The RNA-binding proteomes from yeast to man harbour conserved enigmRBPs

Benedikt M. Beckmann1,*, Rastislav Horos1,*, Bernd Fischer1,†, Alfredo Castello1,†, Katrin Eichelbaum1,†, Anne-Marie Alleaume1, Thomas Schwarzl1, Tomaž Curk1,†, Sophia Foehr1, Wolfgang Huber1, Jeroen Krijgsveld1 & Matthias W. Hentze1

RNA-binding proteins (RBPs) exert a broad range of biological functions. To explore the scope of RBPs across eukaryotic evolution, we determined the in vivo RBP repertoire of the yeast *Saccharomyces cerevisiae* and identified 678 RBPs from yeast and additionally 729 RBPs from human hepatocytic HuH-7 cells. Combined analyses of these and recently published data sets define the core RBP repertoire conserved from yeast to man. Conserved RBPs harbour defined repetitive motifs within disordered regions, which display striking evolutionary expansion. Only 60% of yeast and 73% of the human RBPs have functions assigned to RNA biology or structural motifs known to convey RNA binding, and many intensively studied proteins surprisingly emerge as RBPs (termed ‘enigmRBPs’), including almost all glycolytic enzymes, pointing to emerging connections between gene regulation and metabolism. Analyses of the mitochondrial hydroxysteroid dehydrogenase (HSD17B10) uncover the RNA-binding specificity of an enigmRBP.
RNA-binding proteins (RBPs) mediate pivotal cellular functions such as RNA transport, degradation or translation and represent key effectors of post-transcriptional gene regulation. To fulfil such diverse roles, RBPs vary regarding their RNA-binding modes and specificities. Recently developed unbiased high-content techniques to identify RBPs in vivo yielded information on differences in cell type-specific expression and/or RNA-binding activity of RBPs in mammalian cells. Apart from RBPs with defined functions in RNA biology, many other proteins, among them metabolic enzymes, have been found to bind RNA in vivo. The recent discovery of such unorthodox RBPs using mRNA interactome capture raises the question of whether RBPs using mRNA interactome capture2,3,6 raises the question of whether such RBPs. To answer the first question, we determined the evolutionary conservation and the RNA-binding specificity of such RBPs. We identified a large set of RBPs that are conserved between yeast and human cells. We show that this conserved RNA interactome harbours many proteins without previously assigned roles in RNA biology (enigmRBPs), including surprisingly many metabolic enzymes. We also determined the RNA targets of an RNA-binding mitochondrial enzyme and show its specificity in RNA binding.

Results

The mRNA interactomes of yeast and human HuH-7 cells. HuH-7 liver cells were exposed to ultraviolet light of 254 nm (conventional crosslinking, cCL), or of 365 nm (photo-activatable crosslinking, PAR-CL) after incubation with 4-thio-uridine. We also adapted the mRNA interactome capture protocol to yeast (see Methods and Supplementary Fig. 1) using PAR-CL after incubation with 4-thio-uridine. We identified a large set of RBPs that are conserved between yeast and human cells. We show that this conserved mRNA interactome harbours many proteins without previously assigned roles in RNA biology (enigmRBPs), including surprisingly many metabolic enzymes. We also determined the RNA targets of an RNA-binding mitochondrial enzyme and show its specificity in RNA binding.

Validation experiments corroborate the quality of the data sets (Fig. 1d,e). Comparison of the HuH-7 RBPs with the HeLa and HEK293 data sets begins to define a ‘housekeeping’ human mRNA interactome, while 109 HuH-7 RBPs were previously not found in other human cell types (Fig. 1f; Supplementary Data 2). The latter may be explained by cell-specific expression or culture condition-dependent RNA binding of RBPs. Our data confirm 108 of the 120 yeast RBP candidates from a recent study. Importantly, we identified 570 additional yeast proteins as high-confidence RBPs (Fig. 1g; Supplementary Data 1), and an astounding >10% of the total yeast proteome thus emerge as RBPs (see below).

