Double-Stranded RNA Sensors and Modulators in Innate Immunity

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Abstract

Detection of double-stranded RNAs (dsRNAs) is a central mechanism of innate immune defense in many organisms. We here discuss several families of dsRNA-binding proteins involved in mammalian antiviral innate immunity. These include RIG-I-like receptors, protein kinase R, oligoadenylate synthases, adenosine deaminases acting on RNA, RNA interference systems, and other proteins containing dsRNA-binding domains and helicase domains. Studies suggest that their functions are highly interdependent and that their interdependence could offer keys to understanding the complex regulatory mechanisms for cellular dsRNA homeostasis and antiviral immunity. This review aims to highlight their interconnectivity, as well as their commonalities and differences in their dsRNA recognition mechanisms.

Keywords
dsRNA, RIG-I-like receptor, protein kinase R, oligoadenylate synthase, dsRNA-dependent adenosine deaminase, RNA interference
INTRODUCTION

Detection of foreign nucleic acids is a central mechanism for innate immune defense in most organisms studied to date. Accumulating evidence suggests that the innate immune system can be stimulated not only by pathogen-derived nucleic acids but also by host-derived nucleic acids that signal pathologic cellular states. Such self-derived danger signals include altered structure, location, or loss of interaction partners of cellular DNA or RNA. One of the best-characterized foreign features of RNA is the double-stranded structure. Double-stranded RNAs (dsRNAs) often accumulate during viral infection as a result of viral RNA replication, but they are also present in uninfected cells to varying degrees. Much progress has been made in our understanding of how the innate immune system distinguishes between cellular dsRNA versus viral dsRNAs in normal physiological conditions, and how and when such discrimination breaks down, causing self dsRNA-triggered sterile inflammation. These investigations led to the identification of new sources of viral and cellular dsRNAs and a previously unappreciated network of dsRNA-binding proteins that regulate innate immunity. These proteins include classical innate immune molecules, such as RIG-I-like receptors (RLRs), protein kinase R (PKR), and oligoadenylate synthases (OASes), but also other dsRNA-binding proteins previously known to function in cellular dsRNA homeostasis without a direct connection to innate immunity.

In this review, we discuss several families of cellular dsRNA-binding proteins involved in antiviral innate immunity (Figure 1a, Table 1). We focus on the structural and biochemical mechanisms of how these proteins recognize dsRNAs and how the specific dsRNA binding mode allows discrimination between self and nonself dsRNAs, and between normal and altered self dsRNAs. We also discuss how these proteins influence each other’s functions in the context of antiviral immunity and cellular dsRNA biology. We begin by describing general structural principles of

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Abbreviations: ADAR, adenosine deaminases acting on RNA; dsRBD, dsRNA-binding domain; NTase, nucleotide transferase; OASes, oligoadenylate synthases; PAZ, Piwi Argonaute and Zwille; PKR, protein kinase R; RLR, RIG-I-like receptor; RNase, ribonuclease.
dsRNA-binding proteins in innate immunity. (a) Network of dsRNA sensors and modulators in innate immunity. The diagram below shows their interdependence. Blue arrows and red inhibition lines indicate reported stimulatory and inhibitory relationships, respectively. (b) Structures of dsRNA and dsDNA, which display A-form and B-form double helixes, respectively. (c) Watson-Crick base pair interactions, with edges facing major versus minor grooves indicated. Note that relative locations of hydrogen bond donors and acceptors differ for all four nucleotides in the major groove but are degenerate in the minor groove. Abbreviations: ADAR, adenosine deaminase acting on RNA; PKR, protein kinase R; RLR, RIG-I-like receptor.

dsRNA and dsRNA-protein interactions, and several important points to consider in investigating dsRNA-binding proteins.

**GENERAL STRUCTURAL PRINCIPLES OF dsRNA AND dsRNA-PROTEIN INTERACTIONS**

The structure of dsRNA deviates significantly from that of dsDNA (Figure 1b). dsRNA displays an A-form duplex with deep and narrow major grooves and shallow and wider minor grooves (Figure 1b). dsDNA, on the other hand, forms the B-form duplex, with wide major grooves and narrow minor grooves (1, 2). As such, the major groove of dsDNA is accessible for protein secondary structure, while that of dsRNA is generally not (1, 2). Since the major groove, not the minor groove, displays edges of bases distinct in all four nucleotides (3) (Figure 1c), proteins interacting with the dsDNA major groove have the potential to discriminate among the four nucleotides and bind DNA with high sequence specificity (4). By contrast, dsRNA-binding proteins primarily interact with the RNA backbone and the minor groove and are thus largely sequence independent. While dsRNA sequence can indirectly affect protein binding to a certain extent (5), sequence dependence of dsRNA-binding proteins is generally limited.
CHALLENGES IN STUDYING dsRNA-PROTEIN INTERACTIONS

It is important to recognize common challenges and misconceptions in studying dsRNA-protein interactions. First and foremost, RNA affinity should not be equated to functional relevance. While some of the dsRNA-binding proteins in this review have a significantly higher affinity for dsRNA, many also bind other types of nucleic acids with comparable affinity through non-specific electrostatic interactions. As such, efforts to identify ligands solely based on affinity are inappropriate for these proteins. This was particularly the case for MDA5, an RLR that recognizes dsRNA. MDA5 forms filaments and activates antiviral signaling pathways only upon dsRNA binding (6, 7), but it also binds single-stranded RNA (ssRNA) with equivalent affinity (6). In addition to the high-affinity interaction with ssRNAs, which are abundant in cells, highly cooperative filament formation of MDA5 disfavors enrichment of stimulatory dsRNAs by MDA5 pull-down or by other affinity-based methods (8). By contrast, a conformation-specific method directly utilizing MDA5 filament formation enabled selective isolation of its dsRNA ligands (8).

Another challenge relates to the method of RNA preparation and common presence of RNA by-products or multiple conformers. In vitro transcription with T7 phage RNA polymerase is one of the most common methods for preparing RNAs. But it is known to generate a variety of unintended by-products, one of which is dsRNA formed by intended sense RNA and fully complementary antisense RNA (9). This can lead to misdefining RNA specificity for a protein that is in fact dsRNA specific (10). Even if the RNA of the intended sequence can be obtained without the by-products, whether the prepared RNA adopts the desired RNA structure is of another general concern. For example, hairpin RNAs can open up and pair with the complementary regions in trans rather than in cis, leading to longer dimeric or multimeric dsRNA formation with distinct biological activities. In fact, such dimer formation by hairpin RNAs has caused confusion as to the RNA specificity of PKR, a dsRNA-dependent kinase (11, 12). In many cases, native gel purification, rather than denaturing gel purification, and careful diagnostic analyses are required to determine the precise identity of the RNAs under investigation.

