Genome-wide analysis of Staufen-associated mRNAs identifies secondary structures that confer target specificity

John D. Laver1, Xiao Li1, Kristin Ancevicius2,3, J. Timothy Westwood2,3,*, Craig A. Smibert1,4,*, Quaid D. Morris1,5,* and Howard D. Lipshitz1,*

1Department of Molecular Genetics, University of Toronto, 1 King’s College Circle, Toronto, Ontario, Canada M5S 1A8, 2Department of Cell & Systems Biology, University of Toronto at Mississauga, 3359 Mississauga Road, Mississauga, Ontario, Canada L5L 1C6, 3Department of Biology, University of Toronto at Mississauga, 3359 Mississauga Road, Mississauga, Ontario, Canada L5L 1C6, 4Department of Biochemistry, University of Toronto, 1 King’s College Circle, Toronto, Ontario, Canada M5S 1A8 and 5Banting and Best Department of Medical Research, Terrence Donnelly Centre for Cellular and Biomolecular Research, 160 College Street, Toronto, Ontario, Canada M5S 3E1

Received February 14, 2013; Revised July 17, 2013; Accepted July 18, 2013

ABSTRACT

Despite studies that have investigated the interactions of double-stranded RNA-binding proteins like Staufen with RNA in vitro, how they achieve target specificity in vivo remains uncertain. We performed RNA co-immunoprecipitations followed by microarray analysis to identify Staufen-associated mRNAs in early Drosophila embryos. Analysis of the localization and functions of these transcripts revealed a number of potentially novel roles for Staufen. Using computational methods, we identified two sequence features that distinguish Staufen’s target transcripts from non-targets. First, these Drosophila transcripts, as well as those human transcripts bound by human Staufen1 and 2, have 3’ untranslated regions (UTRs) that are 3–4-fold longer than unbound transcripts. Second, the 3’UTRs of Staufen-bound transcripts are highly enriched for three types of secondary structures. These structures map with high precision to previously identified Staufen-binding regions in Drosophila bicoid and human ARF1 3’UTRs. Our results provide the first systematic genome-wide analysis showing how a double-stranded RNA-binding protein achieves target specificity.

INTRODUCTION

RNA-binding proteins (RBPs) direct many co- and post-transcriptional processes. There are a number of different classes of RBPs that are defined by the presence of different RNA-binding domains (RBDs) (1). One class of RBP is double-stranded RBPs (dsRBPs), defined by the presence of one or more double-stranded RBDs (dsRBDs). dsRBDs are characterized by a conserved \( \text{abba} \) fold (2–4) and bind specifically to double-stranded RNA (dsRNA) (5,6). Proteins containing dsRBPs have roles in diverse processes and include Escherichia coli RNase III, Xenopus laevis Xlrbpa, a dsRBP associated with cellular RNAs and ribosomes, the dsRNA-dependent protein kinase PKR, dsRNA-dependent adenosine deaminases (ADARs), and Dicer, an important component of the RNA interference (RNAi) machinery.

One of the best-characterized dsRBPs is Staufen, an evolutionarily conserved protein first identified in Drosophila (7,8). In Drosophila, Staufen is essential for localization and translation of oskar mRNA at the posterior of the oocyte (9-11), for the anchoring of bicoid mRNA at the anterior of the early embryo (12,13) and for asymmetric localization of prospero mRNA in dividing embryonic neuroblasts (14–17).

Mammals possess two Staufen homologs, Staufen1 and Staufen2, both of which function in developing and adult neurons (18–21). Staufen2 has also been shown to

*To whom correspondence should be addressed. Tel: +416 946 5296; Fax: +416 971 2494; Email: howard.lipshitz@utoronto.ca
Correspondence may also be addressed to J. Timothy Westwood. Tel: +905 828 3894; Fax: +905 828 3792; Email: t.westwood@utoronto.ca
Correspondence may also be addressed to Craig A. Smibert. Tel: +416 946 5538; Fax: +416 978 8548; Email: c.smibert@utoronto.ca
Correspondence may also be addressed to Quaid D. Morris. Tel: +416 978 8568; Fax: +416 978 8287; Email: quaid.morris@utoronto.ca

The authors wish it to be known that, in their opinion, the first three authors should be regarded as Joint First Authors.

© The Author(s) 2013. Published by Oxford University Press.
This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/3.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com
segregate asymmetrically during mammalian neural stem cell divisions and to regulate that lineage (22,23). Staufen1 and 2 have been shown to direct degradation of target RNAs (24,25), and Staufen1 enhances the translation of its targets (26), regulates pre-mRNA splicing (27) and facilitates export of target mRNAs from the nucleus (28). The *Caenorhabditis elegans* Staufen homolog has been implicated in exogenous RNAi (29).

How dsRBPs like Staufen recognize specific mRNA targets *in vivo* is not well understood. *Drosophila* Staufen contains five dsRBDS, three of which (dsRBDD1, dsRBDD3 and dsRBDD4) bind to dsRNA *in vitro* (9), and one of which (dsRBDD3) binds optimally *in vitro* to a stem-loop containing 12 uninterrupted base pairs (bp) when compared against other stem loops (30). However, 12 uninterrupted base pairs are present in only one of the three regions of the bicoid 3' untranslated region (UTR) to which Staufen binds *in vivo* (13,31). In mammals, a 19bp stem is required for Staufen1 binding to *ARF1* mRNA, its best-characterized target (32), but comparable structures have not been detected in other targets of Staufen1 (24,32,33). Intermolecular RNA–RNA interactions may also be important for target recognition by dsRBPs: loop–loop interactions between *bicoid* mRNAs (29) and interactions between long non-coding RNAs and Alu elements in human targets (34) or B elements in rodent targets (35) have been shown to be important for Staufen binding.

To understand how Staufen recognizes its targets *in vivo*, as well as to identify new biological roles for *Drosophila* Staufen, we have performed RNA co-immunoprecipitations (RIPs) followed by microarray analysis (RIP-Chip) to identify Staufen mRNA targets in early *Drosophila* embryos. We used an anti-green fluorescent protein (GFP) antibody to immunoprecipitate transgenic GFP-tagged Staufen (16) as well as a synthetic anti-Staufen antibody (36) to immunoprecipitate endogenous Staufen from wild-type embryos. These experiments identified numerous novel Staufen-associated mRNAs, with a high degree of overlap between the Staufen targets identified by each approach. The functions and localization patterns of these targets support previously known Staufen functions and suggest novel roles for Staufen in early embryos. Using computational methods, we identified secondary structures that are enriched among Staufen targets and are highly specific to Staufen-bound transcripts.

**MATERIALS AND METHODS**

*Drosophila* stocks

*Drosophila* stocks used were as follows: w¹¹¹⁸, GFP-Staufen transgenics (14) (line: GFP311) and staufen mutants with the genotypes w; staufD²³ cn sp/CyO; GFP-Stau2.2FL/TM3 and TM6B, w; stauf⁴⁹⁵/CyO (37), and P[ry11] stauf⁵⁹⁸ cn¹/CyO; ry⁵⁰⁶ (8).

RNA co-immunoprecipitations

For RIPs with synthetic anti-Staufen antibody, synthetic antibodies were expressed and purified as Fabs, and immunoprecipitations were as described (36) with only minor modifications. For anti-GFP-Staufen immunoprecipitations for western blots, RIP-Chip and quantitative PCR (qPCR) validation experiments, protein G magnetic beads (Invitrogen Cat. # 10004D) were first blocked (38), and immunoprecipitations were then performed using a protocol adapted from Invitrogen’s Dynabeads® Protein G protocol and Roche’s immunoprecipitation protocol for anti-GFP (Roche Cat. # 11814460001). The RNA retrieved from these immunoprecipitations was isolated using TRIzol (Invitrogen) following the manufacturer’s protocol and concentrated using RNA clean and concentrator 5 columns (Zymo Research Cat #R1015). For the comparison of synthetic antibody to anti-GFP RIPs (Figure 2C), a slightly different protocol was used, with the anti-GFP RIP protocol modified to be as similar as possible to the synthetic antibody RIP protocol. Details can be found in the Supplementary Materials and Methods.

Microarrays

For microarray analysis, double-stranded cDNA was prepared following the protocol described in the NimbleGen Array User’s Guide (Gene Expression Arrays, version 5.0) with minor modifications. For all samples, 500 ng of double-stranded cDNA was labelled with Cy3- or Cy5-tagged random nonamers following the Roche NimbleGen protocol. Labelled cDNA was then hybridized to custom-designed *Drosophila* 4x72K NimbleGen arrays (GEO platform number: GPL10539). Microarray data were analyzed using the Significance Analysis of Microarrays (39) function available in the MultiExperiment Viewer (MeV) software application (40,41). Details can be found in the Supplementary Materials and Methods, and see Supplementary Figures S1 and S4.

Data access

The data reported in this study have been deposited in NCBI’s Gene Expression Omnibus (42) and are accessible through GEO Series accession number GSE43418 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE43418).

Reverse transcription-quantitative PCR

For reverse transcription-quantitative PCR (RT-qPCR), RNA isolated from immunoprecipitates was reverse transcribed with random hexamer primers and Superscript II reverse transcriptase (Invitrogen). The resulting single-stranded cDNA was subjected to real-time PCR with SYBR green PCR master mix (ABI) using a CFX384 Real-Time System (Bio-Rad). Relative levels of different transcripts were determined using a standard curve.

