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Article

CORE

KH-RNA interactions: back in the groove.

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Abstract

The hnRNP K-homology (KH) domain is a single stranded nucleic acid binding domain that mediates RNA target recognition by a large group of gene regulators. The structure of the KH fold is well characterised and some initial rules for KH-RNA recognition have been drafted. However, recent findings have shown that these rules need to be revisited and have now provided a better understanding of how the domain can recognize a sequence landscape larger than previously thought as well as revealing the diversity of structural expansions to the KH domain. Finally, novel structural and functional data show how multiple KH domains act in a combinatorial fashion to both allow recognition of longer RNA motifs and remodelling of the RNA structure. These advances set the scene for a detailed molecular understanding of KH selection of the cellular targets.

Introduction

The hnRNP K-homology (KH) domain is a single stranded, sequence specific, nucleic acid binding domain present in proteins that regulate gene expression in eukaryotes and prokaryotes. The domain comprises three α -helices that pack onto the surface of a central antiparallel β -sheet (**Figure 1a**). Eukaryotic type I and prokaryotic type II KH domains share a minimal $\beta \alpha \alpha \beta$ core, with two additional α and β elements positioned either C-terminal (type I, eukaryote) or N-terminal (type II, prokaryote) to this core motif **[1-3]** (**Figure 1a**). In all KH domain-nucleic acid structures the nucleic acid backbone interacts with a conserved GxxG loop that links the two helices of the minimal KH core (**Figure 1b**). This orients four nucleic acid bases towards a groove in the protein structure where hydrophobic interactions and a network of mainchain and sidechain hydrogen bonds mediate nucleobase recognition (**Figure 1c**) **[4-6]**. So far, domains with a classical KH fold but lacking the conserved GxxG motif have shown no nucleic acid-binding activity - although they interact with other nucleic acid binding domains and can modulate their RNA binding activity **[7]**.

KH domains are found in arrays of up to 15 repeats (**Figure 1d**) and neighbouring domains can be structurally decoupled or form two-domain structural units. Individual domains often cooperate in the selection of the cellular targets and the combinatorial action of multiple domains allows recognition of longer sequences to increase specificity. Alternatively, longer RNA sequences can be recognised by expanding the classical KH fold with the addition of secondary structure elements and by dimerisation of KH containing proteins, such as the one reported for the STAR family of RNA regulators **[8]**.

A set of rules for KH-nucleic acid recognition has been devised based on the structures of KH domains in complex with RNA and DNA targets and on functional and biochemical information regarding their physiological targets **[9]**. However, recent data suggests that some of these rules should be re-visited. Further, new information has come to light regarding how KH domains can act combinatorially to recognise longer RNA sequences, a necessity to select targets within the large pool of cellular RNAs. Below we review recent key studies on KH-RNA interactions and discuss how these studies have changed our understanding of recognition and of the role played by KH domains in RNA regulation.

An expanded sequence specificity landscape for KH domains

The structure of several KH domain-nucleic acid complexes including, amongst others, Nova-1 KH3 **[4]**, SF1 **[10]**, hnRNPK **[9]**, FBP **[11]**, and PCBP2 **[12]** indicated that, in the protein-RNA complexes, the two central nucleobases of the tetranucleotide recognition sequence (positions 2 and 3) were either adenine or cytosine, while a pyrimidine was preferred in the first and fourth position. The recognition of a cytosine in position 2 is mediated by hydrophobic contacts or stacking interactions and by two H-bonds between an arginine side chain located on the central β -sheet of the domain and the O2 and N3 groups of the base. Instead, when adenine is selected at position 2 the bulky arginine side chain is replaced by a smaller lysine that better accommodates the purine base (**Figure 2a**). In position 3, the Watson-Crick edge of the nucleobase is recognised specifically by two Hydrogen-

bonds with the backbone amide and carboxyl moieties of same amino acid from the second (third for type II domains) strand of the KH β -sheet. Only an A or a C can make the double hydrogen bond, which was thought to be an intrinsic feature of KH-nucleic acid recognition. Discrimination between A and C can be attributed to an additional direct or water-mediated contact between the nucleobase and a protein side chain on $\alpha 2$ (**Figure 2a**).

