

1 ***In silico* analyses on the comparative sensing of SARS-CoV-2 mRNA by**  
2 **intracellular TLRs of human**

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## 1 **1. Introduction**

2           The worldwide outbreak of Coronavirus disease or COVID-19 pandemic caused by the  
3 Severe Acute Respiratory Syndrome Coronavirus-2 (SARS-CoV-2), leads to the infection of  
4 about 0.33% of the total world's population causing the death of 1.24 million people by the  
5 first week of November 2020.<sup>1</sup> The continuous expansion of contagion and pathogenesis, it is  
6 taking the shape of another dark age in the history of mankind, not only to health crises but  
7 also in bankrupting the global socio-economic status. The SARS-CoV-2 belongs to the  $\beta$ -  
8 Coronavirus genus of a 2B group of the Coronaviridae family and is considered the third most  
9 virulent type, leading to the highest fatality rate in humans followed by the SARS-CoV and  
10 MERS-CoV.<sup>2</sup> It transmits from person to person mainly via close physical contact and by  
11 respiratory aerosols that produces during coughing, sneezing, and even talking within  
12 proximity, although some recent studies indicate transmission through fecal matters and  
13 fomite-borne contaminations.<sup>3-5</sup>

14           The proteome of the virus consists of structural proteins, existing in three forms – the  
15 Spike 'S' protein, the Envelope 'E' protein and the Membrane 'M' protein. Along with these,  
16 16 forms of non-structural proteins (NSPs) combining together to generate different catalytic  
17 models.<sup>6</sup> The genome, however, is much simpler and comprises of a 29,903 base long, positive-  
18 sense single-stranded RNA molecule. This (+)ssRNA genome makes it further feasible to be  
19 detectible by the intracellular Toll-like receptors (TLRs) which have a high affinity towards  
20 nucleic acid-related pathogen-associated molecular patterns (PAMPs).<sup>7</sup> Several empirical  
21 studies have presented various propositions relating the intracellular TLR 3, 7, 8, and 9 to the  
22 cytokine storm produced by the virus that majorly owes it the lethality.<sup>8</sup> Cytokine storm is  
23 apparently the incessant extreme activation of cytokine production leading to prolonged and  
24 consistent inflammatory response, which becomes almost continuous due to positive feedback

1 loops in the TLR signaling pathways. The studies are quite successful in relating the role of  
2 TLRs in recognizing oligonucleotide PAMPs and triggering the cytokine storm.<sup>9</sup>

3         The binding of Spike protein with the human ACE2 receptor triggers the pathogenesis  
4 of the SARS-CoV-2, leading to the activation of TLRs to activate the proliferation and  
5 production of pro-inflammatory cytokines causing cytokine storm, those results in  
6 inflammations.<sup>10,11</sup> From previous studies, it has been found that the spike protein shows  
7 binding efficiency with the extracellular domains of TLRs including TLR1, TLR4 and, TLR6,  
8 with the strongest affinity with TLR4.<sup>12</sup> Furthermore, the development of several *in-silico*  
9 multi-epitope-based peptide vaccine candidates against the SARS-CoV-2 has shown to be  
10 effectively binding with TLR3, TLR4 and, TLR5 to regulate the TLR signaling pathways  
11 activation and proliferation.<sup>13</sup> It has been found that targeting human TLRs, in an order to either  
12 block the binding of SARS-CoV-2 or by inhibiting the TLR activation and proliferation that  
13 induce the production of pro-inflammatory cytokines using certain TLR agonists, might be  
14 used as an effective therapeutic strategy against coronavirus disease.<sup>11</sup> Additionally, the  
15 continuous replication and generation of new virus within the host system indicates that there  
16 must be binding efficiency and potency with the intracellular TLRs including TLR3, 7, 8, and  
17 9 in order to induce the severity and pathogenesis of the virus and further introduction to the  
18 new host. TLR3 is well known for its sensing capabilities of viral PAMPs and exists as a  
19 monomer that is attached to the membrane of endosomes, thereby it detects and binds to certain  
20 motifs in the invading viral RNA.<sup>14</sup> TLR7 expressed majorly in the cerebral cortex, lung,  
21 bronchus, breast, kidney, rectum, and smooth muscle tissues, and functions by adhering to the  
22 endosome and thereby binding to the viral RNAs with high guanosine and uridine content <sup>15</sup> to  
23 initiates a MyD88 signal transduction resulting in activation of NF-κB, mitogen-activated  
24 protein kinase (MAPK) cascades, as well as IRF-7 and IRF-5 activation via IL-1 receptor-  
25 associated kinases (IRAK)-1/2/4 and TNF receptor-associated factor-3/6.<sup>16</sup> The signaling

1 finally induces the production of pro-inflammatory cytokines including IL-1 $\beta$ , IL-6, IL-12,  
2 TNF- $\alpha$ , and IFN- $\alpha$ .<sup>17</sup> TLR8 is expressed predominantly in the lung and the peripheral blood  
3 leucocytes and plays a major role in recognizing GU-rich viral ssRNAs including those of  
4 SARS-CoV-2.<sup>18</sup> TLR9 is predominantly expressed in the spleen, lymph node, bone marrow,  
5 and peripheral blood leukocytes and it recognizes CpG motifs in viral DNAs. However, several  
6 studies suggest its role in sensing ssRNA fragments generated by the SARS-CoV-2 genome.<sup>19</sup>

7 However, a lack of precise knowledge regarding the nature of oligonucleotides and  
8 their ligating affinity towards the TLRs still pertains to exist. This in context is abstaining the  
9 medical research community from developing certain therapeutic interventions that would have  
10 been vitally important in this hour of severity. Our study is hoped to rationalize the picture and  
11 provide clues regarding the interaction range of the oligonucleotides towards the intracellular  
12 TLRs, considering the solvent-based force-fields operational in the cytosolic aqueous  
13 microenvironment that predominantly drive these reactions.

## 14 **1. Materials and Methods**

### 15 **2.1 Data mining**

16 Current literatures suggest the predominant expression of ten proteins of SARS-CoV-2.<sup>20</sup> Four  
17 of them are structural proteins – the spike protein subunits S1 and S2, the envelope protein (E),  
18 and the membrane protein (M). While the other six are non-structural proteins viz. NSP7,  
19 NSP8, NSP9, NSP10, papain-like protease (PLpro), and main protease (Mpro). So, first of all,  
20 full-length RNA sequences of the aforementioned proteins were retrieved by extensive  
21 literature research. Further, the respective sequences were subjected to pairwise alignment with  
22 the complete genome of the virus (Accession No. MT438755) available at the GenBank  
23 database of NCBI showing ~100% identity matches and 0 gap penalties. From this, it was  
24 evident that the quality of the sequences obtained was very high and equivalent to the raw

1 original. Thereafter, crystal structures of TLR3 (PDB ID: 1ZIW) and TLR8 (PDB ID: 5WYX)  
2 were obtained from the RCSB PDB database. However, the structures of TLR7 and TLR9 were  
3 obtained by performing homology modelling in the SWISS-MODEL server from ExPASy  
4 (<https://swissmodel.expasy.org/>) by using protein sequences obtained from GenBank with  
5 Accession No. AAZ99026 and AAZ95518.1 respectively. All the crystal surface structures of  
6 TLRs and the whole genome of SARS-CoV-2 were then prepared to visualize (Figure 1).  
7 To perform further studies, PyMOL is used for removing water molecules, aboriginal  
8 heteroatoms as well as any xenobiotic ligands whenever present, and adding polar hydrogens  
9 and Kollman charges to the structures.

