The extended AT-hook is a novel RNA binding motif

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Abbreviations: eAT-hook–extended AT-hook; HMG–high mobility group; TAF1–TATA-binding protein-associated factor 1; AKNA–AT-hook-containing transcription factor; PSR/JMJD6–Phosphatidylinerse receptor–Jmj-domain containing protein D6; CaATL1–Capsicum annuum AT-hook-Like gene 1; TdIF1–Terminal deoxynucleotidyltransferase-interacting factor 1; LEDGF/p75–Lens epithelium-derived growth factor 75; McCP2–Methyl-CpG-binding protein 2; Tip5–TTF-I interacting protein 5; rDNA En–rDNA enhancer; rDNA IGS–rDNA intergenic spacer; PTOV1–Prostate Tumor Overexpressed 1; Med25–Mediator complex subunit 25; GPBP1–GC-rich Promoter Binding Protein 1; VL1–Vascular protein family Vasculin-like 1; RACK1–receptor of protein C kinase 1; Brg1–Brahma related gene 1; Dot1L–Dot1 [Disrupter of telomere silencing protein 1]-like; snRNA–small nuclear RNA; IncRNA–long non-coding RNA

Introduction

Nucleic acid binding domains are essential in protein-mediated regulation of DNA- and RNA-dependent biological processes. The AT-hook belongs to this largely diverse group of protein domains and was originally described as a short, 3 times repeated peptide motif in the HMGA1 (High mobility group AT-hook 1) protein.1 It binds to the minor groove and alters the architecture of DNA, which facilitates binding of other proteins in the opposite major groove.2 The AT-hook motif is a characteristic sequence feature of HMGA proteins and distinguishes them from the HMGB (HMG box) and HMGN (HMG nucleosome-binding domain) proteins.3 Based on the biochemical and structural characterization of HMGA1s AT-hooks,1,2 the G-R-P tripeptide motif has been considered as the core sequence of this short protein domain. Almost two decades ago a systematic search for the G-R-P motif in protein sequence databases led to the discovery of a wide variety of AT-hook proteins, including numerous non-HMG proteins.4 Multiple sequence alignments of all AT-hooks identified a 10–15 amino acids long peptide motif, in which the G-R-P core motif is mainly surrounded by K and R residues. Notably, most of the eukaryotic AT-hook proteins are associated with chromatin regulatory functions, such as chromatin remodelling, histone modifications or chromatin insulator function. Therefore the AT-hooks were proposed to anchor chromatin-modifying proteins to AT-rich DNA sequences.

Keywords: AT-hook, GPBP1, microscale thermophoresis, protein-RNA interaction, PTOV1, RNA-binding domain, Tip5

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Submitted: 03/27/2015; Revised: 06/01/2015; Accepted: 06/04/2015

http://dx.doi.org/10.1080/15476286.2015.1060394

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Importantly, several reports suggest a more relaxed definition for the AT-hook sequence motif than originally proposed. Most strikingly, 2 conserved AT-hook-like motifs have been identified in the β′ subunits of DNA-dependent RNA polymerases (DNGRRGAPVTNP9SDRPLRSL in Thermus thermophilus; the GRP-like motifs are underlined). Based on the ubiquity of this sequence motif in the β′ subunits of a wide variety of species, it has been suggested that AT-hook-like domains function as nucleic acid recognition peptides already in the last universal common ancestor of the present life forms, or even earlier. Moreover, a careful investigation of Drosophila TAF1 (TATA-binding protein-associated factor 1), the only protein in this species with detectable canonical AT-hook, revealed that the replacement of glycine and proline residues of the G-R-P core motif does not interfere markedly with the DNA binding activity of this protein domain. This led to the assumption that AT-hook-like motifs, which do not perfectly match to the consensus AT-hook sequence, might exist in the fruit fly proteome. Indeed, 5 AT-hook-like motifs (KRGRGRARK, EKKPGRKRK, VKRMRRSVVR, PKKRRGRKAK, and PKRKNRRRIK) have been finally identified in the Drosophila homolog of the G9a histone methyltransferase enzyme. Additional examples for non-canonical AT-hook-like domains include the AKNA (AT-hook-containing transcription factor), PSR/JMJD6 (Phosphatidylinerine receptor, Jmj-domain containing protein D6), CaATL1 (Capsicum annum AT-hook-Like gene 1), TdIF1 (Terminal deoxynucleotidyltransferase-interacting factor 1), LEDGF/p75 (Lens epithelium-derived growth factor 75) and MeCP2 (Methyl-CpG-binding protein 2) proteins. Remarkably, LEDGF/p75 and MeCP2 contain also canonical AT-hooks besides their AT-hook-like domains. These studies revealed largely different non-canonical AT-hooks displaying alternative amino acid sequences in the close proximity of the G-R-P core motif, or even within the motif. Notably, these AT-hook-like sequence motifs are similar in size to the canonical AT-hook, i.e. 10–15 amino acids long, and they point to the existence of a large number of so far unrecognized DNA binding domains, which likely perform similar functions to the AT-hooks.