Recently, it was shown that the third KH domain of KSRP recognises a G-rich sequence in the precursor of the tumour suppressor Let-7 miRNA [13] rather than a canonical A/C rich sequence. This recognition is necessary to enhance the processing of the precursor to the mature miRNA. The structural basis of Grecognition was revealed by the solution structure of the KH3-AGGGU complex [14,15]. In KH3 of KSRP the hydrophobic groove is wider than that of other KHnucleic acid complexes (Figure 2b) allowing G bases in position 1, 2 and 3 to enter the groove. Moreover, in positions 2 and 3, guanine is not only tolerated but also selected for. In position 2, lysine 368 makes a specific hydrogen bond with the O6 of Guanine-2. When the lysine is substituted by arginine the specificity of position 2 changes from a purine to a pyrimidine, as predicted, but the RNA binding affinity of the domain remains unchanged [15]. In position 3, the broader groove facilitates a shift of the base edge along $\beta 2$ and the formation of three intermolecular hydrogen bonds between the Watson-Crick edge of the base and the carboxy and amide groups of two residues, isoleucine 356 and phenylalanine 358. A fourth hydrogen bond is formed with the side-chain of glutamine 349, completing a recognition pattern that is very specific, but also different from the canonical one (Figure 2b). Although no other structures of KH domains in complex with G-rich sequences are available, in vitro and in cell data have suggested that the fourth KH domains of the protein IMP1 recognises a G-rich RNA sequence [16]. Overall, the data show that sequence specific recognition in the KH hydrophobic groove relies both on specific contacts and on the overall shape of the groove, which is governed by the conformation of multiple side chains. Several of the key side chain determinants of recognition have been identified and we expect on-going work to result in further advances in deciphering the contact-specific sequence recognition code of KH domains. A future more complex challenge is to understand how the overall side chain composition defines the shape of the binding-groove and provides a template for nucleobase discrimination.

Recognition of longer RNA sequences by expanded KH domains

The hydrophobic groove of canonical KH domains can recognize up to four nucleotides. Specific recognition of longer sequences by individual domains is achieved by expanding the classical KH fold with additional secondary structure elements that elongate the RNA recognition surface **[17,18]**. The best studied of these expanded KH folds is the Signal Transduction and Activation of RNA (STAR) fold, which mediates RNA recognition in a family of proteins important to splicing, mRNA shuttling and translational repression (**Figure 3a**) **[8,19-21]**. The STAR domain includes an N-terminal protein dimerisation motif (the QUA1 motif), a central KH domain and a C-terminal motif involved in RNA binding (the QUA2 motif). The structure of the SF1-RNA complex (**Figure 3b**) has shown that QUA2 comprises a long loop followed by an amphipathic helix that folds back to contact the α 1 and α 3 helices and the GxxG loop of the KH domain. This extends the hydrophobic surface of the KH groove to allow recognition of an additional 2 nucleotides.

Three recent structures, two crystallographic and one in solution, have expanded our understanding of STAR-RNA recognition. The crystal structures of Qk1 and GLD-1

[22,23] in complex with the RNA Recognition element (RRE) target RNA, have provided an overall structural description of STAR domain-RNA interactions and shown that direct contacts with the RNA are made exclusively by the KH and QUA2 domains. Nevertheless, QUA1 dimerization increases the RNA binding affinity of the individual KH-QUA2 units within the homodimer, possibly as a result of QUA1-QUA2 contacts that stabilize the KH-QUA2 arrangement [22,24]. A major factor that determines the KH-QUA2 structural arrangement is the interaction with RNA. In the absence of RNA, the QUA2 helix is poorly ordered, as shown by the analysis of backbone motions of free Qk1 [23,25]. An important role of RNA binding in shaping the protein interaction surface is also suggested by the finding that, despite a network of conserved QUA2-RNA interactions, the orientation of the QUA2 helix varies between two complexes of GLD1 with RNA targets of different length [22,23]. In the solution structure of the GLD-1-RNA complex three nucleotides rather than two make contact with the QUA2 motif and the nucleobase of the third nucleotide docks into a shallow pocket in the QUA2. It has been proposed that this additional contact results in an approximate 10° change in the orientation of the QUA2 helix with respect to the X-ray structure. An important open question is whether the structural flexibility observed in the arrangement of the QUA2 motif is related to a functional flexibility in the recognition of different of RNA targets.