## 10 **2.2 Prediction of mRNA-Protein interactions**

11 The retrieved RNA sequences were subjected to the imRNA tool developed by IIIT-Delhi  
12 (<https://webs.iitd.edu.in/raghava/imrna/>). This tool is based on Motif—Emerging and with  
13 Classes—Identification (MERCIC) program and scans through the RNA sequences for motifs  
14 that are potent to have immunomodulatory properties.<sup>21</sup> Again, conventional high-throughput  
15 experiments for analyzing RNA-protein interactions demand heavier resources and are highly  
16 expensive, where RPISeq tool from Iowa State University  
17 (<http://pridb.gdcb.iastate.edu/RPISeq/>) presents a very inexpensive method of predicting  
18 RNA-protein interactions, simply by analyzing the RNA and protein sequence data.<sup>22</sup> RPISeq  
19 tool operates on machine learning algorithms and generates outputs in two major forms of  
20 RNA-protein interaction prediction parameters – the Random Forest (RF) classifier and  
21 Support Vector Machine (SVM) classifier.<sup>23</sup> Now the protein sequences of TLR3 and TLR8  
22 were extracted from their original PDBs in FASTA format and were used along with the other  
23 sequences for the analysis with this tool.

## 24 **2.3 Molecular docking studies and interactions visualization**

1 The HDock server (<http://hdock.phys.hust.edu.cn/>) presents a novel algorithm which is a  
2 hybrid of template-dependent along with template-independent *ab initio* free docking.  
3 Moreover, it is one of the advance programme which support protein docking against  
4 DNA/RNA molecules.<sup>24</sup> Oligonucleotides have great sizes and demand heavier computing  
5 resources for the rendering of molecular models, which is seldomly feasible for any  
6 supercomputing server to provide. Thus, HDock was a program of choice. The retrieved PDB  
7 structures along with the RNA sequences were used for the purpose, however, the species-  
8 sensitive and intricate nature of the molecules of interest drove us to go with the template-free  
9 docking method. Thereafter, the docked complexes were further visualized with the Protein-  
10 Ligand Interaction Profiler (PLIP) tool (<https://projects.biotec.tu-dresden.de/plip-web/plip/>) it  
11 is a Python-based open-source program that provides a complete analysis and visualization of  
12 the non-covalent protein-ligand interactions even on single-atom level that include seven prime  
13 interaction types *viz.* hydrogen bonds, hydrophobic contacts,  $\pi$ -stacking,  $\pi$ -cation interactions,  
14 salt bridges, water bridges, and halogen bonds.<sup>25</sup>

## 15 **2.4 Simulation studies of the docked complexes**

### 16 **2.4.1 Normal mode analysis (NMA) study**

17 The so retrieved docked structures were then fed into the iMOD webserver from ChaconLab  
18 (<http://imods.chaconlab.org>). This program has a user-friendly GUI and is a well-recognized  
19 tool for performing normal mode analysis (NMA) and simulating various modal trajectories of  
20 protein dynamics. NMA in dihedral coordinates naturally mimics the combined functional  
21 motions of protein molecules modelled as a set of atoms connected by harmonic springs.<sup>26</sup> As  
22 output the server delivers affine modelled arrow, vector field and a modal animation to signifies  
23 the motions. Moreover, study also provide more detail profile about mobility using B-factor  
24 and deformability plots whereas eigenvalue helps to measure the relative modal stiffness of the

1 structure.<sup>27</sup> Besides, covariance matrix implies the variations among the different mode of  
2 motions.

### 3 **2.4.2 Molecular dynamics (MD) simulation study**

4 In-silico molecular docking between receptor and ligand is not sufficient to conclude the nature  
5 of the complex, further analysis viz. MD simulation is desired to validate the structural stability  
6 of that complex. GRONingen MACHine for Chemical Simulations (GROMACS) is the utmost  
7 platform for the MD simulations as it possesses ability to employ different force fields to  
8 generate simulation data not only of proteins complexes but also complexes of DNA-protein  
9 and RNA-protein.<sup>28</sup>

10 In this work, Chemistry at Harvard Macromolecular Mechanics (CHARMM) force field was  
11 accessed initially to generate necessary input files for MD simulation.<sup>29-31</sup> Solution builder  
12 approach under CHARMM-GUI web tool was first add water box (TIP3 216) and then  
13 neutralizing atoms to solvate the system. In order to remove bad contacts and generate more  
14 specific outcomes periodic boundary conditions (PBC) were analysed and minimization steps  
15 were performed sequentially. Server provided outcomes were then applied on GROMACS v.  
16 2020.1 to achieve the equilibrium (NVT- constant volume and temperature) of the system.  
17 Finally, module gmx\_mdrrun was accessed to run the MD simulation of the system and xmgrace  
18 was used to visualize the output of simulation as root mean square deviation (RMSD) plot,  
19 radius of gyration (Rg) plot, solvent accessible surface area (SASA) plot and hydrogen bond  
20 plot.

### 21 **2.5 Analysis of conformational changes of TLRs during and after MD simulation**

22 Protein with conformational changes from its native form suggests it is in bounded state with  
23 any ligand.<sup>32</sup> Thus, to analyse the changes in TLRs, firstly we extract the PDBs from  
24 trajectories and then visualized using PyMOL.

## 1 **3. Results**

### 2 **3.1 Homology modelling**

3 In order to assess the quality of modelled structure of TLR7 and TLR9, we access  
4 Structure Assessment tool under SWISS-MODEL server and found out both the structures are  
5 significant as more than 92% residues from both structures reside in Ramachandran plot  
6 favoured region signifying a stable stereochemical structure (Figure S1).

### 7 **3.2 Motif analysis**

8 Motif analysis played a crucial role in determining the immunoregulatory potency of  
9 the given RNA fragments in interacting with TLR proteins and also their role in inciting pro-  
10 inflammatory cytokine production.<sup>33</sup> According to outcomes from imRNA tool Figure S2A  
11 was modified where the number of motifs was plotted against the sequence length of the  
12 mRNAs to predict the interaction between mRNAs and TLRs. In support RPISeq tool  
13 developed predictions on RF and SVM classifier confirm the interactions as they scored above  
14 0.5 and are visualized in Figure S2B-S2F.

