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CHAPTER SIX

MOLECULAR MECHANICS OF RNA TRANSLOCASES

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Abstract

Historically, research on RNA helicase and translocation enzymes has seemed like a footnote to the extraordinary progress in studies on DNA-remodeling enzymes. However, during the past decade, the rising wave of activity in RNA science has engendered intense interest in the behaviors of specialized motor enzymes that remodel RNA molecules. Functional, mechanistic, and structural investigations of these RNA enzymes have begun to reveal the molecular basis for their key roles in RNA metabolism and signaling. In this chapter, we highlight the structural and mechanistic similarities among monomeric RNA translocase enzymes, while emphasizing the many divergent characteristics that have caused this enzyme family to become one of the most important in metabolism and gene expression.

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1. Introduction

Progress in the field of RNA remodeling has largely resulted from the multidisciplinary nature of experimentation in this area. A combination of genetics, biochemistry, and biophysical studies in many different laboratories has established the biological functions of many RNA remodeling enzymes, the unique structural units upon which the enzymes are built, and the specific chemical and structural determinants by which RNA translocases and helicases identify their nucleic acid targets. As a result, several unifying themes and structural classes have emerged. We have limited the scope of this review to focus on monomeric RNA translocases that are phylogenetically classified as belonging to “helicase superfamily 1” (SF1) or “helicase superfamily 2” (SF2), which are nonetheless quite similar in sequence and morphology.

To date, enzymes in these families translocate unidirectionally along RNA in a processive manner. In order to exemplify the three major types of translocation behaviors that are observed for these enzymes, we focus on three different enzymes that share structural similarities while displaying divergent functional behavior (Fig. 6.1). Specifically, we discuss the following: Nonstructural protein 3 from hepatitis C virus (HCV NS3), a 3' → 5'

![Figure 6.1 Cartoon representation of translocase activities. Depictions of diverse translocase activities are shown. These activities include (A) 3' → 5' translocation on ssRNA, (B) 5' → 3' translocation on ssRNA, and (C) translocation on dsRNA. In the absence of ATP, the helicases are static molecules (red ovals). Upon the addition of ATP, helicases begin to exert mechanical and dynamic movement (green ovals). Translocative behavior can lead to unwinding of RNA duplexes ((A) and (B)). However, it is worth noting that there are occasions in which translocation can occur in the absence of unwinding (C).]
ssRNA translocase; Upf1, a 5′ → 3′ ssRNA translocase; and retinoic acid inducible gene I (RIG-I), a dsRNA translocase. For each enzyme, we describe structural features of the translocative motor domains, and the contributions of additional domains that aid directional movement. We also discuss the molecular determinants for substrate recognition and translocation. These findings will be interpreted in light of the fact that most of these proteins function as components of larger macromolecular machines rather than in isolation (i.e., in the exon junction complex or within replication complexes). We highlight the examples where the presence or absence of additional cofactors alters enzymatic behavior, thereby providing insight into how the cell may use different strategies to regulate helicase activity in vivo.

1.1. The protein domains of an RNA translocase: The RecA motor fold and companion domains

Like their DNA-remodeling cousins, the core of RNA translocase enzymes is composed of two RecA-like domains (α/β domain; a.k.a. Rossman fold) that are linked in tandem (Fairman-Williams et al., 2010; Pyle, 2011; Singleton et al., 2007). ATP binds in the cleft formed between these two domains and conserved amino acid motifs responsible for ATP binding and hydrolysis line up along each RecA-like domain facing inward toward the cleft. Atop these two domains is the platform where RNA binds. Depending upon the structural features of additional domains outside the two RecA-like domains, the geometry of this platform can accommodate either ssRNA or dsRNA (Fig. 6.2).

