Predictable fold switching by the SARS-CoV-2 protein ORF9b

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Abstract

Extant fold-switching proteins remodel their secondary structures and change their functions in response to environmental stimuli. These shapeshifting proteins regulate biological processes and are associated with a number of diseases, including tuberculosis, cancer, Alzheimer’s, and autoimmune disorders. Thus, predictive methods are needed to identify more fold-switching proteins, especially since all naturally occurring instances have been discovered by chance. In response to this need, two high-throughput predictive methods have recently been developed. Here we test them on ORF9b, a newly discovered fold switcher and potential therapeutic target from the Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2). Promisingly, both methods correctly indicate that ORF9b switches folds. We then tested the same two methods on ORF9b1, the ORF9b homolog from SARS-CoV-1. Again, both methods predict that ORF9b1 switches folds, a finding consistent with experimental binding studies. Together, these results (a) demonstrate that protein fold switching can be predicted using high-throughput computational approaches and (b) suggest that fold switching might be a general characteristic of ORF9b homologs.

KEYWORDS

fold-switching proteins, metamorphic proteins, protein folding, SARS-CoV-2

1 | INTRODUCTION

Fold-switching proteins remodel their secondary structures and change their functions in response to environmental stimuli.1 These proteins challenge the long-held paradigm that the amino acid sequence of a globular protein encodes its unique stable structure.2 Furthermore, fold switchers occur in all kingdoms of life, perform over 30 different functions, and are triggered by nearly a dozen stimuli.1 Additionally, the structural transitions of some fold switchers regulate biological processes, such as the expression of bacterial virulence genes3 and the circadian rhythm of cyanobacteria.4

Given the growing amount of evidence suggesting that fold switchers play important regulatory roles,5 it is not surprising that a number of them are associated with different human diseases. For example, PimA, which undergoes an α-helix ↔ β-strand transition, initiates the biosynthetic pathway of virulence factors produced by M. tuberculosis.6 Human lymphotactin (a.k.a. XCL1) iso-energetically populates two β-sheet conformations with completely different hydrogen bonding patterns7,8 and is associated with autoimmune disorders.9
Furthermore, human Chloride Intracellular Channel 1 (CLIC1) remolds its secondary structure and changes its function from a glutathione reductase\textsuperscript{10} to a chloride channel\textsuperscript{11} that balances intracellular chloride levels when cells undergo oxidative stress due to cancer\textsuperscript{12} or Alzheimer’s.\textsuperscript{13}

The biological relevance and increasing number of identified fold switchers have motivated the development of computational methods that predict more. The need for accurate predictive methods is especially acute because, to date, all naturally occurring fold switchers have been discovered by chance. Two years ago, we reported that discrepancies between predicted and experimentally determined protein structures can indicate fold switching.\textsuperscript{14} More recently, we developed a sequence-based method that predicts fold switchers with high levels of statistical significance.\textsuperscript{15}

Here, both predictive approaches are tested on the newly discovered fold switcher, ORF9b\textsuperscript{16} (Figure 1). This protein is from the genome of the Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2), the cause of the current global pandemic. When expressed, ORF9b binds the human mitochondrial protein Tom70, an outer membrane protein that acts as a host-dependency factor for SARS-CoV-2.\textsuperscript{16} In other words, viral titers in Caco-2 cells with \textit{TOMM70} (the gene that encodes Tom70) are significantly higher than in cells without. ORF9b-Tom70 binding has been proposed to have one of two cellular effects:\textsuperscript{16} (a) modulating interferon and apoptosis signaling or (b) decreasing mitochondrial import efficiency, leading to mitophagy.