### 15 **3.3 Molecular docking studies and interactome analysis**

16 Our study focuses principally on the interactions of the intracellular TLRs with the mRNA  
17 fragments, and molecular docking studies play a pivotal role in this study by precisely  
18 determining the most probable binding pocket. The docking algorithms used by the HDOCK  
19 program globally samples all the binding poses of the RNA to the protein using a fast and  
20 flexible Fourier transform (FFT) search strategy with focuses on cavity prediction and shape  
21 complementarity.<sup>24</sup> Further, all the sampled binding modes are analyzed by an iterative  
22 knowledge-based scoring function. The more negative the value, the higher the score. Overall,  
23 40 RNA-protein docked complexes were generated in this study and the scores are furnished

1 in Table S1. However, only the four topmost scoring docked complexes were selected for  
2 further experimentation and analysis.

### 3 ***3.3.1 Interaction of NSP10 RNA fragment with TLR3***

4 In accord to our docking studies, TLR3 binds with a much proficient binding pose with  
5 the RNA fragment encoding NSP10, with a high docking score of -404.77 which is higher than  
6 any other docking operation involving TLR3 like that of the RNA fragment of papain-like  
7 protease which binds with a score of -346.51 or that of the main protease which has a score of  
8 -333.76 (Table S1). While in search of insights of molecular docking PLIP study found the  
9 involvement of 251Arg, 252Asn, 277Thr residues in hydrogen bonding, 326Tyr, 359His in  $\pi$ -  
10 Cation interactions and Salt bridges from 32 and 319His (Figure 2A, Table 1)

### 11 ***3.3.2 Interaction of Envelope-protein RNA fragment with TLR7***

12 In this study, the docking operations determine a good binding pocket for E-protein  
13 mRNA within the TLR7 dimeric cavity, with a binding score of -381.60. PLIP analysis of the  
14 docking complex later reveals that the complex stability is regulated by hydrogen bonding  
15 interactions through residues *viz.* 684Lys, 707Ser, 708His, by  $\pi$ -Cation Interactions through  
16 758(B)Lys, 781His, 782(B)His residues and through Salt Bridges with residues of 627Arg,  
17 736Arg, 758(B)Lys, 820(B)His as shown in Figure 2B and Table 1.

### 18 ***3.3.3 Interaction of NSP8 RNA fragment with TLR8 protein***

19 Herein, the most probable binding pose of NSP8 RNA fragment and TLR8 is built with  
20 a definite score of -416.84, which is much higher than those poses comprising of the RNAs of  
21 PL-pro and Mpro which had scores of -360.49 and -355.80 respectively. PLIP analysis as in  
22 Figure 3A again confirms these results as it shows that there exists a strong concerted  
23 interaction among the complex domains that are 426(B)Ser, 470(B)Phe, 473(B)Pro, 489(B)Ser,

1 513(B)Ser, etc residues in hydrogen bonding, 569Arg, 696(B)Arg, and 699Lys in  $\pi$ -Cation  
2 interactions and Salt bridges from 643(B)Arg, 673(B)Asp, 721His, etc. as shown in Table 1.

### 3 **3.3.4 Interaction of S2 Subunit RNA fragment with TLR9 protein**

4 Among the docked complexes with TLR9 possessing the S2 subunit mRNA fragment  
5 stood at the highest position with a score of -440.33. The molecular insights of this ligation are  
6 well supported by the PLIP analysis as finding involves 224Tyr, 247Arg, 248Val, 292Lys  
7 residues in hydrogen bonding, 311Arg in  $\pi$ -Cation interactions and Salt bridges from 207Lys,  
8 287Glu, and 292Lys (Figure 3B, Table 1).

## 9 **3.4 Simulation analysis**

### 10 **3.4.1 Normal mode analysis (NMA)**

11 iMODs server generates conformational morphing trajectories of the molecule as depicted in  
12 Figure S2G-S2J and enables us to inspect the compatibility and ultimate stability of the  
13 complex. Affine-model-based arrow within the structure represents the nature of domain  
14 dynamics (Figure S3A, S3B, S4A and S4B). While structure mobility and flexibility are  
15 categorized by deformability and B-factor plot (Figure S3C, S3D and Figure S4E, S4F). In  
16 deformability plot hinge regions defined the non-rigid helical contents and in B-factor  
17 distortions are atomic positional fluctuations. Calculations to measure eigenvalue are fully  
18 depended on the energy required to deform the structure, as Figure S4H with very low  
19 eigenvalue direct lower energy can deform the structure and infer greater stability. According  
20 to Abdelli et al. (2020),<sup>34</sup> variance shows a direct and inverse relation with eigenvalue as  
21 depicted in Figure S3I, S3J, S4I and S4J, where red colour represents individual and green to  
22 cumulative variance. In the covariance matrix, motions are categorized in different modes and  
23 visualized in Figure S3K and S3M with red, blue and white colour that signify variations  
24 among correlated, anti-correlated and uncorrelated motions respectively. In the end, the last

1 outcome of NMA study, elastic network map graph is represented by Figure S4L and S4N  
2 where grey dots indicate the stiffness of the motions and springs direct the atomic connections  
3 of the complex.

#### 4 **3.4.2 Molecular dynamics (MD) simulation**

5 Four most active complexes, those were primarily strained out according to their significant  
6 docking score were subjected to further analysis through MD simulation. MD is the  
7 computerized method of simulation that allows mobility of atoms and molecules for a given  
8 time to monitor the interaction and calculate the forces, that ultimately figure out the atomic  
9 level configuration of the complex structure and then visualize using several graphical plots.<sup>35</sup>  
10 Herein, RMSD plot in Figure 4A reflects the structural stability of the backbone of different  
11 receptor TLRs of the four complexes and suggests complex TLR8-NSP8 is less stable among  
12 others. Where Figure 4B and 4C define the compactness and solvent accessible area of that  
13 complexes as  $R_g$  plot and SASA plot, respectively. At last the graphical plot of hydrogen bond  
14 clearly describes the insights of the complexes and supports the postulation reflected on  
15 RMSD,  $R_g$  and SASA plots (Figure 4D).

#### 16 **3.5 mRNA induced conformational changes in TLRs during and after MD simulation**

17 Apart from RMSD,  $R_g$ , SASA and hydrogen bond studies, we also analyse conformational  
18 changes of receptor TLRs arisen during simulation processes. Here, we prepared and visualized  
19 six different conformation on different time steps of each TLRs and revealed mRNA of NSP10,  
20 NSP8, S2 subunit of spike protein and Envelope protein have enough potential to change the  
21 conformation of extracellular domain of TLR3, TLR8, TLR9 and TLR7 respectively (Figure  
22 5).