We have recently analyzed the AT-hook domains of the Tip5 (TTF-I interacting protein 5) chromatin regulator protein, the large subunit of the Nucleolar Remodelling Complex. The DNA binding affinities of the 4 AT-hooks of Tip5 were determined in vitro, and the nucleolar targeting function of the first 2 AT-hooks was reported. While studying Tip5s AT-hooks, we have found an additional G-R-P peptide motif within the Tip5 sequence, which is flanked by basic amino acid patches at a larger distance than the G-R-P core of a canonical AT-hook. Thus, we named the new sequence motif extended AT-hook (eAT-hook). This novel protein domain is approximately 3 times longer than AT-hooks, and the basic amino acids are spaced apart from the G-R-P core in it. Here we show the characterization of the nucleic acid binding activities of Tip5s eAT-hook, define the eAT-hook sequence motif and identify human and mouse proteins with this sequence feature. Finally, we show that the eAT-hooks of 3 different proteins bind RNA with one order of magnitude higher affinity than DNA and therefore suggest a possible function for the eAT-hook in diverse nucleic-acid-dependent cellular processes.

Results

The eAT-hook of Tip5 exhibits DNA binding activity

In addition to the 4 canonical AT-hooks, we have identified in the amino acid sequence of Tip5 another G-R-P tripeptide motif, which resembles the core of the AT-hook domain (Fig. 1A). The additional G-R-P tripeptide is flanked by basic amino acid patches located at a 6–20 amino acids distance, which is larger compared to their typical 1–5 amino acids distance detected in canonical AT-hooks. This observation led us to speculate about the existence of a new kind of DNA binding domain, which we termed extended AT-hook (eAT-hook). This description refers to the patches of basic amino acids located distant from the core motif. To experimentally test that Tip5s eAT-hook is able to bind DNA, we first cloned, expressed and purified a GST-tagged version of this protein domain (eAT), furthermore point mutant (eATmut) and deletion mutant (eATcore) versions of it. In the eATmut construct 2 arginine residues were mutated to aspartic acid. These point mutations should interfere with the DNA binding as described previously for an AT-hook domain.14 The eATcore deletion mutant was truncated at its N- and C-termini, removing the basic amino acid patches of the peptide, but the core motif was not mutated (Fig. 1B). Since Coomassie-stained gels of different purifications of the eAT-hook consistently showed a dominant slower migrating and a secondary faster migrating protein, both proteins were subjected to mass spectrometry analysis prior to performing functional studies. MALDI-TOF measurements identified the larger protein as the full-length GST-tagged eAT-hook, and the smaller one as its C-terminally truncated version (Fig. S1).

After assessing the purity of the isolated protein, microscale thermophoresis (MST) experiments were performed with GST-tagged Tip5 eAT-hook peptides. A DNA fragment from the enhancer region of the mouse rDNA (“rDNA En”), which has been shown to be bound efficiently by the canonical AT-hooks of Tip5, was used as binding partner at a constant concentration of 50 nM. Note that the same oligonucleotide concentration was used in all subsequent MST experiments to allow direct comparison of the results. The equilibrium binding constant concentration value (EC50), i.e., the eAT-hook concentration at which 50% of the DNA is bound by the protein, was calculated from MST data, resulting in 10 ± 0.8 μM. In contrast, none of the eAT-hook mutants did measurably bind to this DNA fragment (Fig. 1C).

To verify the DNA binding activity of Tip5s eAT-hook, gel shift assays were performed. We included here a peptide containing the first and second AT-hook of Tip5 as positive control. Two other AT-rich sites, namely an “rDNA IGS” fragment and the previously characterized HMGA1 binding site of the interferon β (“IFNβ”) promoter were selected in addition to the “rDNA En” fragment. The results clearly show that the eAT-hook binds to all 3 DNA fragments with similar affinity and the observed
supershifts suggest that more than one eAT-hook molecule binds to one DNA molecule at higher peptide concentrations (Fig. 1D).