Additionally, several biochemical and molecular biology tools that are well established for ssRNA and dsDNA are often unsuitable for or difficult to apply to dsRNA. For example, protein-RNA UV cross-linking approaches used for isolating in vivo ligands are often known to be inefficient for dsRNA-binding proteins (13, 14), although successful cases do exist (15, 16). The inefficient cross-linking is likely due to limited contacts between protein side chains and dsRNA bases. Reverse transcription is another example that is difficult to apply to dsRNA. Even with heat denaturation during primer binding, duplex structure can be restored during reverse transcription and can block the activity of the reverse transcriptase, which generally lacks the strand-displacement activity. Knocking down dsRNA by small interfering RNA (siRNA) is also challenging, as the actions of siRNAs require accessibility of the complementary region in the target RNA. Finally, RNase treatment is frequently used to examine whether certain biological events are dependent on RNAs. However, dsRNAs, unlike ssRNAs, are more resistant to RNases, possibly requiring different choices of RNase treatment conditions to examine dsRNA dependence.

The challenges described above, some more serious than others, demand careful examination of dsRNA-protein interactions with the understanding that dsRNAs have different physicochemical properties than ssRNA and dsDNA, and that most molecular biology tools are developed for ssRNA/dsDNA and may not be suitable for dsRNA. With these considerations in mind, we discuss seven families of dsRNA-binding proteins at the interface of innate immunity and cellular dsRNA biology (Figure 1a). We apologize in advance for the studies omitted in this review due to the limited space. More extensive reviews for each family of proteins are available elsewhere.
RIG-I-LIKE RECEPTORS

RLRs consist of RIG-I, MDA5, and LGP2 and are innate immune receptors that recognize viral dsRNAs during infection. Upon viral dsRNA binding, RIG-I and MDA5 activate the common downstream adaptor molecule MAVS. Activated MAVS then recruits multiple signaling molecules, including TRAFs, TBK1, and IRF3/7, leading to the transcriptional upregulation of type I interferons and other proinflammatory cytokines (17, 18). Unlike RIG-I and MDA5, LGP2 does not directly activate MAVS but is thought to positively and negatively regulate MDA5 and RIG-I (19–21), respectively, although additional regulatory activities were also reported (22). Despite sharing the same downstream signaling pathways, RIG-I and MDA5 play nonredundant roles by recognizing largely distinct groups of viral RNAs (17, 18). RIG-I recognizes dsRNA structure in a manner dependent on the dsRNA end structure, in particular the blunt end and 5′-triphosphate group (5′ppp) (10). 5′ppp is present in all nascent transcripts but is generally removed from cellular RNAs through normal 5′ processing (e.g., endoribonuclease cleavage and 7-methyl guanosine cap addition) in the nucleus. By contrast, many (but not all) viral RNAs do not undergo 5′ processing and harbor 5′ppp. When these viral RNAs form dsRNAs with blunt ends, as in copy-back defective interfering particles or in genome panhandles, they can robustly activate RIG-I signaling. By contrast, dsRNA recognition by MDA5 is independent of 5′ppp or dsRNA end structure but instead requires a long (generally ≥0.5–1 kb) duplex stem, as in the replication intermediates of picornaviruses. Note that the dsRNA length threshold for MDA5 is determined by filament assembly and disassembly kinetics and thus can change depending on the level of MDA5 and dsRNA (23, 24).

Detailed structural and biochemical studies revealed how individual RLRs differentially recognize dsRNAs and carry out their respective antiviral functions. We first describe the molecular mechanism of RIG-I, and then those of MDA5 and LGP2. In the absence of viral RNA, RIG-I is in the autorepressed state wherein the signaling domain, the N-terminal tandem CARD (2CARD), is prevented from activating MAVS (25). The autorepressed 2CARD is liberated when dsRNA binds the helicase and C-terminal domains (Hel-CTD) of RIG-I (26, 27), although ATP binding was also proposed to be important (25, 28). Multiple studies showed that a release of 2CARD is not sufficient for MAVS activation; it additionally requires a tetramerization of 2CARD (29). At least two non–mutually exclusive mechanisms have been proposed to stimulate 2CARD tetramerization. First, while RIG-I binds to the end of dsRNA as a monomer (to sense 5′ppp and blunt ends), it assembles a filamentous oligomer on longer dsRNAs through ATP-driven translocation (30–33) (Figure 2a). RIG-I filament formation in turn promotes the 2CARD tetramerization through a proximity-induced mechanism (30) (Figure 2a). Second, K63-linked polyubiquitin (K63-Ubn+), synthesized by the E3 ligase TRIM25 and RIPLET, also plays an important role in tetramerization of RIG-I 2CARD by wrapping around the core tetramer structure of 2CARD (34, 35) (Figure 2a). The 2CARD tetramer then acts as a helical template to nucleate the MAVS filament formation (36), which in turn serves as a signaling platform to recruit downstream signaling molecules (36, 37).

MDA5 has a similar domain architecture and signaling mechanism as RIG-I. MDA5 2CARD activates MAVS by forming an analogous oligomer, which in turn nucleates the MAVS filament formation. MDA5 also binds dsRNA using Hel-CTD and forms filaments along the length of dsRNA (Figure 2a). Unlike RIG-I, however, MDA5 does not bind dsRNA ends, and filament formation is obligatory for stable dsRNA binding and signal activation (6, 23). Furthermore, the MDA5 filament undergoes end disassembly during ATP hydrolysis (23), which results in a selective destabilization of the MDA5 filament on short dsRNA, and preferential accumulation of MDA5 on long dsRNA (23). This leads to selective recognition of long dsRNAs generated during viral
RLR and PKR. (a) Domain architectures and structures of human RIG-I and MDA5 in complex with dsRNA (PDB: 3TMI and 4GL2). RIG-I binds the dsRNA end as a monomer (as shown in the crystal structure), but on long dsRNA, it assembles into filamentous oligomers. Filament formation of RIG-I on dsRNA promotes 2CARD tetramerization through a proximity-induced mechanism. The K63-linked polyubiquitin chain also promotes 2CARD tetramerization. In contrast to RIG-I, MDA5 only binds the dsRNA interior, not the end, and its filament formation is obligatory. As with RIG-I, 2CARD tetramerization (with or without the K63-linked polyubiquitin chain) is required for antiviral signal activation. (b) Domain architecture and structures of PKR. The structure of PKR dsRBDs in complex with dsRNA is not available, but studies suggest that PKR dsRBDs bind dsRNA in a manner similar to other canonical dsRBDs. The RNA-free structure of PKR dsRBD1 (blue) (PDB: 1QU6) is overlaid on the structure of *Aquifex aeolicus* RNase III in complex with dsRNA (green) (PDB: 2EZ6). On the right is the structure of the PKR kinase domain (dark blue) in the active dimeric state and in complex with elf2α (gray) (PDB: 2A19). Analogous to RLRs, PKR utilizes dsRNA binding and the proximity-induced mechanism to dimerize and activate the kinase domain. Abbreviations: CTD, C-terminal domain; dsRBD, dsRNA-binding domain; PKR, protein kinase R; RLR, RIG-I-like receptor; Ub, ubiquitin.