Many helicase proteins belonging to the nonprocessive DEAD-box classification, which also hydrolyze ATP upon RNA binding, only have two RecA-like domains (Linder and Jankowsky, 2011). However, most translocative and processive enzymes have additional domains that potentiate important roles in substrate recognition and helicase function. Some of these domains play positive roles in conferring processive behavior (Domain 3 of NS3), some play autoinhibitory or regulatory roles to prevent processive behavior (Domains CH and 1B in Upf1), and some mediate complex functions such as the transmission of mechanical information over long distances (pincer domain of RIG-I) (Fig. 6.2). Further, some helicases, such as E. coli DbpA, use these extra domains to recognize a specific RNA tertiary structure in order to activate its ATPase activity (Tsu et al., 2001), whereas other helicases lacking these additional domains, such as S. cerevisiae Dbp5, require the binding of auxiliary cofactors (Gle1) and small molecules (inositol hexakisphosphate) to stimulate RNA release (Montpetit et al., 2011). Functional dissection of these additional domains or binding cofactors represents an active area of investigation by many laboratories.
2. Major Examples of Monomeric RNA Translocases

2.1. The NS3/NPH-II family of SF2 proteins: 3′ → 5′ translocation enzymes

Most of our knowledge about these enzymes stems from pioneering work on SF2 helicases of viral origins, primarily from work on the NS3 proteins from the Flaviviridae viruses, Dengue Virus and HCV, and NPH-II from Vaccinia Virus. The NS3 helicases from HCV and Dengue are believed to be important during replication, where they may play a role in unraveling the highly structured single-stranded genomes of these viruses. However, HCV NS3 is now known to play other roles in the life of the virus, including a major function in virus particle packaging and assembly (Jones et al., 2011; Ma et al., 2008). Thus, despite the apparent helicase activity of NS3 enzymes, their actual roles in the viral lifecycles are not firmly established. Similarly, the NPH-II helicases of pox viruses are thought to be required for efficient transcription and inhibition of R-loop formation (Gross and Shuman, 1996). However, there is increasing evidence that they may play a role in release of the polymerase.
The robust unwinding activities of NS3 and NPH-II enzymes have helped reveal the specific structural and chemical features of RNA that are important for translocation and unwinding. Through a combination of ensemble studies in which unwinding was used as a proxy for translocation, together with direct single-molecule studies of translocation, it has been established that these enzymes specifically engage backbone residues on single-stranded regions of RNA (Beran et al., 2006; Dumont et al., 2006; Kawaoka et al., 2004). ATP hydrolysis then promotes directional movement along this bound strand with concomitant displacement of the annealed strands. Indeed, robust helicases like NPH-II strip away annealed strands and even bound proteins with little to no regard for their chemical composition (Jankowsky et al., 2000, 2001). Importantly, translocation by these enzymes requires physical continuity of phosphodiester linkages on this tracking strand, and a basic site along the tracking strand are tolerated, thereby indicating that direct interactions with nucleobases are not essential for unwinding activity (Beran et al., 2006; Kawaoka et al., 2004).

HCV NS3 is a bifunctional enzyme that contains a helicase domain, and an appended serine protease domain that is important during early stages of viral protein processing (Raney et al., 2010). Although the protease and helicase domains are not known to function during the same stages of the viral lifecycle, they strongly influence each other’s functions. That is, the serine protease domain is essential for conferring strong binding to the helicase domain, and the helicase domain enhances proteolysis by the protease domain (Beran and Pyle, 2008; Beran et al., 2007). Thus, as in many helicases, NS3 has evolved a functional dependence on appended domains. The helicase domain of HCV NS3, as well as those for all other Flaviviridae NS3 proteins, is a three-lobed structure; two lobes are the RecA-like domains (HEL1 and HEL2) and the third lobe (Domain 3) is an all α-helical structure (Luo et al., 2008a; Yao et al., 1997; Fig. 6.2A). ssRNA binds in the cavity formed between HEL1 and HEL2 on the bottom and Domain 3 on the top (Appleby et al., 2011; Luo et al., 2008b). The geometry of the RNA binding cavity is such that it can only accommodate ssRNA, a finding that is consistent with direct binding studies demonstrating that NS3 binds tightly to single-stranded substrates and weakly to blunt, duplex substrates (Tai et al., 1996).