The eAT-hook is a novel putative nucleic acid binding domain

In order to identify sequences which are similar to Tip5’s eAT-hook, peptide motifs that contain more than 3 basic amino acid residues on both sides of the G-R-P core motif in a distance of 6–20 amino acids, i.e., (K/R)_{>3}-X_{6–20}-GRP-X_{6–20}-(K/R)_{>3} sequences, were systematically searched in the human and mouse proteome datasets of the Uniprot database (www.uniprot.org). Similar to the sequence homology-based search for AT-hooks, the position of the core G-R-P tripeptide was fixed in our sequence analysis. After removing duplicate hits, 80 and 60 different eAT-hooks were identified in the human and mouse proteomes, respectively (Table S1). Next we characterized these eAT-hooks using the pLogo motif visualization tool, which scales residue heights relative to their statistical significance. Human and mouse eAT-hook sequences were compared to all G-R-P containing peptides of the respective proteomes and the results show a statistically significant overrepresentation of basic amino acids, primarily arginine residues at specific positions. A symmetric appearance of positively charged blocks at a distance of 12–15 amino acids from the G-R-P core, together with an overrepresentation of arginine and lysine amino acids at positions 2, 4, 40 and 42, i.e., 17 and 19 amino acids away from the core motif, were identified as characteristic features of human eAT-hooks (Fig. 2A). Though the overrepresentation of basic amino acids at the aforementioned positions is less prominent, the pLogo visualization of mouse eAT-hooks provides a similar result (Fig. S2A).

As the canonical AT-hook is a short protein motif and only 22 amino acids sequences were analyzed in the original sequence logos, we wondered whether N- and C-terminal extensions of AT-hook peptides show similar sequence patterns to eAT-hooks or differ from them. Notably, the lists of human and mouse AT-hooks slightly vary between different sequence motif repositories, such as InterPro (AT-hook ID: IPR017956), Pfam (AT-hook ID: PF02178) and SMART (AT-hook ID: SM00384). We
selected the human and mouse AT-hook lists of the SMART repository because they contained the most of the sequence entries. The sizes of the AT-hooks were extended to 43 amino acids, which is the predetermined size of eAT-hooks. Additionally we removed duplicated entries from the AT-hook lists, and hypothetical proteins, which also appeared occasionally as redundant sequences. Finally, we generated a set of 44 human and 24 mouse lengthened AT-hook sequences for further sequence analyses (Table S1). Human and mouse AT-hook pLogo images were created, and the sequence visualization clearly shows the expected, characteristic pattern of this sequence motif (Fig. 2B; Fig. S2B). Importantly, the highly significant overrepresentation of basic amino acids in the immediate vicinity of the core motif does not coincide with an overrepresentation of arginine and lysine residues at positions in larger distances from the central G-R-P tripeptide.

The pLogo results and the fact that the eAT-hook lists contain only a few canonical AT-hook sequences already suggested that eAT- and AT-hooks are distinct protein motifs. To confirm this assumption, differences between AT-hook and eAT-hook motifs were investigated using the Two Sample Logo sequence analysis tool. Combined multiple sequence alignments of human and mouse eAT-hooks or AT-hooks were generated, and they were subsequently analyzed by applying 2 different settings (Fig. 2C; Fig. S2C). We observed a highly significant overrepresentation of R residues at positions 8 and 36, as well depletion of the KRKR motif at position 17–20 and the K residue at position 25 in eAT-hooks, when spurious statistical significance is eliminated by Bonferroni correction (P < 0.05, Fig. 2C). Taken together, these results clearly demonstrate that eAT-hooks are not simply elongated canonical AT-hooks, but distinct sequence motifs.

Besides sequence motif analyses, the group of eAT-hook-containing proteins was subjected to functional profiling using the FatiGO web-tool, which is included in the Babelomics suite. In order to obtain statistically significant information, AT-hook containing sequences were removed from the eAT-hook list prior to the analysis. Although nucleic acid metabolism-related gene ontology terms (e.g. DNA binding, RNA binding etc.) associate with approximately half of the proteins, no gene ontology term was found to characterize the eAT-hook proteins with statistical significance. This result

Figure 2. Sequence logo visualizations of eAT-hooks and AT-hooks. (A) pLogo visualization of human eAT-hook peptide sequences. Fixed G-R-P positions are labeled with gray shading. The title indicates that eAT-hooks were used as foreground (FG) and all G-R-P-containing peptides of the human proteome as background (BG) sequences. The n(fg) and n(bg) values at the bottom left of the pLogo indicate the number of aligned foreground and background sequences used to generate the image, respectively. The red horizontal bars on the pLogo correspond to P = 0.05. Note the symmetrical appearance of basic amino acids 12–15 residues away from the G-R-P core motif on both sides. (B) pLogo visualization of human AT-hook peptide sequences. Labels are as described in (A). (C) Two Sample Logo of the differences between eAT-hooks and AT-hooks. The combined groups of human and mouse eAT-hooks vs. AT-hooks were compared. Bonferroni correction (P < 0.05) was applied to eliminate spurious statistical significance. Note the highly significant overrepresentation of R residues at positions 8 and 36, as well depletion of the KRKR motif at position 17–20 and the K residue at position 25 in eAT-hooks.
suggests yet unidentified functions for many of these proteins and implies that their molecular function ontology terms could be further complemented.