Definition of the conserved eukaryotic mRNA interactome. These sets of yeast and human RBPs were analysed for RBP conservation. Comparison of yeast with each of the three human cell lines consistently yields an overlap of >200 orthologous RBPs (Fig. 2a). We used the InParanoid database that assigns RBPs to ‘ortholog groups’ of corresponding biological functions (Methods). Two-hundred and thirty ortholog groups consisting of 243 individual RBPs in yeast and 256 RBPs in human (Fig. 2b; Supplementary Data 1 and 2) constitute the conserved eukaryotic ‘core mRNA interactome’. As expected, it includes many well-studied RBPs with established functions in RNA biology and/or with well-defined RNA-binding domains (Supplementary Fig. 3a,b). Interestingly, some ‘core RBPs’ share lysine [K]- and arginine [R]-rich tripeptide repeat motifs that numerically increase from yeast to human (Fig. 2c,d). Whereas their general occurrence is conserved, their number within orthologous RBPs expands with increasing complexity of the organisms. These expansions may directly interact with RNA and represent an emerging structural property of nucleic acid interactors, possibly to enhance RNA-binding specificity within increasingly complex transcriptomes. The RBPs eIF3a and THOC2 are depicted as examples in Fig. 2c.

Well-studied proteins emerge as conserved enigmRBPs. Merging our data sets with published information on yeast (690 RBPs in total, Supplementary Data 1) and human RBPs (1,217 RBPs in total, Supplementary Table 2), we were surprised to find that 40% (274 yeast proteins) and 27% (326 human proteins), respectively, of the identified RBPs lack both recognizable RBDs and known functions in RNA biology (selection criteria see Methods). These RBPs include many well-studied proteins whose roles in RNA biology remain to be defined (Fig. 3a); we therefore termed these proteins ‘enigmRBPs’. Phosphoglycerate kinase and thioredoxin represent enigmRBPs that we directly validated for RNA binding both in human and yeast cells (Fig. 1d,e and Castello et al., submitted). enigmRBPs cover a wide spectrum of biological functions.

Figure 1 | mRNA interactome capture in yeast and HuH-7 cells. (a) Schematic representation of the mRNA interactome capture protocol in yeast and HuH-7 cells using PAR crosslinking (PAR-CL) or conventional crosslinking (cCL). Quality controls for both yeast and HuH-7 include silver staining (middle upper and lower panel) and western blotting (right upper and lower panel) (M, protein marker, KHD1, KH-domain containing protein fused to eGFP, PUB1, polyU binding protein 1, TUB1, tubulin 1, H4, histone 4, PTBP1, polyprymidine tract binding protein 1, CSDE1, cold shock domain containing protein E1). (b) Volcano plot showing the distribution of proteins according to their enrichment in crosslinked (CL) over non-CL samples. Proteins shown in red (FDR 0.01) represent the mRNA interactome. (c) Overview of mRNA interactome proteins in yeast and HuH-7. (d) Validation of the yeast mRNA interactome using western blotting of input samples and eluate after interactome capture with specific antibodies (ADH1, alcohol dehydrogenase 1, PUB1) or against TAP-tagged proteins (PGK1, phosphoglycerate kinase 1, TDH1, triose phosphate dehydrogenase, TRX2, thioredoxin 2, SHE2, Swhy5-dependent Holo Expression 2). (e) Validation of HuH-7 mRNA interactome RBPs using polynucleotide kinase-mediated 32P labelling after IP with an anti-eGFP antibody (negative control eGFP and MRPP3, mitochondrial RNAseP protein 3; positive control MOVI10, monoloy leukaemia virus 10; and novel RBPs, HSD17B10, hydroxysteroid dehydrogenase 17B 10, c11orf68, chromosome 11 open reading frame 68, CRKL, v-crk avian sarcoma virus CT10 oncogene homologue-like, LLPH, long-term synaptic facilitation homolog, PGK1, phosphoglycerate kinase 1). (f) Comparison of mRNA interactome RBPs from HuH-7 with those from HeLa and HEK293 cells. (g) Comparison of the yeast mRNA interactome RBPs to a published study using cCL.
**a**

in vivo 4thio-U labeling

UV crosslinking

**b**

Yeast

HuH-7

PAR-CL 365 nm

PAR-CL 365 nm

PAR-CL 254 nm

+/– in vivo 4thio-U labeling

UV crosslinking

**c**

PAR-CL 0.72 J/cm²

PAR-CL 7.2 J/cm²

PAR-CL 0.72 J/cm²

PAR-CL 7.2 J/cm²

PAR-CL

PAR-CL

**d**

PAR-CL J cm⁻²

**e**

Phosphoimage

Anti-eGFP

Phosphoimage

Anti-eGFP

**f**

Castello et al.

HuH-7

109

99

84

92

154

860

437

12

**g**

Mitchell et al.