Isolated ORF9b folds into a \(\beta\)-sheet topology that forms a domain-swapped dimer, part of which transforms into a long \(\alpha\)-helix when bound to Tom70 (Figure 1). Interestingly, ORF9b has two infection-driven phosphorylation sites (S50 and S53),\textsuperscript{17} which contact Tom70 directly but are solvent-exposed in the ORF9b dimer. Thus, it has been hypothesized that phosphorylation weakens ORF9b’s interactions with Tom70, causing its homodimeric \(\beta\)-sheet fold to become more energetically favorable.\textsuperscript{16}

Since ORF9b undergoes a large \(\alpha\)-helix \(\leftrightarrow\) \(\beta\)-sheet transition, it is a suitable target for both the sequence- and structure-based predictive methods reported previously.\textsuperscript{15,18} Consistent with experimental observations, both methods indicate that ORF9b switches folds. These methods were then tested on the SARS-CoV-1 ORF9b homolog, hereafter called ORF9b1, which is also binds Tom70 \textit{in situ}\textsuperscript{16} but has not been shown to switch folds. Again, both methods predict that ORF9b1 switches folds. Together, these results (a) corroborate previous work demonstrating that low-resolution, high-throughput methods can predict fold switching and (b) suggest that fold switching might be a general characteristic of ORF9b homologs.
2 | RESULTS

2.1 | Structure-based predictions suggest that SARS-CoV-2 ORF9b switches folds

Previous work has shown that inconsistencies between experimentally determined secondary structures and homology-based secondary structure predictions often indicate protein fold switching. These predictions are homology-based because the algorithms that generate them (in this case PSIPRED, SPIDER2, and JPred4) leverage conservation patterns from homologous sequences to infer secondary structure.

Secondary structure predictions of full-length ORF9b were calculated using PSIPRED, SPIDER2, and JPred4 (Figure 2) and compared with the experimentally determined structures. Interestingly, all three secondary structure predictors suggest helical propensities in the region of ORF9b that forms an α-helix when it binds Tom70 (this has been reported previously for JPred4, but not the other secondary structure predictors). Homology-based predictions of this region uniformly disagree with the experimentally determined secondary structure of the ORF9b dimer, which forms a domain-swapped β-hairpin (Figure 1, green). By contrast, these predictions suggest more helical content than was observed in the cryo-EM structure of the ORF9b-Tom70 complex. Accordingly, low prediction accuracies (Methods) were observed from all three secondary structure predictors. Specifically, prediction accuracies ranged from 48–59% when referenced against the full-length experimentally determined ORF9b dimer and 51–62% when compared with the cryo-EM-resolved region of ORF9b that binds Tom70 (Figure 2). These accuracies fall well below previously benchmarked predictor accuracies, all three of which exceed 80%. Additionally, these accuracies are lower than the mean/median secondary structure prediction accuracies for fold-switching regions of proteins reported previously, which range from 67–68%/68–71%, respectively. Thus, in line with previous findings, these inaccurate secondary structure predictions are consistent with the experimental observation that ORF9b switches folds.

![Figure 2](image_url)

**Figure 2** Three state-of-the-art algorithms inaccurately predict the secondary structures of both experimentally determined forms of ORF9b. Names of the three algorithms lie to the right of their corresponding predictions (three middle secondary structure diagrams, all colored blue). Coils are represented by black lines; α-helices/β-strands are represented by rounded rectangles/arrows. Predictions of α-helices and β-strands that span ≥2 contiguous residues are shown. Experimentally determined secondary structures of the ORF9b Dimer/ORF9b + Tom70 are shown above/below the predictions and are colored as in Figure 1: green secondary structures switch folds; all other secondary structures are shown in orange. The dash in ORF9b’s sequence (above secondary structure diagrams) represents an area of missing electron density in both structures; the gray background corresponds to the region of the ORF9b structure that could be resolved by cryo-EM when in complex with Tom70. The table reports secondary structure prediction accuracies of each algorithm referenced against each experimentally determined structure.
2.2 | Sequence-based predictions suggest that SARS-CoV-2 ORF9b switches folds

Given that the structure-based method correctly inferred that ORF9b switches folds, the next step was to determine whether ORF9b fold switching could be inferred from its sequence alone. It has been shown previously that JPred4 predicts fold switching from sequence more robustly than other secondary structure predictors,18 and α-helix <-> β-strand prediction discrepancies from JPred4 between whole protein sequences and excised sequence fragments can be a robust indicator of fold switching.15 Such a fragment (residues 59–80) was identified in ORF9b (Figure 3). Within the context of its parent protein, this fragment is predicted to form N- and C-terminal α-helices, whereas the excised fragment is predicted to assume an α-helix at the N-terminus and β-strands towards the C-terminus.

