Solution structure and RNA-binding of a minimal ProQ-homolog from Legionella pneumophila (Lpp1663)

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ABSTRACT
Small regulatory RNAs (sRNAs) play an important role for posttranscriptional gene regulation in bacteria. sRNAs recognize their target messenger RNAs (mRNAs) by base-pairing, which is often facilitated by interactions with the bacterial RNA-binding proteins Hfq or ProQ. The FinO/ProQ RNA-binding protein domain was first discovered in the bacterial repressor of conjugation, FinO. Since then, the functional role of FinO/ProQ-like proteins in posttranscriptional gene regulation was extensively studied in particular in the enterobacteria E. coli and Salmonella enterica and a wide range of sRNA-targets was identified for these proteins. In addition, enterobacterial ProQ homologs also recognize and protect the 3′-ends of a number of mRNAs from exonucleolytic degradation. However, the RNA-binding properties of FinO/ProQ proteins with regard to the recognition of different RNA targets are not yet fully understood. Here, we present the solution NMR structure of the so far functionally uncharacterized ProQ homolog Lpp1663 from Legionella pneumophila as a newly confirmed member and a minimal model system of the FinO/ProQ protein family. In addition, we characterize the RNA-binding preferences of Lpp1663 with high resolution NMR spectroscopy and isothermal titration calorimetry (ITC). Our results suggest a binding preference for single-stranded uridine-rich RNAs in the vicinity of stable stem-loop structures. According to chemical shift perturbation experiments, the single-stranded U-rich RNAs interact mainly with a conserved RNA-binding surface on the concave site of Lpp1663.

Keywords: Legionella; NMR structure; ProQ; RNA-binding protein; sRNA

INTRODUCTION
Posttranscriptional regulation of gene expression is an important mechanism for bacteria to quickly adapt to environmental changes. Particularly striking examples in this regard are host-activated bacterial virulence (Westermann et al. 2019), the response to oxidative stress or the regulation of iron metabolism (Altuvia et al. 1997; Massé and Gottesman 2002). Posttranscriptional gene regulation mechanisms in bacteria often depend on the action of transacting small noncoding regulatory RNAs (sRNAs). These sRNAs are 50–300 nt in length and recognize the mRNA of a target gene via a short complementary nucleotide stretch, the so-called “seed” region (Gorski et al. 2017). In principle, one sRNA can regulate different mRNA species, while one mRNA can be targeted by several different sRNAs. Many sRNAs are negative regulators that decrease mRNA stability or inhibit translation initiation (Nitzan et al. 2017). sRNA-stability and efficient complex formation with the target mRNA often depend on the RNA-binding protein (RBP) Hfq (Updegrove et al. 2016). Hfq is a hexameric, “doughnut-shaped” RBP that belongs to the family of Sm-like proteins (Sauter et al. 2003). Communoprecipitation experiments of affinity-tagged Hfq with bound RNAs coupled with next generation RNA sequencing approaches identified numerous sRNA binding-partners for Hfq in a number of different bacterial species (Holmqvist et al. 2016; Heidrich et al. 2017; Chihara et al. 2019). Thus, Hfq is apparently involved in many regulatory pathways. According to structural and biophysical studies of different Hfq proteins and Hfq–RNA complexes it acts as an RNA-interaction platform by providing three distinct RNA-binding surfaces (Schumacher et al. 2002, Link et al. 2009; Mikulecky et al. 2011; Someya et al. 2012; Panja et al. 2013; Dimastrogiovanni et al. 2014) and thereby promotes base-pairing between sRNAs and their mRNA targets.

A chaperoning activity with regard to sRNA–mRNA duplex formation and the stabilization of sRNAs against RNases was also reported as an important function for a
new class of bacterial RNA chaperones, the proteins of the
FinO/ProQ family (Arthur et al. 2003; Chaulk et al. 2010;
Smirnov et al. 2017). In contrast to the hexameric Hfq,
ProQ proteins are monomers and contain a conserved
FinO/ProQ core domain. This domain comprises five char-
acteristic α-helices that are connected by structurally well-
defined loops (Ghetu et al. 2000; Chaulk et al. 2010;
Gonzalez et al. 2017; Olejniczak and Storz 2017). Many
FinO/ProQ family members have additional amino- or car-
boxy-terminal domains and/or unstructured extensions,
which often contain a large number of basic amino acid
residues (Fig. 1; Supplemental Fig. S1A). E. coli FinO fea-
tures long α-helical extensions of the FinO/ProQ core
domain at its amino as well as its carboxyl terminus
(Ghetu et al. 2000). E. coli and Salmonella ProQ contain
an additional carboxy-terminal domain that is structurally
similar to eukaryotic Tudor domains. Tudor domains
form β-barrel-like structures and in eukaryotes interact
with other proteins by recognizing methylated arginine
or lysine residues (Lu and Wang 2013). It is very likely
that the FinO/ProQ-domains harbor the core RNA-binding
activity, but the additional domains/extensions also con-
tribute to RNA-binding or are required for duplex forma-
tion between sRNA and mRNA (Arthur et al. 2003;
Attaiech et al. 2016; Gonzalez et al. 2017; Stein et al.
2020). According to recent genetic experiments a con-
served positively charged area on the so-called “concave”
site of the ProQ/FinO core domain (Pandey et al. 2020)
is a central structural determinant for RNA-binding. How-
ever, how exactly FinO/ProQ proteins recognize their RNA tar-
gets is still not known.