Gene set annotation enrichment analysis

The Database for Annotation, Visualization and Integrated Discovery (DAVID) functional annotation tool web server (43,44) was used. Terms or features...
enriched at a false discovery rate (FDR) of ≤10% and/or a Benjamini P-value of <0.1 were considered significant. Details can be found in the Supplementary Materials and Methods.

Localization pattern enrichment analysis
The subcellular localization of Staufen-associated transcripts as annotated in the Fly-FISH database (http://fly-fish.cebr.utoronto.ca/) was analyzed to ask whether these transcripts were enriched for particular localization patterns (Fly-FISH annotations up to date as of November 2012 were analyzed). Enrichment was determined using Fisher’s exact test, and P-values were adjusted for multiple comparisons using the Benjamini–Hochberg method to estimate FDRs. Localization terms enriched at an FDR ≤10% were considered significant. Details can be found in the Supplementary Materials and Methods.

Source of transcript sequences for assessment of UTR and open reading frame lengths and motif finding
The Drosophila melanogaster (BDGP5) and Homo sapiens (GRCh37.p6) transcript sequences were downloaded from Ensembl using BioMart (http://www.ensembl.org/biomart/martview/eea40c9db7c1002506d5c766e8772c08) in August 2012. We downloaded all cDNA sequences and defined 3’UTRs as the portion of the cDNA 3’ to the coding sequence, as defined by Ensembl. When there were multiple isoforms for a gene, we used the longest isoform to represent its mature mRNA sequence.

Definitions of secondary structure terms
Throughout this article, in reference to a specific secondary structure: we use the term ‘paired’ to refer to RNA bases that participate in a canonical base pair (i.e. a Watson–Crick base pair or a G-U wobble); we use the term ‘mismatch’ to indicate two bases that are found across from one another in a secondary structure but are not canonical base pairs; we use the term ‘unpaired’ to refer to bases that do not have a corresponding partner base on the other strand of a stem. Mismatches only occur in internal loops and unpaired bases occur in either bulge loops or internal loops, although the latter need not contain any unpaired bases. For example, the right-hand internal loop indicated in the schematic in Figure 5A contains two mismatches and one unpaired base, whereas the bulge in Figure 5A contains three unpaired bases and no mismatches.

Defining N of M motif hits
We estimated the ensemble probability that a region of M bases will contain at least N paired bases using the ensemble probabilities that individual nucleotides within these regions will be paired. First, for the entire 3’UTR, we computed the single nucleotide base-pairing probability using RNAplfold (45), with parameter settings $W = 200$, $L = 150$ and $U = 1$ as recommended (46). When folding the 3’UTR, we also included the 150 nt that comprise the 5’ flanking sequence of the 3’UTR (i.e. the 3’-most 150 nt of the coding region) so that the folding window was not truncated at the 5’-end of the 3’UTR. We then estimated the probability that $N$ bases in an $M$ nucleotide region are paired using the $M-N+1^{\text{th}}$ lowest single-nucleotide probability from the region. Specifically, the pairing probability of the $N$ of $M$ motif was estimated using the lowest single-nucleotide probability in the $M$-mer, after removing the nucleotides with the lowest $M$-th single-nucleotide probability among all the nucleotides except the 5’- and 3’-closing bases. When $M = N$, this estimate is an upper bound on the probability that the entire region is paired; for other values of $N$, it is a convenient estimate. For each $N$ of $M$ pair, which we call a ‘motif’ because it corresponds to a contiguous sequence of bases, we deemed ‘hits’ to be those regions in the top 1% of the $N$ of $M$ probabilities across all 3’UTRs. When scoring other regions of the transcript [5’UTR or open reading frame (ORF)], we used the 1% cut-offs defined on 3’UTRs to select motif hits.

Discovery of N of M motifs that predict Staufen binding
To assess how well the $N$ of $M$ motif hits distinguish the Staufen targets from the co-expressed non-targets, we calculated Wilcoxon rank sum P-values and corresponding area under the receiver operating characteristic (ROC) curves (AUROCs) for all values of $N$ and $M$ where $M$ ranged from 1 to 22 and the allowed number of mismatched or unpaired bases ranged from zero to either 4 or $M$ divided by 4, whichever was less. The Wilcoxon rank sum P-values and the AUROCs were computed based on transcript ‘$N$ of $M$ motif hit scores’, which, for a given $N$ and $M$, were the sum of the probabilities of all $N$ of $M$ motif hits in the transcript’s 3’UTR. These Wilcoxon rank sum P-values and AUROCs were compared with those derived from the 3’UTR length alone—as a baseline. Least absolute shrinkage and selection operator (LASSO) sparse logistic regression was performed to further identify the optimal motifs for Staufen binding. The likelihood ratio test was performed to assess the significance of the improved goodness-of-fit of a regression model containing the selected motifs—over 3’UTR length alone—at classifying transcripts according to whether they were bound by Staufen. In some instances, we also used these methods to assess the goodness-of-fit of regression models based on motif hits in 5’UTRs and coding regions; in those cases, we replaced the 3’UTR-based transcript score with summed probabilities of motif hits in the appropriate region.

Defining [12,10] and [19,15] structures
For each 10 of 12 and 15 of 19 motif hit, we input the region and 150 nt of flanking sequence on either side into Sfold (47). We used Sfold to compute both the centroid structure of the input sequence as well as 1000 samples from the structural ensemble. In each structure, we then identified stems that had either a 10 of 12 or a 15 of 19 motif hit as one side of the stem and deemed them [12,10] and [19,15] structures, respectively. For a motif hit to be deemed a valid structure, it had to satisfy three criteria: (i) at least $N$ of its $M$ bases had to be paired, including the first and last bases; (ii) its ‘partner region’, which is the
transcript sequence between the bases that pair with the first and last bases of the \( N \) of \( M \) motif hit, had to pair only with bases in that hit (i.e. contain no hairpins); (iii) the motif hit had to pair only with bases in its partner region. We found that motif hits that corresponded to valid structures in the centroid also corresponded, in nearly every case, to valid structures in the majority of the ensemble samples. We, therefore, used the centroid to represent the ensemble as this simplified subsequent analysis. We removed any motif hit from consideration that did not correspond to an appropriate valid structure in the centroid. Structures in which both sides of the stem corresponded to motif hits were only represented once in subsequent analyses. These steps are diagrammed in Figure 5B and C.

**Identification of additional features of Staufen-recognized \([12,10]\) and \([19,15]\) structures**

We examined additional features of \([12,10]\) and \([19,15]\) structures to determine whether any of these distinguished the target and non-target sets. They were as follows: (i) number of mismatches; (ii) number of unpaired bases; (iii) number of bulge loops; (iv) number of internal loops; (v) maximum loop size—which is the maximum loop size among all bulge and internal loops in the stem spanned by the \([12,10]\) or \([19,15]\) structure where the size of a loop is the length of the longest side of the loop; and (vi) the distance between the two paired regions which, depending on the relative position of the motif hit and its partner region, is either the distance between the 3' end of the motif hit and the 5' end of its partner region or vice versa. We compared the distribution of these feature values in the target and non-target sets at both the level of individual structures and at the transcript level using cumulative distribution function plots and Wilcoxon rank sum tests. The feature value assigned to each transcript was the minimum value for that feature for all corresponding structures in the transcript.

**Using Staufen-recognized structures to predict Staufen targets and non-targets**

The ‘Results’ section defines Type I, II and III Staufen-recognized structures (SRSs), and a Type II SRS is diagrammed in Figure 5C. To assess the predictive value of SRSs, we ranked transcripts based on the presence of any of the three types of SRSs in their 3'UTRs. The relative enrichment for transcripts containing the three SRSs is described in the ‘Results’ section: it decreases from Type I to Type III. We therefore assigned the highest rank to transcripts containing a Type I SRS, the second highest to those with a Type II but not a Type I SRS, the next highest to those that only contained a Type III SRS, and the lowest rank to those without any SRSs. We then plotted an ROC curve that demonstrates the ability of this ranking to distinguish target and non-target transcripts and computed the AUROC.

**Scoring of the precision of motif mapping**

To compute the precision of the mapping of the SRSs, we computed the proportion of nucleotides in motif hits that are in experimentally defined Staufen-binding regions in the 3'UTR-region of interest: the entire 3'UTR in the case of \( \text{bicoid} \) (13,31) and an experimentally defined 300 nt subset of the \( ARF1 \) 3'UTR (32). ‘Baseline precision’ is the proportion of the 3'UTR (\( \text{bicoid} \)) or 3'UTR subset (\( ARF1 \)) that is in experimentally defined Staufen-binding regions. In the case of \( \text{Drosophila} \) Staufen’s binding region in the \( \text{bicoid} \) 3'UTR, the union of the experimentally defined sites (13) was used in the calculation.

**Defining bound and unbound sets for the human Staufen**

To define the Staufen targets and co-expressed non-targets in human cells, we re-analyzed the published human Staufen RIP-Chip data sets (33). Details can be found in the Supplementary Materials and Methods.