PAR-CL

729

108

570

H4

Tubulin

PTBP1

CSDE1

β-actin

PAR-CL

PAR-CL

PAR-CL

PAR-CL

PAR-CL

PAR-CL
functions, including, for example, actin-binding/remodelling, protein folding, ATP-binding and enzymatic functions in classic metabolic pathways (Supplementary Table 1). enigmRBPs resemble orthodox RBPs in terms of typical features (Supplementary Fig. 3c,d) in both yeast and human cells, and constitute a surprisingly large fraction of the conserved core RBPs (42 (17%) yeast and 28 (11%) human RBPs). A few enzymes of classical biochemical pathways that moonlight as RBPs have previously been identified16–18. We thus searched the complete yeast and human RBP data sets (including the HuH-7 candidate RBPs) for ‘classic’ metabolic enzymes (hereafter referred to as enzymes; for selection criteria see Methods). Yeast RBPs (17%) and 9% of the human RBPs belong to this group (Fig. 3b; Supplementary Data 1 and 2); transferases and oxidoreductases constitute more than half of these (Supplementary Fig. 3e).

Strikingly, 9% of the conserved core RBPs are metabolic enzymes (Fig. 3c; Supplementary Fig. 3f), and central carbon metabolism, especially glycolysis, emerges as a hotspot for RNA-binding enzymes (Fig. 3d,e).

**RNA binding of a mitochondrial enzyme.** To explore the binding specificity of metabolic enzymes for RNA, we performed iCLIP19 and determined the interacting RNAs of the mitochondrial enzyme HSD17B10, which is mutated in patients with a mitochondrial cardiomyopathy/neuropathy syndrome (OMIM 300438), and for which a non-enzymatic function has been suspected to account for the disease phenotype 20. HSD17B10 has been described as a subunit of the RNase P complex 21 (together with TRMT10C and MRPP3) that processes mitochondrial tRNAs, which are interspersed within polycistronic mitochondrial transcripts22. Of note, MRPP3 did not appear in any of the human mRNA interactome data sets, nor could we detect ultraviolet-crosslinked RNAs on the protein (Fig. 1e). Thus, the RNA binding presumably resides on TRMT10C (RNA methyltransferase), the tetrameric HSD17B10 (dehydrogenase) or the complex of these two23. We observed enriched binding of HSD17B10 to mitochondrial RNAs (Supplementary Fig. 4a,b), and compared the RNA-protein crosslink sites of wt HSD17B10 to an eGFP background control.
We found that HSD17B10 preferentially binds at the 5' ends of tRNAs (Fig. 4d), especially their D-stem, D-loop and anticodon stem and loop regions, on 15 out of 22 mt tRNAs (Supplementary Fig. 5); suggesting that the mitochondrial RNAse P does not mediate processing of all tRNAs. Indeed, processing of tRNAs that are encoded in clusters (tRNA<sup>His</sup>, tRNA<sup>Ser(AGY)</sup>, tRNA<sup>Leu(CUN)</sup>, Supplementary Fig. 5) was suggested to be mediated by the combination of RNAse P and the ELAC2 complex, which processes 3' end of mt tRNAs<sup>24</sup>. Next, we performed iCLIP on the disease-associated variant, HSD17B10.
R130C, that causes the classical phenotype of HSD10 disease, retains the ability for tetramerization and displays a reduced interaction with TRMT10C in vitro. Of note, the R130C variant exhibits a decreased binding signal to several pre-tRNAs (Fig. 4f; Supplementary Data 4). Our data identify the mitochondrial enzyme HSD17B10 as the RNA-binding subunit of RNaseP in vivo, and reveals that the R130C mutant is deficient in binding of a subset of pre-tRNAs. They also identify an RBP from the dehydrogenase enzyme family with clear RNA-binding specificity.

**Discussion**

Taken together, the data identify a surprisingly high number of RBPs in yeast and humans, including many previously well-characterized proteins that emerge to have conserved RNA-binding activity in vivo (enigmRBPs). Although in vivo RNA binding does not prove physiological function per se, we note that the enzyme β-hydroxysteroid dehydrogenase displays remarkable RNA-binding specificity (Fig. 4). Moreover, the two enzymes aconitase 1 (ref. 18) and GAPDH are known to function as regulatory RNA-binding proteins in vivo, suggesting that other enigmRBPs may also moonlight as post-transcriptional regulators. Alternatively, RNAs could regulate enigmRBPs: by competition with substrates for binding sites within enzymes, as allosteric regulators, or as assembly scaffolds for alignment of enzymes in a biochemical pathway. RNA binding could also influence the folding, assembly or fate of newly synthesized proteins emerging from the ribosome, especially considering the fact that lysine and arginine-rich sequences have a propensity to induce ribosome stalling and protein degradation. The innate immune effectors PKR, TLR3, TLR7, TLR8 and RIG-1 are controlled by pathogen-derived RNAs. We propose that endogenous ‘effectorRNAs’ could serve roles akin to protein–protein interactors for enigmRBPs, and endow the genome with the possibility to affect mature proteins.
Methods