The GLD-1 and Qk1 proteins are dimers in the cell and indeed recognise bi-partite RNA sequences. The structure of a (dimeric) QUA1-KH-QUA2 STAR protein in complex with a complete bi-partite RNA sequence will help clarify how dimerisation is related to the arrangement of the two recognition sequences. Interestingly, the expanded KH domain of FILIA, a protein important in RNA regulation during mouse oogenesis, also creates a strong inter-molecular dimer **[26]**. However, this domain has a different structural arrangement with respect to the STAR domain (Figure 3B) and its specific target sequence is yet to be identified.

Finally, structural extensions have also been reported for type II KH domains. In the ERA-RNA complex an extended type II KH domain interacts with nine nucleotides - five of which make contacts outside the hydrophobic groove. The additional contacts however are made within the core KH fold and not with the structural extension (**Figure 3B**). The structure of the ERA-RNA complex shows that the KH fold can interact with RNA using residues outside the hydrophobic groove and highlights the differences between eukaryotic and prokaryotic KH domains [27].

Combinatorial binding and architectural role by multiple KH domains

As observed for other RNA binding domains, combinatorial binding of multiple KH domains within the same protein is often key to high affinity and high specificity interaction with the RNA target **[28,29]**. Individual KH domains bind their short single stranded RNA target sequences with dissociation constants in the micromolar range (10⁻⁶ to 10⁻⁴ M), while the dissociation constant of multi-KH protein–RNA complexes is in the low nanomolar range **[14,28,29,30]**. The contribution of individual domains to RNA binding and protein function has been evaluated using domain deletions and, more recently, conservative double mutations in the GxxG loop that eliminate RNA binding of individual domains **[30]**. Additionally, structural and biophysical studies have characterized inter-domain interactions. In several instances, flexible linkers join individual domains establishing some degree of domain coupling in the binding, while also allowing the protein to adapt the inter-domain arrangement to different RNA (and DNA) targets. However, other arrangements have been reported where stable inter-domain association of neighbouring KH domains produces a continuous binding site proposed to play a role in the folding of the RNA targets.

The most common inter-domain arrangement in di-domain KH units is a side by side anti-parallel configuration where the three β-strands of each KH orient to form an extended six-stranded β -sheet (**Figure 4a**). This arrangement is stabilised by contacts between the two KH α 3 helices. NMR and X-ray crystallography studies of the KH di-domains of the Nova-1 [31], IMP1 [16,32] and PCBP2 [33,34] proteins, indicate that the dimer does not undergo significant conformational rearrangements upon RNA (or DNA) binding. Instead the two domains form a rigid structural unit and an architectural function has been proposed where binding of the same nucleic acid chain to the anti-parallel hydrophobic grooves of each KH domains results in a 180 ° looping of the RNA [16,32]. In order to validate this observation in a cellular context, the distribution and 5'-to-3' orientation of the cognate sequences of IMP1 KH3 and KH4 was analysed in a set of physiological targets [16]. The analysis showed that both RNA sequences were present in the majority of targets at a distance within the boundaries expected from the structure of the di-domain and in vitro binding data. However, the sequences were found both in a 5-3' and a 3'-5' arrangement, in equal shares, indicating that the RNA chain can loop on either side of the di-domain.

Other inter-domain KH-KH arrangements have also been reported implying that different architectural constraints are imposed on the RNA chain. For example, the orthogonal arrangement of the KH2-KH3 domains of KSRP [35] (Figure 4a) has been proposed to bend the RNA chains with ~90 ° angle while the NusA binding of the type II KH di-domain to one contiguous stretch of 11 nucleotides [36] (Figure 4a) also results in a change in direction of the RNA but with no requirement for looping. Other inter-domain arrangements have been reported, for example for the two KH domains of the Fragile X mental retardation protein (FMRP) but it is unclear whether both domains bind to specific RNA targets [37,38].

KH-domains that do not contain the conserved GxxG motif – and do not interact with RNA – can assemble in multi-domain units where more than one KH-KH interface is observed, exemplified by the structure of the type I KH tetra-domain unit of Bicaudal-C (Bic-C) (**Figure 4b**) [**39**]. Type II KH domains lacking the GxxG motif have also been recently reported in proteins of the metallo- β -lactamase superfamily. As seen for type I KH, type II KH domains lacking the GxxG loop also do not participate in the recognition of the RNA targets [**40-42**].

