synthetic challenges. In addition, potentially mutagenic nucleoside analogues frequently suffer from poor metabolic conversion to the active triphosphate form required by the viral polymerase. The first phosphorylation of the nucleoside analogue is often the rate limiting step to obtain the active mutagenic nucleoside triphosphate used by the viral polymerase [23, 24]. In order to overcome the potential first phosphorylation difficulty, we propose to use nucleobases (purine or pyrimidine base without the ribose or phosphate moieties of a nucleoside). Enzyme mediated condensation of nucleobases with 5-phosphoribosyl-1-pyrophosphate to give the corresponding nucleoside-5’-monophosphate can provide an alternative pathway. Thus, for a nucleoside where the first
phosphorylation is inefficient then using its corresponding nucleobase could allow metabolic conversion to the corresponding nucleoside triphosphate, thereby providing a more efficient metabolic conversion to the triphosphate. Studies on nucleobases have been limited [25], with the notable exceptions of 5-fluorouracil and T-705 (favipiravir) [26–32]. However, in the context of targeting viruses that affect developing countries, mutagenic nucleobases present key advantages over mutagenic nucleosides. In addition to their different metabolic activation pathways, nucleobase analogues are considerably cheaper, more diverse and commercially available in higher numbers compared to corresponding nucleoside analogues. The chemical synthesis of a nucleobase is faster and simpler than the synthesis of the corresponding nucleoside. Similarly to nucleosides, nucleobases possess their own cellular transporters [33, 34].

In this study, we describe the identification of five nucleobases and three corresponding nucleosides that possess potent anti-DENV activity. These compounds have not been previously described to have anti-DENV activity, except for the nucleoside 1b (ribavirin) [35, 36]. We compared the antiviral activities and toxicities of the nucleobases with their corresponding nucleosides. For virus passaged in the presence of a nucleobase 3a (T-1105) or nucleoside 3b (T-1106), we detected an increase in mutations compared to virus passaged in DMSO indicating a possible reduction in virus titer via increased mutagenesis. To our knowledge, our study is the first to fully compare the antiviral mechanisms and efficacies of a nucleobase and its corresponding nucleoside, highlighting the differences, similarities and potential advantages of nucleobases versus nucleosides. Our study also highlights the potential of lethal mutagenesis induction during DENV replication as an alternative to classical antiviral strategies.

**Methods**

**Compounds**

5a (T-705) (CAS# 259793-96-9, 6-fluoro-3-hydroxypyrazine-2-carboxamide) was purchased from ASTA Tech. 3a (T-1105) (CAS# 55321-99-8, 3-Hydroxy-2-pyrazinecarboxamide) was purchased from Alfa Aesar. 3b (T-1106) was synthesized according to known procedures (Preparation of nucleosides with non-natural bases as anti-viral agents Can. Pat. Appl. (2006), 149pp. CODEN:CPXXEB; CA2600359). 1a (ribavirin base) (CAS# 3641-08-5, 1,2,4-Triazole-3-carboxamide) was purchased from Ark Pharm. 1b (ribavirin) (CAS# 36791-04-5) was purchased from Carbosynth. 2a (mizoribine base) (CAS# 56973-26-3, 5-Hydroxy-1H-imidazole-4-carboxamide) was purchased from Ark Pharm. 2b (mizoribine) (CAS# 50924-49-7) was purchased from Carbosynth. 4a (diaminopurine) (CAS# 1904-98-9, 2,6-diaminopurine) was purchased from Sigma-Aldrich. 4b (diaminopurine riboside) (CAS# 2096-10-8, 2-Aminoadenosine) was purchased from Berry and Associates. 6 (mycophenolic acid) (CAS# 24280-93-1) was purchased from Sigma-Aldrich.

**Cell lines and virus**

The hepatocyte-derived cellular carcinoma cell line Huh-7 [37] was used for DENV infection and drug treatment. The African green monkey kidney Vero cell line (ATCC CRL-81) was used to titer DENV via plaque assay. The baby hamster kidney cell line carrying a DENV subgenomic replicon, BHK pD2-hRucPac-2ATG30 [38] (obtained from Dr. M. Diamond, Washington University, School of Medicine), was used for DENV replicon assay. All cell lines were maintained in Dulbecco’s modified Eagle’s (DME) medium supplemented with 10% fetal bovine serum (FBS), 100 IU streptomycin/penicillin per ml and 10 μg/mL plasmocin (InvivoGen) at 37°C in a 5% CO2 incubator. DENV replicon cells were supplemented with 3 μg/mL puromycin (Life Technologies). DENV-2 stocks from New Guinea C strain (ATCC VR-1584) were generated from C6/36 mosquito cell cultures (ATCC CRL-1660) grown in Minimum
Essential Medium (MEM) supplemented with 10% FBS, 1% non-essential amino acids and 1% sodium pyruvate at 28˚C with 5% CO₂. The C6/36 cells on T-150 flasks were inoculated with virus and the supernatant harvested after complete cytopathic effects. Viral stock titers were determined by plaque assay on Vero cells.

**Cell viability assay**

The sensitivity of the cell lines to the compounds was examined using the 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium (MTS)-based tetrazolium reduction CellTiter 96 Aqueous Non-Radioactive cell proliferation assay (Promega G5430). The compounds were initially tested at 10 and 50 μM final concentrations. Each plate also contained DMSO alone, medium alone, and an inhibitory compound, 6. DENV replicon or Huh-7 cells were plated at a density of 1,500 or 8 × 10³ cells, respectively, per well in 96-well plates containing 100 μl of culture medium overnight. Compounds were added to triplicate wells in culture medium and incubated for an additional 72 h. MTS reagent was then added to each well and incubated at 37˚C in a humidified 5% CO₂ atmosphere. The plates were read at various time points at a wavelength of 490 nm using a Molecular Devices M5e plate reader. Mean values of triplicate wells were determined and compared to the mean value for the wells that received DMSO alone. For compounds selected for dose-response experiments, the CC₅₀ was determined by comparing cell viability for eight serial dilutions of the compound and DMSO treated cells using GraphPad Prism software. The CC₅₀ value was defined as the compound concentration resulting in a 50% reduction readout compared with the DMSO.