Interestingly, some members of the FinO/ProQ family
were shown to bind various RNA targets, whereas other
members of the family bind only a small number of target
RNAs with very high specificity. The classical example in
the later regard is the E. coli FinO protein, which is part
of the F-plasmid encoded conjugation repression system
in enteric bacteria. This protein exclusively binds the
FinP sRNA and the complementary region of the corre-
sponding traJ mRNA (van Biesen and Frost 1994).
Similarly, the very recently identified FinO/ProQ-like pro-
tein FopA, which is plasmid encoded, binds primarily to
the Inc sRNA which is involved in plasmid replication in
Salmonella (Gerovac et al. 2020).

The E. coli ProQ protein was initially identified as a spe-
cific regulator of the osmoregulatory transporter ProP
(Kunte et al. 1999). However, decades later it was discov-
ered to be a general sRNA-binding protein interacting
with a large number of different sRNA targets (Smirnov
et al. 2016). In many cases the sRNA targets of ProQ are
not bound by Hfq and vice versa suggesting a functional
complementarity of the two proteins. Some sRNA–
mRNA pairs, however, are bound by both proteins impli-
cating competing roles for ProQ and Hfq in certain regula-
tory pathways (Melamed et al. 2020).

Regarding the function of ProQ/FinO proteins, it is un-
clear at the moment why proteins like FinO or FopA bind
specifically to only a few RNA targets whereas ProQ has
a much wider target range. RNase footprinting experi-
ments showed that both E. coli FinO and ProQ protected
the GC-rich 3′-terminator stem–loops and the uridine-rich
3′-end extensions in their target RNAs upon binding
(Arthur et al. 2011; Smirnov et al. 2017). Furthermore, a sin-
gle-stranded 3′-tail of at least 6 nt was required for high-affi-
ity RNA binding by E. coli FinO (Jerome and Frost 1999).
Recently, it was shown that enterobacterial ProQ also asso-
ciates with stem–loop structures at the 3′ end of mRNAs
and protects them against ribonucleases (Holmqvist
et al. 2018). Furthermore, a number of known ProQ
sRNA targets have single-stranded 3′-overhangs with an
average length of 4–6 nt (Olejniczak
and Storz 2017). A very recent study
aimed at identifying target RNAs for
the N. meningitidis ProQ homolog
NMB1681 by UV CLIP-seq also found
binding sites for this protein at mRNA
or sRNA 3′-ends including stem–loop
structures and U-rich single-stranded
regions (Bauriedl et al. 2020).

Interestingly, Legionella pneumo-
phila contains two ProQ-homologs
(Attaiech et al. 2016). RocC
(Lpp0148—regulator of competence
chaperone) specifically binds the
RocR sRNA and its plasmid-encoded
homolog—the RocRp sRNA—as its
main targets (Attaiech et al. 2016;
Durieux et al. 2019). The RocR sRNA
recognizes mRNAs encoding the pro-
teins of the DNA-uptake system and

FIGURE 1. Domain architecture of selected ProQ/FinO domain proteins. ProQ/FinO-proteins
often carry additional amino- or carboxy-terminal domains or extensions. The core domain
comprises ~100 amino acids and is mainly responsible for RNA-binding. The theoretical pl (iso-
electric point) values of the isolated domains as well as for the full-length proteins are indicat-
ed. (Ec) Escherichia coli; (Se) Salmonella enterica; (Nm) Neisseria meningitidis; (Lp) Legionella
pneumophila.
regulates their expression. The major determinant of RocR sRNA binding to RocC is its 3'-terminal stem–loop which is extended by a 6-nt-long single-stranded uridine-rich tail while the first two stem–loops of the RocR sRNA are involved in interactions with its target mRNAs (Attaiech et al. 2016).

The second ProQ homolog in *L. pneumophila* Lpp1663 (121 amino acids) only consists of a ProQ-domain and very short but rather basic amino- and carboxy-terminal extensions (Fig. 1; Supplemental Fig. S1A). It therefore represents a minimal version of a FinO/ProQ family member protein. So far, this protein has not been structurally and functionally characterized, but the presence of a FinO/ProQ domain suggests a likely function as an RNA-binding protein. Using NMR-spectroscopy, we determined a high-resolution structure of this protein in solution and characterized its general RNA-binding properties. We identified a potential RNA-binding site and could observe a clear preference for binding U-rich single-stranded RNAs. Furthermore, we show that a stem–loop structure in the vicinity of the U-rich region increases the affinity of Lpp1663 to RNA. With this study, we provide new insights into the structural determinants for RNA-binding of ProQ/FinO proteins that will help to understand the functional role of this protein domain in additional bacterial species.

