Comparison of Binding Site of Remdesivir and Its Metabolites with NSP12-NSP7-NSP8, and NSP3 of SARS CoV-2 Virus and Alternative Potential Drugs for COVID-19 Treatment

Lindsey S Jung1 · Tamara M Gund2 · Mahesh Narayan3

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Abstract
Remdesivir was approved by the U.S.A. Food and Drug administration for emergency use to interfere with the replication of SARS CoV-2 virus (the agent that causes COVID-19) in adults and children hospitalized with severe disease. The crystal structure of the metabolite of remdesivir (Monophosphate of GS-441524) and NSP12-NSP8-NSP7 of SARS CoV-2 virus was recently reported. The crystal structures of ADP-Ribose or AMP and NSP3 of SARS CoV-2 virus were also released, recently. This study compared their binding sites and suggests the crystal structure of NSP3 of SARS CoV-2 virus as an alternative binding site of AMP or ADP-ribose to treat COVID-19. We virtually screened 682 FDA-approved compounds, and the top 10 compounds were selected by analysis of docking scores, (G-score, D-score, and Chemscore) and visual analysis using a structure-based docking approach of NSP3 of SARS CoV-2 virus. All immunization approaches are based on the SARS-CoV-2 virus spike protein. A recent study reported that the D614G mutation in the SARS-CoV-2 virus spike protein reduces S1 shedding and increases infectivity of SARS COV-2 virus. Therefore, if there is a severe change in the spike protein of a modified Coronavirus, all developed vaccines can lose their efficacy, necessitating the need for an alternative treatment method. The top 10 compounds (FDA-approved) in this study are selected based on NSP 3 binding site, and therefore are a potential viable treatment because they will show potential activity for all mutations in the SARS-CoV-2 virus spike protein.

Keywords Remdesivir · Non-structural protein 3 (NSP3) of SARS CoV-2 virus · Virtual screening

1 Introduction
Coronaviruses (CoV) were first identified in the 1960s as human pathogens [1]. A new virus, known as the severe acute respiratory syndrome coronavirus-2 (SARS CoV-2 virus) arose in China, rapidly instigating the COVID-19 pandemic. Remdesivir is an antiviral medication developed by Gilead Sciences to treat hepatitis C and was also tested as a potential treatment for Ebola and Coronavirus [2, 3]. Although remdesivir did not originally target SARS CoV-2 virus, remdesivir began to be tested in laboratory by Gilead Sciences in January 2020 and was stated to be active against severe acute respiratory syndrome coronavirus (SARS) and Middle East respiratory syndrome (MERS) in animal models. In 2020, the U.S. Food and Drug Administration issued an emergency use authorization for the investigational antiviral drug remdesivir for the treatment of suspected or laboratory-confirmed SARS CoV-2 virus in adults and children hospitalized with severe disease. The binding mode of remdesivir in SARS CoV-2 virus is not clear, but structures of remdesivir and its parent drug, GS-441524 (Figs. 1 and 2) help understand how they bind in SARS CoV-2 virus and are effective as an antiviral [4]. Recently, Yin reported [5] a structural basis for the inhibition of the RNA-dependent RNA polymerase from SARS-CoV-2 virus by remdesivir.
The actual binding site of the metabolite of remdesivir (the complex of diphosphate-Mg- monophosphate of GS-441524) is disclosed by Yin, and many crystal structures are released for the AMP (Adenosine monophosphate) or ADP-ribose (Adenosine-5-diphosphoribose) binding site of NSP3 [6, 7]. We observe that the structures of AMP (complex of NSP3)
and monophosphate of GS-441524 (complex of NSP12) are very similar. The size of ADP-ribose (complex of NSP3) and the triphosphate of GS-441524 (the active metabolite of remdesivir) [4] are also very similar (ATP-like in Fig. 2); we hypothesize that remdesivir will potentially bind to NSP3 and NSP12-NSP8-NSP7.