**DENV replicon assay**

Compounds were evaluated for antiviral properties using BHK cells containing a DENV-2 viral replicon. 1.5 × 10³ replicon-containing cells per well were plated in white opaque 96-well plates in the absence of antibiotic selection and the next day, compounds dissolved in DMSO were added to triplicate wells in culture medium. The compounds were initially tested at 10 and 50 μM final concentrations and each plate also contained DMSO alone, medium alone, and 6. Three days later, medium was replaced with a 1:1000 dilution of ViVi-Ren Live Cell Substrate (Promega) in DME minus phenol red and 10% FBS. Luminescence was measured with a Molecular Devices M5e plate reader. Mean values of triplicate wells were determined and compared to the mean value for the wells that received DMSO alone. For compounds selected for dose response experiments, the concentration of compound that reduced luciferase activity by 50% was defined as the 50% effective concentration (EC₅₀). The EC₅₀ was determined by comparing luciferase activity for eight serial dilutions of the compound and DMSO treated cells using GraphPad Prism software.

**Titer reduction assay**

Huh-7 cells were seeded in 12-well plates at a density of 4×10⁵ cells per well in 1 mL culture medium. The next day, cells were washed and inoculated with DENV at a multiplicity of infection (m.o.i.) of 0.2 in 500μl infection medium (MEM containing 2% FBS and 10 mM HEPES). The inoculum was removed after 1h, cells were washed with PBS and then incubated in 1 mL MEM, 2% FBS, 1% pen/strep plus compound for 72 h. Viral supernatants were clarified by centrifugation for 5 min at 1500xg and aliquoted and stored at -80˚C. Viral titers were determined using a plaque assay on Vero cells. Briefly, confluent Vero cell monolayers in 24-well plates were incubated at 37˚C for 1 h with duplicate 300 μl samples of 10-fold serial dilutions of viral supernatants. The cells were then washed to remove unbound viral particles and overlaid with 500ul MEM containing 1.3% methylcellulose, 5% FBS and 10mM HEPES. After 5
days of incubation at 37˚C and 5% CO₂, cells were washed with PBS, fixed, and stained using 1% Giemsa. Infectious virus titer (pfu/mL) was determined using the following formula: number of plaques × dilution factor × (1/inoculation volume). The viral titer was presented as the mean of duplicate samples from a dilution yielding approximately 20–50 plaques per well.

**Viral RNA extraction, RT-PCR amplification, quantitative PCR, detection of viral genome mutations**

For qPCR determination of viral genome copy number, viral RNA was isolated from 140 μL of drug-treated or DMSO-treated infected culture supernatant using QIAamp Viral RNA mini kit (Qiagen), following manufacturer’s protocol. Viral RNA was quantified using the TaqMan RNA-to-C₇ 1-Step qPCR Kit (Applied Biosystems). Primers used for qPCR were 5’-CAT-GATGGGAAAGAGAGAAGAAGCT-3’ (forward) and 5’-GGCTCTGCTGCTTTTGCC-3’ (reverse) amplifying a region numbering 8928–8988 in the genome (numbering starting from the beginning of genome, accession number KM204118). The qPCR FAM probe sequence is 5’-TTGCCGAACTCCC-3’. Serial 10-fold dilutions of plasmid containing the NS5 gene of DENV were used to generate a standard curve for the quantification of viral RNA genome copy number based on cycle threshold (C₇) values. The limit of detection for NS5 plasmid dilutions was 30 copies (S1 Fig). One-way ANOVA was performed to determine statistical significance of mean genome copy numbers among treatments at each virus passage and Tukey’s honestly significant difference (HSD) used to determine statistical significant (p < 0.05) between DMSO and 3a or 3b at each passage. Statistical analysis via GraphPad Prism 5 software. To obtain sequence data from viral RNA isolated at each passage, cDNA was generated via M-MLV reverse transcriptase and random hexamers (New England Biolab Inc.) per manufacturer’s instructions. An approximately 1600-base fragment covering membrane protein (prM) and envelop protein gene (E) of DENV-2 viral genome was amplified using PfuUltra II Fusion HS DNA polymerase (Agilent) with primers prMEfor (5’- AACTCAGAATTCTCAAATTGAACCAC-3’) and prMErev (5’- AACTCAGAATTCTCCTTTGAGTTGAGCT-3’). PCR products were purified using Qiaquick PCR purification kit (Qiagen) and then digested with EcoRI and ligated into pcDNA3.1 for sequencing. Sequence for approximately 40–50 individual clones per sample was obtained from the University of Minnesota Genomics Center. Sequences were aligned over a 980-base region that had adequate quality sequencing reads for all clones. Only mutations present in both the forward and reverse reads of a clone were counted. All incidence-based determinants of mutation frequency (Mf min, Mf max, Mfe) were calculated as described [39]. A two-tailed Mann-Whitney U test (GraphPad Prism 5 software) was used to determine if there were statistically significant differences for the mean number of mutations per clone between DMSO-treated and each drug-treated virus passage.