**RESULTS AND DISCUSSION**

**Lpp1663 adopts a ProQ/FinO fold**

The NMR solution structure of Lpp1663 was solved using 1892 NOE-based distance restraints and 172 chemical shift derived torsion angle restraints using protocols described previously (Christ et al. 2012; Hacker et al. 2015). The calculated 20 structures with the lowest energy have an rmsd for the backbone heavy atoms in the structurally ordered regions (residues 12 to 113) of 0.4 Å (Table 1; Fig. 2A). According to $^1$H, $^{15}$N-HetNOE-data both the basic amino (residues 1 to 12) and carboxyl terminus (residues 113 to 120) are flexible (Fig. 2B). In addition, there is no evidence for even a transient interaction of the basic termini with the core of the protein since a deletion of the two termini (Lpp1663tr—aa 10 to 117) causes only minimal chemical shift perturbations for residues in the ProQ core domain (Supplemental Fig. S1B). The fold of Lpp1663 features 5 α-helices (α1–α5) in agreement with previous predictions based on chemical shifts (Immer et al. 2018) which are connected by four structurally very well defined rigid loops (L1–L4) that lack additional regular secondary structure elements. Similar to the other reported ProQ/FinO structures, the surface of Lpp1663 shows two positively charged patches, a larger one on the concave site and a smaller one on the convex site of the protein that might play a role in RNA-binding (Fig. 2C). Furthermore, the flexible amino and carboxyl termini are also rich in positively charged basic residues.

The high resolution structure of Lpp1663 as shown in Figure 2A strongly resembles the prototypic ProQ/FinO domain fold with a Ca- rmsd of 1.8 Å (80 residues) and 1.9 Å (91 residues) to the *E. coli* FinO and *N. meningitidis* NMB1681 structures, respectively (Fig. 3). The sequence identity of Lpp1663 is ~28% to *E. coli* FinO and ~25% to NMB1681. The *E. coli* ProQ amino-terminal domain (lowest energy structure of the NMR-determined structural bundle—Gonzalez et al. 2017) aligned with our Lpp1663 structure with an rmsd of 4.2 Å for 75 residue pairs, despite a higher sequence identity of the two proteins of 34%. The reason for the rather large rmsd between the *E. coli* ProQ amino-terminal domain and Lpp1663 are conformational differences in the loops connecting the core helices of the ProQ domain, as well as differences in the orientation of helices α2 and α5. It should be noted, however, that the structure of the *E. coli* ProQ NTD shows also rather high rmsd values in alignments with the X-ray structures of

| TABLE 1. NMR and refinement statistics of the Lpp1663 solution structure |
|---------------------------------------------------------------|
| **NMR constraints**                                         |
| Total NOE-based distance restraints                          | 1892 |
| Intraresidue                                                   | 458  |
| Sequential                                                     | 570  |
| Medium range                                                  | 400  |
| Long range                                                    | 464  |
| Dihedral angle restraints (φ + ψ) from TALOS+                | 172  |
| Number of restraints per residue                              | 17.2 |
| Number of long-range restraints per residue                  | 3.9  |
| **Residual restraint violations**                             |
| Distance violations/structure >0.5 Å                         | 0.9  |
| RMS of distance violation/constraint                          | 0.03 Å |
| Maximum distance violation                                    | 0.87 Å |
| Dihedral angle violations >10°                                 | 0    |
| **Model quality**                                             |
| Structures in final ensemble                                  | 20   |
| Target function value                                         | 3.76 ± 0.28 |
| Backbone rmsd: all/ordered residues                          | 2.5 Å/0.4 Å |
| Heavy atom rmsd: all/ordered residues                        | 2.7 Å/0.9 Å |
| **Ramachandran plot**                                        |
| Residues in most favored regions                             | 84.6%|
| Allowed regions                                               | 13.8%|
| Generously allowed regions                                    | 1.3% |
| Disallowed regions                                            | 0.3% |
| PDB entry                                                    | 6S10 |

*Ordered residues: 12–30, 33–120.*
E. coli FinO (4.5 Å, 92 residues) and NMB1681 (5.6 Å, 93 residues).

The similarity of the structure and surface properties of Lpp1663 and the proteins of the ProQ/FinO family lends additional support to the notion that Lpp1663 is also a functional homolog of these proteins. Due to the absence of additional domains or extended amino- and carboxy-terminal extension segments, it represents a minimalistic model for further investigations of the RNA-binding properties of the ProQ protein family.

Lpp1663 has a preference for U-rich single-stranded RNAs

Since Lpp1663 has not yet been functionally characterized there are no known target RNAs for this protein. RocC—the other ProQ-like protein in L. pneumophila—interacts specifically with the 3′-terminal stem–loop III of the RocR sRNA and its 3′ single-stranded extension (Supplemental Fig. S2A; Attaiech et al. 2016). An RNA containing stem–loop III and the 3′ single-stranded extension of RocR bound to Lpp1663 with a $K_D$ of 28 µM in ITC experiments (Supplemental Fig. S2B). Thus, the affinity of Lpp1663 for this RNA is $\sim$40-fold lower in comparison to the affinity of the FinO/ProQ core domain of RocC for which this RNA is a specific target (Attaiech et al. 2016). Interestingly, a short single-stranded oligomer with the sequence 5′-CCUUUCU-3′ which corresponds to the isolated single-stranded 3′-end of the RocR sRNA bound to Lpp1663 with a similar $K_D$ of 41 µM (Supplemental Fig. S2C). In contrast, the RocR stem–loop III without the 3′-terminal single-stranded extension did not bind to Lpp1663 (Supplemental Fig. S2D). Thus, the single-stranded 3′-tail of RocR is the major contributor to this interaction.