**RESULTS**

**Genome-wide identification of Staufen-associated mRNAs**

To identify mRNAs associated with Staufen in early \( \text{Drosophila} \) embryos, we performed RIP-Chip using two complementary approaches. First, we carried out RIP-Chip of endogenous Staufen from wild-type 0–3h old embryos using a synthetic antibody, anti-Staufen 2A5, that we previously showed immunoprecipitates Staufen protein along with \( \text{bicoid} \) mRNA (36). As a negative control, we performed immunoprecipitations using a control antibody (C1) derived from the same synthetic antibody library as anti-Staufen 2A5 (36). We identified 46 genes whose mRNAs were enriched at least 2-fold in Staufen immunoprecipitates compared with negative control immunoprecipitates and had an FDR of <5% (Figure 1A and B and Table 1; Supplementary Figures S1 and S2; Supplementary Table S1; see ‘Materials and Methods’ section for details). All three previously identified Staufen target mRNAs, \( \text{bicoid} \), \( \text{oskar} \) and \( \text{prospero} \), were among these 46, and \( \text{bicoid} \) mRNA was the most highly enriched target identified. Validation experiments using RT-qPCR are presented in Supplementary Table S2.

To complement the synthetic antibody RIP-Chip, we also performed RIP-Chip using flies expressing GFP-tagged Staufen (16) immunoprecipitated with a commercially available anti-GFP antibody. Western blotting showed that this antibody successfully immunoprecipitated the fusion protein from transgenic embryo extract (Supplementary Figure S3A) and that GFP-Staufen is present at 1.5–2.0-fold higher levels in GFP-Staufen extract than endogenous Staufen in wild-type extract (Supplementary Figure S3B). This anti-GFP RIP-Chip identified 503 genes (of 6151 expressed) whose mRNAs were enriched at least 2-fold in the anti-GFP immunoprecipitates compared with an anti-FLAG control and had an FDR of <5% (Figure 1C and D; Supplementary Figures S4 and S5; Supplementary Table S3). As with the synthetic antibody, all three previously known Staufen mRNA targets were identified, and \( \text{bicoid} \) was again the most highly enriched target. RT-qPCR validation experiments are presented in Supplementary Table S2.
Figure 1. Enrichment of expressed transcripts in Staufen RIPs using synthetic anti-Staufen and anti-GFP-Staufen antibodies. The average, across three biological replicates, of the log2 microarray signal intensities of each expressed transcript in the anti-Staufen and control RIPs were plotted against each other (A, C, E). Highlighted in red and shown at the individual replicate level in the adjacent boxplots (B, D, F) are the transcripts that were found, through Significance Analysis of Microarray two-class analysis, to be significantly enriched in the anti-Staufen versus the control RIPs. Those transcripts with an FDR < 5% and passing a fold enrichment cut-off of at least two in the synthetic anti-Staufen experiments are shown in panels (A) and (B), and those with an FDR < 5% and passing fold enrichment cut-offs of at least two and five in the transgenic anti-GFP experiments are shown in panels (C, D) and panels (E, F), respectively. The three previously identified targets of Staufen (bicoid, oskar and prospero) are labelled in each scatter plot and further denoted by crosses. In (A, C, E), solid diagonal lines represent no enrichment, and dotted diagonal lines represent 2-fold (A, C) or 5-fold (E) enrichment or depletion.
In all, 41 of the 46 genes (89%) identified as Staufen targets using the synthetic antibody were also identified by the anti-GFP RIP-Chip (Fisher’s exact test, $P < 3 \times 10^{-16}$; Figure 2A). Moreover, 36 of the 46 targets identified by the synthetic antibody lay within the 100 most highly enriched targets in the anti-GFP experiment (Figure 2B). Therefore, there is a high degree of similarity between the results of the two experiments.

However, the anti-GFP RIP-Chip identified 10-fold more targets than the synthetic antibody RIP-Chip. This discrepancy is likely attributable to differences both in antibody affinities and in expression levels of endogenous versus transgenic Staufen. For example, RIPs using anti-GFP antibody from GFP-Staufen extract gave a 3-fold higher enrichment of *bicoid* mRNA than synthetic anti-Staufen antibody from the same extract (Figure 2C), most likely as a result of more efficient pull-down of Staufen by the anti-GFP antibody. However, RIPs using the synthetic antibody from GFP-Staufen extract yielded 1.5–2-fold higher enrichment of *bicoid* mRNA than RIPs using the same antibody from wild-type extract (Figure 2C). As GFP-Staufen is present at 1.5–2-fold higher levels in GFP-Staufen extract than endogenous Staufen in wild-type extract (Supplementary Figure S3B), overexpression of GFP-Staufen also appears to contribute to binding to a larger number of mRNAs.

This effect of the overexpression of GFP-Staufen raises the possibility that a subset of the 503 targets identified by the anti-GFP RIP-Chip may not be natural Staufen targets and may only be bound in the context of its overexpression. We therefore created a high-confidence list of 48 targets by applying a more stringent 5-fold enrichment cut-off to the anti-GFP data (Figure 1E and F; Table 1; Supplementary Table S3). These included *bicoid*, *oskar* and *prospero* and shared a total of 25 transcripts with the synthetic-antibody-identified targets (Figure 2A and B), a highly significant overlap (Fisher’s exact test, $P < 3 \times 10^{-16}$).

To avoid excluding low-affinity but real targets of Staufen identified in the anti-GFP RIP-Chip with a 2-fold enrichment cut-off, while also ensuring particular consideration of the high-confidence targets identified using the synthetic antibody and the anti-GFP RIP-Chip with a 5-fold cut-off, our subsequent analyses were conducted on all three lists of Staufen-associated transcripts.

### Functional analysis of Staufen-associated transcripts

To gain insight into potentially novel functions for Staufen in early embryos, we analyzed Staufen-associated mRNAs by gene set annotation enrichment analysis using the DAVID functional annotation tool (43,44). Two stringencies were applied: the standard FDR cut-off ($\leq 0.1$) or the more stringent ‘Benjamini’ $P$-value ($\leq 0.01$) (Table 2 and Supplementary Tables S4–S6). This revealed a number of functions for Staufen in early embryos.

First, our results suggest a more general role than previously suspected for Staufen in controlling the spatial and/or temporal aspects of transcription in early embryos by binding and regulating mRNAs encoding transcriptional regulators. Along with the previously

---

**Table 1.** Staufen-associated mRNAs identified by synthetic anti-Staufen RIP-Chip (fold enrichment $\geq 2$) and anti-GFP-Staufen RIP-Chip (fold enrichment $\geq 5$)

| Targets identified by synthetic anti-Staufen (FDR < 5%, fold enrichment $\geq 2$) and anti-GFP-Staufen RIP-Chip (FDR < 5%, fold enrichment $\geq 5$) | Additional targets identified by synthetic anti-Staufen RIP-Chip (FDR < 5%, fold enrichment $\geq 2$) | Targets identified by anti-GFP-Staufen RIP-Chip only (FDR < 5%, fold enrichment $\geq 5$) |
|---|---|---|
| *bicoid* | CRI4578 | CG5830 |
| *dacapo* | Phosphoenolpyruvate carboxykinase | ocelliless |
| capping protein beta | Mns19 | CG32756 |
| CR18854 | 825-Oak | no on or off transient A |
| CG4068 | CG12519 | lethal (3) neo38 |
| CG32212 | seqoia | Roughened |
| CG17724 | Vacular H$^{+}$-ATPase 26 kD E subunit | nubbin |
| CG3523 | Histone demethylase 4B | Flotillin-2 |
| falafel | Rubral protein S27 | small wing |
| CR32207 | split ends | hves |
| CG32756 | CG4788 | CG32767 |
| Vacular H$^{+}$-ATPase accessory protein AC45 | viritizer | Sprouty-related protein with EVH-1 domain |
| prospero | kagelkern | Tob |
| Peroxisiredoxin 6005 | vielfaltig/Zelda | CG43736 |
| Vacular H$^{+}$-ATPase subunit PPA1-1 | CG9977 | fusilli |
| partner of paired | Neurofibromin 1 | CG33932 |
| oskar | CG14915 | pasilla |
| CG14100 | Mediator complex subunit 14 | punt |
| Protein phosphatase 19C | CG32267 | CG10777 |
| CG18273 | CG17270 | CG5966 |
| staufen | CG14100 | AMP-dependent protein kinase R1 |
| Dorsal switch protein 1 | CR18166 | vein |
| squeeze | | TFN-receptor-associated factor 6 |
| Polycomb | | |
| CG31688 | | |

---

In all, 41 of the 46 genes (89%) identified as Staufen targets using the synthetic antibody were also identified by the anti-GFP RIP-Chip (Fisher’s exact test, $P < 3 \times 10^{-16}$; Figure 2A). Moreover, 36 of the 46 targets identified by the synthetic antibody lay within the 100 most highly enriched targets in the anti-GFP experiment (Figure 2B). Therefore, there is a high degree of similarity between the results of the two experiments.