#### 23 **4. Discussion**

1 Toll-like receptors (TLRs) are the essential mediators and regulators of host immunity. The  
2 intracellular TLRs play a vital role in sensing the viral RNA and triggers the downstream  
3 activation of signalling cascade of immunomodulation. TLR3 dimerizes upon binding and  
4 recruits signal transfer proteins MyD88, TIRAP, TRAM, or TRIF in the cytoplasmic TIR  
5 domain.<sup>36</sup> Certain kinases alike IRAKs, TBK1, and IKKs are activated as well as TRAF6, as  
6 per different adaptors, thus leading to activation of the MAPK, NF- $\kappa$ B, or JNK-STAT  
7 pathways for promotion of the transcription of inflammatory cytokines and for production of  
8 IFN, IL-1 $\beta$ , IL-6, to coordinate the local or systemic inflammatory responses among which IL-  
9 1 $\beta$  and IL-6 are the major pro-inflammatory cytokines.<sup>37</sup> TLR8 protein has various similarities  
10 with TLR7 including their gene loci on the X chromosome and believed to possess a close  
11 phylogenetic relationship.<sup>38</sup> The binding to TLR7/8 to the RNA, recruits MyD88 which on one  
12 hand activates IRF-7 and on the other hand also activates IRAK-4 resulting in the activation of  
13 the transcription factor NF- $\kappa$ B and subsequent antiviral response.<sup>39</sup> Ligated TLR9 moves from  
14 the endoplasmic reticulum to the Golgi apparatus and lysosomes, recruiting MyD88-dependent  
15 pathway, and resulting in IRF7-mediated IFN production or in proinflammatory responses by  
16 producing IL-6, IL-12, and TNF $\alpha$  through activation of the NF- $\kappa$ B.<sup>40</sup>

17 This study focuses on the establishing the binding and potency of SARS-CoV-2 with the human  
18 intracellular TLRs including TLR 3, 7, 8, and 9. Regarding this crystal structures and templates  
19 are preliminarily obtained via X-Ray Diffraction techniques with resolution of most 2.50Å,  
20 considering good for molecular docking studies. Since the quality of results obtained from  
21 further experiments would rely on the stereochemical quality of modelling, and it showed the  
22 models had >92% residues in favored regions of their respective Ramachandran plots (Figure  
23 S1) indicating standard quality of modelling.

24 The motif analysis recognizes the immunomodulated motifs within the RNA sequence that can  
25 stimulate the immune responses. According to imRNA tool the mRNA fragments are highly

1 capable of interacting with the TLRs and influence immunomodulation (Figure S2A). Though  
2 mRNA of the S2 subunit and NSP8 proteins are almost similar in size and have the highest  
3 potency compared to others. While RPISeq tool strained out best candidates showing maximum  
4 potency in interaction for intracellular TLR 3, 7, 8 and 9. As RPISeq tool predicted a high  
5 interaction coefficient for NSP10 RNA with TLR3 as exhibited in Figure S2B. While the  
6 threshold SVM and RF classifier values for minimum interaction is 0.5 for SVM classifier as  
7 well as for RF classifier, in contrast, the NSP10 mRNA with SVM classifier value of 0.984 and  
8 RF classifier value of 0.8 it stands among the best candidates for interacting with TLR3. While  
9 prediction figure out E-protein RNA fragment is best in interact TLR7 with an RF classifier  
10 value of 0.75 and an SVM classifier value of 0.884 (Figure S2C). Later on, studies also revealed  
11 best interaction between NSP8 mRNA and TLR8 protein having SVM and RF classifier as  
12 high as 0.991 and 0.7 respectively (Figure S2D). Moreover, prediction again showed a higher  
13 probability of interaction of TLR9 with the S2 subunit with SVM classifier value of 0.868 and  
14 RF classifier value of 0.7 (Figure S2E).

15 Further, the molecular docking studies showed a similar trend by reflecting highest docking  
16 score to the complexes selected through RPISeq tool and in turn support the prediction of best  
17 docked structures (Table S1). After getting confirmation on best predicted structure through  
18 HDOCK we forwarded to next step of simulation to analyze the conformational stability and  
19 flexibility of that selected four structures viz. TLR8-NSP8 mRNA complex (Figure 3A),  
20 TLR3- NSP10 mRNA complex as shown in Figure 2A, TLR7-E mRNA complex in Figure 2B  
21 and TLR9-S2 mRNA complex (Figure 3B). Initially, the NMA study reveals all the compounds  
22 have better stability as it reflects quite similar and significant eigenvalue scores, those are  
23  $3.154899 \times 10^{-5}$  for TLR8-NSP8 mRNA complex as shown in Figure S4G,  $1.27056 \times 10^{-5}$  TLR3-  
24 NSP10 mRNA complex (Figure S3G),  $7.042567 \times 10^{-5}$  TLR7-E mRNA complex as in Figure  
25 S3H , and  $9.131746 \times 10^{-5}$  TLR9-S2 mRNA complex (Figure S4H). Later, MD simulation

1 studies figure out postulation of NMA studies is not fully accepted as TLR8-NSP8 mRNA  
2 complex found as unstable throughout the simulation process in the RMSD plot Figure 4A.  
3 The PLIP analysis Table 1 and hydrogen bond plot Figure 4D finally reveals the interactions  
4 between the mRNA ligand and the TLR proteins and the reasons for the stability of the  
5 complexes. At the end, our final experiment on changes of conformation of TLRs after binding  
6 and simulation hypothesized as mRNA of NSP10, S2 and E proteins are potent enough to bind  
7 with TLR3, TLR9 and TLR7 and trigger downstream cascade reactions (Figure 5).

## 8 **5. Conclusion**

9 The intracellular TLRs play a vital role in the recognition of PAMPs and viral pathogens and  
10 it is of utmost necessity to understand the sensing of SARS-CoV-2 viral RNA with the human  
11 intracellular TLRs including TLR3,7,8 and 9. The current study primarily focuses on the  
12 comparative binding of mRNAs of SARS-CoV-2 with the human intracellular TLRs. Later,  
13 NMA study found out stability between complexes with significant values for TLR8-NSP8  
14 mRNA complex, TLR3-NSP10 mRNA complex, TLR7-E mRNA complex, and TLR9-S2  
15 mRNA complex. While, upon MD simulation it is confined that the mRNA of NSP8 has lower  
16 potency to make complex with TLR8. In turn significant binding potency of NSP10, S2, and E  
17 proteins mRNA with TLR3, TLR9, and TLR7 respectively, help us to conclude our study and  
18 we infer complexes of TLR3-NSP10 mRNA, TLR7-E mRNA, and TLR9-S2 mRNA can  
19 potentially trigger the downstream activation and proliferation of immune responses and can  
20 be used in validation for therapeutic action.