Yeast cultures. Yeast colonies (BY4741, TAP-tagged strains) were used to inoculate a 5 ml YPAD pre-culture overnight at 30 °C and 160 r.p.m. The next day, 1 litre SC-medium 2% Ura (YNB, SD-Ura, 120 mM Ura, 1% glucose) cultures were started with an OD600 of 0.01–0.05.

Huh7 cell culture. Cells (kind gift from M. Muckenthaler, MPPU, Heidelberg) were cultured in low glucose (5 mM) DMEM supplemented with 10% heat inactivated FCS (PAA). For the validation procedures, we derived a Huh-7 Flp-In TReX cell line using published protocols (Flp-In T-Rex, Life), and prepared stably expressing tetracycline-inducible cell lines with genes of interest following manufacturer’s instructions. The cells were grown in medium containing blasticidin (5 μg ml⁻¹) and zeocin (100 μg ml⁻¹) or hygromycin (200 μg ml⁻¹). Transfections were done using Lipofectamine (18324-012, Life).

Cloning. Human genes of interest were cloned into pcDNA5_FRT_TO (Life). The detailed cloning strategies and primer sequences are described in Supplementary Table 2.

mRNA interactome capture. For Huh-7 cells, experiments were done with minor modifications in the cell lysis procedure to previously described procedure. The cells were washed twice with PBS on ice before ultraviolet crosslinking. After crosslinking, the cells were lysed directly with lysis buffer on the cell culture plates, scraped and collected into 50 ml tubes. Lysates were sheared through 27G needle and incubated with oligo d(T) beads (volume ratio lysate to beads 15:1) for 1 h at 4 °C. Beads were then washed twice with each wash buffer and pooled elutions from three rounds of purifications were used for RNA treatment and subsequent protein analysis.

For yeast mRNA interactome capture, cells were grown as described above to an OD600 of 0.5 before adding 4-thiouracil (4U, Sigma 440736) to a final concentration of 500 μM. Cells were allowed to grow for 3 h before harvesting by centrifugation (4,000 r.p.m.; 15 min; 4 °C). The cell pellet was dissolved in 40 ml cold water and spread onto two Petri dishes. Ultraviolet crosslinking was performed on ice in a Spectrolinker device (Spectronics, XL1500/F/A emitting 365 nm wavelength using energies from 0.72 to 7.2 J cm⁻²). Cells were re-pelleted (4,000 r.p.m.; 5 min; 4 °C), and pellets were dissolved in 2 ml lysis buffer (20 mM Tris pH 7.5, 500 mM LiCl, 0.5% LDS, 1 mM EDTA, 5 mM diithiothreitol (DTT), 1% protease inhibitor mix (EDTA-free, Roche), 1 mg ml⁻¹ RNasin, 200 mM VRC). Cells were distributed into 1 ml screw-capped tubes containing an equivalent of 300 RNasin, 200 mM VRC). Cells were distributed into 2 ml screw-capped tubes containing an equivalent of 300 RNasin, 200 mM VRC). Cells were washed twice with PBS on ice before ultraviolet crosslinking. After crosslinking, the cells were lysed directly with lysis buffer on the cell culture plates, scraped and collected into 50 ml tubes. Lysates were sheared through 27G needle and incubated with oligo d(T) beads (volume ratio lysate to beads 15:1) for 1 h at 4 °C. Beads were then washed twice with each wash buffer and pooled elutions from three rounds of purifications were used for RNA treatment and subsequent protein analysis.