**Construction of DENV-2 NS5 plasmid**

The DENV-2 viral RNA was isolated from 140 μL of lab stock DENV-2 using QIAamp Viral RNA mini kit (Qiagen). cDNAs corresponding to viral RNAs were generated with random hexamers (New England Biolab Inc.). A 2600-base fragment of NS5 cDNA was amplified using PfuUltra II Fusion HS DNA polymerase (Agilent) with primers NS5for (5’- GCCCGGTGCCAGTTGAATTCCAGCAGCAGC-3’) and NS5Rev (5’- CCGGGGATCTCTAGAATCCAGGACTCTCCTTTGAC-3’). PCR product were purified using Qiaquick PCR purification kit (Qiagen) and then digested with XbaI and HindIII digested pUC18 vector using Infusion HD Cloning Kit (Clontech). A positive clone was identified and the nucleic acid sequence of NS5 confirmed by sequencing at the University of Minnesota Genomics Center.
Virus and compound passage

Virus was passaged on Huh-7 cells supplemented with 200 μM 3a, 500 μM 3b or DMSO (0.5%). Huh-7 cells were seeded in 12-well plates and inoculated with DENV-2 as described for Titer reduction experiments above except an m.o.i. of 0.01 was used. After 3 days of compound treatment, 50 μL of the harvested supernatant was used to inoculate fresh Huh-7 cells in the continued presence of compound. The virus titer in the harvested supernatant was determined by plating ten-fold serial dilutions onto single wells of a 24-well plate. The wells were washed and overlaid as described in the Titer reduction assay above. If no plaques were obtained in any of the harvested supernatant dilutions, the undiluted supernatant was used. If plaques were not detectable, the virus titer was considered to be at the limit of detection, 1 plaque (3.3 pfu/mL). Supernatant titer was determined from a single well where there were approximately 20–50 plaques when possible. This experiment was repeated three times at the compound concentrations listed.

Results

Antiviral nucleoside identification can be hindered by difficulties in chemical synthesis and poor conversion of the nucleoside to the active triphosphate form. To circumvent these issues, we propose to use nucleobases in our initial screen for antiviral agents because of their different activation pathway to the active nucleotide (Fig 1), their low cost and ready commercial availability. Phosphoribosyltransferases of the cellular nucleotide salvage pathway directly convert some nucleobases to the corresponding nucleoside monophosphate and therefore the corresponding nucleoside analogue need not be an efficient substrate for a nucleoside kinase (Fig 1) [40, 41]. In that regard, 3a and analogue 5a (Fig 2) are substrates of human phosphoribosyl transferases and are converted in one step to the corresponding nucleoside monophosphate [41].

Our strategy to identify nucleobase and nucleoside DENV inhibitors was to screen for activity and toxicity of selected compounds at 10 μM and 50 μM using a luciferase-reporting DENV replicon cell line, BHK pD2-hRucPac-2ATG30 [38]. Compounds that demonstrated inhibitory activity against the replicon cell line were used in dose-response analysis to assign EC₅₀ and CC₅₀ values. The nucleobases were generally more active with a higher tissue culture therapeutic index (CC₅₀/EC₅₀) than their corresponding nucleosides (Fig 2). The EC₅₀ values of the active nucleobases range from 2.4 to 110 μM, comparable to the EC₅₀ values of the active nucleosides that range from 1.3 to 113 μM (Fig 2). Nucleobase 3a is 5 times more active than nucleobase 5a (favipiravir). The CC₅₀ values of the nucleobases 1a, 3a and 5a were beyond 665 μM (Fig 2). Remarkably, nucleobase 1a did not show cytotoxicity at 1000 μM compared to 1b nucleoside (Fig 2) where the CC₅₀ was 20 μM. 2a nucleobase displayed a clear antiviral effect at 2.4 μM. Representative EC₅₀ curves for 2a plus corresponding nucleoside 2b and for 3a plus corresponding nucleoside 3b are shown in Fig 3. Inactive nucleobases at initial screening are listed in S1 Table in the supplementary material.

We focused on the antiviral mechanism of nucleobase 3a and corresponding nucleoside 3b for the following reasons: 1) among the 5 active nucleobases, 5a was already known to induce viral lethal mutagenesis but its corresponding nucleoside was not available and unable to be synthesized for our study; 2) 1a and 2a are likely to possess a complex mechanism of action due to their probable inhibitory effect of inosine monophosphate dehydrogenase (IMPDH) after conversion to the nucleotide form; 3) 4b was too toxic to perform a full study comparing the nucleobase to the nucleoside (Fig 2). Therefore, we chose 3a, known to be substrate of human phosphoribosyl transferase [41], and its corresponding nucleoside 3b to be the best
candidates for antiviral mechanism of action studies and to compare the effects of nucleobase and corresponding nucleoside.

Our approach to study mechanism of action and possible lethal mutagenesis was to passage virus in Huh-7 cells in the presence of a compound and determine the compound’s effect on virus titer, genome copy number and genome sequence. We had initially identified inhibitors using a DENV replicon BHK cell line so we wanted to verify inhibitory activity in Huh-7 cells.

Fig 1. Activation pathways of antiviral nucleobases and nucleosides.

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using replication competent DENV. We chose Huh-7 cells because they are a human cell line, they are susceptible to DENV infection and they produce readily detectable infectious virus. The BHK replicon cells are a convenient tool to identify initial DENV inhibitors. We used human cells for more detailed studies for these compounds that require conversion to the active form by cellular enzymes. We repeated the dose-response experiments for 3a and 3b using a titer-reduction assay with Huh-7 cells as previously described [42]. The values obtained for the compounds (Table 1) were consistent with those from the replicon assay. Based upon the data in Table 1, we used non-toxic levels of 3a (200 μM) and 3b (500 μM) that we empirically determined (S2 Fig) would reduce virus replication steadily during multiple passages and ultimately lead to undetectable levels of infectious virus. The results for virus passage in the presence of 3a (200 μM), 3b (500 μM) or DMSO (0.5%) are shown in Fig 4. Compounds were added to Huh-7 cells shortly after inoculation. After 3 days incubation in compound, the supernatant was collected and a fixed volume (50 μL) of supernatant was used to inoculate fresh cells. The level of infectious virus present in the supernatant for each passage was determined by plaque assay and the number of viral genomes by RT-qPCR. For the RT-qPCR, we