It was reported that the D614G mutation reduces S1 shedding and increases infectivity [8]. All immunization approaches against SARS CoV-2 virus are currently based on this spike protein. Therefore, if there is a severe change in the spike protein of a modified Coronavirus, there is a possibility that the developed vaccines will lose their efficacy. While there is no direct evidence of the effect of the G614 mutant on developing vaccines, this mutation serves as an example of possible future mutations that may cause developing vaccines to lose their efficacy. Unlike immunization approaches, the approach of developing a drug through remdesivir with its NSP binding sites is a promising choice to treat modified SARS CoV-2 viruses.

Clinical trials of remdesivir are still progress, rendering the possible side effects of remdesivir to be unknown. Therefore, in addition to remdesivir, this study searches for alternative treatment against mutations in the spike protein. We perform virtual screening of 682 FDA approved compounds based on the binding modes of remdesivir with NSP3. The top 10 compounds from virtual screening are selected as potential treatments for Coronavirus. We search for additional alternatives in hope that the compounds could potentially test to be more active against Coronavirus in the future.

2 Material and Methods

2.1 General Methods

Visual analysis was performed by observing the hydrogen bonding, molecular surface (MOLCAD) of cavity depth, and lipophilic and electrostatic potentials. The binding cavity was based on each crystal structure (6W6Y, 6WOJ, 6 M71, 7BV2). Superimposition was calculated in “match” function. A commercially available 682 FDA approved data set from Enamine was used for virtual screening through the Surflex-Docking (GEOM) setting. Scoring functions were calculated by CSCORE. All calculations and simulations were done using SYBYL-X2.1 [9].

2.2 Binding Site of NSP3 and NSP12-NSP8-NSP7 of SARS CoV-2 Virus

Two crystal structures were selected for NSP3: AMP (PDB: 6W6Y) and ADP-ribose (PDB: 6WOJ). Using SYBYL-X2.1 [9], structures of the ligand complex were identified, and H-bonds were analyzed. In addition, the lipophilic and electrostatic interactions were validated for each complex. Two crystal structures (PDB: 6W6Y, 6WOJ) were superimposed and analyzed based on the binding site of AMP and ADP-ribose [6, 7]. An apo structure [10] and a bound structure with the metabolite of remdesivir (monophosphate of GS-441524) [5] were available for NSP12-NSP8-NSP7. Both structures (PDB: 6 M71, 7BV2) were superimposed and compared in the binding site by SYBYL-X2.1.

2.3 Validation of Previously Reported Experimental Data for NSP3 or NSP12 of Coronavirus

This study targets ADP-ribose as a potential binding site as a result of Frick’s report [7] of the molecular basis ADP-ribose binding to the Mac1 domain of SARS CoV-2 virus, NSP3. It was suggested that NSP3 binds to both SARS-CoV virus and SARS-CoV-2 virus, despite the two amino acid sequences varying by 26%. When testing the ADP-ribose binding in the two Coronaviruses, it binds to SARS-CoV-2 virus protein more tightly (Kd = 10 μM) than the SARS-CoV virus protein (Kd = 24 μM). Thus, he encourages the use of ADP-ribose to design antiviral drugs targeting this NSP3 region in SARS-CoV-2 virus. This study superimposes the crystal structures of SARS-CoV [11] and SARS CoV-2 viruses using SYBYL-X2.1 to explain the different binding affinities of ADP-ribose to both Coronaviruses. Tchesnokov reported [12] that as shown through steady-state kinetic parameters, the efficiency of ATP incorporation decreases threefold when with the mutant enzyme. To validate these experimental data, the crystal structure of NSP12 (wild type) and V557L were generated and superimposed by SYBYL-X2.1.

2.4 Selected Binding Sites of Virtual Screening in This Study

The crystal structure of NSP3 with ADP-ribose (PDB: 6WOJ) was selected to search potential inhibitors for SARS CoV-2 virus using virtual screening of FDA approved 682 drugs (commercially available from Enamine).