This UV254-induced RNA damage represents the likely cause of low RBP recovery after limited irradiation with ultraviolet light at 254 nm (Supplementary Fig. 1). (see above) using an RNA Pico Chip (Agilent BioAnalyzer 2100) was found to be RBP crosslinking, or the integrity of total RNA and RNA after oligo-d(T) selection observed contaminants as reversed sequences of all entries had been added. Initial maximal allowed mass tolerance was set to 20 p.p.m. for peptide masses, followed by 6 p.p.m. in the main search, and 0.5 Da for fragment ion masses. The minimum peptide length was set to six amino-acid residues, and three labelled amino-acid residues were allowed. A 1% FDR was required at both the protein and peptide level. In addition to the FDR threshold, proteins were considered identified if they had at least one unique peptide. Protein identifications were transferred between matching runs, based on the retention time (2 min window) and the accurate peptide masses determined in the Orbitrap analyzer. Protein identification was reported as an indistinguishable ‘protein group’ if no unique peptide sequence to a single database entry was identified. Protein quantification was based on unique and razor peptides.

Definition of mRNA interactome proteins. Peptide UniProt accession numbers were converted into ENSEMBL gene IDs. Where multiple ENSEMBL gene IDs applied, the peptide group was not considered. Statistical analysis was performed using an empirical Bayes moderated t-test within the Limma package in R/Bioconductor. P values were adjusted for multiple testing using the method of Benjamini and Hochberg. Proteins within FDR 1% were considered as mRNA interactome proteins.

SDS-PAGE, western blotting and silver staining. The procedures were performed according to the standard protocols. Antibodies used were anti-eGFP (1:3,000, H9, Chromotek), PUB1 (1:3,000, kind gift from Maurice Swanson), tubulin, histone H4 (1:4,000, Abcam), Calnexin (1:4,000, Abcam), PTBP1 (1:1,000, 572581M, Sigma), β-actin (1:2,000, A5441, Sigma) and CSD1E1 (1:12,000, 13319-1-AP, PTG). The TAP-tagged yeast proteins were visualized by staining with anti-rabbit secondary conjugates against HRP (1:10,000, NA934V, GE). Uncropped blots are shown in the Supplementary Fig. 6.

Polynucleotide kinase assay. Cells expressing tagged genes of interest were ultraviolet crosslinked, lysed (100 mM KCl, 5 mM MgCl₂, 10 mM Tris pH 7.5, 0.5% NP40, 1 mM DTT, protease inhibitors) and homogenized with ultrason (3 × 10 s, 50% amplitude) on ice. Cleared lysates were treated with 50 μl⁻¹ DNase1 (Takara) and 8 ng μl⁻¹ RNase A (Sigma) for 15 min at 37 °C, and used for immunoprecipitation with anti-eGFP coupled to magnetic beads (Chromotek) for 2 h at 4 °C. Beads were washed 4 × with washing buffer (500 mM NaCl, 20 mM Tris pH 7.5, 1 mM MgCl₂, 0.05% NP40, protease inhibitors) and 2 × with polynucleotide kinase (PK) buffer (50 mM Tris pH 7.5, 50 mM NaCl, 10 mM MgCl₂, 0.5% NP40, 5 mM DTT). Beads were resuspended in PK buffer with 0.1 μg ml⁻¹ γ-32P [ATP (Hartmann) and 1 μl µl⁻¹ T4 PNK (NEB) and labelled for 15 min at 37 °C. After five washes with PK buffer without DTT, beads were boiled and loaded onto SDS-PAGE gels. The blot was then autoradiographed exposed to x-ray film. IP efficiency was controlled by anti-eGFP antibody (1:3,000, H9, Chromotek). Uncropped blots and phosphorimages are shown in the Supplementary Fig. 6.

RBPs classifications. We used the Gene Ontology (GO) database to curate the protein list (ENSEMBL gene identifiers). We assigned proteins as ‘linked mRNA biology’ if their associated GO terms contained at least one of the following terms:

- ‘mRNA’
- ‘splice’
- ‘RNA binding’
- ‘RNA’
Some of the above terms were shortened from original words on purpose to capture multiple variations of these. If the protein did not fall in this category, we assigned it as ‘unknown’ in RNA biology. Domain classification based on Interpro and Pfam domains was described in ref. 2. We devised new category for RBD domains ‘recognized’ that encompasses both ‘classical’ and ‘non-classical’ categories from ref. 2.

**Complete human mRNA interactome data set.** For the complete human mRNA interactome data set, we used the combination of proteins found in this study (FDR 1%), proteins from HeLa mRNA interactome and HKE293 cell RNA-binding proteins (Supplementary Data 1).14

**Complete yeast mRNA interactome data set.** For the complete yeast mRNA interactome data set, we combined the proteins of this study (FDR 1%) and the proteins from ref. 9.