However, the anti-GFP RIP-Chip identified 10-fold more targets than the synthetic antibody RIP-Chip. This discrepancy is likely attributable to differences both in antibody affinities and in expression levels of endogenous versus transgenic Staufen. For example, RIPs using anti-GFP antibody from GFP-Staufen extract gave a 3-fold higher enrichment of *bicoid* mRNA than synthetic anti-Staufen antibody from the same extract (Figure 2C), most likely as a result of more efficient pull-down of Staufen by the anti-GFP antibody. However, RIPs using the synthetic antibody from GFP-Staufen extract yielded 1.5–2-fold higher enrichment of *bicoid* mRNA than RIPs using the same antibody from wild-type extract (Figure 2C). As GFP-Staufen is present at 1.5–2-fold higher levels in GFP-Staufen extract than endogenous Staufen in wild-type extract (Supplementary Figure S3B), overexpression of GFP-Staufen also appears to contribute to binding to a larger number of mRNAs.

This effect of the overexpression of GFP-Staufen raises the possibility that a subset of the 503 targets identified by the anti-GFP RIP-Chip may not be natural Staufen targets and may only be bound in the context of its overexpression. We therefore created a high-confidence list of 48 targets by applying a more stringent 5-fold enrichment cut-off to the anti-GFP data (Figure 1E and F; Table 1; Supplementary Table S3). These included *bicoid*, *oskar* and *prospero* and shared a total of 25 transcripts with the synthetic-antibody-identified targets (Figure 2A and B), a highly significant overlap (Fisher’s exact test, $P < 3 \times 10^{-16}$).

To avoid excluding low-affinity but real targets of Staufen identified in the anti-GFP RIP-Chip with a 2-fold enrichment cut-off, while also ensuring particular consideration of the high-confidence targets identified using the synthetic antibody and the anti-GFP RIP-Chip with a 5-fold cut-off, our subsequent analyses were conducted on all three lists of Staufen-associated transcripts.

### Functional analysis of Staufen-associated transcripts

To gain insight into potentially novel functions for Staufen in early embryos, we analyzed Staufen-associated mRNAs by gene set annotation enrichment analysis using the DAVID functional annotation tool (43,44). Two stringencies were applied: the standard FDR cut-off ($\leq 0.1$) or the more stringent ‘Benjamini’ $P$-value ($\leq 0.01$) (Table 2 and Supplementary Tables S4–S6). This revealed a number of functions for Staufen in early embryos.

First, our results suggest a more general role than previously suspected for Staufen in controlling the spatial and/or temporal aspects of transcription in early embryos by binding and regulating mRNAs encoding transcriptional regulators. Along with the previously
known targets, bicoïd and prospero, which encode homeodomain-containing transcriptional regulators, the newly identified targets included two additional homeodomain-encoding transcripts (ocelliless and nubbin), Mediator complex subunit 14 of the basal transcription machinery and several known or potential epigenetic regulators (Polycomb, Dorsal switch protein 1 and Histone demethylase 4B). Consistent with this more general role in transcriptional regulation, annotation terms related to ‘transcription regulator’ or the keyword ‘homeobox’ were significantly enriched among targets (Table 2 and Supplementary Tables S4–S6).

Second, all three of Staufen’s previously identified targets control cell fate either via transcriptional (bicoïd, prospero) or post-transcriptional (oskar) regulation. Our results suggest a more general role for Staufen in regulation of cell fate and patterning at these levels as well as at the level of signal transduction and the cell cycle. Staufen’s newly identified targets include mRNAs encoding signaling molecules, cell-cycle regulators, additional transcriptional regulators and additional post-transcriptional regulators with roles in cell fate determination. Accordingly, GO terms related to cell fate were significantly enriched among all three lists of targets (Table 2 and Supplementary Tables S4–S6).

Third, Staufen is expressed and functions both during oogenesis and in the early embryo. Many of its targets are maternal mRNAs that encode proteins that are likely to function at both of these developmental stages. Thus, GO terms related to ‘gamete generation’, ‘oogenesis’ and various aspects of ‘reproductive process’ were significantly enriched among the targets (Table 2 and Supplementary Tables S4–S6). The fact that GO terms related to ‘reproductive process’ are also enriched in the targets recently identified for a Staufen homolog in C. elegans (29), suggests a conserved role during oogenesis and in early embryos.

Fourth, the SwissProt keyword ‘alternative splicing’ and/or the UniProt sequence feature ‘splice variant’, both of which denote genes with alternatively spliced isoforms, were significantly enriched among Staufen targets (Table 2 and Supplementary Tables S4–S6). This suggests a novel role for Drosophila Staufen: binding to transcripts that are alternatively spliced and/or regulation of alternative splicing. The latter function would be consistent with a recently reported role for mammalian Staufen1 in regulation of alternative splicing (27).

Finally, GO terms related to nervous system development (i.e. ‘neuroblast differentiation’, ‘neurogenesis’) and function (‘cognition’) were significantly enriched among Staufen targets (Table 2 and Supplementary Tables S4–S6). Ninety-five percent (42/44) of the genes from the anti-GFP 2-fold list in the categories ‘neuron

Figure 2. Continued
and Supplementary Tables S7–S12). These categories are the stage 4–5 category ‘apical enrichment’ (Figure 3 and Supplementary Tables S7–S12), consistent with the striking enrichment for categories related to posterior localization patterns of Staufen targets showed a marginally longer (Bonferroni-corrected Rank sum P = 0.04), but this was not true for the other two lists.

Staufen targets were also significantly enriched for the stage 1–3 localization category ‘cell division apparatus’ and the stage 4–5 category ‘apical enrichment’ (Figure 3 and Supplementary Tables S7–S12). These categories are intriguing, as they are consistent with the previous observation that on injection into early embryos of in vitro synthesized bicoid 3′UTR RNA, Staufen localizes to mitotic spindle poles during mitosis and to the cortex of the embryo in association with the migrating syncytial nuclei (13).

Finally, analysis of the stage 4–5 patterns of Staufen-associated mRNAs showed highly significant enrichment (FDRs ranging from 0.004 to 8.3% for anti-GFP 2-fold targets) for categories describing zygotic transcription (Figure 3 and Supplementary Tables S7–S12). Enrichment for targets in the process of being transcribed is intriguing in light of the enrichment among Staufen’s targets for alternatively spliced transcripts (discussed earlier in the text) and the reported role of mammalian Staufen1 in regulation of alternative splicing (27).

Drosophila and human Staufen targets have unusually long 3′UTRs

We next assessed features of the transcript sequence that might distinguish Staufen-associated mRNAs from co-expressed non-target mRNAs (hereafter referred to simply as ‘non-targets’). First, we compared 5′UTR, ORF and 3′UTR lengths and found that the median 3′UTR length of targets was 3–4-fold greater than that of the non-targets for all three target lists (synthetic anti-Staufen, anti-GFP-Staufen 2-fold and anti-GFP-Staufen 5-fold) with a Bonferroni-corrected Wilcoxon rank sum of P < 10^{-3} (Figure 4A; Supplementary Table S13). The 3′UTRs of targets on both anti-GFP lists were also significantly longer than non-targets, although to a lesser extent (2–3-fold; Bonferroni-corrected Rank sum P < 10^{-3}), and this was not the case for the synthetic anti-Staufen, anti-GFP-Staufen 2-fold and anti-GFP-Staufen 5-fold list.

Analysis of the embryonic stage 1–3 as well as the stage 4–5 localization patterns of Staufen targets showed a striking enrichment for categories related to posterior localization (FDR < 1% for anti-GFP 2-fold targets; Figure 3 and Supplementary Tables S7–S12), consistent with the fact that Staufen is concentrated in the posterior of the early embryo (8). The ~30 posterior-localized Staufen targets include nanos, oskar, arrest, lost, IGF-II mRNA-binding protein (Imp) and oo18 RNA-binding protein (orb) mRNAs, all of which encode well-known post-transcriptional regulators, and several of which are known to function in the germ plasm.

Staufen targets were also significantly enriched for the stage 1–3 localization category ‘cell division apparatus’ and the stage 4–5 category ‘apical enrichment’ (Figure 3 and Supplementary Tables S7–S12). These categories are intriguing, as they are consistent with the previous observation that on injection into early embryos of in vitro synthesized bicoid 3′UTR RNA, Staufen localizes to mitotic spindle poles during mitosis and to the cortex of the embryo in association with the migrating syncytial nuclei (13).