## 21 **Acknowledgment**

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23 and researchers currently working endlessly against coronavirus disease worldwide. We have  
24 kept selected studies as reference due to the limitation in space, but we appreciate all the uncited

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### 3 **Conflict of Interest**

4 The authors declare that there is no conflict of interest.

### 5 **Authors' Contributions**

6 AC, NCD performed all experiments; AC, NCD, and RP write the manuscript; MB and SM  
7 drafted and edited the manuscript; SM design the experiments, analyses the result and supervise  
8 the study.

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## 16 Captions

### 17 Figure captions

18 **Figure 1.** A. Conceptual representation of the SARS-CoV-2 genome. B. TLR3 C. TLR7 D.  
19 TLR8 E. TLR9.

20 **Figure 2.** A. Molecular interaction between NSP10 RNA fragment and TLR3 protein. (i)  
21 Zoomed in configuration (ii) Detailed bonding interactions determined by PLIP. Solid blue  
22 lines represent Hydrogen bonds, dotted orange lines represent  $\pi$ -Cation interactions, while  
23 dotted yellow lines denote Salt bridges. B. Molecular interaction between E-protein RNA

1 fragment and TLR7 protein. (i) Zoomed in configuration (ii) Detailed bonding interactions  
2 determined by PLIP. Solid blue lines represent Hydrogen bonds, dotted orange lines represent  
3  $\pi$ -Cation interactions, while dotted yellow lines denote Salt bridges. It is to be noted that for  
4 the sake of perceptual clarity some residues have been intentionally omitted from the diagram,  
5 however the same have been furnished in table 1.

6 **Figure 3.** A. Molecular interaction between NSP8 RNA fragment and TLR8 protein. (i)  
7 Zoomed in configuration (ii) Detailed bonding interactions determined by PLIP. Solid blue  
8 lines represent Hydrogen bonds, dotted orange lines represent  $\pi$ -Cation interactions, while  
9 dotted yellow lines denote Salt bridges. B. Molecular interaction between S2-subunit RNA  
10 fragment and TLR9 protein. (i) Zoomed in configuration (ii) Detailed bonding interactions  
11 determined by PLIP. Solid blue lines represent Hydrogen bonds, dotted orange lines represent  
12  $\pi$ -Cation interactions, while dotted yellow lines denote Salt bridges. It is to be noted that for  
13 the sake of perceptual clarity some residues have been intentionally omitted from the diagram,  
14 however the same have been furnished in table 1.

15 **Figure 4. Molecular dynamic (MD) simulation study.** Reflecting outcomes generated during  
16 MD simulation studies of TLR3-NSP10 mRNA complex, TLR7-E protein mRNA complex,  
17 TLR8-NSP8 mRNA complex and TLR9-S2 protein mRNA complex. A. RMSD plots. B. Rg  
18 (Radius of gyration) plots. C. SASA plots. D. Graphical presentation of Hydrogen bonds.  
19 (TLR3-NSP10 mRNA complex shown in red, TLR8-NSP8 mRNA complex shown in violet,  
20 TLR9-S2 protein mRNA complex shown in black and TLR7-E protein mRNA complex in  
21 green).

22 **Figure 5. mRNA induced conformational changes in TLR3, TLR7, TLR8 and TLR9.**  
23 Representing changes in helix, sheet, and loop in the structure of TLRs at various time points  
24 of molecular dynamic simulation. A represents changes in TLR3 after NSP10 mRNA

1 induction. B signifies envelope (E) protein mRNA induced alteration of TLR7. C represents  
2 changes in TLR9 after S2 subunit of spike protein mRNA induction. While D exhibits NSP8  
3 mRNA induced alteration of TLR8. Red is depicting the helix, yellows are the sheets, and loops  
4 are given as green.

## 5 **Supplementary Figures**

6 **Figure S1. Validation of homology modelled structures.** A. Ramachandran plot for modelled  
7 TLR7 protein, showing 92.11% residues within favored regions. B. Ramachandran plot for  
8 modelled TLR9 protein, showing 93.09% residues within favored regions.

9 **Figure S2. Motif analysis.** A. imRNA Motif analysis graph of different RNAs. B. RPISeq  
10 graph for RNAs & TLR3 C. RPISeq graph for RNAs & TLR7 D. RPISeq graph for RNAs &  
11 TLR8 E. RPISeq graph for RNAs & TLR9 F. Legend for graphs. G. H. I. & J. Exhibit different  
12 conformations of TLR3, -7, -8, & -9 respectively in a given mode, analyzed by iMOD tool.

13 **Figure S3. Normal mode analysis (NMA) depicting stability and flexibility of complexes**  
14 **as computed by iMODS webserver.** Results of NMA of TLR3-NSP10 mRNA and TLR7-E  
15 mRNA complex respectively (A,B) NMA mobility (arrow field), (C,D) deformability, (E,F)  
16 B-factor, (G,H) eigenvalue, (I,J) variance (red colour represent individual variances and green  
17 colour indicates cumulative variances), (K,M) co-variance map (red- correlate, white-  
18 uncorrelated and blue- anti-correlated) and (L,N) elastic network (darker gray regions indicates  
19 more stiffer regions) of the complex.

20 **Figure S4. Normal mode analysis of the TLR-mRNA interactions.** iMODS webserver  
21 depicted outcome of NMA of TLR8-NSP8 mRNA and TLR9-S2 mRNA complex respectively,  
22 (A,B) NMA mobility (arrow field), (C,D) deformability, (E,F) B-factor, (G,H) eigenvalue, (I,J)  
23 variance (red colour represent individual variances and green colour indicates cumulative  
24 variances), (K,M) co-variance map (red- correlate, white- uncorrelated and blue- anti-

1 correlated) and (L,N) elastic network (darker gray regions indicates more stiffer regions) of the  
2 complex.

### 3 **Table captions**

4 **Table 1.** Residues involved in bonding interactions among the given docked complexes.

### 5 **Supplementary table**

6 **Table S1.** Docking scores of different RNA fragments docking with different TLRs.

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**Table 1.** Residues involved in bonding interactions among the given docked complexes.