Based on these results as well as on the previously reported RNA-binding preferences for other members of the ProQ/FinO protein family we hypothesized that short single-stranded U-rich oligomers might be suitable model systems to derive initial insights into the RNA-binding mechanism of Lpp1663.

Initially, we tested the binding of a single-stranded oligo $U_6$ RNA to Lpp1663 using isothermal titration calo-

![NMR solution structure and electrostatic surface features of Lpp1663. (A) Lpp1663 has a ProQ/FinO fold with five central helices that are connected by four structurally well-defined loops. Shown is an overlay of the 10 final structures with an rmsd of 0.4 Å (ordered residues) that were deposited in the PDB under accession number 6S10. (B) The $^{1}H$, $^{15}N$ heteronuclear NOE of $^{15}N$-labeled Lpp1663 indicates the flexible amino- and carboxyl termini of the protein. Secondary structure elements are depicted by blue boxes. (C) The electrostatic surface potential of the lowest energy structure reveals two positively charged patches on either side of the protein, the concave and the convex site. Positive and negative charges are colored in blue and red. Amino- and carboxyl termini are indicated as N and C, respectively.](image-url)
there is any sequence preference in single-stranded RNA-binding we titrated Lpp1663 with oligo A₆, C₆ and a G-rich hexameric RNA (sequence GGAGGA) (Fig. 4). We avoided using an oligo G₆ RNA since such a sequence could form G-quadruplex structures. The affinity of Lpp1663 to oligo A₆ was decreased 2.5-fold (∼70 µM) compared to oligo U₆. The interaction of oligo C₆ and the G-rich RNA with Lpp1663 was either very weak or absent and therefore yielded no interpretable binding curves in the corresponding ITC experiments (Fig. 4). Since Lpp1663 has a clear preference for oligo U, and Lpp1663 seems to be able to discriminate between the two pyrimidine bases uridine and cytosine, the interaction of the two binding partners is not only due to electrostatic interactions between the positively charged residues of the protein and the negatively charged RNA phosphate backbone.

**The concave site of Lpp1663 harbors a potential RNA binding region**

To investigate which parts of the protein are involved in binding RNA by NMR-spectroscopy we titrated ¹⁵N labeled Lpp1663 with up to five equivalents of the unlabeled oligo N₆ RNAs and recorded changes in the chemical shifts of the amide group signals in ¹H, ¹⁵N HSQC spectra. (Fig. 5; Supplemental Fig. S3A). NMR titration experiments are carried out at significantly higher protein concentrations compared to our ITC experiments. Thus, they also yield information about interactions that were too weak to be analyzed reliably by ITC and consequently, spectral changes were observed in titrations with all four oligo N₆ RNAs.

In all titration experiments, a number of Lpp1663 amide group signals gradually changed their chemical shifts upon titration with the RNAs. For these signals, the difference in chemical shifts of the apo and the RNA-bound state is small compared to the rate of interconversion between the RNA-bound and the RNA-free state of the protein—they are in fast exchange on the NMR time scale. Other signals disappeared in the course of the titration (Fig. 5A). For these signals, the difference of their chemical shift in the apo and the RNA-bound state is larger and similar to the rate of interconversion between the RNA-bound and RNA-free state of the protein—they are in intermediate exchange on the NMR time scale. In order to map the observed NMR spectral changes on the surface of Lpp1663 we colored residues with disappearing signals red, signals with shift changes blue and those with signals unaffected by RNA addition gray. Residues with signals that could not be analyzed due to overlap with other signals and proline residues were colored black (Fig. 5B).

The most dramatic effects on the ¹H, ¹⁵N HSQC spectrum of Lpp1663 were observed upon titration with oligo U₆ as expected based on the dissociation constants measured by ITC. For many amino acid residues on a continuous surface at the concave site of the protein, the signals of the amide protons gradually weakened and disappeared upon increasing the RNA concentration indicating binding in intermediate exchange. This surface includes the amino-terminal half of helix α₂, helices α₃, α₄, and α₅, as well as loop L1 (Fig. 5B) and is surrounded by additional residues showing gradual chemical shift changes. Gradual chemical shift changes in agreement with fast exchange are also observed for the convex site of the protein upon titration with oligo U₆. However, since the concave and the convex surface of Lpp1663 are only separated by a single layer of α-helices it is unclear if these changes report on weak and probably unspecific binding to this surface or if they are due to indirect allosteric effects of RNA-binding to the concave surface. The titration experiments with oligo A₆ also reveal the concave surface of the protein as a hotspot for RNA-induced spectral changes. In contrast, the effects induced by addition of the G-rich oligomer and oligo C₆ are much more limited (Fig. 5B). In particular, for the G-rich oligomer the induced spectral changes are very minor
in agreement with a very weak and unspecific RNA–protein interaction.