2.5 Selection of Reference Compounds and Top 10 Compounds

Reference compounds are needed for post-docking analysis to select potential compounds for alternative treatments of COVID-19 (via the etiological SARS CoV-2 virus). The reference compounds consisted of structures of ATP, ADP, and AMP-like (size) compounds (all ten compounds from Figs. 1 and 2), along with remdesivir and GS-441524. GS-441524 was added because Eastman and et al. reported how remdesivir is metabolized to GS-441524 [4]. 682
FDA approved compounds were prepared and minimized in MMFF94s (Force Fields) and MMFF94 (Charges) and docked to the NSP3 binding site of the SARS CoV-2 virus crystal structure (PDB: 6WOJ). Scoring functions have been used to estimate and predict which compounds would have the best actual binding affinities in an experimental test. Surflex-score, G-score, D-score, and Chemscore were scoring functions used to predict the best compounds using virtual screening [13–19]. Although scoring functions are reliable mathematical functions, there is still a possibility for a protein and ligand to clash and protrude [9]. Therefore, visual analysis is crucial in predicting the best potential compounds. Every docked compound was visually analyzed, and compounds with protruding ligands and proteins were eliminated. The scoring functions in this study were compared and analyzed using SYBYL-X2.1 [9] (Table 1).

### 3 Results

#### 3.1 Binding Site of NSP3, NSP12-NSP8-NSP7

Figures 3 and 4 each display hydrophilic interactions of AMP and ADP-ribose with their hydrogen bonds and NSP3 binding site. Figures 3 and 4 also display that the size of the NSP3 binding site cavity fits well with the AMP and ADP-ribose complex ligands in each crystal structure. The lipophilic and electrostatic molecular surfaces are shown in Figs. 5 and 6 for AMP and ADP-ribose with NSP3 of SARS CoV-2 virus respectively. Both crystal structures (PDB: 6W6Y, 6WOJ) were superimposed and the binding cavities were compared (Fig. 7). Two crystal structures of NSP12-NSP8-NSP7 were superimposed and analyzed (Fig. 8); one is an apo structure [8] (PDB: 6M71) and another is a bound structure (PDB: 7BV2) with a metabolite of remdesivir (monophosphate of GS-441524) [5].

| Table 1 Scoring functions to select top 10 compounds in this study [9–15] |
| Scoring functions | Descriptions |
|-------------------|-------------|
| **General scoring** [9] | A linear empirical scoring function can be written as a sum of independent terms such as 
\[
\Delta G_{\text{bind}} = c_0 + c_1 \Delta G_{\text{vdw}} + c_2 \Delta G_{\text{hbond}} + c_3 \Delta G_{\text{entropy}}
\]
where \(c_i\) is the weighting coefficients of the respective \(\Delta G_{\text{bind}}\) terms, adjusted to reproduce affinity data based on the training set. In the example, \(\Delta G_{\text{vdw}}\) is a Van der Waals potential, \(\Delta G_{\text{hbond}}\) is a specific term accounting for hydrogen bonds, and \(\Delta G_{\text{entropy}}\) is related to the ligand entropic loss upon binding |
| **Surflex-score** [11] | Surflex scoring function includes an entropic penalty term that is linear in the number of rotatable bonds in the ligand, intended to model the entropic cost of fixation of these bonds, and a term that is linearly related to the log of the molecular weight of the ligand, intended to generate putative poses of ligand fragments |
| **G-score** [12] | This scoring function is from GOLD program. It is basically a force field-based scoring function, which consists of terms for the hydrogen bonds and van der Waals interactions between protein and ligand and the internal steric energies of the ligand |
| **PMF-score** [13] | This knowledge-based scoring function was developed by Muegge et al. It sums up pairwise knowledge-based interaction potentials between protein and ligand |
| **D-score** [14] | This scoring function is adopted by the DOCK program by Kuntz et al., including Van der Waals and electrostatic interactions between protein and ligand |
| **Chemscore** [15] | This empirical scoring function is based on the work of Eldridge et al. which includes terms for hydrogen bonds, metal-ligand interactions, lipophilic contacts, and conformational entropies |