Finally, analysis of the stage 4–5 patterns of Staufen-associated mRNAs showed highly significant enrichment (FDRs ranging from 0.004 to 8.3% for anti-GFP 2-fold targets) for categories describing zygotic transcription (Figure 3 and Supplementary Tables S7–S12). Enrichment for targets in the process of being transcribed is intriguing in light of the enrichment among Staufen’s targets for alternatively spliced transcripts (discussed earlier in the text) and the reported role of mammalian Staufen1 in regulation of alternative splicing (27).

| Enriched term or feature | Enriched with FDR ≤ 10% among targets identified by \( ^b \) | Enriched with ‘Benjamini’ adjusted \( P ≤ 0.1 \) among targets identified by \( ^b \) |
|-------------------------|-------------------------------------------------|----------------------------------|
|                         | Synthetic anti-Staufen | Anti-GFP 5-fold | Anti-GFP 2-fold | Synthetic anti-Staufen | Anti-GFP 5-fold | Anti-GFP 2-fold |
| Cell fate determination/commitment | \( ^\checkmark \) | \( ^\checkmark \) | \( ^\checkmark \) | | | |
| Alternatively spliced | \( ^\checkmark \) | \( ^\checkmark \) | \( ^\checkmark \) | | | |
| Transcription regulator activity/Homeodomain-containing | \( ^\checkmark \) | \( ^\checkmark \) | \( ^\checkmark \) | | | |
| Reproductive process-related/Oogenesis | | \( ^\checkmark \) | | \( ^\checkmark \) | | |
| Pattern specification process | | \( ^\checkmark \) | | \( ^\checkmark \) | | |
| Cognition | | \( ^\checkmark \) | | \( ^\checkmark \) | | |
| Neuroblast differentiation/Neurogenesis | | | | | | |

\( ^a \)Only terms or features enriched in at least two of the three Staufen target lists (synthetic anti-Staufen, anti-GFP 2-fold, anti-GFP 5-fold) are shown.

\( ^b \)Enrichment analysis was performed using the DAVID functional annotation tool.

Sub-cellular localization of Staufen-associated transcripts

Given Staufen’s well-established and conserved role in mRNA localization, we analyzed the Staufen-associated mRNAs for enrichment of subcellular localization patterns reported by the Fly-FISH database (49), the BDGP in situ database (50,51) and/or the modENCODE temporal expression data (52) available on FlyBase as well as later in the developing or adult nervous system. Given that Staufen is expressed in the developing and adult nervous system, our data suggest that Staufen may regulate these targets at two different stages of development.

Enriched term or feature | Enriched with FDR ≤ 10% among targets identified by \( ^b \) | Enriched with ‘Benjamini’ adjusted \( P ≤ 0.1 \) among targets identified by \( ^b \) |
|-------------------------|-------------------------------------------------|----------------------------------|
|                         | Synthetic anti-Staufen | Anti-GFP 5-fold | Anti-GFP 2-fold | Synthetic anti-Staufen | Anti-GFP 5-fold | Anti-GFP 2-fold |
| Cell fate determination/commitment | \( ^\checkmark \) | \( ^\checkmark \) | \( ^\checkmark \) | | | |
| Alternatively spliced | \( ^\checkmark \) | \( ^\checkmark \) | \( ^\checkmark \) | | | |
| Transcription regulator activity/Homeodomain-containing | \( ^\checkmark \) | \( ^\checkmark \) | \( ^\checkmark \) | | | |
| Reproductive process-related/Oogenesis | | \( ^\checkmark \) | | \( ^\checkmark \) | | |
| Pattern specification process | | \( ^\checkmark \) | | \( ^\checkmark \) | | |
| Cognition | | \( ^\checkmark \) | | \( ^\checkmark \) | | |
| Neuroblast differentiation/Neurogenesis | | | | | | |

\( ^a \)Only terms or features enriched in at least two of the three Staufen target lists (synthetic anti-Staufen, anti-GFP 2-fold, anti-GFP 5-fold) are shown.

\( ^b \)Enrichment analysis was performed using the DAVID functional annotation tool.
have long 3′UTRs. We calculated the ratios of median 3′UTR lengths of single-stranded RBP targets to the median lengths of the 3′UTRs of co-expressed non-targets identified using RIP-Chip for Drosophila PUM (53), human ELAVL1 (54), human PTB (55) and human PUM1 (56). In all cases, the mRNAs in these target sets also had long 3′UTRs; however, the fold increase was less than for Staufen targets: 1.5–2.5-fold rather than 3.0–4.5-fold (Supplementary Table S13 and Supplementary Figure S6).

Thus, although long 3′UTRs are a feature of the targets of both double-stranded and single-stranded RBPs, Staufen targets exhibit particularly long 3′UTRs.

High-confidence Drosophila Staufen target mRNAs are enriched for paired regions of specific lengths in their 3′UTRs

We next asked whether we could identify specific structural motifs in the 3′UTRs of Staufen targets that would distinguish them from non-targets. Specifically, we searched for double-stranded stems of varying length, ranging from 1 to 22 bp, and with varying degrees of imperfect pairing, and asked whether any such structures were enriched among Staufen targets compared with non-targets. Local folding of mRNA is a better predictor of its secondary structure and protein interaction than

Figure 3. Staufen targets are enriched for specific annotated embryonic localization patterns. A search of the Fly-FISH database was performed to identify localization categories that are enriched among Staufen target genes for (A) embryonic stages 1–3 and (B) embryonic stages 4–5. For both (A) and (B), categories in closed boxes were tested for enrichment. The grey shading indicates the localization categories that were enriched among anti-GFP 2-fold targets, FDR < 10%. No categories were enriched among synthetic antibody or anti-GFP 5-fold targets with an FDR < 10%, likely due to the small sizes of these lists. See Supplementary Tables S7–S12 for detailed results.
global folding (46, 57) but local folding tools, such as RNAplfold, only output the probability that a single base is paired. As such, we adopted a hybrid strategy to identify the characteristics of stems bound by Staufen: we first identified regions of the mRNA sequence likely to be paired; then, we assessed Staufen’s preferences for paired regions of various sizes (1–22 nt) and, finally, filtered the preferred regions for those that were actually stems. Figure 5 diagrams the steps in this process.

To identify 3'UTR regions likely to be in stems, for all M-mers (M = 1 to 22 contiguous nucleotides), we estimated the probability that a number, N, of these M contiguous bases was paired using RNAplfold. This is a necessary but not sufficient condition for the M-mer to be in a stem with N of its bases paired. For each M-mer, we set a range for the allowed number of mismatched or unpaired bases from zero to either four or M divided by four, whichever was less (e.g. where M-N = 0 to min(4, M/4)). As an example, based on this definition, an M-mer of 19 with from 15 to 19 paired bases (where the lowest number of paired bases is designated N) is given the designation 15 of 19 (i.e. N of M). We did not allow unpaired bases at the first and last position of the M-mer (see ‘Materials and Methods’ section). We then scanned each 3'UTR for each N of M combination and scored a region as containing a particular N of M ‘hit’ if the probability that the region contained N of M paired bases was found in the top 1% of the N of M probabilities across all 3'UTRs (see Supplementary Table S14 for the cut-offs). We refer to each N of M as a different ‘motif’ because each corresponds to a contiguous sequence of bases. Figure 5B diagrams several motif hits. We assigned each transcript an N of M ‘motif hit score’ equal to the sum of the probabilities of all hits in that transcript’s 3'UTR (see ‘Materials and Methods’ section for details) and asked whether these scores were significantly higher in the Staufen target transcripts versus the non-targets using a one-sided Wilcoxon rank sum test. In addition, to ensure that any significant increase in N of M motif scores of Staufen targets was not solely a result of the differences in 3'UTR length, we set the Rank sum P-value of the 3'UTR length as the baseline. We used AUROC to indicate the effective size of the enrichment, as it has a linear relationship with the Wilcoxon rank sum test statistic; in the vast majority of cases, the feature (i.e. N of M motif score or 3'UTR length) associated with a higher AUROC on a data set also has a more significant P-value. AUROC is also a measure of classification accuracy; in this context, if a feature has a higher AUROC, its scores are better predictors of Staufen binding in these data.

We performed this analysis on each of the three sets of Drosophila Staufen targets described earlier in the text as well as, for comparison, Pumilio targets (53). Analysis of the Staufen targets identified by anti-GFP RIP-Chip with a 2-fold enrichment cut-off and of the Pumilio targets did not reveal any N of M motifs with higher AUROC than 3'UTR length (Supplementary Table S15). In contrast, analyses of the targets identified using either synthetic anti-Staufen or anti-GFP-Staufen with 5-fold cut-off revealed AUROCs above baseline for several N of M motifs (Figure 6; Supplementary Table S15): for the synthetic antibody targets, these included values of M ranging from 4 to 21, with the highest peak at the 10 of 12 motif (Figure 6A, Supplementary Table S15); for the anti-GFP 5-fold targets, these included values of M ranging from 8 to 22, with the highest peak at the 16 of 19 motif (Figure 6C, Supplementary Table S15).

Many of these motifs differed by only one in M or in N, raising the possibility that some of the motifs performed well simply because they were imperfect predictors of the presence of a related motif. Indeed, the values of the transcript scores for similar motifs were extremely highly...
correlated (Supplementary Figure S7). Therefore, to select a core non-redundant set of No fM motifs that collectively explained Staufen’s binding preferences, we performed LASSO logistic regression on the synthetic antibody and anti-GFP 5-fold lists. As potential features to distinguish Staufen targets from non-targets, we used 30 UTR length and all of the No fM motifs identified by us as individually having AUROCs above baseline. This sparse regression analysis assigns weights to each feature, with the most important features being assigned the greatest weights and the least important features receiving weights of zero. For both data sets, LASSO assigned non-zero weights to 10 of 12 (which had the greatest weights: 0.34 and 0.43) and to 15 of 19 (which had a weight of 0.13 for both target sets) (Figure 6B and D). Non-zero weights were also given to 9 of 10, 11 of 11, 14 of 16 and 18 of 20. The logistic regression models containing these collections of motifs had a significantly better fit to the Staufen-binding data from their corresponding sets than ones based on 3’UTR length alone (likelihood ratio test P < 10^{-11} for synthetic antibody and P < 10^{-11} for the anti-GFP, 5-fold; Figure 6 and Supplementary Table S15), indicating that the presence of these motifs explains Staufen’s binding preferences better than the strong bias in 3’UTR length noted earlier.