The eAT-hook domains of PTOV1 and GPBP1 display similar DNA binding properties to Tip5s eAT-hook

After classifying the new eAT-hook domain as a functional DNA binding motif in Tip5 and identifying a large number of eAT-hooks in the human and mouse proteomes, we set out to test the DNA binding activity of eAT-hook domains found in other proteins. Two candidate proteins, which are involved in nucleic acid metabolism but do not contain known nucleic acid binding domains, were selected. The PTOV1 (Prostate Tumor Overexpressed 1) protein was initially reported as a factor overexpressed in prostate carcinoma cells. In addition, it has been suggested to have some regulatory function in cell cycle progression and was observed to shuttle between the cytoplasm and the nucleus.\textsuperscript{19,20} The human PTOV1 protein consists of 416 amino acids, of which the first 43 residues represent the eAT-hook. In addition, the protein possesses two Med25 (Mediator complex subunit 25; PTOV activation and synapsin 2; Pfam ID: PF11232) motifs, which are predicted transactivation domains. The other selected eAT-hook protein, GPBP1 (GC-rich Promoter Binding Protein 1, also known as Vasculin, vascular wall–subunit 25, PTOV activation and synapsin 2; Pfam ID: PF15337) at the C-terminus (\textsuperscript{21,22} Fig. 3A). The VL1 motif (Vascular protein family Vasculin-like 1; Pfam ID: PF15337) at the C-terminus (Fig. 3A). The VL1 motif is linked to transcription factor function, but its exact role has not been revealed so far.

To analyze the DNA binding activity of the eAT-hooks of PTOV1 and GPBP1, GST-tagged wild-type (eAT), point mutant (eATmut) and deletion mutant (eATcore) versions of the protein domain were cloned. In the eATmut constructs the R-G-R-P and G-R-P cores of PTOV1s and GPBP1s eAT-hook were changed to D-G-D-P and G-D-P, respectively. The eATcore deletion mutants were truncated at their N- and C- termini, removing the basic amino acid patches of the peptides (Fig. 3B). GST-tagged eAT-hook proteins were expressed, affinity purified and analyzed on Coomassie gels prior to functional analyses (Fig. 3C). MST experiments were performed using the “rDNA En” DNA fragment and the same experimental setup as described for Tip5s eAT-hook. The analysis of the MST data yielded 4.0 ± 0.7 \mu M and 7.0 ± 1.7 \mu M equilibrium binding constant concentration values for the eAT-hooks of PTOV1 and GPBP1, respectively. Similar to Tip5, none of the eAT-hook mutants did measurably bind to this DNA fragment (Fig. 3D, E).

In order to confirm the results of MST analyses, gel shift assays were performed using PTOV1s eAT-hook peptides (Fig. 3F). The different eAT-hook proteins were incubated with the AT-rich “rDNA En” sequence under identical experimental conditions and the DNA binding properties of the different GST-eAT proteins were monitored on native polyacrylamide gels. The results clearly show binding of the target DNA by PTOV1s eAT-hook. This is illustrated by the appearance of high molecular weight protein-DNA complexes in lanes 5 and 6. Apparently, with an increase of the protein amount, all DNA was bound to the protein and the DNA was supershifted in lanes 7 and 8. As expected the eATmut and eATcore peptides displayed no significant DNA binding activity in these experiments. In summary PTOV1s and GPBP1s eAT-hooks showed very similar DNA binding behavior to Tip5s eAT-hook.