Figure 2

replication and robust discrimination against short cellular dsRNAs during normal physiological conditions (24, 38). The third RLR member, LGP2, contains the helicase domain and CTD but lacks 2CARD and is thus unable to activate MAVS. Structural studies, however, suggest that LGP2 binds dsRNA in a manner similar to RIG-I (39). It preferentially binds dsRNA ends (albeit in a 5′ppp-independent manner), and at the same time forms filamentous oligomers upon ATP hydrolysis (39–41). Previous studies suggest that LGP2 sensitizes MDA5 to dsRNA ligands and facilitates MDA5 filament
formation by assisting the otherwise slow and inefficient nucleation process (42). Considering the high dsRNA affinity of LGP2 and its structural similarity to MDA5, LGP2 may serve as a nucleator for the MDA5 filament assembly, although direct interaction between LGP2 and MDA5 has not been demonstrated.

In all cases of RLRs, they recognize dsRNA structure and additional features, such as 5′ppp and dsRNA length, for robust discrimination between self and nonself dsRNAs. Despite these multilayered discriminatory mechanisms, accumulating evidence suggests that RLRs misrecognize self RNAs under various pathologic conditions. In mice, a single point mutation (G821S) in MDA5 leads to constitutive activation of its signaling activity and development of a multiple-organ autoimmune disease (43). In humans, mutations in at least seven different residues of MDA5 were shown to cause a broad spectrum of autoinflammatory diseases, such as Aicardi-Goutières syndrome and systemic lupus erythematosus (44–46). While some studies have suggested the possibility of ligand-independent activation of MDA5 (47), others have reported the role of endogenous dsRNAs, in particular those formed by inverted repeat Alu elements (IR-Alus), in constitutive activation of MDA5 (8). Similar IR-Alu-mediated activation was observed for wild-type MDA5 under the condition where the dsRNA-modifying enzyme ADAR1 (adenosine deaminase acting on RNA 1) is deficient (discussed below) (8, 48). Recent studies showed that a chemotherapeutic agent, 5-aza-2′-deoxycytidine, also activates MDA5 by upregulating endogenous retrovirus dsRNAs, leading to sterile inflammation that is beneficial for the subsequent cancer immunotherapies (49, 50). A more recent study showed that mitochondrial dsRNAs accumulated under pathologic conditions can be released to the cytosol and function as an additional source of endogenous ligands for MDA5 (51).

Several point mutations in RIG-I were also found to constitutively activate antiviral signaling, causing another autoinflammatory disease, Singleton-Merten syndrome (52, 53). Misrecognition of endogenous RNAs was also proposed to underlie aberrant activation of RIG-I, but the identity of the endogenous RNA ligand is less clear. Wild-type RIG-I can also be activated by ionizing radiation (IR) cancer therapy (54). IR was proposed to induce mislocalization of U1 and U2 small nuclear RNAs (snRNAs) and subsequent misrecognition by RIG-I. However, the precise nature of the RIG-I-stimulatory elements in U1/U2 snRNAs is yet unclear, as U1 and U2 snRNAs harbor tri-methyl-guanosine cap, not 5′ppp. In the context of breast cancer cells, RIG-I was found to be activated by 5′ppp-harboring transcripts of RNA polymerase III (RNAP III). It was proposed that cancer cell–stromal cell interactions upregulate RNAP III activity, leading to abnormal accumulation of RNAP III transcripts, such as RNA7SL1, without corresponding upregulation of its protein partner that normally shields 5′ppp (55, 56). Similar unshielding of 5′ppp-containing RNAP III transcripts was proposed for the activation of RIG-I during infection with herpes simplex virus 1, Epstein-Barr virus, and influenza A virus (57). Additionally, multiple other conditions were also shown to generate altered self RNAs, by the action of RNase L (58, 59) or inaction of the RNA exosome (RNase) component, SKIV2L (60), thereby triggering RIG-I signaling. In most cases, however, the proposed RNA ligands are not known to harbor canonical RIG-I-stimulatory elements, i.e., 5′ppp and blunt-ended dsRNA structure. Mechanistic details of how these RNAs stimulate RIG-I remain to be further investigated.

RLRs can also function beyond antiviral signaling pathways and impact the functions of other dsRNA-binding proteins. RIG-I and MDA5 were found to remodel dsRNA-protein complexes (dsRNPs) in an ATP-dependent manner (61). This remodeling activity can result in displacement of viral proteins from dsRNAs, exposing otherwise sequestered dsRNA to the host immune system, such as RLRs and PKR (61). In line with this observation, RIG-I and MDA5 were shown to have a signaling-independent antiviral activity against a broad range of viruses (61–64). Whether LGP2 displays a similar dsRNP-remodeling activity is as yet unclear, but recent studies have also
suggested LGP2 activities beyond antiviral immunity. LGP2 inhibits Dicer-dependent processing of pre-microRNA (miRNA) and long dsRNA, thereby suppressing biogenesis of miRNA and siRNA, respectively (65, 66). These observations together suggest that RLRs have multiple functions in antiviral immunity and host dsRNA biology beyond simply activating the antiviral signaling pathway.

**PROTEIN KINASE R**

PKR is a dsRNA-dependent protein kinase and is transcriptionally upregulated by interferon. PKR normally exists in the latent form and is activated by dsRNA binding (67). One of the best-characterized targets of PKR is eIF2α, the α subunit of the eukaryotic translational initiation factor eIF2. eIF2 delivers the initiator tRNA to the small ribosomal subunit and is essential for cap-dependent translation. Phosphorylation of eIF2α prevents recycling of eIF2 and suppresses cap-dependent translational initiation. Thus, activation of PKR leads to the global shutdown of protein synthesis and inhibition of viral replication, cell cycle progression, and cell growth.