Interestingly, the concave surface region containing the amino acid residues with the most pronounced spectral changes upon oligo U₆ titration contains many highly conserved residues. This includes L38, G41 and the partially conserved positively charged residue K39 in loop L1, Y76 and L77 in helix α4, and R86 in loop L4 (Olejniczak and Storz 2017) reinforcing the notion that this area represents the primary RNA-binding site of Lpp1663 (Supplemental Fig. S4A). Another residue in loop L1 with a signal disappearing upon titration with oligo U₆ RNA is the partially conserved K36. In contrast to the other affected residues in loop L1, the backbone amide group and the sidechain of this residue point toward the convex surface of ProQ. However, as mentioned above, this might indicate an allosteric conformational change in loop L1 in response to RNA binding rather than a direct interaction. An RNA binding surface on the concave site of the E. coli ProQ domain very similar to our observations was recently proposed by Berry and coworkers based on results from a bacterial three-hybrid mutational screen aimed at identifying amino acids important for RNA-binding in vivo (Pandey et al. 2020). Substitution of conserved residues with alanine as well as random mutagenesis revealed an essential role in RNA-binding for, e.g., G37, K54, R58, Y70, L71, R80, and D82 (E. coli ProQ numbering). These amino acids correspond to G41, K60, R64, Y76, L77, R86, and D88 in Lpp1663, which are all affected by oligo U₆ binding in the NMR titration experiments, respectively. Residue R80 (E. coli ProQ numbering)/R86 (Lpp1663 numbering) from this list warrants a more detailed discussion.

In the NMR solution structure of E. coli ProQ, the side chain of R80 points toward the convex site of the protein (Gonzalez et al. 2017). Since its mutation to A lead to a decrease in RNA-binding in the bacterial three-hybrid experiments Berry and coworkers assigned this residue to an RNA-binding site on the convex surface of the protein (Pandey et al. 2020). Since its mutation to A lead to a decrease in RNA-binding in the bacterial three-hybrid experiments Berry and coworkers assigned this residue to an RNA-binding site on the convex surface of the protein (Pandey et al. 2020). In our structural bundle representing the NMR solution structure of Lpp1663, the equivalent R86 side chain points toward the concave face of the protein in all structures (Supplemental Fig. S4B,C). Furthermore, in the two crystal structures of E. coli FinO and N. meningitidis NMB1681, the arginine side chain points toward the concave site as well, suggesting that this residue is part of the RNA binding surface on the concave face in the majority of ProQ-like proteins (Supplemental Fig. S4D).
We also tested the influence of oligonucleotide length on the binding affinity and the affected binding surface. Therefore, we titrated Lpp1663 with uridine-5′-monophosphate (UMP), oligo U₄ and oligo U₈ in ITC and/or NMR experiments. NMR titrations did not show evidence for even a transient interaction of UMP with Lpp1663 (Supplemental Fig. S3B). On the other hand, according to ITC experiments oligo U₄ with a $K_D$ of $31 \pm 3$ µM has almost the same affinity for Lpp1663 as oligo U₆ (Fig. 6). In contrast, extending the RNA by two additional uridine nucleotides to oligo U₈ resulted in a 3.5-fold increase in the affinity ($K_D = 8 \pm 1.0$ µM, Fig. 6). However, NMR-based titration experiments with the oligo U₄ and oligo U₈ RNAs and $^{15}$N- labeled Lpp1663 (Supplemental Fig. S3B) identified the same surface area on the concave site of the protein as the major RNA binding site (Supplemental Fig. S5).

To further support the importance of the positively charged surface patch on the concave face of the protein for single-stranded oligo U binding we mutated residues Y76 or R86 (Lpp1663 numbering) to A and characterized the affinity of the two mutant proteins for oligo U₈ by ITC (Supplemental Fig. S6A). Remarkably, both single point mutations Y76A and R86A completely abolished RNA binding. Importantly, both mutant proteins are folded properly according to their 1D $^1$H NMR spectra (Supplemental Fig. S6B). Our results agree well with the in vivo results of Berry and coworkers (Pandey et al. 2020) for E. coli ProQ, where the equivalent point mutations also abrogated RNA binding.

