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To sense or not to sense viral RNA — essentials of coronavirus innate immune evasion

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An essential function of innate immunity is to distinguish self from non-self and receptors have evolved to specifically recognize viral components and initiate the expression of antiviral proteins to restrict viral replication. Coronaviruses are RNA viruses that replicate in the host cytoplasm and evade innate immune sensing in most cell types, either passively by hiding their viral signatures and limiting exposure to sensors or actively, by encoding viral antagonists to counteract the effects of interferons. Since many cytoplasmic viruses exploit similar mechanisms of innate immune evasion, mechanistic insight into the direct interplay between viral RNA, viral RNA-processing enzymes, cellular sensors and antiviral proteins will be highly relevant to develop novel antiviral targets and to restrict important animal and human infections.

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Introduction
Our immune system functions as a collaborative network of cells, lymphoid organs, cytokines and humoral factors, to combat infections caused by pathogenic agents such as bacteria, parasites and viruses [1]. The first line of host defence consists of innate immunity, which is evolutionary conserved, and has the main task to limit replication and spread of pathogens [2]. This response is then complemented by adaptive immunity, which develops more gradually and relies on lymphocytes that recognize antigenic structures on the surface of pathogens that managed to pass through the epithelium [1].

The emphasis of this present review lies on Coronaviruses (CoVs), which are positive-stranded RNA viruses of both veterinary and medical importance causing mainly respiratory and enteric diseases. Human CoV-infections usually present as mild respiratory tract disease (common cold) that may be more severe in elderly or immune-compromised individuals. However, the epidemic in 2002/03 caused by the severe acute respiratory syndrome-associated CoV (SARS-CoV) and the recently discovered Middle East Respiratory Syndrome coronavirus (MERS-CoV) demonstrate the zoonotic potential of CoVs and their ability to seriously impact human health [3,4,5].

Coronavirus innate immune evasion
CoVs belong to the order Nidovirales and are characterized by a large single-stranded (ss) RNA genome of positive polarity and a complex replication cycle [3,6**]. Upon entry into host cells the viral genome is uncoated and primary translation results in the synthesis of two large polyproteins pp1a and pp1ab that are cotranslationally and post-translationally cleaved by viral proteases into 16 non-structural proteins (nsp 1–16) (Figure 1). These CoV nsps form the replication/transcription complex (RTC) that is responsible for the synthesis the viral RNA, the major CoV pathogen-associated molecular pattern (PAMP) recognized by host cell pathogen-recognition receptors (PRRs). Most PRRs responsible of sensing viruses are either located in the endosomes (e.g. Toll-like receptors; TLRs) [7] or in the cytosol (e.g. retinoic acid inducible gene I (RIG-I)-like receptors; RLRs) [8,9], suggesting that viruses are primarily detected by the presence of their nucleic acids. Sensing of PAMPs by PRRs results in their activation and ultimate expression of numerous cytokines, including interferons (IFNs). IFNs in turn induce the expression of a broad array of IFN-stimulated genes (ISGs) to establish an antiviral state.

Much of our current understanding concerning CoV-induced innate immune responses results from studies using mouse hepatitis virus (MHV) in the context of a murine model of infection. Upon MHV infection, plasmacytoid dendritic cells (pDCs) rapidly secrete high amounts of IFN-α mediated through TLR7 stimulation and constitutively expressed IFN-regulatory factor (IRF) 7 [10]. Considerably lower levels of type-I IFNs are produced by other cell types, such as macrophages, microglia and oligodendrocytes [11–13]. pDC-derived Type-I IFN is particularly important to protect cells that fail to raise a significant IFN-response during CoV-infections and thus secures their integrity and efficiently reduces virus replication and spread [14]. Remarkably,
human CoVs (HCoVs) infection in human airway epithelial cells (HAEs), the primary target tissue at the entry port of respiratory pathogens, did not result in upregulation of IFNs or inflammatory cytokines, although these cells readily respond to exogenous type-I and type-III IFN [4*]. In line, SARS-CoV and MERS-CoV did not induce a pronounced IFN-response in polarized airway epithelial cells (Calu-3) [15,16], in A549 cells (human alveolar adenocarcinoma cell line) and in human monocyte-derived macrophages [17]. Collectively, these data suggest that CoVs efficiently evade innate immune sensing in other cell types than pDCs, either passively, by hiding PAMPs, or actively, by encoding viral antagonists to counteract induction of IFN expression.