PKR consists of two tandem repeats of dsRNA-binding domains (dsRBDs) and a kinase domain (68, 69) (Figure 2b). dsRBDs are one of the most common dsRNA-binding protein motifs. Multiple structures of dsRBDs with and without dsRNA are available (5, 70) that show that dsRBDs have a conserved protein core structure and a common dsRNA-binding mode. dsRBD binds to one face of dsRNA primarily through phosphate and ribose backbones of two adjacent minor grooves (Figure 2b). Consistent with the backbone-binding mode, most dsRBDs, including PKR dsRBDs, interact with dsRNA in a sequence-independent manner (71). Studies showed that while the first dsRBD acts as the primary dsRNA binder on short dsRNAs (72, 73), both dsRBDs are required for high-affinity interaction with longer dsRNAs and dsRNA-dependent activation of the kinase domain (see below) (73, 74).

Earlier studies suggested that PKR exists in the autorepressed state in the absence of dsRNA, and dsRNA binding releases autorepression, thereby activating the kinase activity (75, 76). The second dsRBD was proposed to mediate autorepression by directly binding to the kinase domain. However, other studies have argued against the autorepression model and supported the role of dsRNA-induced dimerization as a mechanism for PKR activation. For example, biophysical analysis showed that latent PKR is in the extended monomeric conformation (77), inconsistent with the folded autoinhibited conformational state. Deletion of dsRBDs does not constitutively activate PKR, while replacement of dsRBDs by an artificial dimerizing domain does (78). The minimal dsRNA length (∼33 bp) required for PKR activation also coincides with the minimal length stably recruiting two PKR molecules (74). Additionally, an excess amount of stimulatory dsRNA, which promotes monomeric PKR binding but suppresses PKR dimerization, was found to inhibit PKR activity (79).

Structures of the kinase domain further showed a detailed picture of the dimerization-induced activation of PKR (Figure 2b). Although the structure of inactive, monomeric PKR is not available, the structure of the active, dimeric PKR kinase domain suggested that the dimerization of the N-terminal lobe of the kinase domain likely induces conformational changes in the αC helix and the activation segments in a manner similar to other well-characterized kinases (69, 80) (Figure 2b). The structure also showed how the autophosphorylation of Thr446 in the activation segment further helps stabilize the active site conformation (80, 81) (Figure 2b). Altogether, these observations support the model where dsRNA binding brings two or more PKR molecules in close proximity to allow dimerization of the kinase domain (Figure 2b). Once dimerized, the kinase domain autophosphorylates in αC, which further stabilizes the dimeric structure even in the absence of dsRNA and fully activates kinase activity (78).
Accumulating evidence suggests that PKR can tolerate a certain level of structural irregularities (e.g., mismatches, bulges) in dsRNA (79) and can be activated by cellular dsRNAs under various physiological conditions. For example, in the absence of ADAR1, cellular dsRNAs formed by unmodified IR-Alu were proposed to activate PKR (48). Similarly, breakdown of the nuclear membrane during mitosis and consequent exposure of an excess amount of nuclear IR-Alus were also proposed to activate PKR (82). Additional sources of endogenous ligands for PKR include cellular small nucleolar RNAs (snoRNAs) (83) and dsRNAs formed by mitochondrial sense and antisense transcripts (84), although mechanistic details of how these RNAs access cytosolic PKR remain unclear. Intriguingly, RNAs with short stem loops were also shown to activate PKR in a 5′ppp-dependent manner (85). This observation raises the possibility that PKR could be activated by a broader range of cellular RNAs, for example 5′ppp-containing RNAP III transcripts, as proposed for RIG-I (55, 57).

Innate immune functions of PKR appear to be complex. Several studies showed that PKR enhances the antiviral signaling activities of RLRs against certain viruses (86–88). Multiple, mutually nonexclusive mechanisms have been proposed. In one mechanism, PKR phosphorylates molecules involved in the signaling pathway, such as IkB (89), thereby altering their activities and synergizing with RLR signaling. However, it remains controversial whether PKR directly phosphorylates those signaling molecules or whether the effect is indirect (90, 91). Another proposal is that PKR amplifies RLR signaling by inducing formation of stress granules, the cytosolic nonmembranous granules formed upon dsRNA stimulation and enriched for RLRs and other signaling molecules (88, 92). However, there are cases where RIG-I and MDA5 are recruited to stress granules without activating antiviral signaling (93), suggesting that stress granule localization is not sufficient. PKR was also found to promote IFN-α/β production by stabilizing their mRNAs, rather than by upregulating their transcription (86), although the underlying mechanism for this observation is also unclear. Finally, PKR was proposed to exert its impact on RLR through translational suppression (94, 95). This proposal is supported by the observation that chemical inhibition of protein synthesis by cycloheximide also amplifies RLR signaling. All in all, PKR clearly plays an important role in antiviral immunity and cellular dsRNA biology, not only by linking dsRNA detection to translational inhibition but also by stimulating the antiviral signaling pathways.

**Oligoadenylate Synthases and RNase L**

OASes are a family of interferon-inducible enzymes that synthesize 2′−5′ phosphodiester–linked oligoadenylates (2−5An) upon binding to dsRNA (96, 97). 2−5An function as a second messenger activating latent endoribonuclease RNase L by inducing its homodimerization or oligomerization (98). Activated RNase L cleaves unpaired RNA with minimal sequence specificity, targeting a wide range of RNAs, including rRNAs, tRNAs, other noncoding RNAs, and mRNAs, from both viruses and host. Accordingly, activation of the OAS–RNase L system leads to global suppression of protein synthesis, inhibition of cell growth, and induction of cell death, effects analogous to those of PKR, but more rapid and possibly with a broader impact (96, 97).