**Disordered regions and low complexity.** The intrinsic disorder of proteins is computed with IUPred20. Disordered amino-acid residues are defined by a IUPred score of >0.4. For each protein, the fraction of disordered amino-acid residues is computed. To assess complexity, shannon entropy is computed for each amino acid position within a window of ±10 residues. Positions with an entropy <3 bits are considered as low complexity. For each protein, the fraction of amino-acid residues in low-complexity regions is computed.

**Ortholog definition.** To define orthologs of yeast and human, ENSEMBLE gene ID were converted to UniProt IDs and used for the InParanoid ortholog groups database (Release 7.0, June 2009)27. There were 2,041 InParanoid ortholog groups in total, covering 3,670 human and 2,586 yeast proteins. InParanoid clusters were categorized according to their mRNA-binding behaviour. An InParanoid cluster was regarded as mRNA binding, if at least one of the contained proteins is included in the complete human or yeast mRNA interactome data set. This categorization resulted in three groups of InParanoid clusters: some showed mRNA binding in human and yeast, some showed mRNA binding only in human and some did not show mRNA binding either in yeast or in human. There was not a single InParanoid cluster that shows mRNA binding in yeast, but not in human.

**K-mers motif evolution across species.** The InParanoid database was used to find clusters from *Caenorhabditis elegans* (C. elegans), *Drosophila melanogaster* (D. melanogaster) and *Danio rerio* (D. rerio), which contain orthologs to the 2,041 ortholog groups described above. Within each InParanoid cluster, the protein with the longest (isofrom) sequence was chosen as the representative for the cluster. For each yeast–human orthology group, one orthologous protein in fish (respectively, *D. melanogaster*) was chosen. A list of K-mers was computed for each organism, protein in yeast and the representative protein in human. If there were multiple orthologous protein in fish (respectively, *D. melanogaster*) of the same yeast–human orthology group, one orthologous protein in fish (respectively, *D. melanogaster*) was selected. The K-mers contained an ‘X’ or ‘U’ were excluded. A table counting the repeat number of each K-mer for each protein in each organism. Next, we created a vector of K-mers for each protein in each organism. If there were multiple orthologous protein in fish (respectively, *D. melanogaster*) of the same yeast–human orthology group, one orthologous protein in fish (respectively, *D. melanogaster*) was selected. A table counting the repeat number of each K-mer for each protein in each organism. We then used the fraction of disordered amino-acid residues in low-complexity regions is computed.

**GO enrichments analysis.** For GO enrichment analysis, the DAVID database (version 6.7)28,29 was used. As a background, the total human or yeast proteomes, ‘classic’ metabolic enzymes were used the IntEnz database (http://www.ebi.ac.uk/ intenz/) to classify all enzymes in the mRNA interactomes. For the purpose of this study and to exclude obvious RNA-related enzymes, we did not consider the following as classic metabolic enzymes:

- RNA/DNA helicases
- RNA polymerases
- tRNA modification enzymes (tRNA methyltransferase, pseudouridyllases and so on)
- nucleases (RNA or DNA)
- tRNA aminoacylsynthetases
- RNA/DNA polymerases
- topoisomerases
- proteosome subunits
- regulatory subunits (of any enzyme)

**iCLIP and data analysis.** iCLIP was performed with following modifications. Stably expressing HuH-7 cells were induced overnight with 100 ng/ml - 1 of tetracycline, ultraviolet-crosslinked and lysed on plate. Lysates were homogenized using Branson sonifier (3 × 10 s, 50% amplitude) and cleared at 13,000 r.p.m. for 10 min. IP with anti-GFP magnetic beads was performed as described above and following washes were applied twice each: high-salt wash (500 mM NaCl; 20 mM Tris pH 7.5; 1 mM MgCl₂; 0.05% NP40; 1% SDS), medium-salt wash (250 mM NaCl; 20 mM Tris pH 7.5; 1 mM MgCl₂; 0.05% NP40) and low-salt wash (150 mM NaCl; 20 mM Tris pH 7.5; 1 mM MgCl₂; 0.01% NP40). After RNA treatment and dephosphorylation of 5’ ends, RNA linker was ligated overnight at 850 r.p.m. at 16°C. Beads were treated with proteinase K and cluates were used for RNA isolation, cDNA production and sequencing following published protocol30. Low-FDR crosslink sites read values were then used for the DESeq analysis40.