Complex	Interacting residues		
	Hydrogen bonding residues	$\pi$ -Cation Interactions	Salt Bridges
TLR3 with NSP10 RNA	251ARG, 252ASN, 277THR, 322PHE	326TYR, 359HIS	32HIS, 319HIS
TLR7 with E protein RNA	684LYS, 707SER, 708HIS, 710(B)GLN, 731LYS, 732ASN, 734(B)GLN, 736ARG, 755SER, 756SER, 758LYS, 782HIS, 784(B)ARG	758(B)LYS, 781HIS, 782(B)HIS	627ARG, 736ARG, 758(B)LYS, 820(B)HIS
TLR8 with NSP8 RNA	426(B)SER, 470(B)PHE, 473(B)PRO, 489(B)SER, 513(B)SER, 538(B)THR, 562(B)SER, 563TYR, 569ARG, 590(B)ASN, 592(B)SER, 672(B)ASN, 696(B)ARG, 699LYS, 723ARG, 779ASP, 797(B)ARG, 810ARG	569ARG, 696(B)ARG, 699LYS	375(B)ARG, 643(B)ARG, 673(B)ASP, 721HIS, 723(B)ARG, 749(B)LYS
TLR9 with S2 subunit RNA	224TYR, 247ARG, 248VAL, 292LYS, 311ARG, 312VAL, 335GLN, 337ARG, 340ASN, 397ARG, 399GLN, 464GLU	311ARG	207LYS, 287GLU, 292LYS, 337ARG, 368GLU, 467ARG

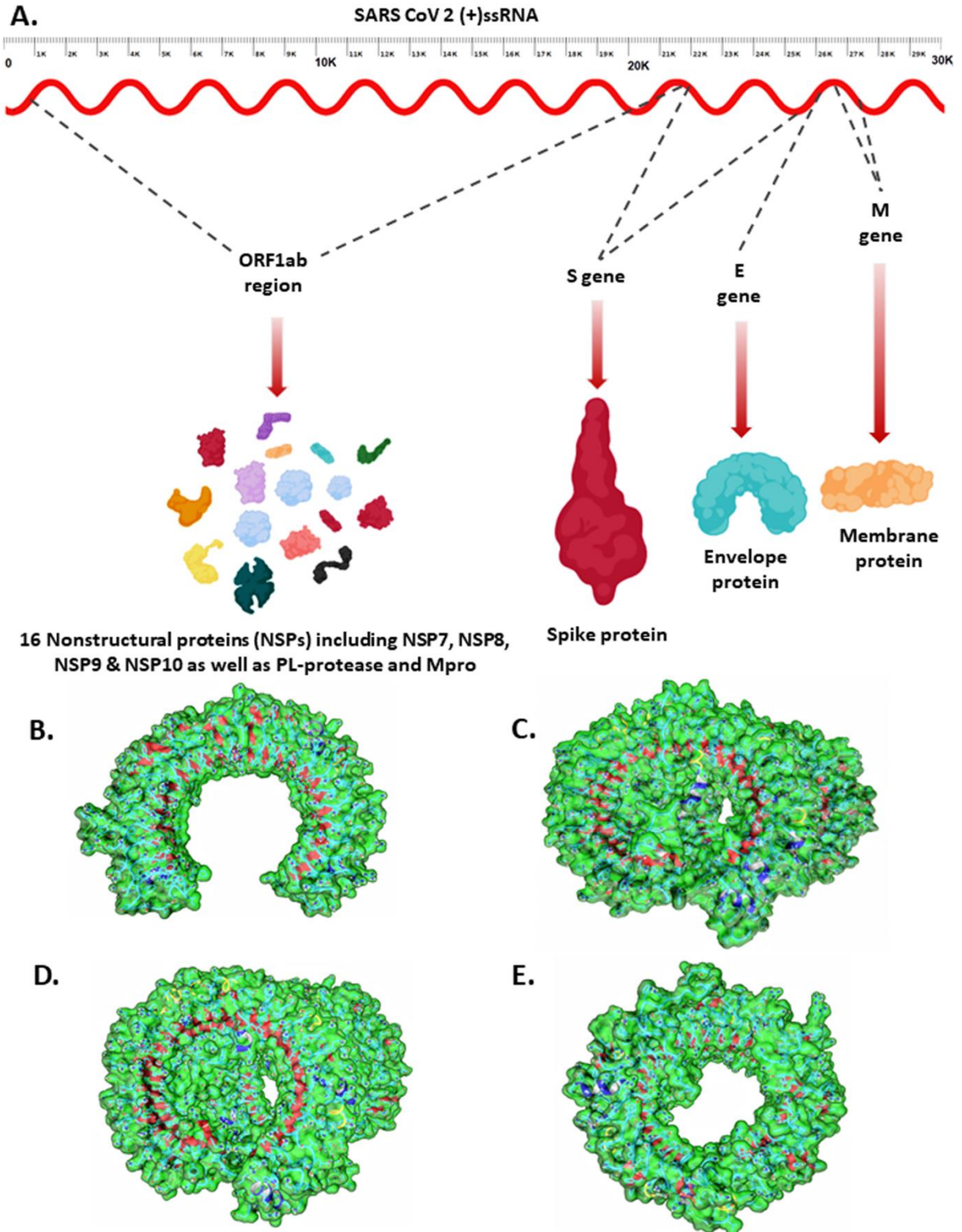
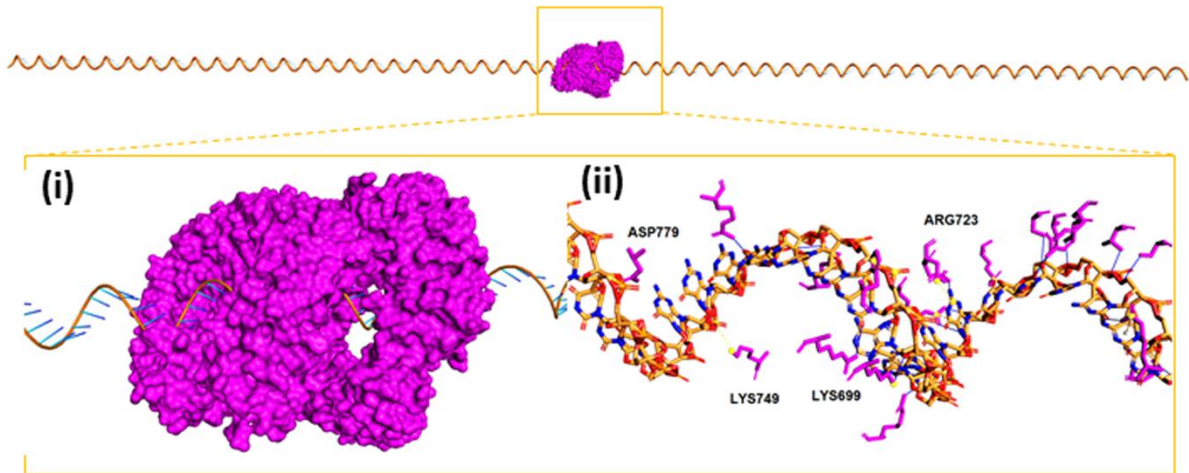
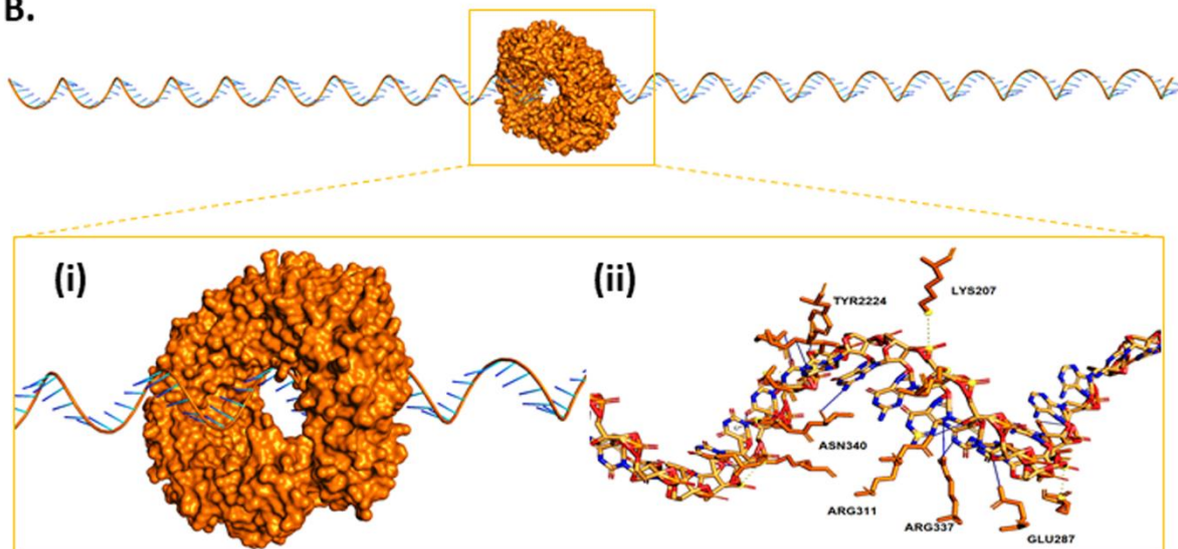