In all NMR-based titration experiments, we noticed that the chemical shifts of the amino acids in the positively charged flexible amino and carboxyl termini were not affected upon RNA addition and are therefore apparently not important for interactions with single-stranded RNAs. ITC titration experiments with the amino- and carboxy-terminally truncated protein variant (Lpp1663tr aa 10–117) confirmed that the affinity for oligo U₈ is virtually the same as for the full-length protein (Supplemental Fig. S7; $K_D = 10$ µM). These results also suggest that the

![Figure 5](https://example.com/figure5.png)

**Figure 5.** Lpp1663 interacts with RNA through its core domain and mainly with the concave face of the protein. (A) Sections of $^1$H, $^{15}$N HSQC spectra of $^{15}$N-labeled Lpp1663 titrated with oligo U₆, A₆, C₆, or a G-rich hexameric RNA. NMR signal assignments are given for residues undergoing substantial changes in signal intensity or chemical shift in the oligo U₆ titration. (B) Results of the NMR-titrations (three equivalents of RNA) were plotted on the surface of one of the lowest energy structures of Lpp1663. Changes in the chemical shift are indicated as the following: (red) peak disappeared, (blue) small shift, (gray) no change, (black) unclear. The most affected region is the concave face of Lp1663 as can be seen for oligo U₆ and oligo A₆. For oligo C₆ and the G-rich RNA, mostly minor chemical shift changes can be observed. The amino- and carboxyl termini are not involved in RNA-binding. Amino- and carboxyl termini are indicated as N and C, respectively.
unstructured termini do not contribute with unspecific electrostatic interactions to the binding of single-stranded RNA oligonucleotides.

The influence of RNA stem–loops on Lpp1663 RNA-binding

While Lpp1663 preferentially binds to single-stranded oligo U rather than to oligo A, C or a G-rich RNA and oligo U₈ binds Lpp1663 with a Kₐ in the low µM range this Kₐ is still ~10-fold higher than the Kₐ for the RocC/RocR stem–loop III interaction and significantly higher than the Kₐs reported for other ProQ/FinO like proteins for their targets (Attaiiech et al. 2016; Smirnov et al. 2016). This suggests that in biologically relevant Lpp1663–RNA interactions additional RNA elements might be needed to further contribute to the affinity. Thus, we tested the influence of adding a 3′-terminal stem–loop to oligo U on the affinity of the protein–RNA interaction. As there is no known biological target of Lpp1663, we chose the 3′-terminator hairpin of the RaiZ sRNA which is a verified target of Salmonella ProQ (Smirnov et al. 2017). We tested an RNA construct containing only the isolated RaiZ 3′-terminator hairpin (RaiZ hp) and hairpins with either a single-stranded 3′ oligo U₈ (RaiZ hp U₈) or A₆ tail (RaiZ hp A₆). The native RaiZ terminator hairpin stem is predicted to feature two terminal A:U Watson–Crick base pairs and is succeeded by an oligo U₈ 3′-single-stranded tail. The RNAs were folded properly according to their 1D 1H NMR spectra (Supplemental Fig. S8). In ITC experiments the isolated RaiZ hairpin (RaiZ hp) did not show binding to Lpp1663 (Fig. 7A). However, the RaiZ hp U₈ RNA bound Lpp1663 with a Kₐ of ~800 nM (Fig. 7B) which is ~10-fold lower than the Kₐ for single-stranded oligo U₈ and similar to the Kₐ observed for the biologically relevant RocC/RocR interaction (Attaiiech et al. 2016). In contrast, the RaiZ hp A₆ RNA did not show binding in ITC titrations (Fig. 7C). Thus, both the presence of the structured stem–loop element and the sequence of the 3′-terminal single-stranded tail are important determinants of RNA-binding by Lpp1663. In agreement with our results the importance of a structured hairpin element for RNA-binding to enterobacterial ProQ was postulated recently (Westermann et al. 2019).

For the E. coli FinO protein which binds specifically to its RNA-target—stem–loop II of the FinP sRNA—the presence of a 3′-single-stranded tail improved the affinity of this interaction ~10-fold compared to an RNA construct consisting only of stem–loop II (Jerome and Frost 1999). Interestingly, however, in this case the relative importance of the two RNA structural elements—the stem–loop and the single-stranded 3′-tail—for the binding affinity seems to be reversed compared to the interaction between Lpp1663 and RaiZ hp U₈. This might be due to the presence of the long amino-terminal α-helical extension of the ProQ core fold in FinO and its postulated interaction with the stem of the FinP sRNA stem–loop II (Arthur et al. 2011).

The results of the NMR titrations for RaiZ hp U₈ are very similar in comparison to the titrations with oligo U₈ (Supplemental Fig. S3C). The previously identified RNA binding surface on the concave face of Lpp1663 was again affected strongly in the titration with the RaiZ hp U₈ RNA. However, the surface area on the concave face of Lpp1663 affected by the presence of the RaiZ hp U₈ RNA (Supplemental Fig. S9) is slightly increased compared with the binding surface for oligo U₈. This suggests that the single-stranded oligo U stretch of RaiZ hp U₈ is bound similar to the isolated single-stranded oligo U RNA. The stem–loop region of the RaiZ hp U₈ RNA apparently only supplements the interactions with the single-stranded 3′-tail of the RNA. A computational model for the complex of FinO and FinP stem–loop II RNA based on SAXS, enzymatic probing and mutational data also suggested that the single-stranded 3′-tail of the FinP RNA is recognized by the concave face of the core domain, whereas the stem forms additional contacts to the long amino-terminal helical
extension of FinO (Arthur et al. 2011). For Lpp1663 the absence of chemical shift perturbations for residues in the flexible amino- and carboxy-terminal extensions in titration experiments with the RaiZ hp U₆ RNA suggest that here the short extensions do not form additional interactions with the stem part of the RNA. This is borne out by the results of an ITC titration experiment where the RaiZ hp U₆ RNA is added to the amino- and carboxy-terminally truncated variant Lpp1663tr (amino acids 10–117). The $K_D$ for this interaction is essentially the same as the one involving the WT protein ($K_D = 1.2$ µM, Supplemental Fig. S10). Thus, the flexible basic tails of Lpp1663 might be important for chaperoning RNA duplex formation as shown, for example, for the amino-terminal helix of FinO or the carboxy-terminal domain of *E. coli* ProQ (Arthur et al. 2003; Chaulk et al. 2011).