In the context of a model unwinding substrate, which usually consists of a “tailed duplex,” NS3 binds the single-stranded region, which serves as a launch pad for ATP-dependent tracking through the duplex portion of the substrate. NS3 binds ssRNA in an orientation whereby the phosphodiester backbone faces HEL1 and HEL2, and the nucleobases are displayed in the opposite direction toward Domain 3 (Fig. 6.2A). Enzymological studies and crystallographic investigations show that NS3 tracks along the phosphodiester backbone by forming two types of interactions: hydrogen bonds mediated by main-chain amide groups and contacts formed through polar
side chains (Appleby et al., 2011; Beran et al., 2006). Examples of interactions between the RNA backbone and main-chain amide groups include Val232 and Gly255 that extend from HEL1, and Lys371 and Arg393 that extend from HEL2. The use of polar side chains to form contacts is best illustrated through two evolutionarily conserved threonine residues (Thr269 from HEL1 and Thr411 from HEL2). For SF2 helicases, these two residues project from each RecA-like domain and act as dynamic pincers by forming hydrogen bonds with the nonbridging phosphoryl oxygen atoms (Fig. 6.3B). An additional hydrogen bond is established with the RNA backbone using the side chain of Thr416, a residue within motif V. However, this residue only engages the ssRNA backbone in the absence of ATP and disengages from the ssRNA in the presence of ATP. In the absence of ATP, Thr269 and Thr411 are spaced three nucleotides apart, allowing the protein to adopt a more open conformation. ATP binding induces closure of HEL1 and HEL2 by bringing them together in closer proximity such that the two threonine residues are now separated by two nucleotides.

Figure 6.3 Dynamic motions of HCV NS3, a 3′ → 5′ ssRNA translocase. (A) Domain motions of HCV NS3 in response to RNA and ATP binding. Upon binding ssRNA, HEL2 undergoes a small conformational rearrangement that allows for optimal positioning. In the presence of ATP, HEL1, and HEL2 move closer in proximity to each other to enclose the cleft. Domain motions were interpreted based upon aligning all structures to HEL1. (B) In the absence of ssRNA, the side chains of three threonine residues hydrogen bonds form direct contacts with the phosphodiester backbone of the ssRNA, whereas a sole tryptophan residue makes a stacking interaction with a nucleobase. Note that Thr269 and Thr411 are positioned three nucleotides apart. Upon binding of ATP and closure of the two RecA-like domains, Thr416 disengages from the backbone, and Thr269 and Thr411 are now spaced by only one nucleotide. Trp501 remains stacked with the nucleobase.
nucleotides (Fig. 6.3A). Upon ATP hydrolysis, the products ADP and inorganic phosphate dissociate from the ATP binding cavity, and HEL2 relaxes to move away from HEL1, thus effectively advancing the helicase by one nucleotide toward the 5′ direction of the ssRNA.

Cycles of ATP binding and hydrolysis directly influence the number of hydrogen bonds formed between the helicase and RNA substrate, thus resulting in alternating high- and low-affinity states. Most interactions are observed in the ATP-free state, which represents the high-affinity state. Upon binding ATP, the number of contacts between helicase and the RNA decreases (notably, Thr416 disengages from the RNA backbone) and resulting in a low-affinity state, which is consistent with previous direct binding studies demonstrating the helicase has a lower affinity for RNA in the presence of nucleotide.

The 3′ → 5′ directionality of movement is achieved in part through conserved hydrophobic residue Trp501, which extends from Domain 3 (Fig. 6.3B). This residue stacks upon the nucleobase at the 3′ end of the ssRNA and effectively serves as an anchor point for the helicase to move unidirectionally toward the 5′ end of the ssRNA rather than slipping backward. Interestingly, through the cycles of ATP binding and hydrolysis, the orientation of Trp501 remains stacked with the nucleobase (Appleby et al., 2011). Maintaining this interaction with the nucleobase is functionally significant for the translocation and possibly for unwinding mechanisms of NS3. Indeed, single-molecule experiments have suggested that Trp501 acts as a plowshare, dragging behind the translocating helicase (which moves along the tracking strand in one nucleotide increments as described above) (Myong et al., 2007). Specifically, the data supports a model where Trp501 maintains its interactions with the same nucleobase, while the helicase undergoes two forward cycles of ATP binding and hydrolysis; during the third cycle, the helicase accumulates enough tension in the system and causes Trp501 to spring forward along the tracking strand by three nucleotides concomitant unwinding of three base pairs. Importantly, no sequence–specific interactions are formed between the nucleobases on the ssRNA strand and residues from Domain 3. This lack of specificity likely contributes to the ability of the helicase to unwind substrates in a sequence-independent fashion.