**References**

1. Gerstberger, S., Hafner, M. & Tuschl, T. A census of human RNA-binding proteins. *Nat. Rev. Genet.* **15**, 829–845 (2014).
2. Castello, A. et al. Insights into RNA biology from an atlas of mammalian mRNA-binding proteins. *Cell* **149**, 1393–1406 (2012).
3. Baltz, A. G. et al. The RNA-binding proteome and its global occupancy profile on protein-coding transcripts. *Mol. Cell* **46**, 674–690 (2012).
4. Cech, T. R. & Steitz, J. A. The noncoding RNA revolution-trashing old rules to forge new ones. *Cell* **157**, 77–94 (2014).
5. Hentze, M. W. & Preiss, T. The REM phase of gene regulation. *Trends. Biochem. Sci.* **35**, 433–446 (2010).
6. Kwon, S. C. et al. The RNA-binding protein repertoire of embryonic stem cells. *Nat. Struct. Mol. Biol.* **20**, 1122–1130 (2013).
7. Castello, A. et al. System-wide identification of RNA-binding proteins by interactome capture. *Nat. Protoc.* **8**, 491–500 (2013).
8. Creamer, T. J. et al. Transcriptome-wide binding sites for components of the Saccharomyces cerevisiae non-poly(A) termination pathway: Nrd1, Nab3, and Sen1. *PLoS Genet.* **7**, e1002329 (2011).
9. Mitchell, S. F., Jain, S., She, M. & Parker, R. Global analysis of yeast mRNPs. *Nat. Struct. Mol. Biol.* **20**, 127–133 (2013).
10. Picotti, P. et al. A complete mass-spectrometric map of the yeast proteome applied to quantitative trait analysis. *Nature* **494**, 266–270 (2013).
11. Kato, M. et al. Cell-free formation of RNA granules: low complexity sequence domains form dynamic fibers within hydrogels. *Cell* **149**, 753–767 (2012).
12. Jonas, S. & Izaurrealde, E. The role of disordered protein regions in the assembly of decapping complexes and RNP granules. *Genes Dev.* **27**, 2628–2641 (2013).
13. Han, T. W. et al. Cell-free formation of RNA granules: bound RNAs identify features and components of cellular assemblies. *Cell* **149**, 768–779 (2012).
14. Wu, C. et al. Identification of novel nuclear targets of human thioredoxin 1. *Mol. Cell Proteomics* **13**, 3507–3518 (2014).
15. Hakimi, H. et al. Plasmodium knowlesi thioredoxin peroxidase 1 binds to nucleic acids and has RNA chaperone activity. *Parasitol. Res.* **113**, 3957–3962 (2014).
16. Chang, C. H. et al. Posttranscriptional control of T cell effector function by aerobic glycolysis. *Cell* **153**, 1239–1251 (2013).
17. Tsvetanova, N. G., Klass, D. M., Salzmann, J. & Brown, P. O. Proteome-wide search reveals unexplored RNA-binding proteins in Saccharomyces cerevisiae. *PloS ONE* **5**, e12671 (2010).
18. Hentze, M. W., Muckenthaler, M. U., Galy, B. & Camaschella, C. Two to tango: regulation of Mammalian iron metabolism. *Cell* **142**, 24–38 (2010).
19. Konig, J. et al. iCLIP—transcriptome-wide mapping of protein-RNA interactions with individual nucleotide resolution. *J. Vis. Exp.* e2638, doi:10.3791/2638 (2011).
20. Rauschenberger, K. et al. A non-enzymatic function of 17beta-hydroxysteroid dehydrogenase type 10 is required for mitochondrial integrity and cell survival. *EMBO Mol. Med.* **2**, 51–62 (2010).
21. Holzmann, J. et al. RNAse P without RNA: identification and functional reconstitution of the human mitochondrial RNA processing enzyme. *Cell* **135**, 462–474 (2008).
22. Ojala, D., Montoya, J. & Attardi, G. RNA punctation model of RNA processing in human mitochondria. *Nature* **290**, 470–474 (1981).
23. Vilard, E. et al. A subclass of human mitochondrial RNase P is a bifunctional methyltransferase—extensive moonlighting in mitochondrial RNA biogenesis. *Nucleic Acids Res.* **40**, 11583–11593 (2012).
24. Rossmanath, W. Processing of human mitochondrial RNA(Ser(AGY))GCU: a new pathway in RNA biogenesis. *J. Mol. Biol.* **265**, 365–371 (1997).
25. Vilard, E. & Rossmanath, W. Molecular insights into HSD10 disease: impact of SORCS1 mutations on the human mitochondrial RNase P complex. *Nucleic Acids Res.* **43**, 5112–5119 (2015).