In conclusion, we show here that Lpp1663 structurally belongs to the ProQ/FinO family and is a genuine RNA-binding protein with a preference for RNA-substrates containing single-stranded U-rich regions in the vicinity of stem–loop elements. The primary RNA-binding surface is located on the concave face of the protein and includes a number of highly conserved amino acid residues such as Y76 and R86. Apparently, this RNA-binding site is present in other members of the FinO/ProQ protein family (Ghetu et al. 2002; Chaulk et al. 2010; Pandey et al. 2020). In particular, it was suggested to be the binding site for the 3’ single-stranded terminal tail in the interaction between FinO and the stem–loop II of the FinP sRNA (Jerome and Frost 1999). On the other hand, the comparison of the available structures and sequences for functionally characterized FinO/ProQ family members suggest that the presence of large structured or unstructured amino- or carboxy-terminal extensions of the ProQ core domain as observed in FinO, RocC or FopA is related to a high substrate specificity in these proteins. These extensions could either directly contribute to the recognition of specific RNA targets as suggested for FinO (Arthur et al. 2011) or they could sterically restrict the access to the core binding site by transiently folding back on the concave surface of the core domain. However, since no high-resolution structural information for an RNA/protein complex nor data about their intrinsic dynamics are available for FinO/ProQ-family members with specific RNA-substrate recognition, this currently remains speculation.

The genome of *L. pneumophila* harbors two proteins that are members of the FinO/ProQ family—RocC and Lpp1663. The function of RocC was demonstrated to be rather specific (Attaiech et al. 2016) and here we show that the specific sRNA target of RocC—the sRNA RocR—

![FIGURE 7. A hairpin containing RNA binds to Lpp1663 with enhanced affinity. (A–C) Secondary structure of the RaiZ hairpin variants used in this study and ITC thermograms of Lpp1663 titrated with these constructs. The six U and A residues that are not present in the RaiZ hp construct are labeled in red.](image-url)
is bound only weakly by Lpp1663. On the other hand, Lpp1663 is apparently able to bind with a biologically significant affinity to RNAs containing single-stranded U-rich elements next to structured stem–loops. Thus, Lpp1663 might therefore functionally resemble those FinO/ProQ family members with a broad substrate binding ability and a more general mRNA-protection and RNA chaperone function. However, a final assessment of the function of Lpp1663 will have to await the identification of its native RNA targets in vivo.

**MATERIALS AND METHODS**

**Cloning of Lpp1663 plasmids**

The Lpp1663 coding sequence was cloned into a pET11a vector with an amino-terminal hexahistidine tag and a TEV cleavage site as a synthetic gene (GenScript) with the codon usage optimized for *E. coli* expression.

Lpp1663 point mutations Y76A and R86A were obtained by site-directed mutagenesis as described previously (Liu and Naismith 2008). For the PCR reactions the following primer pairs were used: R86A_forward (CCGGCCGTGATTTATTGTTAAA CGAGGTTGATGTGGTT) and R86A_reverse (TCCACCGCCGGTGCAGTCCGTCTTGGA); Y76A_forward (CCGGCCGTGAGTGCCAGAAGCCGGAC) and Y76A_reverse (CTCAGGCGCGCCGG TAGCGGGCTATA). The introduced point mutation is underlined in the complementary region, the nonoverlapping primer pairs are highlighted in italic.

The PCR products were treated with DpnI (1 U/1 ng DNA) in cut smart buffer for 2 h at 37°C followed by 20 min of heat inactivation at 80°C. A total of 10 μL of the reaction mix was used for transformation of chemically competent *E. coli* DH5α cells. Plasmids were isolated from grown colonies and verified by sequencing.

**Protein expression and purification**

The full-length Lpp1663 protein, Lpp1663tr (residues 10 to 117), and the amino acid point mutants Lpp1663 Y76A and R86A, were expressed and purified as described in detail previously (Immer et al. 2018).

Briefly, protein expression was induced at OD600 ∼0.8 with 1 mM IPTG at 20°C for ~16 h in *E. coli* BL21(DE3) Gold (Agilent Technologies/Stratagene). Media were supplemented with ampicillin (100 μg/mL). For uniformly 15N- or 15N,13C-labeling, the protein was expressed in M9 minimal medium as described in detail previously (Cambridge Isotope Laboratories). Amino acid (0.13 g/L); and U-13C proline (0.1 g/L).