eAT-hooks bind to RNA with higher affinity than to DNA

Recent publications already suggested a role in RNA interactions for canonical AT-hooks of the HMGA1 and Tip5 proteins.\textsuperscript{23,24} Importantly, it was shown that a stem-loop structure is necessary for the proper RNA binding of the first AT-hook domain of HMGA1.\textsuperscript{25} These observations in combination with a recent report implicating an RNA component in PTOV1 function\textsuperscript{26} made it tempting to speculate about a possible RNA binding activity for the eAT-hook domains. To test this hypothesis, MST analyses with the described GST eAT-hook peptides of Tip5, PTOV1 and GPBP1 were performed. For these experiments two RNA oligonucleotides, which were already used in\textsuperscript{23,24} in vitro protein–RNA interaction studies in our lab, were selected\textsuperscript{26} (Fig. 4A). As predicted by the RNAfold web tool,\textsuperscript{27} oligonucleotide “RNA1” harbours a large stem-loop structure with a long double-stranded region, while “RNA2” comprises only a small hairpin structure and a large loop region. Thus, they provided an interesting structural diversity for the intended assay. For the MST measurements, both RNAs were incubated with dilution series of the GST-tagged Tip5, PTOV1 and GPBP1 eAT-hooks under identical experimental conditions. The analysis was conducted as described previously,\textsuperscript{13} and all data points were normalized to the bound fraction of the RNA molecules and plotted against the protein concentration. The results revealed a strong RNA binding activity for all 3 eAT-hook domains tested. Interestingly, all proteins display a higher affinity for the “RNA1” molecule, with EC50 values of 187 ± 10 nM for Tip5s (Fig. 4B), 203 ± 6 nM for PTOV1s (Fig. 4C) and 378± 18 nM for GPBP1s eAT-hook (Fig. 4D). All three proteins also bind to the “RNA2” molecule, however a 2- to 3-fold decrease in binding affinity was detected. The EC50 values increased to 569 ± 40 nM for Tip5s, 600 ± 30 nM for PTOV1s and 938 ± 92 nM for GPBP1s eAT-hook. In summary the results of the MST analyses show that all tested eAT-hook domains bind to RNA with approximately one order of magnitude better than to DNA, and the secondary structure of RNA influences the binding affinity.

eAT-hooks bind preferentially to RNA in competitive nucleic acid binding assays

To further dissect the function of the eAT-hook domains in nucleic acid binding, competitive MST analyses were performed and the preference for RNA versus DNA was monitored. For this purpose, a FAM-labeled “RNA1” oligonucleotide and a Cy5-labeled “rDNA En” DNA oligonucleotide were mixed in equimolar ratio and MST measurements were carried out and
analyzed as described above. The results strongly support the observed preference for RNA binding of the 3 different eAT-hook domains. Tip5s eAT-hook binds under the competitive conditions with an EC50 of 277 ± 8 nM to the RNA (Fig. 4E), whereas PTOV1s eAT-hook displays an EC50 of 252 ± 10 nM (Fig. 4F), and GPBP1s eAT-hook reveals an EC50 of 373 ± 19 nM (Fig. 4G), which are comparable to the binding affinities measured in the absence of competitor DNA. In all 3 cases no interaction with the DNA was detected, the proteins bound exclusively to RNA.

Figure 3. For figure legend, see page 870.
Deletion of the eAT-hook domain impairs nucleocytoplasmic shuttling of PTOV1

Previous studies revealed that PTOV1 shuttles between the cytoplasm and the nucleus, further it associates with ribonucleoprotein particles which also shuttle. Because eAT-hooks displayed a strong RNA binding activity in the MST experiments, we decided to test the influence of the eAT-hook on PTOV1 shuttling in vivo. For this 2 mammalian expression vector constructs were cloned, encoding a GFP-tagged, full-length PTOV1 fusion protein (PTOV1 WT), as well as an N-terminal deletion mutant lacking the eAT-hook comprising first 48 amino acids (PTOV1 ΔeAT). Since the eAT-hook presents the only characterized nucleic acid interaction domain of PTOV1, this deletion was likely to interfere with its functions in RNA- or DNA-related processes. The two constructs were transfected in HeLa or HCT116 cells and their expression was monitored on western blots, further sub-cellular localization of the expressed GFP fusion proteins was examined by fluorescence microscopy (Fig. 5; Fig. S3). Strikingly, PTOV1 WT and PTOV1 ΔeAT revealed a clear difference in their localization patterns. PTOV1 WT expressing cells show 3 distinct patterns of GFP signal distribution with cells displaying mainly nuclear localization (1), cells revealing evenly dispersed signals in the nucleus and the cytoplasm (2), and cells showing a higher amount of GFP signal in the cytoplasm (3) (Fig. 5A; Fig. S3A). These results were in good agreement with previously published reports, describing similar patterns, caused by the shuttling mechanism of PTOV1. In contrast, PTOV1 ΔeAT shows one particular localization pattern with the GFP signals predominantly located in the nucleus and displaying only very faint cytoplasmic signals (Fig. 5A; Fig. S3A). To get a more detailed view on the differential distribution of the GFP fusion proteins throughout the cells, the transfected cells were examined with a higher magnification, too (Fig. 5B,C). In addition to the 3 different localization patterns of PTOV1 WT, an accumulation of the GFP signal can be observed around the nucleus in cells that contain higher amounts of the protein in the cytoplasm (Fig. 5B). The PTOV1 ΔeAT expressing cells exhibit the primary nuclear localization as seen before (Fig. 5C). The possibility that this effect is simply due to the loss of a nuclear export signal present in the deleted region is unlikely, because a search for this motif using a web based in silico prediction tool delivered a negative result. These observations clearly suggest a role for the eAT-hook in nucleocytoplasmic shuttling and indicate its importance in PTOV1 function.