### Table 1: Dose-response results for selected compounds evaluated in DENV replicon cells

| Structure | Nucleobase | EC₅₀ (μM) | CC₅₀ (μM) | TTI  |
|-----------|------------|-----------|-----------|------|
| ribavirin | 1a         | 4.9 ±1.6  | >1000     | 204  |
| mizoribine| 2a         | 2.4 ±0.1  | 23 ±4.2   | 9.6  |
| T-1105    | 3a         | 21 ±0.7   | >665      | 32   |
| diamino-purine | 4a    | 3.6 ±1.3  | 13 ±2.1   | 3.6  |
| T-705     | 5a         | 110 ±30   | >1000     | 9.1  |
| Control   | myco-phenolic acid | 0.3 ±0.03 | 1.4 ±0.4 | 4.7  |

N/A = not available.
chose primers within the NS5 gene to increase likelihood of detecting full-length genomes. Both compounds induced a steady decline in infectious virus production and supernatant genome copy number over repeated passages (Fig 4). In the presence of 3b, infectious virus

Table 1. Titer reduction assay dose-response.

| Compound | EC_{50} (μM) | CC_{50} (μM) |
|----------|--------------|--------------|
| 3a       | 20 ± 11      | >1000        |
| 3b       | 60 ± 22      | >1000        |

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was undetectable at passages 4 and 5 and genomic RNA was undetectable at passage 5 (Fig 4). For 3a, no infectious virus was detected at passages 5 and 6 and no genomic RNA was detected at passage 6 (Fig 4). The mean genome copy numbers (Fig 4B) for the three treatments were statistically different at each passage (p < 0.001, One Way ANOVA) with statistically significant differences between 3a and DMSO and between 3b and DMSO at each passage (p < 0.05, Tukey’s HSD). Both compounds displayed a reduction in the ratio of infectious virus to RNA genome copy number (sometimes referred to as RNA specific infectivity) when compared to DMSO (Fig 4). This delay in reduction of genomic RNA compared to infectious virus is a hallmark of a mutagenesis-based antiviral activity [43–46]. These observations, along with the knowledge that 5a (analogue of 3a and 3b) induces mutagenesis in influenza A, norovirus and the flavivirus West Nile virus [27–29], motivated us to determine if a mechanism of action for 3a or 3b included an enhanced mutagenesis of the viral genome.

We hypothesized that an increase in mutations induced by a particular nucleobase or nucleoside would be detected by analyzing the viral genome sequence at a passage near where the titer was significantly reduced or undetectable. Therefore, we amplified a region containing the pre-membrane (prM) and envelope (E) genes from viral genomic cDNA derived from passage 3b-treated cells and passage 4 3a-treated cells because those passages were just before viral titer was undetectable. We amplified the prM/E region of the viral RNAs instead of the downstream NS5 gene to increase the likelihood we would obtain PCR products when viral titers were greatly reduced. The presence of the NS5 gene would require an almost complete genomic RNA whereas the prM/E region was closer to the 5’ end of the viral genomic RNA its presence would not require a complete viral genome. The amplified products were inserted into a cloning vector and the nucleotide sequence was analyzed for the resulting 35–50 independent clones. We sequenced at least 31,000 nucleotides for each set of clones similar to other published studies of viral lethal mutagenesis [27, 43] and compared the sequences to that of the consensus sequence obtained from DMSO-treated cell supernatants at passages 3 and 4.

An initial plot and analysis of the number of mutations per clone for 3a treatment passage 4 and 3b passage 3 clearly indicate there was a significant difference (p < 0.0001) in the number of mutations per clone for sequence obtained from compound-passaged virus compared to
DMSO (Fig 5A). We next calculated a number of incidence-based mutation indices to better determine possible heightened mutagenesis as recommended in a recent review [39]. There was an increase in every index of mutation frequency and sequence diversity for 3a- and 3b-treated clones compared to those for DMSO (Table 2). These include, the minimum mutation frequency (Mf min, mutation at a given nucleotide counted only once), maximum mutation frequency (Mf max, all mutations counted) and the entity level mutation frequency (Mfe, mutations with respect to sequence of dominant haplotype of drug-treated clones). In addition, the number of haplotypes, number of different mutations, number of total mutations and number of clones with a mutation (Table 2) were all higher for 3a and 3b passage sequences. We noted fewer mutations in the DMSO passage 4 sequence than seen for other DMSO passages (Table 2 and Fig 5B). That does not change our interpretation of the 3a passage 4 results. The mean number of mutations per clone for 3a passage 4 was significantly higher than the mean mutations per clone for any DMSO passage (ranging from $p < 0.01$ to $p < 0.03$). In addition, all advanced indices of mutation frequency and sequence diversity were greater for 3a passage 4 when compared to all DMSO passages (Table 2). Lastly, we examined sequence data from other 3a- and 3b-treated passages. Sequence from passage 2 also showed higher numbers of mutations per clone and an increase in all advanced mutation indices for 3a- and 3b-passaged virus compared to DMSO (Fig 5 and Table 2). The sequence data from 3a passage 3 yielded mixed results with certain measures of mutation frequency similar to DMSO and certain others higher than those for DMSO (Table 2). We were unable to obtain sequence data for 3b passage 4 or 5. Taken together, these results indicate an enhanced mutagenesis as a likely significant contributor to the antiviral mechanism of action for 3a and 3b.