Currently, there are no crystal structures available of bona fide helicases in complex with duplex RNA substrates (there are structures of RIG-I with RNA duplexes, but it is not a helicase, vide infra). However, unwinding is likely achieved by the steric pressure from a β-hairpin that projects from HEL2. This conserved structural element is present in other related helicases and is proposed to function like a wedge to splay the two composite strands apart (Buttner et al., 2007; Luo et al., 2008b). While the tracking strand is fed through the ssRNA channel described above, the displaced strand is directed away. However, the trajectory of the displaced strand is currently unknown and represents a potential area of research.
2.2. The UPf1 family of SF1 proteins: 5′ → 3′ translocation enzymes

Much less is known about the translocation and unwinding mechanisms of RNA helicases that move with 5′–3′ directionality. This is unfortunate because members of this class include helicases of viral origin such as SARS coronavirus, which had been shown to be unusually processive. The SARS helicase can efficiently unwind RNA duplexes of several hundred base pairs under single-cycle conditions (Ivanov and Ziebuhr, 2004). However, the most structurally and functionally characterized 5′ → 3′ translocation enzymes of the SF1 protein Upf1. Upf1 is a core component of the nonsense-mediated decay (NMD) machinery that detects and rapidly degrades aberrant mRNA transcripts (Conti and Izaurralde, 2005). Intriguingly, Upf1 is phylogenetically related to well-characterized DNA helicases such as Rep and PcrA, rather than to other SF2 RNA helicases.

Upf1 exhibits low ATPase and unwinding activities in isolation and requires the binding of additional NMD factors Upf2 and Upf3 to stimulate these activities (Chamieh et al., 2008). The structural basis for these functional observations was recently illustrated in a series of crystal structures of Upf1 in isolation and with various cofactors that alter its enzymatic activities (Chakrabarti et al., 2011; Cheng et al., 2007; Clerici et al., 2009). Like the NS3 helicase, Upf1 contains a core motor domain composed of two RecA-like domains (HEL1 and HEL2) (Fig. 6.2B). The N-terminus of Upf1 contains a zinc-knuckle domain (CH domain) that exerts an inhibitory effect in cis on the enzymatic activities of the protein. Between the CH domain and HEL1 is a domain composed of six antiparallel β-strands (Domain 1B), and a stalk domain composed of two α-helices. Projecting from HEL1 is an additional domain also composed of two α-helices (Domain 1C). Such modular domains embedded within the RecA-like domains are unique features that typify members belonging to SF1. Interestingly, ssRNA binds in an orientation atop the HEL1 and HEL2 platform of Upf1 similar to that seen for NS3. This is a surprising finding, given that one would expect that a helicase that translocates and unwinds with an opposite polarity as NS3 to position itself on the RNA in a reversed orientation. However, this similarity in binding polarity, despite opposite translocation directionality, has also been observed in comparisons of the structures of DNA helicases, RecD2 and PcrA (Saikrishnan et al., 2009; Velankar et al., 1999), and in the hexameric helicases, Rho and E1 (Enemark and Joshua–Tor, 2006; Thomsen and Berger, 2009). Results with Upf1 are therefore consistent with the notion that translocases do not “turn around” to run backward. Rather, they reverse their gears (Pyle, 2009).

At a molecular level, Upf1 contacts the ssRNA using main–chain amide groups and side–chain interactions that project from HEL1, HEL2, the stalk,
Domain 1B, and Domain 1C. The CH domain does not directly interact with the RNA but rather exerts an allosteric effect on RNA binding by impinging upon Domain 1B to contact two additional nucleotides at the 3' end of the RNA (Fig. 6.4B). Impinging upon Domain 1B likely prevents forward translocative movement and may account for the intrinsically low ATPase and unwinding activities of the full-length protein. Interestingly, the crystal structure of a Upf1 construct lacking its CH domain shows that Domain 1B rotates away from the 3' end of the nucleic acid. This rearrangement allows Upf1 to translocate processively with 5'→3' directionality. Domain motions were interpreted based upon aligning all structures to HEL1. (B) In the presence of the CH domain (green), Upf1 adopts a compact, globular conformation that prevents processive translocation activity. Inhibition can be partially relieved by the binding of the cofactor Upf2 (orange). Binding leads to a large conformational rearrangement of the CH domain. Whereas the CH domain was once positioned above HEL2, binding of Upf2 repositions the CH domain behind HEL1.