Having identified this core set of motifs through our analysis of 3’UTRs, we also asked whether there was enrichment for these motifs in the 5’UTRs or ORFs of Staufen-bound mRNAs. To do this, we asked whether the motif scores computed for the 5’UTR or ORF had significantly higher AUROCs than just the length of the corresponding region at the task of classifying transcripts according to Staufen binding, on either the synthetic antibody or anti-GFP 5-fold target sets (Supplementary Table S16). For both 5’UTR and ORF, none of the motifs was a better predictor than length. In addition, notably, the motif hits in the 3’UTR had consistently higher AUROCs than those in either the ORF or 5’UTR (e.g. AUROCs of 0.73–0.90 for those in the 3’UTR versus 0.48–0.63 for 5’UTR and ORF; see Supplementary Table S16).

Having identified several potential motifs that are enriched in the 3’UTRs of Staufen’s targets, we combined the Staufen-associated transcripts from the synthetic antibody and GFP 5-fold data sets to further refine our model of Staufen binding. This created a new set of Staufen target transcripts consisting of the union of the Staufen targets from the two data sets, and a new set of co-expressed non-targets consisting of the intersection of the non-targets from the two data sets. Repeating our regression analysis on these new sets, we found that the fit of the model containing only the 10 of 12 and 15 of 19 motifs (and 3’UTR length) was statistically indistinguishable from one that included all six motifs (and 3’UTR length).
Furthermore, the model containing only the 10 of 12 and 15 of 19 motifs had a significantly better fit than one that only contained one of these two motifs ($P < 0.05$ for both, one-tailed LRT, Bonferroni-corrected). As such, for further analysis, we considered only 10 of 12 and 15 of 19.

**Figure 6.** Specific double-stranded structures are enriched in Staufen target transcript 3’UTRs. (A, C) Wilcoxon rank sum $P$-values were used to assess how well a particular double-stranded stem could distinguish between Staufen targets and co-expressed non-targets that were defined by (A) synthetic antibody RIP-Chip (with 2-fold enrichment cut-off, FDR $\leq 5\%$) or (C) anti-GFP RIP-Chip (with 5-fold enrichment cut-off, FDR $\leq 5\%$). The test was performed on stems of varying length ranging from 1 to 22 base pairs (indicated on the x-axis) and with varying degrees of imperfect pairing ranging from 0 to 4 mismatches (indicated by the different colours and symbols). Among these, only the stems with more significant $P$-value than 3’UTR length (the baseline) are shown and were tested in the analysis shown in (B) and (D). (B, D) Using the stems identified in (A) and (B) as well as the 3’UTR length as the features, LASSO regression was trained to select the features most relevant to Staufen target prediction (i.e. features with non-zero weights). Compared with the training model using 3’UTR length only, the LASSO-selected features significantly improved the prediction of Staufen binding: likelihood ratio test $P < 10^{-7}$ (B) and $P < 10^{-11}$ (D). Exact numbers are given in Supplementary Table S15.

Further refine our analysis, we next sought to identify properties of the dsRNA stems that corresponded to the 10 of 12 and 15 of 19 motif hits. To do so, we used Sfold (47) to predict the secondary structure of a region of ~300 nt centred on each of the 10 of 12 and 15 of 19 motif hits. Sfold outputs two results, both of which we considered in our analysis; a set of 1000 structures sampled from the ensemble of all possible secondary structures, and the ‘centroid’ structure, which is the single structure with the highest total agreement with all 1000 samples. Using the Sfold results, we then identified those centroids where the motif hit satisfied the following three criteria, thus placing it in a dsRNA stem: (i) at least $N$ of its $M$ bases had to be paired, including the first and last bases; (ii) its ‘partner region’, which is the transcript sequence between the

[<P>0.05, one-tailed likelihood ratio test (LRT)]. Furthermore, the model containing only the 10 of 12 and 15 of 19 motifs had a significantly better fit than one that only contained one of these two motifs ($P < 0.05$ for both, one-tailed LRT, Bonferroni-corrected). As such, for further analysis, we considered only 10 of 12 and 15 of 19.

**Computational analysis of the properties of dsRNA stems bound by Staufen**

The aforementioned analysis provides information for only one side of a potential dsRNA structure (i.e. the black strands illustrated in Figure 5B). As such, a single $N$ of $M$ designation describes a large number of potential secondary structures (for some examples, see the combinations of black and grey strands in Figure 5B). Thus, to

---

**Figure 6.** Specific double-stranded structures are enriched in Staufen target transcript 3’UTRs. (A, C) Wilcoxon rank sum $P$-values were used to assess how well a particular double-stranded stem could distinguish between Staufen targets and co-expressed non-targets that were defined by (A) synthetic antibody RIP-Chip (with 2-fold enrichment cut-off, FDR $\leq 5\%$) or (C) anti-GFP RIP-Chip (with 5-fold enrichment cut-off, FDR $\leq 5\%$). The test was performed on stems of varying length ranging from 1 to 22 base pairs (indicated on the x-axis) and with varying degrees of imperfect pairing ranging from 0 to 4 mismatches (indicated by the different colours and symbols). Among these, only the stems with more significant $P$-value than 3’UTR length (the baseline) are shown and were tested in the analysis shown in (B) and (D). (B, D) Using the stems identified in (A) and (B) as well as the 3’UTR length as the features, LASSO regression was trained to select the features most relevant to Staufen target prediction (i.e. features with non-zero weights). Compared with the training model using 3’UTR length only, the LASSO-selected features significantly improved the prediction of Staufen binding: likelihood ratio test $P < 10^{-7}$ (B) and $P < 10^{-11}$ (D). Exact numbers are given in Supplementary Table S15.
basis that pair with the first and last bases of the \( N \) of \( M \) hit, had to pair only with bases in the hit (i.e. contain no hairpins); (iii) the motif hit had to pair only with bases in its partner region (compare Figure 5B and C to see examples of structures that are filtered out, or retained, using these criteria). Based on these analyses, we found that a significantly higher percentage of motif hits were in dsRNA stems in the Staufen targets versus non-targets (Table 3, \( P < 10^{-10} \)). We then removed hits that did not correspond to dsRNA stems and combined hits that were on either side of the same dsRNA stem into a single structure consisting of the motif hit and the sequence with which it pairs. We call these structures [12,10] and [19,15] where the first number indicates that at least one side of the dsRNA stem spans exactly 12 (or 19) nt, and the second number indicates that at least 10 (or 15) bases are paired on that side (e.g. see Figure 5C). The partner region could, in principle, span a different number of nucleotides and have fewer or more unpaired bases.

On further analysis of the [12,10] and the [19,15] structures, we found clear differences between the Staufen targets and non-targets for both structures. Specifically, we assessed the presence of various structural features: (i) unpaired bases, which refer to those bases that do not have a corresponding partner base on the other strand of a stem, (ii) mismatches, which refer to two bases that are found across from one another in a secondary structure but are not canonical base pairs, (iii) internal loops, which are loops emanating from within the dsRNA stem that contain at least one mismatch and, possibly, some unpaired bases, and (iv) bulge loops, which are loops emanating from within one strand of the stem and that contain only one or more unpaired bases. These are diagrammed in Figure 5A.

Comparing Staufen targets with non-targets, our analysis revealed that the [12,10] and [19,15] structures in the targets had significantly fewer unpaired bases (Supplementary Figure S8) as well as a weak preference against mismatches. The bias against unpaired bases also manifested as a significant depletion for bulges in the target set (Supplementary Figure S8). In addition, there was a significant preference for smaller internal loops or bulges in the Staufen targets than the non-targets (Supplementary Figure S9). Taken together, these results suggest an overall model in which Staufen binds [12,10] and [19,15] dsRNA stems that have a small number of mismatches, zero or few unpaired bases and short internal loops.

Many transcripts in both the target and non-target sets contain multiple [12,10] and [19,15] structures; as such, we next considered the same criteria at the transcript level to determine features associated with the ‘best’ Staufen site in each transcript. We did this by assigning each transcript a mismatch, unpaired base, and loop count as well as loop length equal to the minimum of those values across all [12,10] and [19,15] structures in that transcript’s 3‘UTR. This analysis also revealed a significantly lower number of unpaired bases in the ‘best’ structure in Staufen target transcripts compared with non-targets, which also manifested as an even stronger preference for balanced stems (i.e. those containing only paired or mismatched bases and no unpaired bases) (Figure 7). In particular, 82% of Staufen targets that had at least one [12,10] structure contained a balanced [12,10] structure (compared with 47% in non-targets). We use the designation [12,10,0] to describe these balanced structures where the last number refers to the maximum number of unpaired bases in the structure. The enrichment among the Staufen targets for transcripts containing balanced [19,15] structures (i.e. [19,15,0]) was even more striking (67 versus 12%). Because many more Staufen target transcripts have [19,15] structures than do non-targets, if we define a Staufen-bound transcript as one containing a [19,15] that is balanced (i.e. [19,15,0]), this rule would be satisfied by 44% (n = 24) of the target transcripts but only 3% of the non-target transcripts (n = 42).