There are four isoforms of OASes in humans (Figure 3a). OAS1–3 are enzymatically active, while OASL is not. OAS1, 2, and 3 respectively harbor one, two, and three tandem repeats of a nucleotidyl transferase (NTase) domain that belongs to the family of template-independent nucleotide polymerases (96). Structural studies showed that OAS NTases bind one side of dsRNA (99, 100) in a manner analogous to that of dsRBDs (Figure 3a). Unlike dsRBDs, however, OAS NTases undergo a conformational change upon dsRNA binding, which activates its enzymatic activity. More specifically, dsRNA binding organizes the three conserved aspartic acids in the active
OASes and ADARs. (a) Domain architectures and structures of human OAS1–3 and OASL. The structure of OAS1 in complex with dsRNA (PDB: 4IG8). The OAS1 structure is also superposed onto cGAS in complex with dsDNA (PDB: 6CT9) and OAS3 (first pseudo-NTase domain) in complex with dsRNA (PDB: 4S3N). The second UbL domain of OASL (green, PDB: 1WH3) is superposed onto the structure of Ub (dark blue) on the right. (b) Schematic of adenosine-to-inosine modification and domain architectures of human ADAR1–3. The structure of dsRBD1–2 of ADAR2 in complex with dsRNA (PDB: 2L3J). Structures of ADAR1 ZBD1 in complex with dsRNA in the Z-form (PDB: 2GXB) and the deaminase domain of ADAR2 (PDB: 5HP2) are also shown on the right. Abbreviations: ADAR, adenosine deaminase acting on RNA 1; dsRBD, dsRNA-binding domain; NTase, nucleotidyl transferase; OAS, oligoadenylate synthase; PDB, Protein Data Bank; Ub, ubiquitin; ZBD, Z-DNA-binding domain.

site for coordinating the magnesium ions required for catalysis (99). The mechanism by which OASes couple dsRNA binding to 2–5′ AN synthesis is similar to that of cGAS (Figure 3a), a cytosolic DNA sensor that catalyzes synthesis of 2′–3′ cGAMP upon dsDNA binding (101, 102). This suggests a conserved mechanism for double-stranded nucleic acid recognition and antiviral signaling.

OAS2 and OAS3 have two and three copies of NTase domains, but only one domain is catalytically active; the other domains (pseudo-NTase) lack the catalytic triad in the active site (103, 104). The pseudo-NTase domains, however, appear to have important functions. For example, the first pseudo-NTase domain in OAS3 binds dsRNA using the same binding mode as the catalytically active NTase domain of OAS1 (105, 106) (Figure 3a). It is possible that the second pseudo-NTase domain of OAS3 also binds dsRNA, thereby increasing dsRNA affinity of OAS3. Intriguingly, OAS3 responds to dsRNA more efficiently than OAS1 and OAS2 and serves as the primary upstream receptor for RNase L during viral infection (107). OAS3 was also found to preferentially recognize long (>50 bp) dsRNAs, perhaps reflecting linear arrangements of the three NTase domains along dsRNA (103).
OASL, an enzymatically inactive member of the OAS family, was also shown to display broad antiviral activity (108). OASL contains a single pseudo-NTase (Figure 3a), which also appears to bind dsRNA in a manner similar to OAS1 (109). Following the pseudo-NTase domain are the two tandem repeats of ubiquitin-like domains (UbLs, Figure 3a) (110). Both pseudo-NTase and UbLs are required for the observed antiviral activity of OASL (111). A previous study suggested that UbLs of OASL can functionally substitute K63-Ubn in inducing RIG-I 2CARD tetramerization (112), although the nature of this interaction remains to be further characterized. In mice, Oasl1 and Oasl2 are related to human OASL, with 70% and 50% sequence identity, respectively. While Oasl2 is necessary for efficient RIG-I signaling, as with human OASL (112), Oasl1 binds the 5′ untranslated region (UTR) of IRF7 mRNA, inhibits its translation, and negatively regulates antiviral immunity (113). This observation raises the possibility that human OASL plays a more complex role than simply assisting RIG-I.

Accumulating evidence suggests that the OAS–RNase L system has virus-independent activities. RNase L was found to regulate the steady-state mRNA levels of certain genes involved in cell growth, proliferation, and differentiation (114–116). Whether this basal activity of RNase L requires 2–5A and upstream OAS enzymes is unclear. The lack of dsRNA modification by ADAR1 (see the next section) was also shown to activate RNase L (117), causing cytotoxicity in an RNase L–dependent manner. In this case, RNase L activation is most likely mediated by upstream OASes (in particular OAS3) and their activation by unmodified cellular dsRNAs (e.g., IR-Alus). Intriguingly, Oas1 was found to be involved in normal developmental process (118), and Oas1 binds the 5′ untranslated region (UTR) of IRF7 mRNA, inhibits its translation, and negatively regulates antiviral immunity (113). This observation raises the possibility that human OASL plays a more complex role than simply assisting RIG-I.

ADENOSINE DEAMINASES ACTING ON RNA

ADARs are a class of enzymes that modify adenosines (A) to inosines (I) in dsRNAs (120–122) (Figure 3b). Since inosine is recognized as guanosine by reverse transcriptase and ribosomes, ADARs can be mutagenic when the modification occurs in coding regions. In noncoding duplexed regions, A-to-I modification replaces the A:U base pair by the weaker A:I base pair interaction (Figure 3b), resulting in weakening or partial melting of the dsRNA structure (120–122). Both the mutagenic activity and the duplex-melting activity have a profound impact on dsRNA biology and innate immunity.

Humans have three ADAR proteins, ADAR1–3, all of which have tandem repeats of RNA-binding domains [Z-DNA-binding domain (ZBD), dsRBD, and/or arginine-rich domain] followed by the deaminase domain (Figure 3b). ZBDs and arginine-rich domains (R-domains) are specific to ADAR1 and ADAR3, respectively, while dsRBD repeats are common in all three ADAR proteins. While dsRBDs are thought to bind dsRNAs with little or no sequence specificity, structures of ADAR2 dsRBDs with a hairpin dsRNA (Figure 3b) suggested that ADAR2 dsRBDs can recognize limited sequence information both through the minor groove contact and through indirect sensing of structural irregularities embedded within the hairpin dsRNA (5, 123). ZBDs of ADAR1 are also known to bind dsRNA in a sequence-dependent manner, displaying higher affinity for purine-pyrimidine repeat sequences that readily form left-handed Z-forms (124) (Figure 3b). The R-domain, on the other hand, appears to allow sequence- and structure-independent electrostatic interaction with nucleic acids (125).

The deaminase domain also has an intrinsic affinity for dsRNA with limited sequence preference. The crystal structures of ADAR2 showed that the deaminase domain binds dsRNA with a
highly localized distortion in RNA, causing flipping of the target adenosine base out of the RNA duplex and its insertion into the active site (126, 127) (Figure 3b). While the mechanism for base flipping appears somewhat analogous to those of DNA methyltransferases (MTases) (128), ADAR2 and the DNA MTases flip their target bases through minor and major grooves, respectively, reflecting their differences in dsRNA- and dsDNA-binding modes. As the sequence nearby the target site would likely affect the efficiency of target base flipping, the deaminase domain, in addition to the other RNA-binding domains, was proposed to influence the site selectivity of ADARs (127).