**Figure 6.4** Dynamic motions of Upf1, a 5'→3' ssRNA translocase. (A) Similar to HCV NS3, the two RecA-like domains of Upf1 close upon binding ATP. Domain 1B undergoes a small conformational change and rotates away from the interface of the two RecA-like domains. Upon binding ssRNA, Domain1B makes another conformational change to position itself away from the 3' end of the nucleic acid. This rearrangement allows Upf1 to translocate processively with 5'→3' directionality. Domain motions were interpreted based upon aligning all structures to HEL1. (B) In the presence of the CH domain (green), Upf1 adopts a compact, globular conformation that prevents processive translocation activity. Inhibition can be partially relieved by the binding of the cofactor Upf2 (orange). Binding leads to a large conformational rearrangement of the CH domain. Whereas the CH domain was once positioned above HEL2, binding of Upf2 repositions the CH domain behind HEL1.
In a cellular context, preventing the CH domain from impinging on Domain 1B is accomplished by the addition of Upf2 (Fig. 6.4B). The C-terminus of Upf2 binds directly to the CH domain of Upf1, and this interaction is sufficient to enhance the ATPase and unwinding activities of Upf1. From a structural viewpoint, the binding of Upf2 to Upf1 induces a large conformational change in the CH domain of Upf1. Whereas the CH domain was originally positioned atop the HEL2 domain, binding of Upf2 causes the CH domain to occupy a new position distal to HEL1. By occupying this new position, the CH domain can no longer impinge upon Domain 1B to interact with the 3’ end of the ssRNA. The repositioning of the CH domain is likely the structural basis for the functional observation that Upf2 enhances the biochemical properties of Upf1.

For future studies, it will be interesting to understand the coordinated domain motions of Upf1 in the presence of full-length Upf2 and Upf3 proteins or with an RNA unwinding substrate. Having such a global view may help uncover the biological significance of translocation and unwinding by Upf1 in the context of the NMD pathway. Further, it would be valuable to conduct experiments similar to those performed using NPH-II and NS3 proteins probing mechanistic features of substrate recognition. These studies would provide a basis to compare the similarities and differences between helicases that operate on the similar nucleic acid substrates but move in different directions.

2.3. Double-stranded translocation by monomeric RNA helicases: The RIG-I/Dicer family

While members of this class have been characterized biochemically, a structural view of any member of this class had long been elusive until recently. This was especially unfortunate given that members constituting this class of enzymes are involved in important functions such as innate immune signaling (RIG-I, and related proteins MDA5 and LGP2) and RNA interference (Dicer) (Bernstein et al., 2001; Yoneyama et al., 2004, 2005). In the absence of a crystal structure, it was difficult to understand how a helicase recognizes dsRNA, which remained a lingering question in the field. For example, does the dsRNA rest on top of the two RecA-like domains in an orientation similar to those seen for helicases that bind ssRNA? If so, is the binding groove sufficiently deep to discriminate dsRNA from ssRNA? For recognizing and binding a dsRNA substrate, is one strand preferred over the other strand? These questions were especially important to answer given that the RIG-I and Dicer proteins are phylogenetically related SF2 helicases, but function in strikingly different pathways: RIG-I translocates along dsRNA and functions in innate immune signaling, whereas Dicer binds and cleaves long double-stranded pre-microRNAs into short, double-stranded siRNAs. Though both proteins
are powered by the same helicase core motor, their divergent functions are
conferred by accessory domains flanking the N- and C-termini of the
respective protein. Fortunately, the structural basis for recognizing and bind-
ing dsRNA by RIG-I in the presence and absence of bound nucleotide was
recently shown in a series of publications from numerous groups (Civril et al.,
2011; Jiang et al., 2011; Kowalinski et al., 2011; Luo et al., 2011).