Discussion

Characterization of the eAT-hook domain

When investigating the function of Tip5s AT-hooks, we have identified an AT-hook-like domain, which was termed extended AT-hook due to the relatively large distances of basic amino acid patches from the core G-R-P motif. Microscale thermophoresis and gel shift assays revealed a similar DNA binding activity of the eAT-hook to the canonical AT-hooks of Tip5. The characterization of the novel Tip5 eAT-hook domain as a functional non-canonical AT-hook set the stage for the identification of a large group of proteins holding this novel putative DNA-binding domain. The widespread distribution of the (K/R)$_{>3}$X$_{6-20}$-GRP-X$_{6-20}$-(K/R)$_{>3}$ sequence throughout the human and mouse proteomes with 80 and 60 different candidate proteins, resembling about twice the number of already mapped AT-hooks, further underlines the relevance of this new motif. Bioinformatics analyses of sequence alignments using the pLogo and Two Sample Logo web tools demonstrate that eAT-hook and AT-hook sequence motifs are clearly distinct from each other. Notably, the central sequence region of some eAT-hooks can also be identified as canonical AT-hook (see Table S1), suggesting that these domains may exhibit the functions of both eAT- and AT-hooks.

The results gained from gel shift and MST analysis of the eAT-hooks of the PTOV1 and GPBP1 proteins emphasize the observations made for the Tip5 eAT-hook motif. Remarkably, all 3 domains display DNA binding affinities in a range comparable to canonical AT-hooks. Moreover, deletion and point mutants of these eAT-hook peptides lose DNA binding activity, further validating the nucleic acid binding capacity of this protein domain. Surprisingly, additional investigations hinted at a strong RNA-binding activity for eAT-hooks. Therefore the data presented here indicates that the eAT-hook is a new functional nucleic acid binding domain, which likely mediates RNA-dependent functions in many proteins. We would like to note that several bioinformatically-identified eAT-hooks may still represent...
RNA oligos used for MST

RNA 1 5’FAM - AGUUCCAUAGUGUUUUCACCCUAUUACCUCACUAUUUGG
RNA 2 5’CY5 - GAGAACGUGUAAAACAAUACAACUGGAGGG

RNAfold structure prediction

B
Microscale thermophoresis
Tip5 - RNA

C
Microscale thermophoresis
PTOV1 - RNA

D
Microscale thermophoresis
GPBP1 - RNA

E
Microscale thermophoresis
Tip5 - RNA vs DNA

F
Microscale thermophoresis
PTOV1 - RNA vs DNA

G
Microscale thermophoresis
GPBP1 - RNA vs DNA

Figure 4. For figure legend, see page 872.
peptides that do not possess nucleic acid binding activity. In contrast, eAT-hooks with variations in the G-R-P core motif may also exist, and such sequences would represent false negatives of our eAT-hook search.

The importance of the eAT-hook for proper protein function in vivo is shown for a selected eAT-hook protein called PTOV1, which contains an N-terminal eAT-hook and no other known DNA- or RNA-binding domain. PTOV1 was initially described as a factor overexpressed in prostate carcinoma cells. The exact function of the protein is so far not known, however a role for PTOV1 in regulation of cell cycle progression was implicated and the shuttling of the protein between the cytoplasm and nucleus has been revealed. Interestingly, a very recent report described the association of PTOV1 with 40 S ribosomes through interaction with RACK1 (receptor of protein C kinase 1), which in turn could control translation initiation of c-Jun. The authors furthermore suggested a function for PTOV1 in ribonucleoprotein shuttling. We showed that PTOV1 accumulates in the nucleus upon deletion of the eAT-hook, which demonstrates the importance of this domain in nucleocytoplasmic shuttling of the protein, and implies that it may have a regulatory role in the aforementioned biological process.

The relationship of eAT-hooks to other short K/R-rich protein sequence motifs

The discovery of various non-canonical AT-hooks already emphasized that it is not possible to present a single sequence motif, which allows the identification of all short peptides with AT-hook-like function. The analysis of the β’ subunits of DNA-dependent RNA polymerases suggested that such sequence motifs appeared very early in evolution possibly playing a role in nucleic acid – protein interactions. Since AT-hook-like motifs show high sequence variability, we have not intended to search for all possible eAT-hook-like sequences, but rather focused on the characterization of several eAT-hooks and generated a sequence logo for human and mouse (K/R)$_{3-1}$X$_{3-20}$-GRP-X$_{20-20}$- (K/R)$_{3-3}$-type eAT-hooks. This sequence logo was significantly different from the sequence logo of 43-amino-acids-sized AT-hooks.