ADAR1–3 have largely distinct groups of dsRNA substrates and expression patterns (129). ADAR1 primarily modifies dsRNAs in noncoding regions, such as ∼300-bp IR-Alus in introns and 3′ UTRs of mRNAs (129, 130). By contrast, ADAR2 is responsible for modifying small hairpins in coding regions (129). ADAR1 and ADAR2 also have largely distinct expression patterns; ADAR1 is ubiquitously expressed, while ADAR2 is restricted to the central nervous system. The observed difference in their target sites, however, appears to reflect their intrinsic difference in RNA specificity rather than expression patterns. In contrast to ADAR1 and ADAR2, ADAR3 lacks enzymatic activity despite harboring catalytic residues, and was proposed to function primarily as an inhibitor of ADAR2 (129, 131).

Consistent with the distinct modification targets and expression patterns, ADAR1 and ADAR2 have different biological functions. ADAR2 knockout causes postnatal seizures and early death in mouse, but this phenotype can be rescued by a single A-to-G substitution in one of the ADAR2 target sites in the Gria2 gene (132). By contrast, ADAR1 deletion is embryonically lethal (133–135). ADAR1 has two isoforms, constitutively expressed p110 in the nucleus and interferon-inducible p150, that are in both the cytosol and the nucleus (Figure 3b). Similar embryonic lethality was observed with selective deletion of p150 (136) or mice with a catalytically deficient knock-in mutation (137). The embryonic lethality of ADAR1 knockout in mouse was largely rescued by double knockout of MDA5 or MAVS, but not RIG-I (136–138). In humans, loss-of-function mutations in ADAR1 cause Aicardi-Goutières syndrome (139), the inflammatory disease that can also be caused by gain-of-function mutations in MDA5 (44). Further analysis showed that lack of A-to-I modification leads to stabilization of cellular dsRNAs (in particular IR-Alus), which in turn activates MDA5 (8), PKR (48), and OASes/RNase L (117). Consistent with the role of ADAR1 in suppressing these dsRNA-dependent antiviral molecules, ADAR1 has been shown to be a broadly acting proviral factor (108, 122). However, ADAR1 can also display antiviral activities against a subset of viruses or under certain circumstances, likely by directly modifying viral RNAs and accumulating deleterious mutations in the viral genomes (140, 141).

Besides the canonical antiviral pathways, ADAR1 also participates in the RNA interference (RNAi) pathway (142). ADAR1 was shown to modify certain precursors of miRNA and siRNA, thereby impairing their proper maturation (143) or altering their target mRNAs (144). However, more recent studies found that ADAR1, in particular the p110 isoform, can also have a positive effect on Dicer activity (145). ADAR1 forms a complex with Dicer and stimulates its activity independent of the deaminase activity (145). It was proposed that ADAR1 carries out different functions depending on the oligomeric state and interaction partners (145); the full picture remains to be further examined.

Thus, the primary functions of ADAR1 appear to regulate other dsRNA-binding proteins in the host innate immunity and the RNAi system. Whether ADARs are reciprocally regulated by other cellular dsRNA-binding proteins is yet unclear. While DHX9 and Staufen 1 were shown to modulate the metabolism of IR-Alus, the primary targets of ADAR1, the level of A-to-I modification of IR-Alus was found to be largely unaffected by DHX9 and Staufen 1 (146, 147). Future studies aimed at identifying regulatory factors for ADAR1 would be important.
RNA INTERFERENCE PATHWAY

Drosha, Dicer, and Argonautes are conserved RNases involved in RNAi pathways. Drosha and Dicer are responsible for biogenesis of miRNA and siRNA, while Argonaute proteins silence miRNAs harboring the sequence complementary to the miRNA or siRNA (148–150). miRNAs are synthesized by RNAP II as primary transcripts (pri-miRNAs). Drosha processes pri-miRNAs to ∼70-nt-long pre-miRNAs in the nucleus (148, 150). Pre-miRNAs are exported to the cytosol and processed by Dicer, producing mature miRNAs harboring an ∼21-bp duplexed region with 2-nt 3′ overhangs at both ends. Dicer can also process exogenously introduced long dsRNAs, such as viral replication intermediates, to generate siRNAs (148, 150). Mature miRNA and siRNA duplexes are then loaded onto Argonaute proteins, at which point the guide strand is selected, while the passenger strand is degraded. The resultant complex of Argonaute and guide-strand RNA, termed the RNA-induced silencing complex (RISC), binds to a target mRNA for gene silencing. Humans have four Argonaute proteins (Ago1–4), of which only Ago2 has nuclease activity to slice target mRNAs. However, all four Ago proteins can silence target genes by recruiting effector molecules for translational suppression, deadenylation or mRNA degradation (149).

Previous structural and biochemical analyses of Drosha, Dicer, and Argonautes offer detailed pictures of how these enzymes function. Drosha and Dicer have dsRNA-specific RNase III domains and share similar structural organization and dsRNA recognition mechanisms (151–153) (Figure 4a). Both enzymes utilize the PAZ domain to fix one end of pri-miRNA/pre-miRNA and position dsRNA for accurate cleavage by the RNase III domains (Figure 4a). Structural analysis of Dicer revealed that the PAZ domain interacts with dsRNA in at least two conformations representing the preslicing and slicing modes (153, 154) (Figure 4a). Additional accessory dsRBDs, both within Drosha/Dicer and from their interaction partners (e.g., DGCR8 and TRBP, respectively), were proposed to further facilitate dsRNA binding and dsRNA positioning. Dicer also contains the helicase domain homologous to those of RLRs (Figure 4a). The structures of human and Drosophila Dicers suggest that the helicase domain can also interact with dsRNA in at least two distinct modes (Figure 4a), one similarly to RLR (155) and the other distinct from RLR (153). The latter was observed in the preslicing conformation. How the helicase domain interacts with dsRNA in the slicing mode and whether it adopts the conformation similar to RLR remain to be investigated. Unlike Drosha and Dicer, Argonautes have an RNase H–like domain, rather than RNase III, and display vastly different domain architectures. Structures showed that Ago2 binds guide RNA through the phosphate backbone and extends the guide RNA to facilitate target miRNA recognition (149).

Accumulating evidence suggests that Drosha and Dicer can function beyond biogenesis of siRNAs/miRNAs (156). For example, Drosha has been shown to regulate mRNA stability and processing of a select group of RNAs (157). Dicer was shown to process the Huntingtin mRNA when the CAG repeat sequence expands and forms a stable duplex hairpin (158). Dicer was also shown to cleave Alu RNAs and prevent excessive accumulation of Alu RNAs that can trigger NLRP3-mediated inflammation (159, 160). Interestingly, the immune-stimulatory RNAs in this case were reported to be individual Alu RNAs synthesized by RNAP III, rather than IR-Alus synthesized by RNAP II. Whether Dicer can also process IR-Alus and perturb the dynamics of other dsRNA-binding proteins, such as MDA5, PKR, OASes, and ADARs, remains unknown.