RIG-I recognizes and binds dsRNA using its central SF2 helicase motor
core and displays dsRNA-dependent ATPase activity (Gack et al., 2008). At
the N-terminus, RIG-I contains two tandem caspase activation and recruit-
ment domains (CARDs; CARD1 and CARD2) for signaling, and at the
C-terminus, it contains a domain important for recognizing 5′-triphosphate
moieties (the CTD) (Cui et al., 2008) (Fig. 6.2C). The CARD fold is
typically found in proteins that function in apoptosis and inflammatory
signaling pathways (Hofmann et al., 1997). Therefore, it was highly unusual
for a helicase to have such an accessory domain that functions outside the
boundaries of the Central Dogma. In the context of RIG-I, it was later
determined that the CARDs forms a platform to interact with other
CARD-containing proteins to propagate a signaling cascade (Gack et al.,
2008). The CTD recognizes the 5′ end of either ssRNA or dsRNA, though
it binds with the tightest affinity to duplexed substrates bearing a triphos-
phate group (Wang et al., 2010).

The recognition of triphosphorylated substrates is a defining feature of
RIG-I (Hornung et al., 2006; Pichlmair et al., 2006). As most cellular RNAs
are capped or modified at their 5′ ends immediately following transcription,
the chemical nature of free 5′ triphosphate groups are immediately recog-
nized by RIG-I as “nonself” RNA. It is generally accepted that such
“nonself” RNAs are generated when viruses hijack the cellular machinery
to create new strands of viral genomic RNA de novo using its viral polymer-
ase. This recognition event activates RIG-I to initiate a signaling cascade
through the CARDs to ward off the invading virus to maintain an antiviral
cellular state.

Translocation along RNA duplex by the helicase domain was demon-
strated using a single-molecule fluorescence approach (Myong et al., 2009).
Using a wide spectrum of protein and RNA constructs, the authors demon-
strated that RIG-I exhibits robust and repetitive translocation along duplex
RNA substrates. Notably, translocation activity occurred in the absence of
unwinding. This was consistent with previous studies showing that neither
RIG-I, nor its cousin MDA5 display strong helicase activity. Both of these
enzymes display ATPase activity that is specifically activated by duplex
RNA rather than single strands (Gee et al., 2008; Kang et al., 2002).
Functional dissection of the role for the CARD domains indicates that they
play an inhibitory role by negatively regulating translocation activity (Myong
et al., 2009) (Fig. 6.5B). Binding of duplex RNA partially alleviates this
inhibitory role to allow translocation. Interestingly, maximal translocation
activity was observed using triphosphorylated duplex RNA, thereby highlighting the specific structural and chemical features of the nucleic acid substrate requirement for RIG-I. Taken together, RIG-I uses its helicase domain to power translocation along dsRNA, potentially disrupting the viral replication machinery or assisting in the loading of additional RIG-I molecules.

From the collection of available recent RIG-I structures, we can now directly observe the coordinated motions of its multiple domains, which has led to new and unexpected structural findings. In addition to the CARDs, the canonical motor domains HEL1 and HEL2, and the CTD, RIG-I contains two domains that had never been appreciated previously. The
first domain is composed of five antiparallel \( \alpha \)-helices and is termed as the insertion domain (HEL2i). HEL2i exists as an independent structure projecting above from HEL2 and forms direct contacts with the backbone dsRNA. The second domain is composed of two \( \alpha \)-helices that form a V-shape, which we refer to as the pincer domain, which wraps around multiple domains in order to transduce mechanical signals throughout the protein.

In the absence of dsRNA, RIG-I adopts an open, extended conformation. CARD1 and CARD2 stack upon one another, and CARD2 forms extensive interactions with HEL2i that are stabilized by salt bridges and hydrophobic contacts (Kowalinski et al., 2011) (Fig. 6.5B). The interactions between CARD2 and HEL2i may partially explain the inhibitory role of the CARDs observed in the single-molecule fluorescence studies. In the presence of dsRNA, however, RIG-I collapses upon and encircles the dsRNA (Jiang et al., 2011; Luo et al., 2011). This compaction event is consistent with small-angle X-ray scattering (SAXS) data measuring the dimensions of these molecules in solution in the presence and absence of dsRNA. Though a structure of the full-length RIG-I protein with dsRNA is unavailable, SAXS measurements indicate that such a complex would place the CARDs away in solution and distal to the dsRNA binding interface. In fact, such a conformation might even be optimal for the CARDs to interact with other protein cofactors.