In the sequence analysis, we excluded nucleotides 952–4 (numbering starting from the beginning of genome, accession number KM204118) in analysis of mutations. This codon, located in the E gene, changes in response to repeated DENV passaging in cultured cells [47, 48]. The sequence for nucleotides 952–4 of our stock virus was ATA. However, nearly all the clones from virus passaged with DMSO, 3a and 3b displayed changes in that sequence (S2 Table). We also excluded mutations that were present in a majority of clones and therefore were consensus sequence changes. These included one nucleotide position (1043) in the 3a clones and three positions (960, 1057 and 1493) in the 3b clones (S2 Table). A hallmark of lethal mutagenesis is an unchanged consensus sequence despite an increase in single mutations [44, 45] and so it is possible those changes were introduced during the RT-PCR amplification of the viral RNA genomes.

An examination of the types of mutations induced by nucleoside 3b revealed an almost exclusive increase in transition mutations for purines and pyrimidines (Table 2). For nucleobase 3a, a similar increase in mostly transition mutations was observed (Table 2). These results are also consistent with data observed with related compound 5a [27, 28].

**Discussion**

Effective and inexpensive drug therapies for DENV infection are urgently needed. So far, classical antiviral strategies have failed to identify small molecules to treat DENV infection. In addition, developing broad antiviral strategies is critical to face future outbreaks of emerging viruses. The adaptability of viruses to changing environments results in the generation of a heterogeneous population of closely related yet different viral variants during infection. The diversity present within this population may result in viruses escaping drug treatments or host immune defenses. Lethal mutagenesis aims to generate deleterious viral mutations that would prevent viral adaptation and drive the viral population to collapse. Induction of lethal mutagenesis has largely been based on using nucleoside mutagens to target the error prone viral
polymerase, a main source of viral mutations. However, limitations in the chemical synthesis of nucleosides and poor metabolism to the active triphosphate form have limited the discovery of new viral mutagens. Here, we have identified nucleobases and nucleosides with significant antiviral activity against DENV, some of which appear to act through lethal mutagenesis. Along with nucleosides, nucleobases may be valuable molecules to induce viral lethal
In this study, we have identified five nucleobases and three nucleosides that possess anti-DENV activity. Nucleobase 3a and corresponding nucleoside 3b were selected for antiviral mechanism-of-action studies. 3a and 3b both increased mutations. Although the antiviral mechanisms of 5a, 1b and 4b have not been explored in this study, these compounds are known antiviral mutagens [22, 25, 28] supporting our result that viral lethal mutagenesis may be an effective antiviral strategy against DENV. The strong antiviral effects of 2b (EC_{50} 15 μM) and its nucleobase 2a (EC_{50} 2.4 μM) suggest a critical role of IMPDH for DENV replication, yet the fact that 2a and 2b possess a rotatable amide bond (theoretically able to mimic either adenosine or guanosine) indicates a component of the antiviral effect could occur through lethal mutagenesis. To our knowledge, this is the first description of 2a as an antiviral agent.

Interestingly, in our study, the active nucleobases and nucleosides (Fig 2) share common structural characteristics such as 1) a purine analog structure and/or 2) a rotatable amide bond. Some of these nucleobases have been described to be converted metabolically to the active nucleotide form [41]. Our study is the first to compare a nucleobase side by side with its corresponding nucleoside for their abilities to induce viral genome mutagenesis as an antiviral mechanism of action. The way by which mutagenic nucleobases/nucleosides and resulting mutagenic nucleotides induce mutations is usually due to ambiguous base pairing

| Passage (P) | P 2 | P 3 | P 4 | P 5 |
|------------|-----|-----|-----|-----|
|            | DMSO | 3a\(^b\) | 3b\(^b\) | DMSO | 3a | 3b | DMSO | 3a | DMSO |
| Transition |      |       |      |      |      | 3 | 6 | 1 | 3 | 1 | 3 |
| G→A       | 3    | 3    | 3    | 1    | 4    | 1 |
| A→G       | 2    | 3    | 2    | 1    | 4    | 1 |
| U→C       | 1    | 14   | 4    | 2    | 3    | 4 |
| C→U       | 2    | 4    | 1    | 3    | 6    | 3 |
| Transversion |      |       |      |      |      | 1 | 2 | 1 | 2 |
| U→A       | 1    | 2    | 1    | 2    |
| G→U       | 1    | 1    | 1    |
| U→G       | 1    | 1    | 1    |
| C→A       |      |       |      |      | 2 | 1 |
| A→U       |      |       |      | 1    |

Number of different mutations: 7, 17, 33, 9, 9, 15, 1, 16, 7
Number of total mutations: 11, 80, 47, 10, 19, 50, 1, 27, 8
Number of nucleotides sequenced: 38,829, 32,508, 30,702, 36,120, 33,411, 31,605, 40,635, 40,635, 35,217
Number of haplotypes: 7, 17, 22, 9, 9, 17, 2, 15, 8
Number of clones with mutation/total: 7/43, 26/36, 22/34, 9/40, 12/37, 22/35, 1/45, 19/45, 8/39
Mf min\(^a\), per 10,000 nucleotides: 1.8, 5.2, 10.7, 2.5, 2.7, 4.7, 0.2, 3.9, 2.0
Mf max\(^a\), per 10,000 nucleotides: 2.8, 24.6, 15.3, 2.8, 5.7, 15.8, 0.2, 6.6, 2.3
Mfe\(^a\) (×10^{-3}): 1.4, 2.5, 2.4, 1.1, 1.4, 2.2, 0.6, 1.6, 1.0