How does the RNAi system affect viral replication and antiviral immunity? While the importance of RNAi in antiviral defense has been widely documented in plants and invertebrates (161), its potential antiviral function in mammals has been controversial. On the one hand, Dicer-mediated cleavage of viral dsRNAs could lead to suppression of viral replication by knocking down viral genes. On the other hand, it could also suppress dsRNA-dependent antiviral immunity by
Figure 4
Drosha/Dicer/Ago2 and PACT/TRBP. (a) Domain architectures and structures of human Drosha, Dicer, and Ago2. The structures of human Dicer (PDB: 5ZAL and 4NHA) in complex with dsRNA showed two distinct conformations representing the preslicing mode (where dsRNA is far from the slicing center) and the slicing mode (where dsRNA contacts the slicing center). dsRNA in the slicing mode was modeled based on the structure of Dicer platform-PAZ in complex with dsRNA (PDB: 4NHA). Note that in the preslicing conformation, the helicase domain (green) interacts with dsRNA in a manner distinct from the RLR helicase domains. By contrast, the structure of Drosophila Dicer-2 (PDB: 6BU9, right) shows that the Dicer helicase domain partially encircles dsRNA, similarly to the RLR helicase domains. (b) Domain architectures and structures of PACT and TRBP. The structures of canonical dsRBD1–2 of TRBP in complex with dsRNA (PDB: 5N8M) and noncanonical dsRBD3 in complex with Dicer Hel2i (PDB: 4WYQ). The overlay of the two structures on the right shows that noncanonical dsRBD3 is structurally similar to canonical dsRBDs. Abbreviations: dsRBD, dsRNA-binding domain; RLR, RIG-I-like receptor.
eliminating dsRNA ligands for RLR, PKR, and OASes. Multiple lines of evidence have suggested that RNAi and the mammalian antiviral signaling pathways are mutually inhibitory (65, 66, 162). Interferon signaling blocks RISC by ADP-ribosylation, while several interferon-stimulated genes (ISGs) were found to be targeted by RISC (162). LGP2, a CARD-less RLR and an ISG, was also found to inhibit Dicer-dependent processing of pre-miRNA and long dsRNA, thereby suppressing miRNA/siRNA biogenesis (65, 66).

Other studies, however, suggest that Drosha, Dicer, and Argonaute proteins can exert antiviral activities in both RISC-dependent and -independent manners. It was initially thought that Dicer and Ago2 had antiviral functions in only a few undifferentiated tissues where interferon response was minimal or lacking (163, 164); a more recent study has suggested that Ago2 has antiviral activity in a wider range of cell types, including differentiated and interferon-competent cells (165). Furthermore, Drosha was also shown to suppress several positive-strand viruses by binding structural elements in viral RNAs and blocking actions of the RNA-dependent RNAP (166). These studies together offer multiple mechanisms by which RNAi machineries can exert antiviral activity. However, future studies are required to fully understand how these antiviral activities of RNAi interplay with mutually suppressive interferon signaling.

PACT AND TRBP

PACT and TRBP are paralogous dsRNA-binding proteins involved in many branches of dsRNA biology, including functions of RLRs, PKR, and Dicer (167). Both PACT and TRBP consist of three tandem repeats of dsRBDs (Figure 4b). As described for PKR and ADARs, dsRBDs have a conserved structure and dsRNA-binding mode. In PACT and TRBP, the first two (dsRBD1 and 2) are canonical dsRBDs with demonstrated dsRNA-binding activity (similar to those of PKR and ADAR1), while the third domain (dsRBD3) is a noncanonical dsRBD that has the canonical dsRBD structure but lacks dsRNA-binding activity (167) (Figure 4b). Instead, dsRBD3s of PACT and TRBP are involved in a broad range of protein-protein interactions. They were shown to form homodimers as well as heterodimers with each other, other dsRBDs (70, 167–169), and a helicase domain (170) (Figure 4b). To make matters more complex, dsRBD1 and 2 of PACT and TRBP were also proposed to interact with other proteins in dsRNA-dependent and -independent manners (171). As such, despite having a simple repeat of dsRBDs, PACT and TRBP are involved in multiple aspects of dsRNA biology through various protein-protein interactions as well as protein-dsRNA interactions.

One of the first reported functions of PACT and TRBP is to modulate the PKR activity. Earlier studies proposed that PACT directly binds PKR and activates the PKR kinase activity in the absence of dsRNA (172–174), while TRBP inhibits PKR by directly binding to PKR and/or PACT (175–177). A homozygous missense mutation (P222L) in PACT was shown to cause hyperactivation of PKR and early-onset dystonia DYT16 (178), supporting the role of PACT in PKR modulation. However, the exact nature of the PKR-modulatory activity of PACT (and the impact of the mutation P222L) has been controversial, with more recent studies supporting the role of PACT as a negative regulator of PKR, rather than a positive regulator. Mice deficient in the PACT homolog, RAX, display developmental and reproductive defects, and these defects were rescued by double knockout of PKR or expression of a kinase-dead mutant of PKR (i.e., a dominant negative mutant) (179). These observations indicate that the primary function of PACT is to suppress the aberrant activation of PKR. The PKR-suppressive role of PACT has also been observed in the context of HIV infection (180) and the deficiency of the splicing factor TIA1/TIAL1 (181). The reason for the discrepancy between the earlier and more recent studies is yet unclear—it might reflect the context-dependent role of PACT (e.g., cell stress state,
posttranslational modifications) (180, 182, 183) or the accidental mutation in the PACT construct reported to be widely used in the earlier studies (167, 182). More detailed understanding of the structural and biochemical mechanism is required to fully define the role of PACT and TRBP in PKR regulation.

PACT and TRBP were also proposed to be involved in biogenesis of miRNAs and siRNAs. They interact directly with Dicer and fine-tune its RNA cleavage sites (167, 170, 184–186). Detailed structural and biochemical studies revealed that TRBP utilizes dsRBD3 to bind the helicase domain of Dicer (in particular the Hel2i subdomain, Figure 4b) and promotes Dicer activity (170, 184). Although a similar binding mode was proposed for PACT, PACT and TRBP differentially affect Dicer substrate specificity, cleavage kinetics and accuracy, isomer production, strand selection, and RISC assembly (184, 185, 187–189). The structural basis for the differential effects of PACT and TRBP remains to be further examined.