When binding an RNA duplex, it is important to distinguish one strand from the other. We refer to the RNA strand that lies across the HEL1 and HEL2 motor domains as the tracking strand (Fig. 6.2C). Interestingly, the tracking strand is in the same polarity to those seen for NS3 and Upf1, whereby the 5’ end of the tracking strand is located above HEL2, while the 3’ end is located above HEL1. We refer to the complementary sister strand as the activating strand since the 5’ end of this strand directly interacts with the CTD and would carry the triphosphate group that activates RIG-I translocation activity. The duplex RNA is oriented perpendicular to the plane of HEL1 and HEL2, and numerous polar interactions are formed between the 2’-hydroxyl groups of the ribose sugars and the phosphodiester backbone using main-chain amides and side-chain residues; hydrophobic interactions are only observed between the CTD and the 5’ end of the activating strand. The lack of base specificity likely allows RIG-I to translocate on diverse dsRNAs without sequence bias.

The bound nucleotide state directly influences the number of interactions made between RIG-I and the dsRNA. When ATP occupies the cleft formed by HEL1 and HEL2, these two domains move in close proximity to one another in order to prime the catalytic residues lining the cleft for ATP binding and hydrolysis. In this closed conformation, HEL1 and HEL2 form contacts with both the tracking and activating strands. However, in the absence of nucleotide, RIG-I adopts a more open conformation where
HEL2 moves away from HEL1 and disengages from contacting the dsRNA. Intriguingly, HEL1 maintains its grip with the dsRNA regardless of the nucleotide bound state. Similar domain movements are also observed with HEL2i. Using the face of one of its helices, HEL2i forms extensive contacts with both strands of the dsRNA in the presence of nucleotide; these contacts are lost, except one, in the absence of bound nucleotide (Fig. 6.5B).

The act of engaging and then disengaging from the dsRNA by HEL2 and HEL2i in response to the bound nucleotide state may directly reflect the RIG-I translocation mechanism. By maintaining a constant point of contact with the dsRNA through HEL1, RIG-I may use HEL2 and HEL2i to inch the protein forward along one strand of the dsRNA. Upon nucleotide binding, RIG-I maintains a tight grip on the dsRNA by forming direct contacts through HEL1, HEL2, and HEL2i. Following, ATP hydrolysis may permit RIG-I to loosen its grip on the dsRNA by allowing HEL2 and HEL2i to disengage from the dsRNA. Each cycle of ATP binding and hydrolysis would then propel RIG-I forward by one base-pair toward the 5' end of the tracking strand. Indeed, such directional bias along the tracking strand had been observed in a single-molecule setting. In a biological context, it is tempting to speculate that directional translocation along the tracking strand may serve to dislodge the viral replication machinery that would be actively synthesizing new viral RNA (i.e., the activating strand). However, whether RIG-I can remodel protein–RNA complexes, as had been shown for other helicases previously, remains the subject of future inquiry.

Despite having the numerous structural views of RIG-I now available, several outstanding questions remain. For example, given that the CTD has a tight affinity for the triphosphorylated duplex substrates, how does RIG-I disengage from that position to translocate along dsRNA? What are the roles of phosphorylation, ubiquitination, or tetraubiquitin binding in altering the biochemical activities of RIG-I? How do the CARDs interact with other proteins that have CARD domains, and what are the functional outcomes? Is there a role for ATP hydrolysis during the presentation of the CARD domains and in signaling?

3. Concluding Remarks

We now have complete structural views of three RNA translocases that move in different directions and function on different types of RNA. Despite the diversity in the directionalities and substrate specificities of these molecular motors, these enzymes share a common theme: they bind and hydrolyze ATP in the presence of a stimulating nucleic acid substrate. While a main research question is to decipher what ATP binding and hydrolysis
accomplish, an emerging theme appears to be that this orchestrated motion advances the protein forward by one nucleotide. It is thus tempting to speculate that perhaps any helicase containing these two RecA-like modules may be able to move along nucleic acid strands, regardless of the number of additional flanking domains. Nevertheless, there is still much to uncover what is underneath the hoods of Nature’s molecular engines, and future research will undoubtedly shed light on their diverse mechanical properties.

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