The K/R richness of the eAT-hook motif also raised the question about its possible similarity to short K/R-rich protein sequence motifs, i.e. nuclear localization signals (NLSs) and RGG/RG motifs. Although NLSs frequently overlap nucleic acid binding domains, and eAT-hooks are rich in basic amino acids, they do not represent NLSs. A characteristic feature of NLSs is lysine clustering, but the eAT-hook sequence logo illustrates an arginine-rich sequence motif. The arginine-richness, on the other hand, clearly supports the occasional occurrence of RGG/RG motifs within eAT-hooks. In the 80 human eAT-hooks altogether 1 di-RGG, 2 tri-RG, and 8 di-RG motifs can be identified. Remarkably, the role of RGG/RG motifs in mediating nucleic acid - protein interactions was highlighted recently.

Possible roles for the RNA-binding activity of eAT-hook proteins

Surprisingly, the tested eAT-hook domains of Tip5, PTOV1 and GPBP1 displayed a strong RNA binding activity in MST experiments. The affinities of the 3 different domains measured for RNA are in the hundred nanomolar range and approximately 10- to 20-fold higher than their affinity for DNA molecules. The results of nucleic acid binding analyses also suggest a structural preference of RNA binding. All three tested eAT-hooks bind to the RNA oligonucleotide comprising a helix-loop structure with an almost 3-fold higher affinity than to the RNA featuring a small stem – large loop structure. Noteworthy, this behavior parallels the efficient binding of the first AT-hook of HMGA1 to the double-stranded helix-loop structure of 7SK snRNA. Moreover, in competitive MST assays the presence of DNA does not interfere with the RNA interactions, further validating the preferred RNA binding of the eAT-hook proteins.

In recent years the discovery of an ever-growing number of functional ncRNAs, working as guides, regulators, and scaffolds for the assembly and targeting of chromatin modifying complexes, revealed a completely new layer of gene regulatory processes. Now that numerous novel non-coding roles have been discovered for various RNA classes, it is becoming increasingly important to map RNA-dependent regulatory networks and to further identify and characterize the underlying individual RNA-protein interactions in detail. Strikingly, 2 studies involving HMGA1 and Tip5 already illustrated the capability of AT-hook domains to mediate RNA interactions in regulatory complex formation.
results in a ribonucleoprotein complex consisting of the ncRNA 7SK, HMGA1 and P-TEFb playing a role in the transcriptional regulation of target genes.\textsuperscript{23,33} In Tip5 the deacetylation of a lysine residue within the second AT-hook leads to increased binding of the evolutionarily conserved non-coding pRNA, and to enhanced heterochromatin formation at the rDNA locus.\textsuperscript{24,34} Additionally, ncRNA binding of the insulator protein CTCF was also described recently and the RNA-binding domain was mapped to the C-terminus of the protein.\textsuperscript{35} This chromatin regulator protein does not possess known RNA-binding domains but it contains an AT-hook at its C-terminus.\textsuperscript{36} In the light of these facts, the RNA binding of the eAT-hook could suggest a role for eAT-hooks in RNA-mediated processes within the cell. It is tempting to speculate that eAT-hooks and AT-hooks of some proteins may not only tether and anchor them to particular DNA structures and sequences as originally predicted for AT-hook proteins,\textsuperscript{4} but may have a function as RNA-binding factors. Concerning eAT-hooks, the chromatin-dependent regulatory functions of
Brg1 (Broma related gene 1), Dot1L (Dot1 [Disruptor of telomere silencing protein 1]-like), and Tip5 could involve as well eAT-hook-RNA interactions, since these proteins have been shown to associate with specific IncRNAs. The identification of eAT-hooks in the Rps8, Mtp-S12, Rrp12 and Srsf10 proteins suggests a possible involvement of this domain also in the function of specific ribonucleoprotein complexes, for instance in ribosome and spliceosome function. In addition, 19 eAT-hook proteins can be found in current catalogs of human RNA-binding proteins, which have been identified mostly by their ability to bind mRNA (Table S1). Although the aforementioned eAT-hook proteins can already be classified as factors that bind different RNA molecules, many other eAT-hook proteins are yet uncharacterized. The discovery of the presented novel sequence motif and the characterization of selected eAT-hooks set the ground to explore the function of eAT-hook proteins, and to better understand their role in RNA- and chromatin-dependent biological processes.