PACT and TRBP were also found to regulate RLRs. Earlier studies suggested that PACT activates RIG-I and MDA5 through a mechanism independent of PKR and Dicer (190–194). TRBP, on the other hand, was proposed to interact with LGP2, assisting MDA5 detection of cardioviruses (195). Despite having a helicase domain (including Hel2i) closely related to that of Dicer, PACT was proposed to bind RIG-I CTD, not Hel2i, and to stimulate the ATPase activity of RIG-I (190). However, in an apparent contradiction to these findings, more recent studies have shown that PACT is specifically involved in MDA5-mediated antiviral signaling, not RIG-I-mediated signaling (196, 197). It was also proposed that PACT interacts with LGP2 in a dsRNA-dependent manner and that this interaction is required for PACT-mediated MDA5 stimulation (197).

All in all, it is clear that PACT and TRBP function as a hub for dsRNA biology, not only by sharing common dsRNA substrates but also by interacting with various dsRNA-binding proteins in the host. However, the specific roles of PACT and TRBP in functions of PKR, Dicer, and RLRs remain controversial, and their molecular mechanisms unclear. These limitations likely reflect the complex network of interactions among dsRNA-binding proteins, and the challenges in dissecting mechanisms that connect many of these dsRNA-binding proteins.

HELICASES (BESIDES RLRs AND DICER)

Helicases are involved in nearly all nucleic acid biology, from DNA replication, transcription, and translation to dsRNA-dependent innate immunity. While the term helicase implies duplex-unwinding activity, many proteins in the helicase superfamily do not display such activity (198, 199). In fact, helicases are classified based on not duplex-unwinding activity but the presence of conserved sequence motifs (198, 199). Helicases often contain one or two RecA-like domains, and utilize the interface of two RecA-like domains (either within a polypeptide or in an oligomer) to bind ATP and cognate nucleic acids. Nucleic acid binds one side of the interface of the RecA-like domains, which in turn organizes the active site on the other side for ATP binding and/or hydrolysis (198, 199). The ATP binding or hydrolysis, in turn, triggers additional conformational changes in the helicase protein, often resulting in remodeling of the bound nucleic acid or other neighboring proteins.

A small group of helicases recognize dsRNAs and display dsRNA-dependent ATPase activity without duplex unwinding; they are named dsRNA-activated ATPases (DRAs) (200). They include RLRs and Dicer, which commonly share the two RecA-like domains and an insertion domain (Hel2i) between the two. DRAs belong to helicase superfamily 2, one of the largest families of helicases. Recent studies, however, suggest that several other helicases in superfamily 2 are also involved in sensing foreign dsRNAs and regulating cellular dsRNA homeostasis.
One of these helicases is DHX9 (also known as RHA), a highly abundant and multifunctional RNA/DNA helicase involved in DNA replication, transcription, and posttranscriptional and translational regulation (201). It contains two tandem repeats of dsRBDs followed by the DExH-motif helicase domain, and it displays an ATP-driven dsRNA-unwinding activity (201). Earlier studies showed that DHX9 is packaged into HIV-1 virions and is a host factor necessary for HIV-1 replication (202). More recently, DHX9 was shown to play key roles in cellular dsRNA homeostasis by binding IR-Alus and cooperating with ADAR1 in resolving cellular dsRNA structures (146). In apparent contradiction to its role in dsRNA unwinding and resolution, DHX9 was also proposed to function as a dsRNA sensor upstream of MAVS (203) and the Nlrp9b inflammasome (204). The interplay between the activities of DHX9 in resolving dsRNA and sensing dsRNA is yet unclear.

Several other helicases in superfamily 2 have also been proposed to activate antiviral and inflammatory signaling pathways in response to foreign dsRNAs. For example, DDX60 is an interferon-inducible helicase and was found to perform antiviral functions in at least two distinct manners (108, 205, 206). DDX60 assists RLRs in sensing foreign dsRNAs in a tissue-dependent manner (205, 206), but it also degrades viral RNAs, thereby conferring RLR- and interferon-independent antiviral activities (206). However, a more recent study failed to confirm the antiviral function of DDX60 against a broad range of viruses (207), demanding more detailed studies to understand the precise role of DDX60. Additional examples of helicases reported to function in innate immune signaling in response to dsRNA include DHX15 (208), DHX33 (209, 210), DHX29 (211, 212), DDX3 (213, 214), and the complex of DDX1-DDX21-DHX36 (215). Some helicases, such as DDX17, were shown to function as antiviral effectors independent of immune signaling pathways (16), analogous to the reported effector functions of RLRs (61) and DDX60 (206). The antiviral activity of DDX17, however, appears dependent on the virus; DDX17 inhibits bunyavirus replication (16) and promotes replication of HIV-1 (216) and influenza A (217). Although more detailed studies are required to mechanistically define antiviral functions of many of these helicases, these studies place helicases as additional factors that further increase the complexity of the dsRNA biology network and antiviral immunity.

CONCLUDING REMARKS

We discussed seven families of dsRNA-binding proteins involved in mammalian antiviral innate immunity. The main goal was not necessarily to cover each family in depth, but rather to highlight their commonality, differences, and interdependence in their functions. Structures of many of these proteins show that they commonly recognize phosphate backbones and minor grooves of dsRNA, the binding mode largely incompatible with a high degree of sequence dependence. They also utilize several common mechanisms to couple dsRNA binding to their respective downstream functions. These include dsRNA-mediated, proximity-induced oligomerization (RLRs and PKR); dsRNA-mediated allosteric regulation (OASes and helicases); and simple recruitment of effector domains to dsRNA (ADARs, Drosha, and Dicer).

Their functional interdependence, however, appears more complex and is further confounded by seemingly conflicting reports in some of these cases (Figure 1a, Table 1). On the one hand, their functions are inevitably linked through the shared dsRNA substrate—both pathogen-derived and cellular dsRNAs. In particular, various sources of cellular dsRNAs have been identified to play increasingly important roles in human dsRNA biology and pathology, including Alu retroelements, triple-repeat expansions, and mitochondrial dsRNAs. However, it is also clear that the interdependence of these dsRNA-binding proteins goes far beyond simply sharing dsRNA substrates and is further defined by protein-protein interactions. Much remains to be investigated to
structurally and biochemically define their interactions and interdependence. Such system-level understanding of cellular dsRNA biology is likely required for comprehensive understanding of dsRNA-mediated antiviral immunity and regulation.

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