**Mimicking cellular mRNAs**

To distinguish host-cell-derived from viral mRNAs, PPRs have evolved to specifically recognize foreign, or ‘non-self’, RNAs. The cap structure present at 5’-termini of eukaryotic mRNAs has a pivotal role in this context, as uncapped RNA bearing a 5’ triphosphate terminus, and RNA with a non-methylated or incompletely methylated cap structure are now considered as viral PAMPs. In addition, double-stranded (ds) RNA is a long known viral PAMP, since replication of RNA virus genomes involves so-called replicative intermediates consisting of dsRNA. The cytosolic RLRs RIG-I and melanoma-differentiation-associated protein 5 (MDA5) sense viral RNAs, however RIG-I preferentially senses short blunt-end or 5’ triphosphate RNAs [8,9], while MDA5 recognizes cap0-structures (i.e. cap structures lacking ribose 2’O methyl-ation) [18**] and long dsRNAs [8,19]. Host cell mRNAs are usually not recognized by RLRs, since already in the nucleus their nascent 5’ triphosphate-end is co-transcriptionally modified to form a N7-methylated and 2’O-methylated 5’ cap structure [20**].

Many viruses that replicate in the cytoplasm have thus evolved mechanisms to provide a cap structure at the 5’ ends of their mRNAs in order to evade RLR recognition and to ensure eukaryotic translation. The acquisition of a cap structure is most efficiently done by either hijacking the cellular capping machinery, by snatching caps from cellular mRNAs or by encoding viral capping enzymes [20**]. CoV RNAs contain a 5’ cap structure [18**] and the CoV-RTC harbours capping enzymes such as a
RNA-triphosphatase encoded by nsp13 [21], a N7-Methytransferase (N7-MTase) residing in nsp14 [22] and a 2'O-MTase encoded by nsp16 [23].

Detailed analyses including 2'O-MTase-deficient Coronaviruses, Flaviviruses and Poxviruses congruently demonstrated that 2'O-methylation of viral RNA increases virulence through evasion from innate immune responses (Figure 2). Recombinant 2'O-MTase-deficient MHV elicited an elevated type-I IFN-response in wildtype, but not in MDA5-deficient, macrophages, indicating that 2'O-methylation of viral RNA protects from recognition by MDA5 [18**]. Additionally, 2'O-MTase deficient viruses are more sensitive to the antiviral actions of IFN, suggesting that specific ISG(s) may explicitly restrict replication of viruses lacking 2'O-methylation [24]. Indeed, IFIT1 (IFN-induced-protein-with-tetratriopeptide-repeats-1) was shown to preferentially bind to cap0-RNA (i.e. lacking 2'O-methylation) and thereby impairs the attachment of eukaryotic initiation factors.
(eIFs) to the viral template, which will ultimately result in translation arrest [25**]. IFIT1 apparently has only marginal effects on replication and virulence of wildtype viruses [18**,24,25**]. However, replication of MHV-mutants, West-Nile-Virus-mutants, and vaccinia virus-mutants lacking 2′O-methylation was restricted in wildtype macrophages, but restored in IFIT1-deficient macrophages [18**,24,25**]. Interestingly, the ability of IFIT1 to restrict replication of West Nile Viruses lacking 2′O-methylation was highly cell-type specific and revealed that additional mechanisms independent of IFIT1 are involved in control of the infection [26].