Perspectives

It is an exciting time for the study of KH domains. Recent findings have highlighted their capacity to recognize a diverse range of sequences and to expand and combine sequence recognition of individual domains to select longer RNA targets. In addition, di-domain KH units likely play a general role in re-shaping the structure of the bound RNA. It is now important to understand how the architectural role of di-domain units relates to the structure and function of large protein-RNA particles. A further set of questions concerns the complex relationship between nucleobase specificity of individual KH units, the use of multiple KH domains and the targeting of specific RNAs in the cell. A combined structural and cell biology approach, including the genomic analysis of the targets, will be required to understand not only KH recognition of high affinity sequences but, also target selection in a cellular context where binding to intermediate affinity targets is functionally relevant for many KH domains.

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Figure 1.

KH domain structure and global features of KH-RNA interactions.

(a) Cartoon representation of the secondary structure and fold of type I and type II KH domains. Core secondary structure elements are shown in green, additional elements in grey. (b) Sequence alignment of the minimal $\beta\alpha\alpha\beta$ core of type I and type II KH domains. Secondary structure elements are drawn above the alignment. (c) Molecular surfaces of the Type I Nova2 KH3 (left) and Type II NusA KH2 (right) in complex with RNA. Hydrophobic residues are shown in yellow, while the conserved GxxG loop is highlighted in red. The four nucleotides recognized in the KH3 hydrophobic groove are displayed as sticks, color-coded by atom type. In the middle is a ribbon representation of the Nova-2 KH3-RNA complex. The four nucleotides are displayed by sticks, colour-coded by atom type, with carbon atoms in yellow. The equivalent nucleotides in the superimposed structure of NusA-KH2-RNA complex are also similarly displayed, but with carbon atoms in orange. (d) Schematic representation of the multiplicity and arrangement KH domains (light blue boxes) in a selection of eukaryotic proteins.

Figure 2.

RNA sequence recognition in the KH domain hydrophobic groove.

(a) KH domain canonical and non-canonical (far right) recognition of nucleobases in positions 2 and 3 of the target tetra-nucleotide. The two nucleotides are displayed by sticks, colour-coded by atom type, with carbon atoms in green, while interacting amino acid are similarly colour coded but with carbon atoms in magenta. Hydrogen bonds are shown as black dashed lines and the red sphere represents a water molecule. (b) Molecular surfaces of the four RNA-bound KH domains shown above. Hydrophobic residues are coloured red. The bound nucleic acids are drawn using a stick representation and are coloured yellow.



Figure 3. Extended KH domains: structure, dimerisation and RNA binding. (a) Schematic of the arrangement of secondary structure elements of the extended KH domains. Elements belonging to the common KH $\beta\alpha\alpha\beta$ motif are shown in green, the type I/type II specific secondary structures are shown in white and the secondary structure elements that extended KH domains are shown in pink and red. (b) Ribbon representations of expanded KH domains, either free or in complex with RNA. The core KH domain is in grey and the RNA is displayed as sticks, colour coded by atom type with the carbon atoms in green (for the four nucleotides bound in positions 1-to-4 of the hydrophobic groove) or yellow (for the additional nucleotides that make contact with the QUA2 element). The nucleotides making contact with the QUA2 element are numbered starting from the nucleobase 5' to position 1 of the hydrophobic groove (position 0). The RNA bound to the ERA protein has all of the carbon atoms in green, as the contacts made are different than in canonical KH domains. Nucleotides that are not making contacts with the protein are not displayed in any of the structures. Additional secondary structure elements are colour-coded as in a, the QUA1 dimerisation region of Qk1 in pink and QUA2 extra RNA binding elements in red.



(b)

(a)





Figure 4.

Multi-KH structural units.

Ribbon representation of the multi-KH units of (**a**) Bicaudal-C and (**b**) IMP1, KSRP and NusA-RNA. The NusA bound RNA is displayed as sticks, colour coded by atom type. For each structure a schematic arrangement of the two domains is drawn, with the β -sheet face shaded black area and the helical face hatched, colour coded as the corresponding domain.