\(^a\)Defined in the text
\(^b\)3a 200 μM and 3b 500 μM for all passages

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misinterpreted by the viral polymerase when replicating the viral genome [49]. Typically, ambiguous base pairing originates from rotational or tautomeric forms of the base that result in the mutagenic nucleoside triphosphate resembling more than one natural nucleoside triphosphate. In that matter, 5a, an analogue of 3a and 3b, was found to be converted to the active triphosphate form and then used by influenza polymerase as an A or G mimic, likely by rotation of its amide bond [50]. In a similar fashion, 1b (ribavirin) possesses a rotatable amide bond and is widely accepted as an A or G mimic. Overall, 3a and 3b in this study and 5a and 1b in previous studies all essentially induce transition mutations [22, 27], a result compatible with these molecules resembling A or G. The 3b triphosphate would manifest as an ATP or GTP mimic and the rotation of its amide bond would allow base pairing with a U or C, respectively (Fig 6). We did not observe a clear preference for A to G and U to C transition mutations versus the G to A and C to U mutations indicating that the DENV polymerase may not have a preference for using 3b triphosphate. The proportions of A to G or G to A transitions induced by these mutagens can vary from virus to virus [27, 28]. The affinity of the different viral polymerases for one conformation of a mutagen rather than another may explain the observed variance. A mutagen such as 1b also inhibits IMPDH, an enzyme involved in nucleotide biosynthesis, resulting in an imbalance in endogenous nucleotide pools which may impact the induced mutation spectrum compared to a purely mutagenic compound. In our case, we did not observe major trends for one particular transition mutation versus another. Exploring molecules capable of inducing transversion mutations may be of interest to induce a stronger antiviral effect.

Fig 6. Potential base pairing of nucleoside triphosphate forms of 3a and 3b. Once converted to the active nucleoside triphosphate forms, mutagens 3a and 3b may adopt different conformations due to the rotation of the amide bond allowing these compounds to mimic GTP and base pair with C or mimic ATP and base pair with U.

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Supporting information

S1 Table. Inactive molecules. (PDF)

S2 Table. List of sequence changes for prM/E region cDNA. (XLSX)

S1 Fig. Standard curve for DENV NS5 qPCR. (PDF)

S2 Fig. Passage of virus in different concentrations of 3a and 3b. (PDF)

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References

1. Bhatt S, Gething PW, Brady OJ, Messina JP, Farlow AW, Moyes CL, et al. The global distribution and burden of dengue. Nature. 2013; 496(7446):504–7. https://doi.org/10.1038/nature12060 PMID: 23563266; PubMed Central PMCID: PMCPMC3651993.

2. Halstead SB. Neutralization and antibody-dependent enhancement of dengue viruses. Adv Virus Res. 2003; 60:421–67. PMID: 14689700.

3. Simmons CP, Farrar JJ, van Vinh Chau N, Wills B. Dengue. New England Journal of Medicine. 2012; 366(15):1423–32. https://doi.org/10.1056/NEJMc1110265 PMID: 22494122.

4. Hadinegoro SR, Arredondo-Garcia JL, Capeding MR, Deseda C, Chotpitayasunondh T, Dietze R, et al. Efficacy and Long-Term Safety of a Dengue Vaccine in Regions of Endemic Disease. New England Journal of Medicine. 2015; 373(13):1195–206. https://doi.org/10.1056/NEJMoa1506223 PMID: 26214039.

5. Simmons CP. A Candidate Dengue Vaccine Walks a Tightrope. New England Journal of Medicine. 2015; 373(13):1263–4. https://doi.org/10.1056/NEJMc1509442 PMID: 26214040.
6. Chen YL, Yokokawa F, Shi PY. The search for nucleoside/nucleotide analog inhibitors of dengue virus. Antiviral Res. 2015; 122:12–9. https://doi.org/10.1016/j.antiviral.2015.07.010 PMID: 26241002.

7. Lim SP, Wang QY, Noble CG, Chen YL, Dong H, Zou B, et al. Ten years of dengue drug discovery: progress and prospects. Antiviral Res. 2013; 100(2):500–19. https://doi.org/10.1016/j.antiviral.2013.09.013 PMID: 24076358.

8. Low JG, Ooi EE, Vasudevan SG. Current Status of Dengue Therapeutics Research and Development. J Infect Dis. 2017; 215(suppl_2):S96–S102. https://doi.org/10.1093/infdis/jiw423 PMID: 28403438; PubMed Central PMCID: PMCPMC5388029.

9. Loeb LA, Essigmann JM, Kazazi F, Zhang J, Rose KD, Mullins JI. Lethal mutagenesis of HIV with mutagenic nucleoside analogs. Proc Natl Acad Sci U S A. 1999; 96(4):1492–7. PMID: 9990051; PubMed Central PMCID: PMCPMC15492.

10. Perales C, Domingo E. Antiviral Strategies Based on Lethal Mutagenesis and Error Threshold. Curr Top Microbiol. 2016; 392:323–39. https://doi.org/10.1007/82_2015_459 PubMed PMID: WOS:000385415200016. PMID: 26294225.

11. Sanjuan R, Nebot MR, Chirico N, Mansky LM, Belshaw R. Viral mutation rates. Journal of virology. 2010; 84(19):9733–48. Epub 2010/07/28. https://doi.org/10.1128/JVI.00694-10 PMID: 20660197; PubMed Central PMCID: PMCPMC2937809.

12. Drake JW, Holland JJ. Mutation rates among RNA viruses. Proc Natl Acad Sci U S A. 1999; 96(24):13910–3. Epub 1999/11/26. PMID: 10570172; PubMed Central PMCID: PMCPMC24164.

13. Twiddy SS, Holmes EC, Rambaut A. Inferring the rate and time-scale of dengue virus evolution. Mol Biol Evol. 2003; 20(1):122–9. Epub 2003/01/10. https://doi.org/10.1093/molbev/msg010 PMID: 12519914.