We note that Staufen sites defined by previous structural, in vitro and/or in vivo studies are not nearly as predictive of its binding as the [12,10] and [19,15] structures: using the criteria described earlier in the text, only 31% of all targets and 4% of the non-targets contain a perfect 12 bp region of dsRNA (i.e. [12,12,0]), which was defined as an optimal structure for Staufen dsRBD3 binding (30), and none of the target transcripts and only two of the non-target transcripts (0.1%) contain a perfect

| Motif hit | Staufen targets | Co-expressed non-targets | Hypergeometric \( P \)-value |
|-----------|-----------------|--------------------------|-----------------------------|
| 10 of 12  |                 |                          |                             |
| No. of motif hits in dsRNA stem | 672 | 3272 | 1.83 \times 10^{-10} |
| No. of motif hits | 954 | 5469 |                         |
| Percentage motif hits in dsRNA stem | 70.44 | 59.83 | 4.41 \times 10^{-11} |
| No. of genes with at least one hit in dsRNA stem | 45 | 572 |                         |
| Percentage genes with at least one hit in dsRNA stem | 81.82 | 37.43 |                         |
| 15 of 19  |                 |                          |                             |
| No. of motif hits in dsRNA stem | 487 | 1626 | 1.71 \times 10^{-26} |
| No. of motif hits | 954 | 5469 |                         |
| Percentage motif hits in dsRNA stem | 51.05 | 29.73 | 3.54 \times 10^{-11} |
| No. of genes with at least one hit in dsRNA stem | 36 | 344 |                         |
| Percentage genes with at least one hit in dsRNA stem | 55 | 1528 |                         |
| No. of genes with at least one hit | 6.545 | 22.51 |                         |
dsRNA stems bound by Staufen, we next mapped these structures onto: (i) the 3′UTRs of the Staufen target transcripts; (ii) length-matched non-target transcript 3′UTRs; and (iii) a random subset of non-target 3′UTRs (Supplementary Figure S10). This led us to note a substantial overlap of [19,15,4] and [12,10,2] structures in the Staufen target set. As such, we next assessed whether the [12,10,2] structures are independently associated with Staufen binding or if they are simply features of more predictive [19,15,4] structures. To do so, we first assessed whether the presence of [12,10,2] in the absence of [19,15,4] was predictive of Staufen binding by comparing Staufen target and non-target transcripts that did not contain a [19,15,4] structure. We found that 47% (9 of 19) of such transcripts in the Staufen target set have a [12,10,2] structure, whereas only 22% (279 of 1245) of the non-targets have one (P = 0.02, Hypergeometric test). We then asked whether Staufen target transcripts were more likely to have a [19,15,4] structure that contained a [12,10,2] structure than non-targets. Here, we found a highly significant difference: 94% (34 of 36) of Staufen target transcripts with a [19,15,4] structure had one containing a [12,10,2] structure but only 61% (174 of 283) of such non-target transcripts did (P < 2 × 10⁻⁹, Hypergeometric test). We also noted a similar, but not statistically significant, preference for [19,15,0] structures containing a [12,10,0] structure in the Staufen targets: 91.7% (22 of 24) versus 69% (29 of 42). These results indicate that [12,10,0] structures are not only independent predictors of Staufen binding—although the statistical significance of this association is weak—but 19bp region of dsRNA (i.e. [19,19,0]) defined as an optimal site for human Staufen1 binding to ARF1’s 3′UTR (32).

That said, the [19,15,0] rule does not explain all instances of Staufen binding; in particular, the bicoid 3′UTR does not contain one of these high-confidence sites. However, we can slightly refine our model of the [12,10] and [19,15] structures to [12,10,2] and [19,15,4], respectively (Figure 5). A model including these latter structures is better at distinguishing Staufen targets from non-targets than the [12,10] or [19,15] structures. Although this model removes individual sites from the Staufen target set, all Staufen targets with a [12,10] or a [19,15] structure contain at least one [12,10,2] or [19,15,4] structure, respectively. On the other hand, this rule reduces the number of non-target transcripts with these structures by removing those that have a large number of unpaired bases in the partner region (e.g. one [19,15] structure in the non-target set had >100 unpaired bases). Note that all [19,15,0] structures are also [19,15,4] structures because the latter includes structures with three, two, one and zero unpaired bases. In total, 65% of Staufen targets and 18% of non-targets contain a [19,15,4] structure, whereas 82% of Staufen targets and 33% of non-targets contain a [12,10,2] structure.

**Definition of Staufen-recognized structures (SRSs)**

Having defined the structural characteristics of the dsRNA stems bound by Staufen, we next mapped these

---

**Figure 7.** Characteristics of the stems bound by Staufen. We compared the 10 of 12 (A) and 15 of 19 (B) structures in the Staufen targets versus non-targets, by characterizing the structural features: (i) number of mismatches; (ii) number of unpaired bases; (iii) number of bulges; (iv) number of internal loops. If there was more than one structure in a transcript’s 3′UTR, the feature with the minimal score was used to represent the gene. The Wilcoxon rank sum test was used to assess the feature in Staufen targets (the solid line) versus non-targets (the dashed line). pos: targets; neg: non-targets.
are also more likely to be found in Staufen targets as features of \{19,15\} structures.

Based on these results, we define three classes of SRSs: Type I is a \{19,15,0\} containing a \{12,10,0\}, which we abbreviate as \{19,15,0\}&\{12,10,0\}; Type II is a \{19,15,4\} containing a \{12,10,2\}, abbreviated \{19,15,4\}&\{12,10,2\}; and Type III is a \{12,10,2\}. An example of a Type II SRS is shown in Figure 5D.

Figure 8 individually compares the true- and false-positive rates for each of the aforementioned structures. If we classify transcripts based on which type of SRS they contain, we achieve an AUROC of 81.6% by training the whole data set (i.e. as described earlier in the text, the union of the Staufen targets from the synthetic and GFP 5-fold data sets, and the set of co-expressed non-targets consisting of the intersection of the non-targets from the two data sets), which is comparable with that achieved on held-out data by a logistic regression classifier that uses presence or absence of all the structures mentioned in the previous paragraph plus 3'UTR length as features (data not shown). Because we used the training set to define these structures and the SRSs, the AUROC is likely to over-estimate the classification performance we should expect on new data; however, we suspect that this over-estimate is small because (i) we used a simple classification model; (ii) the enrichment P-values we used to define these features were all highly significant; and (iii) we only made a small number of feature selection choices. We also note that \{12,10,0\} lies directly on the curve; therefore, it has no added classification power in our training set above the three types of SRSs, and, as such, we do not identify it as an additional SRS. \{19,15,0\} lies slightly to the left of the ROC curve formed by the three SRSs; therefore, it does have added classification power; however, the increase is small, and, as such, we omit it as a separate feature to ensure that the SRSs are nested, thereby forming the basis of our transcript scoring scheme (see ‘Materials and Methods’ section).

The crystal structures of dsRBDs from Drosophila Staufen or yeast Rnt1p endonuclease bound to dsRNA show interactions between the RBD and the hairpin loop (30,58). We, therefore, next asked whether there was any preference for the location of the three types of SRSs relative to hairpin loops. We scored transcripts based on the linear distance on the mRNA transcript between the two paired regions in the SRS and found that this minimum distance was significantly larger in target transcripts for Type I SRSs but shorter for Type II and Type III SRSs (Supplementary Figure S11). These results were, therefore, inconclusive with respect to the preferred location of SRSs relative to hairpin loops.

Mapping of SRSs in Drosophila and human mRNAs

We next mapped the locations of the three types of SRSs within the 3'UTRs of all Drosophila mRNAs (Supplementary Table S17). Figure 9 shows the locations of SRSs in the 3'UTRs of the Staufen target transcripts, length-matched non-target transcript 3'UTRs and a random subset of non-target 3'UTRs.

We then asked whether the identified Drosophila SRSs predict human Staufen targets better than 3'UTR length alone and found that none had better AUROC than 3'UTR length for either Staufen1 or Staufen2 (Supplementary Table S18). These results, as well as the failure to predict a motif for Drosophila Staufen from the GFP 2-fold data set, may be a consequence of these lists containing a significant fraction of weakly associated targets or co-expressed transcripts that are not normally bound by Staufen but are scored as bound in these data sets due to overexpression of the tagged Staufen protein used in the RIP-Chip experiments. As such, those transcripts may not be enriched for SRSs (see ‘Discussion’ section).