Based on previously reported thresholds,15 these prediction discrepancies between parent and excised sequence fragments are significant, consistent with the observation that ORF9b switches folds. Specifically, both the total predicted secondary structure content (50%) and the 75% discrepancy between α-helix <-> β-strand predictions exceed the minimum parameters15 (secondary structure content ≥35% and α-helix <-> β-strand prediction discrepancies ≥50%). Although the region with α-helix <-> β-strand discrepancy begins 5-residues C-terminal to the region experimentally observed to switch from α-helix <-> β-strand, we note that the β-strand predictions of the excised fragment overlap with some regions of the ORF9b dimer that also fold into β-strands. Furthermore, as reported previously,15,18 this method uses the differences in predicted α-helix and β-strand to infer fold switching. Predictions need not adhere exactly to a solved protein structure to make this inference.

Indeed, as shown in the previous section and in previous work,14 there is generally poor correspondence between the predicted and experimentally determined secondary structures of fold-switching proteins. Furthermore, slightly off-register α-helix <-> β-strand prediction discrepancies have been observed in other fold switchers such as RfaH and Ovalbumin, but such discrepancies are very rare in proteins expected not to switch folds.15

2.3 | Both predictive methods suggest that ORF9b’s homolog from SARS-CoV-1 also switches folds

Since both of our predictive methods indicate that ORF9b switches folds, they were used to assess whether its homolog from SARS-CoV-1 (hereafter called ORF9b1) might switch also. ORF9b1 is the only ORF9b homolog with a solved crystal structure in the PDB. Its experimentally determined structure assumes the same dimeric fold as ORF9b (RMSD = 0.94 Å using PyMOL22), and its sequence is 69% identical to ORF9b’s (Figure 4a).

As with ORF9b, both predictive approaches suggest that ORF9b1 switches folds (Figure 4b-d). Specifically, all three secondary structure predictors suggest that the region of ORF9b1 analogous to the ORF9b region that interacts with human Tom70 is helical, even though it folds into a β-hairpin in the experimentally determined ORF9b1 structure (Figure 4a,b). The prediction that this region can form an α-helix is plausible given that ORF9b1 (a) coimmunoprecipitates with Tom70 in both HEK293T and A549 cells and (b) colocalizes with Tom70 in HeLaM cells.16 Additionally, predictions from all three algorithms had lower-than-expected accuracies,19–21 ranging from 51–59% (Figure 4c), similar to the prediction accuracies the ORF9b dimer. Furthermore, JPred4 predictions of the ORF9b1 sequence fragment analogous to the ORF9b sequence in Figure 3 differ significantly depending on whether the fragment is excised or contextualized within its parent sequence (Figure 4d): 41% predicted secondary structure content, 56% α <-> β discrepancies, both of which, again, exceed the significance thresholds for fold switching reported previously.15

![FIGURE 3](image-url) JPred4 predicts significantly different secondary structures for a fold-switching ORF9b sequence fragment depending on its context. When within its parent sequence, the fragment is predicted to be α-helical (above), but when excised from its parent, it is predicted to be a mix of α-helix and β-strand (below). The sequence is shown above both predictions with residue numbers shown above for reference. Coils are represented by black lines; α-helices/β-strands are represented by rounded rectangles/arrows.

3 | DISCUSSION

Extant fold-switching proteins remodel their secondary structures and change their functions in response to environmental stimuli.1 These shapeshifting proteins regulate biological processes2 and are associated with a number of human diseases.6,9,12 In December 2020, ORF9b, a protein from the SARS-CoV-2 genome and a possible...
therapeutic target for coronaviruses, was reported to switch between a homodimeric β-sheet fold and an α-helix that binds human Tom70.