![Fig. 3](image-url)  
Fig. 3 Analysis Molecular Surfaces AMP with crystal structure of NSP3 (PDB: 6W6Y) (a) Hydrogen Bond between NSP3 and AMP, (b) Molecular Electrical Potential Surfaces of AMP, and (c) Molecular Surface displayed by Cavity Depth for NSP3
Fig. 4 Analysis Molecular Surfaces ADP-ribose with crystal structure of NSP3 (PDB: 6WOJ) (a) Hydrogen Bond between NSP3 and ADP-ribose, (b) Molecular Electrical Potential Surfaces of ADP-ribose, and (c) Molecular Surface displayed by Cavity Depth for NSP3

Fig. 5 Analysis Molecular Surfaces AMP with crystal structure of NSP3 (PDB: 6W6Y) (a) Molecular Lipophilic Potential Surfaces of NSP3 and AMP, (b) Molecular Electrical Potential Surfaces of NSP3 and AMP

Fig. 6 Analysis Molecular Surfaces ADP-ribose with crystal structure of NSP3 (PDB: 6WOJ) (a) Molecular Lipophilic Potential Surfaces of NSP3 and ADP-ribose, (b) Molecular Electrical Potential Surfaces of NSP3 and ADP-ribose
Fig. 7 Superimposed NSP3 of SARS COV-2 with AMP and ADP-ribose (PDB: 6WOJ) (a) Molecular Surface by Cavity Depth for NSP3 with AMP (PDB: 6W6Y), (b) Superimposed NSP3 with AMP and ADP-ribose (PDB: 6W6Y, 6WOJ), (c) Molecular Surface by Cavity Depth for NSP3 with ADP-ribose (PDB: 6WOJ)

Fig. 8 Superimposed NSP12 of SARS COV-2 with Monophosphate of GS-441524; (a) Molecular Surface of NSP12 (PDB: 7BV2), (b) Superimposed NSP12 (PDB: 7BV2, 6M71), (c) Molecular Surface of NSP12 (PDB: 6M71)

Fig. 9 (a) Superimposition, and (b) Sequence Alignment of NSP3 in SARS COV-2 (purple, PDBL 6WOJ) and SARS-CoV (orange, PDB: 2FAV) (Color figure online)
3.2 Validation of Previously Reported Experimental Data for NSP3 and NSP12 of Coronavirus

The crystal structures of SARS-CoV [11] and SARS-CoV-2 viruses were superimposed (PDB id: 6WOJ, 2FAV) in Fig. 9; sequence alignment of two crystal structures of SARS-CoV and SARS-CoV-2 viruses was 73.2% in 168 residues. However, in active sites, they were nearly identical, with the exception in only one amino acid of PHE156 for SARS-CoV-2, and ASN157 for SARS-CoV virus. Adenosine showed hydrophobic interactions with the phenyl group in PHE156 and had no hydrophobic interactions with ASN157 for SARS-CoV (Fig. 9). It explained why the SARS-CoV-2 virus protein binds to ADP-ribose somewhat more tightly (Kd = 10 μM) than the SARS-CoV virus protein (Kd = 24 μM) [7]. The crystal structure of NSP12 (wild type) and its mutation of V557L were generated and superimposed in Fig. 10. The complex of NSP12 and metabolite of remdesivir displayed that it metabolized in monophosphate of GS-441524, bonding with magnesium with diphosphate in Fig. 10. Overall, the size of the structure looks like the main active metabolite of remdesivir, triphosphate of GS-441524, (compound d in Fig. 1) reported by Eastman and Gordon [4, 20]. VAL557 is located in between ARG555 and U10 (Uridine phosphate); the metabolite of remdesivir has H-bonds with ARG555 (NSP12) and U10 (UTP), explaining the counteractive nature of the V557L mutation, in regard to the inhibitory effects of remdesivir; the threefold reduction in the efficiency of ATP incorporation with the mutant enzyme [12]. The H-bond interaction was lost due the H-bond distance between pyrrolo[2,1-f][1,2,4]triazine, and ARG555 increasing from 2.72 to 3.13 Armstrong (from purple 557 Valine to orange 557 Leucine) in Fig. 10.