26. Mitchell, S. F. & Parker, R. Principles and properties of eukaryotic mRNPs. *Mol. Cell* **54**, 547–558 (2014).
27. Hentze, M. W. & Preiss, T. Circular RNAs: splicing’s enigma variations. *EMBO J.* **32**, 923–925 (2013).
28. Lu, J. & Deutsch, C. Electrostatics in the ribosomal tunnel modulate chain elongation rates. *J. Mol. Biol.* **384**, 73–86 (2008).
29. Charneresi, C. A. & Hurst, L. D. Positively charged residues are the major determinants of ribosomal velocity. *PLoS Biol.* **11**, e1001508 (2013).
30. Kim, Y. et al. PKR is activated by cellular dsRNAs during mitosis and acts as a mitotic regulator. *Genes Dev.* **28**, 1310–1322 (2014).
31. Yu, M. & Levine, S. J. Toll-like receptor, RIG-I-like receptors and the NLRP3 inflammasome: key modulators of innate immune responses to double-stranded RNA viruses. *Cytokine Growth Factor Rev.* **22**, 63–72 (2011).
32. Boersema, P. J., Raijmakers, R. & Lemeer, S. & Mohammed S, Heck AJ. Multiplex peptide stable isotope dimethyl labeling for quantitative proteomics. *Nat. Protoc.* **4**, 484–494 (2009).
33. Rappsilber, J., Mann, M. & Ishihama, Y. Protocol for micro-purification, enrichment, pre-fractionation and storage of peptides for proteomics using StageTips. *Nat. Protoc.* **2**, 1896–1906 (2007).
34. Cox, J. & Mann, M. MaxQuant enables high peptide identification rates, individualized p.p.b.-range mass accuracies and proteome-wide protein quantification. *Nat. Biotechnol.* **26**, 1367–1372 (2008).
35. Smyth, G. K. Linear models and empirical bayes methods for assessing differential expression in microarray experiments. *Stat. Appl. Genet. Mol. Biol.* **3**, 1–25 (2004).
36. Dosztáni, Z., Csizmok, V., Tompa, P. & Simon, I. IUPred: web server for the prediction of intrinsically unstructured regions of proteins based on estimated energy content. *Bioinformatics* **21**, 3433–3434 (2005).
37. Ostlund, G. et al. InParanoid 7: new algorithms and tools for eukaryotic orthology analysis. *Nucleic Acids Res.* **38**, D196–D203 (2010).
38. Huang, D. W., Sherman, B. T. & Lempicki, R. A. Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. *Nat. Protoc.* **4**, 44–57 (2009).
39. Huang, D. W. et al. DAVID Bioinformatics Resources: expanded annotation database and novel algorithms to better extract biology from large gene lists. *Nucleic Acids Res.* **35**, W169–W175 (2007).
40. Love, M. I., Huber, W. & Anders, S. Moderated estimation of fold change and dispersion for RNA-seq data with DESeq2. *Genome Biol.* **15**, 550 (2014).

**Acknowledgements**

We thank C. Strein for valuable assistance, and T. Preiss as well as A. Lamond for insightful discussions. We are grateful to A.-C. Gavin for TAP-tagged yeast strains and M. Swanson for antibodies. This work was supported by a Marie Curie Fellowship (FP7/2007-2013/MC-IEF-301031 to B.M.B.), Virtual Liver Consortium (German Ministry for Education and Research to M.W.H.) and European Research Council Advanced Grant (to M.W.H.).

**Author contributions**

B.M.B, R.H., W.H., J.K. and M.W.H. designed the project; B.M.B, R.H., A.C., K.E., A.-M.A. and S.F. performed the research; B.F., T.S., T.C., B.M.B, R.H., W.H., J.K. and M.W.H. analysed data; B.M.B, R.H., J.K. and M.W.H. wrote the paper with input from all authors.

**Additional information**

Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

Competing financial interests: The authors declare no competing financial interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

How to cite this article: Beckmann, B. M. et al. The RNA-binding proteomes from yeast to man harbour conserved enigmRBPs. *Nat. Commun.* 6:10127 doi: 10.1038/ncomms10127 (2015).

This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/