14. Park AW, Daly JM, Lewis NS, Smith DJ, Wood JL, Grenfell BT. Quantifying the impact of immune escape on transmission dynamics of influenza. Science. 2009; 326(5953):726–8. Epub 2009/11/11. https://doi.org/10.1126/science.1175980 PMID: 19900931; PubMed Central PMCID: PMCPMC3800096.

15. Holland JJ, Domingo E, de la Torre JC, Steinhauer DA. Mutation frequencies at defined single codon sites in vesicular stomatitis virus and poliovirus can be increased only slightly by chemical mutagenesis. Journal of virology. 1990; 64(8):3960–2. Epub 1990/08/01. PMID: 1695258; PubMed Central PMCID: PMCPMC249691.

16. Anderson JP, Daifuku R, Loeb LA. Viral error catastrophe by mutagenic nucleosides. Annual review of microbiology. 2004; 58:183–205. Epub 2004/10/19. https://doi.org/10.1146/annurev.micro.58.030603.123649 PMID: 15487935.

17. Bull JJ, Sanjuan R, Wilke CO. Theory of lethal mutagenesis for viruses. Journal of virology. 2007; 81(6):2930–9. Epub 2007/01/05. https://doi.org/10.1128/JVI.01624-06 PMID: 17202214; PubMed Central PMCID: PMCPMC1859999.

18. Domingo E, Holland JJ. RNA virus mutations and fitness for survival. Annual review of microbiology. 1997; 51:151–78. Epub 1997/01/01. https://doi.org/10.1146/annurev.micro.51.1.151 PMID: 9343347.

19. Crotty S, Maag D, Arnold JJ, Zhong W, Lau JY, Hong Z, et al. The broad-spectrum antiviral ribonucleoside ribavirin is an RNA virus mutagen. Nat Med. 2000; 6(12):1375–9. Epub 2000/12/02. https://doi.org/10.1038/82191 PMID: 11100123.

20. Day CW, Smee DF, Jalander JG, Yamshchikov VF, Sidwell RW, Morrey JD. Error-prone replication of West Nile virus caused by ribavirin. Antiviral Res. 2005; 67(1):38–45. Epub 2005/05/28. https://doi.org/10.1016/j.antiviral.2005.04.002 PMID: 15919121.

21. Dietz J, Schelhorn SE, Fitting D, Mihm U, Susser S, Welker MW, et al. Deep sequencing reveals mutagenic effects of ribavirin during monotherapy of hepatitis C virus genotype 1-infected patients. Journal of virology. 2013; 87(11):6172–81. Epub 2013/03/29. https://doi.org/10.1128/JVI.02778-12 PMID: 23536652; PubMed Central PMCID: PMCPMC3648094.

22. Moreno H, Gallego I, Sevilla N, de la Torre JC, Domingo E, Martin V. Ribavirin can be mutagenic for arenaviruses. Journal of virology. 2011; 85(14):7246–55. Epub 2011/05/13. https://doi.org/10.1128/JVI.00614-11 PMID: 21561907; PubMed Central PMCID: PMCPMC3126590.

23. Van Rompay AR, Johansson M, Karlsson A. Phosphorylation of nucleosides and nucleoside analogs by mammalian nucleoside monophosphate kinases. Pharmacol Ther. 2000; 87(2–3):189–98. Epub 2000/09/29. PMID: 11008000.

24. Van Rompay AR, Johansson M, Karlsson A. Substrate specificity and phosphorylation of antiviral and anticancer nucleoside analogues by human deoxyribonucleoside kinases and ribonucleoside kinases. Pharmacol Ther. 2003; 100(2):119–39. Epub 2003/11/12. PMID: 14609716.

25. Negishi K, Bessho T, Hayatsu H. Nucleoside and nucleobase analog mutagens. Mutation research. 1994; 318(3):227–38. Epub 1994/12/01. PMID: 7527486.
26. Agudo R, Arias A, Domingo E. 5-fluorouracil in lethal mutagenesis of foot-and-mouth disease virus. Future Med Chem. 2009; 1(3):529–39. Epub 2009/06/01. https://doi.org/10.4155/fmc.09.26 PMID: 21426129.

27. Arias A, Thorne L, Goodfellow I. Favipiravir elicits antiviral mutagenesis during virus replication in vivo. eLife. 2014; 3:e03679. Epub 2014/10/22. https://doi.org/10.7554/eLife.03679 PMID: 25333492; PubMed Central PMCID: PMC4204012.

28. Baranovich T, Wong SS, Armstrong J, Marjuki H, Weberry RG, et al. T-705 (favipiravir) induces lethal mutagenesis in influenza A H1N1 viruses in vitro. Journal of virology. 2013; 87(7):3741–51. Epub 2013/01/18. https://doi.org/10.1128/JVI.02346-12 PMID: 23325689; PubMed Central PMCID: PMC3624194.

29. Escribano-Romero E, Jimenez de Oya N, Domingo E, Saiz JC. Extinction of West Nile Virus by Favipiravir through Lethal Mutagenesis. Antimicrobial agents and chemotherapy. 2017; 61(11):e01400–17. https://doi.org/10.1128/AAC.01400-17 PMID: 28848019; PubMed Central PMCID: PMCPMC5655064.

30. Liu WJ, Aaskov JG. Fitness peaks of dengue virus populations. PloS one. 2018; 13(1):e0189554. doi: ARTN e0189554. PubMed PMID: WOS:000419101600022. https://doi.org/10.1371/journal.pone.0189554 PMID: 29293524.

31. Li W, Aaskov JG. Fitness peaks of dengue virus populations. PLoS one. 2018; 13(1):e0189554. doi: ARTN e0189554. PubMed PMID: WOS:000419101600022. https://doi.org/10.1371/journal.pone.0189554 PMID: 29293524.