**Materials and Methods**

**Plasmid DNA constructs**

GST-eAT-hook constructs were prepared by inserting the coding sequences for the peptides listed in Figures 1B and 3B into BamHI/Sall digested pGEX4T3 plasmid DNA. The pGEX4T3-Tip5-AT1+2 2 plasmid encoding the GST-tagged GKGRGPRRNKANKEPKVGRGPGPRPPKIKMP peptide of Tip5 has been previously described. Synthetic DNA oligonucleotides with flanking BamHI/Sall restriction sites were used for the cloning of eATcore constructs of Tip5, PTOV1 and GPBP1. The Tip5 eAT coding DNA fragment was PCR amplified using primers with flanking BamHI/Sall restriction sites. The DNA inserts for the cloning of eATmut and eAT constructs of PTOV1 and GPBP1 were obtained from synthetic plasmid DNA (GeneArt Gene Synthesis, Life Technologies). The pEGFP-PTOV1 plasmid DNA was cloned by inserting the full-length cDNA of PTOV1 into EcoRI/XbaI digested pEGFP-C2. The insert was PCR amplified from the clone 5928807 (IMAGE ID) of the NIH_MGC_47 library (Source Bioscience). To obtain pEGFP-PTOV1-ΔeAT, the pEGFP-PTOV1 construct was EcoRI/BamHI digested, the ends refilled with the Klenow enzyme and re-ligated. Novel plasmid DNA sequences were analyzed by sequencing.

**Expression and purification of recombinant proteins**

The GST fusion proteins were expressed in the *E. coli* Rosetta 2 strain. Transformed bacteria were induced at OD600 = 0.6 by adding 0.5 mM IPTG and incubated for 3 hours at 37°C. Cells were then harvested and affinity purified on Glutathione Sepharose 4B beads (Pharmacia) in batch according to the manufacturer’s protocol. Protein concentrations of the elution fractions were measured by using a Qubit fluorometer (Invitrogen).

**Mass spectrometry analysis**

GST-eAT proteins were separated on SDS–polyacrylamide gels. After electrophoresis, gels were stained with Coomassie brilliant blue G-250 and the 2 bands appearing in the GST-Tip5-eAT elution fraction were excised. Gel bands were destained, digested with trypsin, and the digestion reactions were analyzed by MALDI-TOF mass spectrometry as previously described. Spectra were manually interpreted.

**Protein – nucleic acid binding assays**

Gel shift assays and microscale thermophoresis (MST) measurements were performed as described in our previous studies. To analyze the nucleic acid binding activity of recombinant eAT-hooks, the following oligonucleotides were used: double-stranded synthetic DNA oligonucleotides (sense strand sequences): “rDNA En” (gel shift): 5'-TTG ATC TTT TTT TTT TTT TTT TTT TTT TTT TTT TTT TTT TTT TTT TTT TTT TTT TTT TTT GAA TTA AAA TTT TTA
Antibodies

To detect proteins by protein gel blot or immunofluorescence, rabbit polyclonal α-actin (Sigma A-2066), rat monoclonal α-GFP (3H, Chromotek) and the GFP-Booster-Atto488 (Chromotek) antibodies were used at appropriate dilutions.

Immunofluorescence

Transfected HeLa and HCT116 cells were grown on coverslips for 24 and 48 hours, respectively, and fixed in 4% paraformaldehyde in PBS for 10 min. During the last minute a few drops of 0.5% Triton X-100 were added. After washes the cells were permeabilized in PBS with 0.5% Triton X-100 for 10 min. Cells were then washed with PBS-T (PBS with 0.1% Tween20) 3 times for 5 min. Subsequently they were incubated with the GFP-Booster-Atto488 (Chromotek) in 4% BSA in PBS-T for 1 h in a humidified chamber followed by 3 times 5 min washes in PBS-T. The second washing solution was supplemented with 50 ng/ml DAPI to counterstain the DNA. Alternatively, only DAPI staining was performed on fixed cells, and GFP fluorescence was directly visualized. Slides were mounted in Vectashield (Vector). Images were taken on a Zeiss Axiovert 200 inverted fluorescence microscope using the AxiosVision software and processed with ImageJ and Adobe Photoshop.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Acknowledgments

The authors thank Regina Gröbner-Ferreira, Verena Kutenberger and David Boymanns for technical help, and Axel Imhof for supporting the MALDI analysis at the Protein Analysis Unit of the LMU Munich.

Funding

This work was supported by the DFG SFB960 program to G. L. and A.N., and the Elite Network of Bavaria to K.Z.

Supplemental Material

Supplemental data for this article can be accessed on the publisher’s website.

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