Biochemical analyses revealed that 2′O-MTases of CoVs and Flaviviruses specifically methylate N7-methylated RNA structures, suggesting that N7-methylation is a prerequisite for 2′O-methylation to occur, and thus, an inactive N7-MTase would prevent the 2′O-MTase from proceeding with methylation [27,28]. It has been shown that WNVs lacking N7-methylation are non-replicative and that upon transfection of RNA encoding only weak N7-MTase activity, viruses with a number of compensatory mutations were rescued [29]. The coronaviral N7-MTase was only recently identified to reside in nsp14 [22,27] and it is currently unknown how the inactivation of the CoV-N7-MTase will affect viral replication and innate immune recognition (Figure 2). It can only be hypothesized that non-methylated RNA will be preferentially recognized by IFIT1, resulting most likely in a translational arrest similar to RNA lacking 2′O-methylation [25**]. It has not yet been described if CoV-MTases also methylate internal nucleotides of nascent mRNAs as it is known for Flaviviruses [30] or if they employ a similar mechanism as demonstrated for the Marburg Virus (Filoviruses), where a multifunctional protein VP35 not only caps dsRNA-termini but also fully coats the backbone to physically protect the RNA from length-dependent sensing [31,32].

**Ribonucleases remove RNA-PAMPs?**

Efficient removal of RNA-PAMPs prevents the activation of the innate immune system and is thus facilitating successful replication in the host cell. Infections by Arenaviruses, including the Lassa fever virus (LASV), are characterized by a potent mechanism of immune suppression, which is partially mediated by the 3′-to-5′ exoribonuclease (ExoN) which is encoded in the LASV nucleoprotein and structurally very similar to members of the DEDDh family of ExoNs [33]. The LASV-ExoN specifically digests viral dsRNAs and consequently removes RNA PAMPs which otherwise would activate cellular receptors and trigger an IFN-response [31,33,34].

Coronaviruses also encode a 3′-to-5′ ExoN of the DEDDh superfamily within the N-terminal part of nsp14. The enzyme hydrolyses ssRNAs and dsRNAs, and excises single nucleotide mismatches at the 3′-end of dsRNAs [35*]. Replacement of catalytic residues in the ExoN domain of MHV and SARS-CoV resulted in viable mutants. However, mutations accumulated throughout their genomes albeit without any obvious fitness costs [36], indicating that ExoN functions as an unique RNA-proofreading enzyme and thereby confers very high replication fidelity, which is required for stable propagation of large viral RNA genomes exceeding 20 kb [6**,37]. Obviously, RNase-activities are expected to be very target-specific and tightly regulated to prevent unwanted cleavage of viral or cellular RNAs. It has been shown that DNA, ribose-2′O-methylated RNA substrates [38], diribonucleotide, triribonucleotide or longer unpaired ribonucleotide stretches as well as 3′-modified RNAs [35*] are resistant to ExoN-mediated cleavage. Furthermore, CoV-nsp10 has been described to allosterically regulate the 2′O-MTase activity [27,39] and interestingly, also stimulates the CoV-ExoN activity [35*]. This suggests an important regulatory role of nsp10 in immune suppression through its dual impact on ExoN and 2′O-MTase [35*], proposing that the ExoN is not only involved in RNA-proofreading but also in the escape of host innate immunity. As IFN-induction is efficiently inhibited during CoV-infection, a role of ExoN in degrading RNA-PAMPs might be indeed conceivable and would complement known IFN-antagonistic activities during CoV-infection.