32. de Koning H, Diallinas G. Nucleobase transporters (review). Mol Membr Biol. 2000; 17(2):75–94. Epub 2000/09/16. PMID: 10989458.

33. Griffith DA, Jarvis SM. Nucleoside and nucleobase transport systems of mammalian cells. Biochim Biophys Acta. 1996; 1286(3):153–81. Epub 1996/10/29. PMID: 8982282.

34. Koff WC, Elm JL Jr., Halsted SB. Antiviral effects if ribavirin and 6-mercaptop-9-tetrahydropyrimine against dengue viruses in vitro. Antimicrobial agents and chemotherapy. 2005; 49(3):981–6. https://doi.org/10.1128/AAC.49.3.981-986.2005 PMID: 15728892; PubMed Central PMCID: PMCPMC549233.

35. Markland W, McQuaid TJ, Jain J, Kwong AD. Broad-spectrum antiviral activity of the IMP dehydrogenase inhibitor VX-497: a comparison with ribavirin and demonstration of antiviral additivity with alpha interferon. Antimicrobial agents and chemotherapy. 2000; 44(4):859–66. PMID: 10722482; PubMed Central PMCID: PMCPMC89783.

36. Nakabayashi H, Takekita M, Miyano K, Yamane T, Sato J. Growth of human hepatoma cells lines with differentiated functions in chemically defined medium. Cancer research. 1982; 42(9):3858–63. Epub 1982/09/01. PMID: 6286115.

37. Whitby K, Pierson TC, Geiss B, Lane K, Engle M, Zhou Y, et al. Castanospermine, a potent inhibitor of dengue virus infection in vitro and in vivo. Journal of virology. 2005; 79(14):8698–706. Epub 2005/07/05. https://doi.org/10.1128/JVI.79.14.8698-8706.2005 PMID: 15994763; PubMed Central PMCID: PMCPMC1168722.

38. Gregori J, Perales C, Rodriguez-Frias F, Esteban JL, Quer J, Domingo E. Viral quasispecies complexity measures. Virology. 2016; 493:227–37. Epub 2016/04/10. https://doi.org/10.1016/j.virol.2016.03.017 PMID: 27080566.

39. Sinha SC, Smith JL. The PRT protein family. Curr Opin Struct Biol. 2001; 11(6):733–9. Epub 2001/12/26. PMID: 11751055.

40. Naesens L, Guddat LW, Keough DT, van Kuilenburg AB, Meijer J, Vande Voorde J, et al. Role of human hypoxanthine guanine phosphoribosyltransferase in activation of the antiviral agent T-705 (favipiravir). Mol Pharmacol. 2013; 84(4):615–29. Epub 2013/08/03. https://doi.org/10.1121/mol.113.087247 PMID: 23907213.

41. Vernekar SK, Liu L, Zhang J, Kankanala J, Li H, Geraghty RJ, et al. 5'-Silylated 3'-1,2,3-triazolyl Thymidine Analogues as Inhibitors of West Nile Virus and Dengue Virus. Journal of medicinal chemistry. 2015; 58(9):4016–28. Epub 2015/04/23. https://doi.org/10.1021/jm500327w PMID: 25909386; PubMed Central PMCID: PMC4465584.

42. de Avila AI, Gallego I, Soria ME, Gregori J, Quer J, Esteban JL, et al. Lethal Mutagenesis of Hepatitis C Virus Induced by Favipiravir. PLoS one. 2016; 11(10):e0164691. Epub 2016/10/19. https://doi.org/10.1371/journal.pone.0164691 PMID: 27755573; PubMed Central PMCID: PMC5068794.
45. Grande-Perez A, Gomez-Mariano G, Lowenstein PR, Domingo E. Mutagenesis-induced, large fitness variations with an invariant arenavirus consensus genomic nucleotide sequence. Journal of virology. 2005; 79(16):10451–9. Epub 2005/07/30. https://doi.org/10.1128/JVI.79.16.10451-10459.2005 PMID: 16051837; PubMed Central PMCID: PMCPMC1182645.

46. Pariente N, Airaksinen A, Domingo E. Mutagenesis versus inhibition in the efficiency of extinction of foot-and-mouth disease virus. Journal of virology. 2003; 77(12):7131–8. https://doi.org/10.1128/JVI.77.12.7131-7138.2003 PMID: 12768034; PubMed Central PMCID: PMCPMC156209.

47. Guirakhoo F, Hunt AR, Lewis JG, Roehrig JT. Selection and partial characterization of dengue 2 virus mutants that induce fusion at elevated pH. Virology. 1993; 194(1):219–23. Epub 1993/05/01. https://doi.org/10.1006/viro.1993.1252 PMID: 8480420.

48. Pierro DJ, Salazar MI, Beaty BJ, Olson KE. Infectious clone construction of dengue virus type 2, strain Jamaican 1409, and characterization of a conditional E6 mutation. The Journal of general virology. 2006; 87(Pt 8):2263–8. Epub 2006/07/19. https://doi.org/10.1099/vir.0.81958-0 PMID: 16847122.

49. Bonnac LF, Mansky LM, Patterson SE. Structure-activity relationships and design of viral mutagens and application to lethal mutagenesis. Journal of medicinal chemistry. 2013; 56(23):9403–14. Epub 2013/08/07. https://doi.org/10.1021/jm400653j PMID: 23914828.

50. Jin Z, Smith LK, Rajwanshi VK, Kim B, Deval J. The ambiguous base-pairing and high substrate efficiency of T-705 (Favipiravir) Ribofuranosyl 5’-triphosphate towards influenza A virus polymerase. PloS one. 2013; 8(7):e68347. Epub 2013/07/23. https://doi.org/10.1371/journal.pone.0068347 PMID: 23874596; PubMed Central PMCID: PMCPMC3707847.