**NMR measurements and structure calculation**

NMR measurements were performed as described previously (Immer et al. 2018). All NMR spectra were recorded on Bruker AVANCE III HD 600, 700, and 800 MHz spectrometers equipped with cryogenic triple resonance probes at 298 K. 1H chemical shifts were internally referenced with 50 μM DSS (4,4-dimethyl-4-silapentane-1-sulfonic acid). The heteronuclear 13C and 15N chemical shifts were referenced indirectly with the appropriate conversion factors (Markley et al. 1998). All protein samples (400–500 μM protein concentration) were supplemented with 10% (v/v) D2O in NMR buffer (50 mM sodium phosphate buffer pH 6.5, 100 mM NaCl, 2 mM β-mercaptoethanol).

The backbone assignment was obtained using the following BEST-TROSY-based triple resonance 3D-experiments with 13C,15N uniformly labeled Lpp1663: HNCO, HN(CA)CO, HNCACB (Favier and Brutscher 2011); 1H,15N-HSQC and 1H,13C-HSQC spectra were recorded for α-15N-lysine, α-15N-tyrosine, α-15N-phenylalanine, and 13C-proline-labeled protein to verify the assigned backbone amide signals. Side chain signals were assigned using 3D HBHA(CO)NH, (H)CCH-TOCSY, H(C)CH-TOCSY, (H)C(CO)NH, and H(C)CO(NH) and H(C)CO(NH)NH. Distance restraints for the structure calculation were obtained by analyzing cross peaks in 13C-NOESY-HSQC (with the 13C offset and delays optimized for aliphatic and aromatic carbon nuclei in two separate experiments, mixing time 150 msec) and 15N-NOESY-HSQC (mixing time 150 msec) experiments (Sattler et al. 1999). All 15N heteronuclear NOE experiment of the amide resonances was recorded to identify flexible regions of the protein (Farrow et al. 1994).

Spectra were processed using the Bruker TOPSPIN 3.2 software and analyzed with CARA (Keller 2004). Torsion angle restraints were calculated with TALOS+ (Shen and Bax 2013) based on the chemical shift assignments. NOE distance restraints were automatically picked with the ATNOS/CANDID module from UNIO (Herrmann et al. 2002) and manually curated with CCPNmr Analysis (Vranken et al. 2005). The structure was calculated with CYANA (Güntert 2009), performing seven iterations with 100 initial structures and 10 final structures. Restrained energy refinement of the NMR structure was performed with the AMBER force field (Ponder and Case 2003) using the OPAL module (Luginbühl et al. 1996). The structure quality was assessed with the Protein Structure Validation Software (PSVS) (Bhattacharya et al. 2007). The electrostatic surface potential was calculated with the PDB2PQR web server (Dolinsky et al. 2004) and visualized with the APBS plug-in for PyMOL (Baker et al. 2001). All figures of the structures were prepared with PyMOL (The PyMOL Molecular Graphics System, Version 2.0, Schrödinger, LLC).

The NMR assignments and structure are deposited in the BMRB and the PDB under accession numbers 27453 and 6510, respectively.
Preparation of RNA

The following RNA sequences were obtained commercially from Dharmacon: RocR 3′ single-stranded tail (5′-CCUUCGGC-3′), oligo U5, U6, and U8, oligo A6, G-rich (5′-GGAGGA-3′), RaiZ hp (5′-AACCGCGCCUUUGGGCCGCUU-3′), RaiZ hp U5 (5′-AACCGCGCCUUUGGGCCGCUU-3′) and without (5′-GGUCAAUUGGCGACACACUGAUUGGCC-3′) the 3′ single-stranded tail were prepared by in vitro transcription with T7-RNA polymerase and purified as described in detail previously (Duchardt-Ferner et al. 2016).

NMR titrations

For NMR titrations, 1H,15N-HSQC of 80 µM uniformly 15N-labeled Lpp1663 were recorded in NMR buffer and titrated with up to three or five equivalents of RNA at 298 K. Chemical shift perturbations were analyzed manually with Cara (Keller 2004). Shift changes were categorized as the following: peak disappeared (intermediate exchange regime), peak shifted (fast exchange regime), no change, no analysis possibly due to signal overlap or residue is a proline and cannot be identified in a 1H,15N-HSQC spectrum. Residues that showed the same type of chemical shift change were highlighted on the surface of Lpp1663 in the same color.

ITC measurements

ITC measurements were performed at 298 K in NMR buffer using a MicroCal ITC200 calorimeter (Malvern Panalytical). Twenty-five, 50, or 80 µM protein samples were titrated with 0.375, 0.5, 1, or 2 mM RNA (obtained from Dharmacon as described above) by 19 serial injections of 2 µL at a stirring speed of 750 rpm. Data processing and determination of binding constants were performed with an Origin7 (OriginLab) based software provided commercially (Malvern Panalytical) using a one-site binding model. Experiments were performed in triplicates. The results are summarized in Supplemental Table 1.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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