Fig. 10 Superimposition of the crystal structure of the metabolite of Remdesivir (PDB: 7BV2) in wild type (purple) and V557L (orange) with UTP (Uridine phosphate, green) (Color figure online)

Table 2 Top 10 candidates with selected ten reference compounds in different Scoring Functions; ID (compounds a–h from Fig. 2, remdesivir, and GS-441524 from Fig. 1), All ID is from Enamine database. Surflex (Surflex Score), SO (Order of Surflex-score from all 692 compounds; references compounds a–h in Fig. 2, Remdesivir, GS-441524, and FDA approved 682 compounds from Enamine), GOLD (G-Score), GO (Order of Gold-score from all 692 compounds), PMF (PMF-Score), PO (Order of PMF-score from all 692 compounds), DOCK (D-Score), DO (Order of D-score from all 692 compounds), CHEMS (Chemscore), CO (Order of Chemscore from all 692 compounds), SGPD (Sum of SO, GO, PO, and DO)

| ID    | Surflex | SO   | GOLD | GO  | PMF   | PO   | DO    | CHEMS  | CO | SGPD |
|-------|---------|------|------|-----|-------|------|-------|--------|----|------|
| d     | 15.91   | 1    | −397.75 | 8   | −48.28 | 15   | −198.07 | 7     | −11.26 | 621  | 31   |
| remdesivir | 11.84  | 4    | −404.94 | 6   | −32.20 | 49   | −236.21 | 1     | −28.58 | 127  | 60   |
| a     | 12.71   | 2    | −339.33 | 34  | −53.00 | 9    | −164.38 | 54    | −8.37  | 653  | 99   |
| g     | 8.82    | 42   | −343.20 | 26  | −64.14 | 4    | −173.29 | 35    | −7.18  | 663  | 107  |
| h     | 8.71    | 49   | −318.54 | 50  | −55.7  | 7    | −169.56 | 40    | −10.25 | 632  | 146  |
| b     | 10.16   | 13   | −258.21 | 132 | −64.65 | 34   | −136.85 | 149   | −10.21 | 633  | 297  |
| e     | 8.38    | 62   | −294.63 | 80  | −12.89 | 132  | −168.14 | 44    | −23.73 | 282  | 183  |
| c     | 8.47    | 58   | −221.29 | 241 | −37.78 | 35   | −117.39 | 241   | −7.81  | 655  | 574  |
| f     | 6.57    | 225  | −241.91 | 178 | −25.34 | 86   | −133.51 | 149   | −16.52 | 528  | 657  |
| GS-441524 | 7.09   | 158  | −167.95 | 447 | −39.38 | 30   | −114.53 | 262   | −16.58 | 527  | 897  |
| BRC1213 (1)| 10.63  | 9    | −283.71 | 99  | −38.25 | 31   | −168.14 | 44    | −23.73 | 282  | 183  |
| BRC0679 (2)| 8.06   | 81   | −320.23 | 39  | −27.13 | 66   | −181.62 | 25    | −38.12 | 18   | 211  |
| BRC0089 (3)| 9.54   | 23   | −321.76 | 47  | −20.11 | 119  | −161.91 | 60    | −23.73 | 281  | 249  |
| BRC0206 (4)| 10.31  | 12   | −374.95 | 15  | −3.80  | 304  | −202.36 | 5     | −28.56 | 128  | 336  |
| BRC0332 (5)| 7.56   | 126  | −295.68 | 76  | −22.78 | 104  | −196.57 | 30    | −34.34 | 38   | 336  |
| BRC0205 (6)| 8.39   | 61   | −263.83 | 126 | −27.78 | 59   | −147.56 | 105   | −25.65 | 218  | 351  |
| BRC0901 (7)| 8.65   | 51   | −340.55 | 32  | −8.52  | 255  | −185.98 | 19    | −35.77 | 25   | 357  |
| BRC0853 (8)| 8.89   | 38   | −249.50 | 155 | −32.91 | 47   | −144.79 | 120   | −35.13 | 29   | 360  |
| BRC0377 (9)| 7.46   | 130  | −266.64 | 122 | −37.35 | 35   | −155.35 | 77    | −29.57 | 101  | 364  |
| BRC1908 (10)| 7.62  | 120  | −297.43 | 74  | −25.32 | 87   | −152.27 | 88    | −23.60 | 287  | 369  |
3.3 Top 10 Compounds from Virtual Screening of FDA Approved 682 Drugs to NSP3 Binding Site of SARS CoV-2 Virus