Figure 1

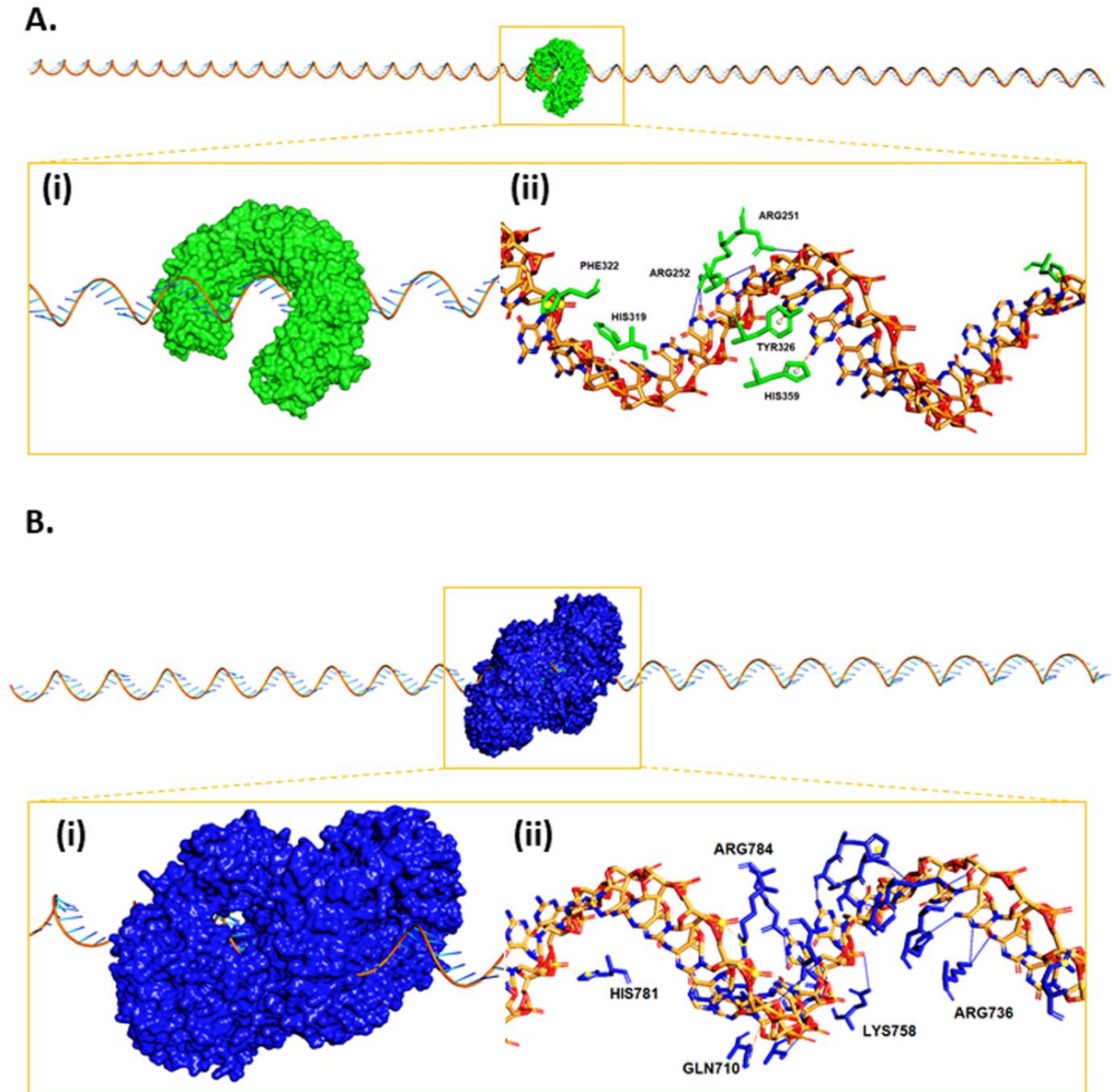
**A.**



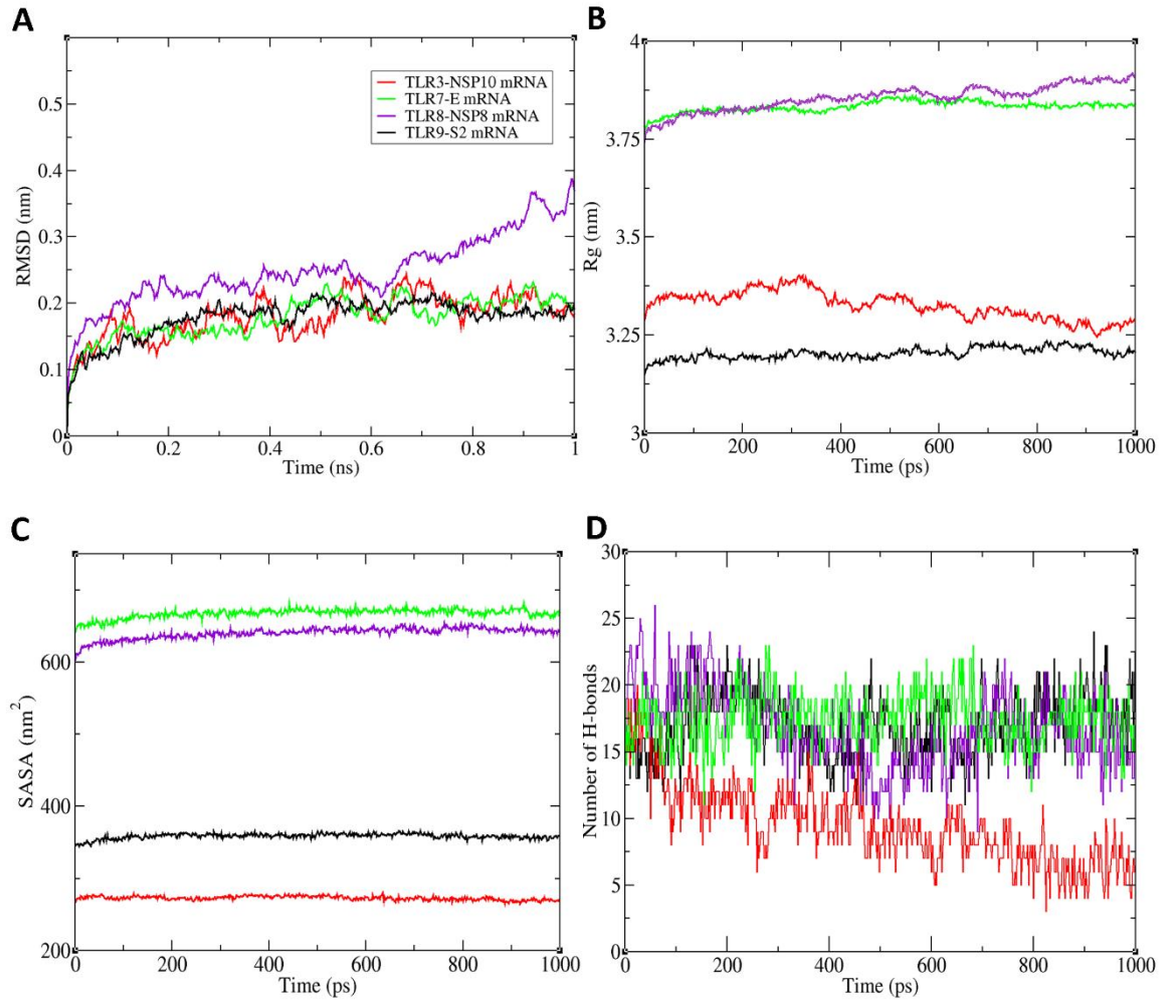