We tested two recently developed predictive methods on ORF9b to assess whether they could identify it as a fold switcher. Both methods—one structure-based, the other sequence-based—were successful. The structure-based method identified the region experimentally observed to assume α-helix in one structure and β-strand in the other, while the sequence-based prediction began within five residues of this fold-switching region. These results suggest that structure-based predictions might identify the precise locations of fold-switching regions more accurately than sequence-based predictions. Nevertheless, the previously reported robustness of the sequence-based method suggests that it is an adequate binary classifier for fold switchers/single-fold proteins. Interestingly, sequence-based predictions identify the part of the domain-swapped β-strands of ORF9b and ORF9b1 as helices in both whole-sequence and excised sequence fragments. This finding suggests that this region of the sequence might have intrinsic helical propensities that are energetically outweighed by favorable interactions with neighboring β-sheets, a possibility...
consistent with other studies suggesting that protein secondary structure formation can be context-dependent.\textsuperscript{23,24}

Our predictions suggest that ORF9b1, the ORF9b homolog from SARS-CoV-1, also switches folds. This finding is consistent with a couple lines of experimental evidence demonstrating that ORF9b1, like ORF9b, binds Tom70.\textsuperscript{16} Based on the ORF9b-Tom70 structure, it is not obvious how ORF9b1 could bind to Tom70 without assuming an α-helical fold, even though its only experimentally determined structure suggests that it folds into a β-sheet topology. The sequences of ORF9b1 and ORF9b are 69% identical. Reconstructed ancestors with similar levels of sequence identity to the metamorphic protein XCL1 have also been shown to switch folds,\textsuperscript{7} demonstrating that fold switching can be conserved among protein homologs, though this is not always the case.\textsuperscript{25}

The folds of ORF9b differ from those of the fold switchers used to develop our predictive methods.\textsuperscript{14,15} Since these methods correctly infer that ORF9b switches folds, they have the potential to identify other proteins that undergo novel fold-switching transitions. Given that these methods are high-throughput, they can be tested on numerous sequences and structures. We are optimistic that they will reveal new fold switchers in future work.

4  |  METHODS

4.1  |  Secondary structure predictions of ORF9b and ORF9b1

Secondary structure predictions of ORF9b (PDB\textsuperscript{26} IDs: 6Z4U, chain A and 7KTD, chain B) and ORF9b1 (PDB ID 2CME, chain A) sequences (both parent and fragments) were determined using the JPred4 (http://www.compbio.dundee.ac.uk/jpred/) and PSIPRED webservers (http://bioinf.cs.ucl.ac.uk/psipred/). SPIDER2 predictions were determined from position-specific scoring matrices generated locally by running three rounds of PSI-BLAST\textsuperscript{27} on the UniRef90\textsuperscript{28} database from 7/2014 with up to 10,000 alignments with a maximum e-value of 0.05. For consistency, each residue from all three predictors were assigned one of three secondary structure annotations: “H” for α-helix, “E” for extended β-strand, and “C” for coil.

4.2  |  Secondary structure prediction accuracy calculations

Experimentally determined secondary structures of ORF9b and ORF9b1 were calculated locally using DSSP\textsuperscript{29} and simplified to the same three-state classification system as the secondary structure predictors: “H” for α-helix, “E” for extended β-strand, and “C” for all other DSSP annotations. Additionally, chain breaks were annotated as “-”. Sequences were aligned using the pairwise2.align.localxs function from Biopython\textsuperscript{30} with a gap-forming score of −1 and gap-elongation score of −0.5. Predicted/calculated secondary structures were then re-registered according to the Biopython sequence alignments. Secondary structure prediction accuracies were calculated using the \(Q_{\text{total}}\) (or \(Q_3\)) metric,\textsuperscript{31} in which experimentally determined and predicted secondary structures are compared one-by-one, residue-by-residue and normalized by the length of the sequences compared. Chain breaks were excluded from both scoring and normalization.

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AUTHOR CONTRIBUTIONS

Lauren Porter: Conceptualization; formal analysis; funding acquisition; investigation; methodology; visualization; writing-original draft; writing-review & editing.

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