Table 2 shows how reference compounds were listed and how the top 10 compounds were selected based on scoring functions [13–19]. The top 10 compounds, remdesivir, and its parent compound (GS-441524) were each displayed in a docking pose in the cavity depth of molecular surfaces (Fig. 11) and a zoomed out figures of the top 9 compounds were displayed in active sites of amino acids, generating H-binds (Fig. 12) to explain where the ligands are binding relative to the natural substrate (Table 3).

4 Discussion

From analyzing the crystal structure of NSP12-NSP7-NSP8, we discovered that the main interaction of remdesivir’s metabolite (monophosphate of GS-441524) was with NSP12. This binding site was superimposed using two available NSP12-NSP7-NSP8 crystal structures. The two structures differ greatly, as one is an apo structure and other is a bound structure, and both are extremely flexible depending on the conditions (Fig. 8). Contrastingly, the superimposed ADP-ribose and AMP (NSP3) binding sites (PDB: 6W6Y, 6WOJ) are visually consistent and stable. Therefore, the NSP3 binding site was selected to virtually screen 692 compounds (10 reference compounds in Table 2 and FDA approved 682 compounds). In this study, the docking scores revealed vitamin B9, Folic acid, as the number one candidate to treat COVID-19 (via SARS CoV-2 virus replication inhibition). The docking pose arranged well with NSP3, and the binding pose looked considerably similar with remdesivir and its metabolite (Fig. 13); Both compounds had Hydrogen bonds with ASP22, PHE132 with NSP3. A strong salt bridge interaction (aspartic acid and amine) in ASP22 was formed for both compounds. It has been reported that maternal folic acid supplementation is commonly used to prevent neural tube defects (NTDs). In addition, it may play a role in preventing pregnancy complications [21–23] and increasing anti-HBs titers in the primary response, together with the fact that antibody levels after primary vaccination are strongly associated with the persistence of the protective antibody [24]. Recently Serseg reported that Hispidin and Lepidine E: two natural compounds and folic acid as potential inhibitors of 2019-novel coronavirus main protease (2019-nCoVMpro) using molecular docking and SAR study [25]. It was noted that folic acid is able to inhibit furin, preventing binding by the SARS-CoV-2 virus spike protein, preventing cell entry and virus turnover [26]. In this study by the structure-based approach of NSP3 binding domain, folic acid is a promising candidate to treat and prevent COVID-19 infections. Therefore, folic acid may be a potential treatment for the management of COVID-19.
Fig. 12 Docking poses in the cavity of NSP3 for top 9 compounds with active sites, generating H-bonds: (a) (top 1) Folic acid, (b) (top 2) Telmisartann, (c) (top 3) Methotrexate, (d) (top 4) Bosentan, (e) (top 5) Lapatinib, (f) (top 6) Gefitinib, (g) (top 7) Ketoconazole, (h) (top 8) Carvedilol, (i) (top 9) Glyburide
| ID          | Name             | Therapeutic agent                  | Structures |
|-------------|------------------|------------------------------------|------------|
| BRC1213     | Folic acid (Vit. B9) | Dietary supplements, Hematologic, vitamins | ![Structure Image] |
| BRC0679     | Telmisartan      | Antihypertensive                   | ![Structure Image] |
| BRC0089     | Methotrexate     | Antineoplastic                     | ![Structure Image] |
| BRC0206     | Bosentan         | Antihypertensive                   | ![Structure Image] |
| BRC0332     | Lapatinib        | Anticancer                         | ![Structure Image] |
| BRC0205     | Gefitinib        | Antineoplastic, BCRP/ABCG2 inhibitors | ![Structure Image] |
| BRC0901     | Ketoconazole     | Anti-infective                      | ![Structure Image] |
### Table 3 (continued)