The second ribonuclease of CoVs, an endoribonuclease (EndoU) is encoded by nsp15 and was until recently considered a Nidovirus-specific feature. However, the discovery of an insect-born Nidovirus and the re-evaluation of invertebrate Ronivirus genomes revealed that they lack an EndoU domain, indicating that the utilization of an EndoU domain is restricted to vertebrate Nidoviruses [6**]. The CoV-EndoU domain is structurally very conserved but exhibits only moderate sequence conservation [40,41]. It hydrolyses ssRNA and dsRNA substrates, preferably 3′ of uridylylates [41,42] and cleaves similar as XendoU and RNase A, as the reaction products possess 2′,3′-cyclic phosphate ends [42]. Ectopic expression studies demonstrated that CoV-nsp15 is not only co-localized with the RTC, but also distributed throughout the cytoplasm [40], thus an additional function besides its involvement in viral replication has been proposed [43]. Overexpression studies in the absence of CoV-infection suggested a role of the SARS-CoV EndoU as a robust IFN-antagonist [44] and it seems plausible that CoVs might use the RNase activity of nsp15 to digest RNA-PAMPs and thus avoid activation of the innate immune system, similar as described for the Pestivirus ribonuclease activity of the structural protein E [45*,46]. However it should be kept in mind that during virus infection, Arterivirus and CoV EndoUs co-localize mainly with the RTC, and it still remains to be determined if the proposed mechanism of immune evasion is actually taking place at Arterivirus and CoV RTCs. Further it remains
elusive if the EndoU substrate is RNA of viral or cellular origin, and under which circumstances the utilization of an EndoU is required, as the enzyme is absent in invertebrate Nidoviruses [6**].

Shielding dsRNA from innate immune sensors
A hallmark of CoV replication is the induction of ER-derived double membrane vesicles (DMVs) that are part of a reticulovesicular network in the host cytoplasm. Particular proteins involved in replication (nsp3, nsp5, nsp8) are found predominantly at the outer membrane of DMVs [47], whereas dsRNA was mostly detected within the inner lumen of the vesicles [48,49]. Interestingly, no connection of the DMV lumen to the cytoplasm could be identified so far. Thus, assuming that the dsRNA within CoV-induced DMVs is of viral origin, it remains enigmatic how viral mRNAs may leave the vesicles and become accessible for translation. It furthermore raises the question if dsRNA within DMVs represents replicative intermediates [47] or rather dead-end products that are shielded from innate immune sensing by cytoplasmic receptors. It thus remains to be determined if DMVs are the actual site of RNA synthesis and additionally, how transcripts are transported to the sites of assembly. Moreover, given the assumption that dsRNA may simply represent potent PAMPs that are hidden from innate sensors, it would be crucial to evaluate the strength of the IFN-response in the absence of DMVs and also, which receptors may play a role in this context.

Significance and perspectives
CoVs raised considerable concerns as zoonotic pathogens. However, they have long been known as important animal pathogens and our knowledge on molecular details of their replication strategy has significantly increased during the past decades. CoVs have evolved versatile mechanisms to counteract host innate immune responses, including mimicking cellular mRNAs, engagement of virus-encoded ribonucleases, and shielding of dsRNA from innate immune sensing. There are a number of parallels described for other viruses, suggesting that those mechanisms target evolutionary conserved innate immune pathways. Nevertheless, mechanistic details of CoV innate immune evasion are still sparse. It will be important in future studies to address the overarching theme ‘how, where and when CoV-RNA is sensed during infection’. For example, it is important to clarify where exactly CoV-RNA synthesis takes place and which role(s) CoV-induced DMVs play in the context of viral replication and innate immune evasion. How and where do cytosolic PRRs access CoV-RNA and which mechanisms for self–non-self distinction of RNA exist. Important consideration are (i) when and where during the infection do viral RNA and PRRs interact, (ii) is there a particular sensing complex assisting RLRs to assess molecular signatures, such as 2′O-methylation, in order to distinguish self from non-self RNA, and (iii) is there a particular sensing complex to detect ‘mis-localized’ RNA, for example viral RNA outside DMVs. A better understanding of these basic questions will be crucial to design novel strategies of antiviral intervention that will be applicable to a wide spectrum of virus-induced diseases in animals and humans.

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