**B.**



**Figure 2**



**Figure 3**



**Figure 4**

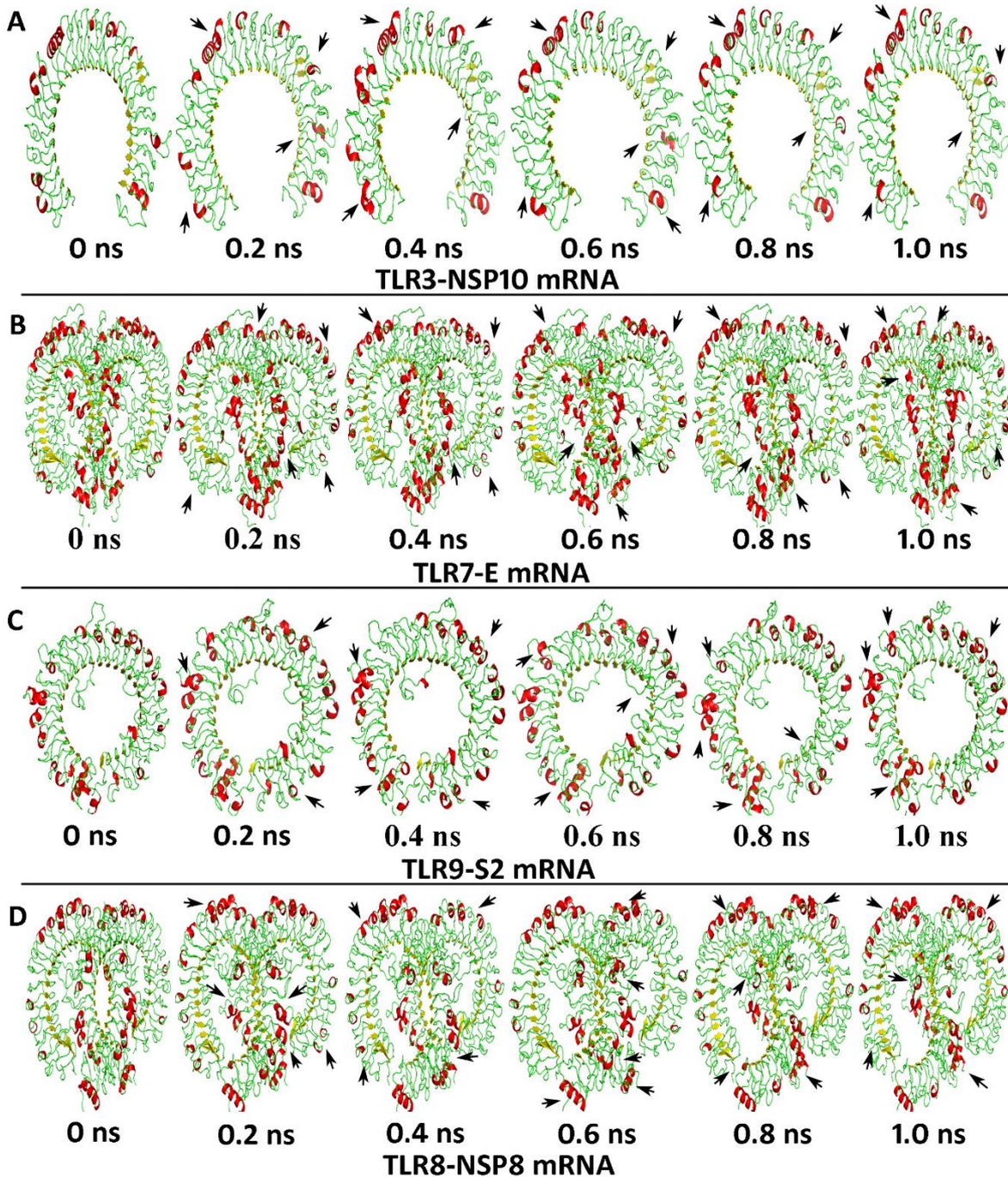


Figure 5