| ID            | Name       | Therapeutic agent                      | Structures |
|---------------|------------|----------------------------------------|------------|
| BRC0853 (top 8) | Carvedilol | Antihypertensive vasodilator agent     | ![Carvedilol Structure] |
| BRC0377 (top 9) | Glyburide  | Antihyperglycemic                      | ![Glyburide Structure] |
| BRC1908 (top 10) | Avanafil   | Urologicals Vasodilating               | ![Avanafil Structure] |

**Fig. 13** Docking poses of Remdesivir and Folic Acid (top 1) with NSP3 of SARS COV-2; (a) H-bonds of NSP3 with Remdesivir (green) and Folic acid, (b) The cavity of NSP3 with Remdesivir (green) and Folic acid (Color figure online)
Author Contributions LSJ, TMG, and MN conceptualized and designed the experiments and analyzed the data. MN reviewed the data and manuscript.

Compliance with Ethical Standards

Conflict of interest The authors declare no competing financial interests.

References

1. Killery ME, Biggs HM, Haynes A, Dahl RM, Mustaquim D, Gerber SI, Watson JT (2018) Human coronavirus circulation in the United States 2014–2017. J Clin Virol 101:52–56
2. Stephens B (2020) The story of remdesivir. The New York Times https://www.nytimes.com/2020/04/17/opinion/remdesivir-coronavirus.html. Accessed 21 Sept 2020
3. Warren TK, Jordan R, Lo MK, Ray AS, Mackman RL, Soloveva A, Gruez A, Campanacci V, Cambillau C, Ziebuhr J, Ahola T, Canard B (2006) Structural and functional basis for ADP-ribose and poly(ADP-ribose) binding by viral macro domains. J Virol 80:8493–8502
4. Tchesnokov EP, Gordon CJ, Woolner E, Kocincova D, Perry JK, Feng JY, Porter DP, Götte M (2020) Template-dependent inhibition of coronavirus RNA-dependent RNA polymerase by remdesivir reveals a second mechanism of action. J Biol Chem. jbc.AC120.015720. https://doi.org/10.1074/jbc.AC120.015720
5. Guedes IA, Pereira FSS, Dardene LE (2018) Empirical scoring functions for structure-based virtual screening: applications, critical aspects, and challenges. Front Pharmacol 9:1089–1106
6. Li Y, Han L, Liu Z, Wang R (2014) Comparative assessment of scoring functions on an updated benchmark: 2. Evaluation methods and general results. J Chem Inf Model 54:1717–1736
7. Jain AN (2003) Surflex: fully automatic flexible molecular docking using a molecular similarity-based search engine. J Med Chem 46:499–511
8. Moeij WT, Verdonk ML (2005) General and targeted statistical potentials for protein-ligand interactions. Proteins 61:272–287
9. Fernández A (2020) Structural impact of mutation D614G in SARS-CoV-2 spike protein: enhanced infectivity and therapeutic opportunity. ACS Med Chem Lett 11:1667–1670
10. Fernández A (2020) Structural impact of mutation D614G in SARS-CoV-2 spike protein: enhanced infectivity and therapeutic opportunity. ACS Med Chem Lett 11:1667–1670
11. Schonberger K, Riedel C, Mansmann U, Jilg W, Scherf M, Elstner E, Stähler S0378-5122(20):30348-0. https://doi.org/10.1016/j.maturitas.2020.08.007