Finally, to assess whether SRSs map to the experimentally determined in vivo Staufen-binding sites, we focused on Drosophila bicoid (13,31) and human ARF1 (32). In the bicoid 3'UTR, the SRSs mapped almost exclusively to the three experimentally determined Staufen-binding regions (Figure 10A): for Type II SRSs, the precision was 1.0, and for Type III SRSs, the precision was 0.94 (baseline precision = 0.47; there were no Type I SRSs). For human ARF1's 3'UTR, again, the SRSs mapped almost exclusively to the known Staufen1-binding regions (Figure 10B): for all three types of SRSs, the precision was 1.0 (baseline precision = 0.13). Taken together, these results provide strong evidence that the
Figure 9. Mapping of SRSs in the 3'UTRs of Staufen targets and non-targets. The three types of SRSs were mapped in the 3'UTRs of Staufen targets (A), length-matched non-targets (B) and a random subset of non-targets (C). The x-axis represents the 3'UTR in nucleotides, starting from the first nucleotide after the stop codon. Each 3' UTR is represented by a grey bar, within which the predicted SRS hits are represented by coloured bars (Type I: dark blue; Type II: light blue; Type III: red; Type III embedded in or overlapping with Type I: black; Type III embedded in or overlapping with Type II: magenta). For each SRS, the 5'-most nucleotide in the corresponding 15 of 19 or 10 of 12 motif is connected to the paired nucleotide in the partner arm by a line of the same colour as the SRS (Type I: dark blue, Type II: light blue, Type III: red).
computationally identified structures correspond to bona fide, in vivo Staufen-binding sites.

DISCUSSION

Here, we have performed RIP-Chip to identify mRNA targets of Staufen in vivo in early Drosophila embryos, using a synthetic antibody to immunoprecipitate endogenous Staufen and an anti-GFP antibody to immunoprecipitate transgenic GFP-Staufen. In silico analyses of the functions and subcellular localization of these bound transcripts suggested novel roles for Staufen in early embryos. In addition, computational analyses identified dsRNA structures that are highly specific to Staufen’s in vivo targets.

Figure 9. Continued.
Localization and functions of Staufen-associated transcripts

The localization patterns of the Staufen-associated transcripts that we have identified here are largely consistent with known localization of Staufen protein in early Drosophila embryos. We found strong enrichment for posteriorly localized mRNAs among Staufen’s targets, consistent with the fact that Staufen is concentrated at the posterior of early embryos (8). The 30 newly identified posterior-localized Staufen targets include mRNAs that encode additional post-transcriptional regulators, several of which are known to function in the germ plasm (nanos, arrest, lost, Imp and orb). Thus, Staufen may play a more general role in localization and regulation of germ-plasm transcripts than previously suspected.

Our gene set annotation enrichment analyses suggest that Staufen also plays a more general role in cell fate specification than previously thought, via regulation of mRNAs encoding intra- and inter-cellular signalling molecules, as well as transcriptional and post-transcriptional regulators. This more general role may not be limited to early embryos but might extend into the developing nervous system as well, as Staufen targets were enriched for transcripts that confer neuroblast and neuronal fates.
Although *Drosophila* Staufen was previously known to localize *prospero* transcripts in the neuroblast lineage (14–17), a more general role for *Drosophila* Staufen in neuronal development and function is consistent with the previously reported role of mammalian Staufens in these processes (18–23).

Our gene set annotation enrichment analyses also suggest that Staufen may serve a more general role in regulating transcription in early embryos via the basal transcriptional machinery, additional homeodomain-encoding transcripts and several known epigenetic regulators. One of the new targets encodes Zelda (also known as Vifaltig), an important regulator of zygotic genome activation during the maternal-to-zygotic transition (59–62). Thus, Staufen may regulate both spatial and temporal aspects of zygotic transcription.

Finally, our analyses suggest a potential role for *Drosophila* Staufen in regulation of alternative splicing and/or a preference for binding to alternatively spliced transcripts. Although *Drosophila* Staufen has not been observed to localize to nuclei, it contains at least one putative nuclear localization signal as predicted by NLStradamus (63) and NucPred (64) web tools. Furthermore, mammalian Staufen1 contains a nuclear localization signal, has been observed in nuclei (27,28,65) and has been shown to function in splicing regulation (26). Posterior localization of one of *Drosophila* Staufen’s targets, *oskar*, requires splicing of its first intron and deposition of the exon junction complex (9,66–71). This raises the possibility that Staufen’s role in posterior localization of its targets could, in part, be related to a function in splicing regulation.

**The role of 3’UTR length in Staufen’s targets**

We have shown that 3’UTR length is a major feature that distinguishes both the *Drosophila* and human Staufen targets from non-targets. Our analyses also indicated that the targets of single-stranded RBPs had long 3’UTRs relative to co-expressed but unbound transcripts. However, the fold-increase was less than for fly and human Staufen targets: 1.5–2.5-fold rather 3.0–4.5-fold. Thus, although Staufen targets have particularly long 3’UTRs because they undergo a great deal of such regulation.

An alternative explanation is that the 3’UTR length derives from a particular mechanism that is specific to this set of targets. For example, studies of human Staufen suggest that Staufen-mediated decay (SMD) is dependent on the presence of an upstream termination codon, and therefore Staufen binding sites that direct SMD must be located in the 3’UTR of targets (23). In addition, in yeast, *C. elegans*, *Drosophila* and mammals, transcripts with longer 3’UTRs show increased susceptibility to nonsense-mediated decay, which shares many mechanistic similarities with SMD (74–79). It is therefore possible that the longer 3’UTRs of Staufen targets may be important for SMD. There is not, however, currently any evidence for SMD in *Drosophila* embryos. If SMD were a major role of *Drosophila* Staufen, then a large fraction of its targets should be degraded and, although 19 to 25% of the Staufen targets we have identified are degraded in early embryos, this number is similar to the overall percentage of maternal mRNAs that is cleared (72,78).

**Staufen-recognized secondary structures**

Previous *in vitro* studies on the binding of dsRBD3 from *Drosophila* Staufen to artificial RNA substrates (30) and *in vivo* studies on the binding of mammalian Staufen1 to ARFI mRNA (32) have identified 12 and 19 bp Watson–Crick (or G-U) paired stems, respectively, as binding sites for Staufen. In addition, studies of two other dsRBPs have shown that 16 bp stems can also act as binding sites for RBPs containing two or more dsRBDS in *in vitro* (6,81). Our computational analysis of *Drosophila* Staufen’s *in vivo* targets has revealed enrichment for three types of SRSs in their 3’UTRs: Type I [19,15,0]&[12,10,0], Type II [19,15,4]&[12,10,2] and Type III [12,10,2]. Together, the Type I, II and III SRSs map to the previously identified Staufen-binding regions in *Drosophila bicoid* and human ARFI mRNAs with a precision of ≥0.94.

To our knowledge, this is the first report that uses genome-wide *in vivo* binding data to define structural preferences—and thus specificity—for a double-stranded RBP. Although *in vitro* studies have suggested that mismatches or unpaired bases in dsRNA stems reduce the degree of dsRBD-RNA binding (6,30,82), all three SRSs predicted from our *in vivo* data allow one or more mismatches, and Type II and III SRSs also allow unpaired bases. Consistent with this finding, *Drosophila*, the experimentally identified regions within the 3’UTR of *bicoid* RNA that are required for Staufen binding contain imperfect stems of the length and mismatch number predicted by our computational analysis (13). Moreover, in other cases, where the binding of dsRBDS to endogenous substrates has been studied, bulges or mismatches are often present. For example, mammalian ADAR2 binds a structure in GluR-2 mRNA referred to as the R/G stem-loop, which contains mismatches required for binding (83,84). In addition, mammalian PKR binds to a variety of cellular and viral RNAs that contain mismatches (28,85–87). Thus, a perfect Watson–Crick/G-U dsRNA helix is not a prerequisite for dsRBD binding; indeed, we argue that imperfect stems provide the major specificity of Staufen’s binding to its targets. Experimental analyses will be required to test this proposal. We note that our definition of SRSs permits non-canonical base pairs (or the rotation of unpaired bases) that lead to stable A-form helical stems in the absence of perfect canonical base pairing. Indeed, the strong
preferençe for mismatches over unpaired bases supports this possibility.

How does Staufen recognize and bind stems of different lengths?

In addition to being imperfect stems, the SRSs we have identified are comprised of two major substructures of length 19 and 12, with the latter often contained within the former. One possibility to explain how Staufen binds structures with different lengths is that the shorter stems mediate the binding of a single dsRBD and the longer stems mediate the binding of two or more dsRBDS, either housed in the same dsRBP or in two simultaneously bound dsRBPs molecules. Indeed, it has recently been shown that mammalian Staufen 1 and 2 can form both homo- and heterodimers in vitro as well as in tissue-culture cells (25,88). Consistent with the possibility that the longer stems mediate the binding of two or more dsRBDS, three of Staufen’s dsRBDS bind RNA in vitro (9) and structural studies of Xenopus Xlrpa and mammalian ADAR2 have shown that the binding of a single dsRBD involves interactions with only one face of a dsRNA helix, thereby allowing a second dsRBD to bind a different face of the same helix (81,83,84).

A second, not necessarily mutually exclusive, possibility is that the short Type III SRS represents the binding site for a dsRBD only in the context of a stem-loop structure, whereas the longer SRSs represent binding to stems with no adjacent hairpin loop. With respect to the former, in vitro studies of the binding of dsRBD3 of Drosophila Staufen to RNA stem loops have shown that a stem length of 12 bp with a loop is optimal for binding, and that, in this context, the dsRBD makes contacts with the loop as well as the stem (30). Similarly, the structure of the dsRBD from the yeast Rnt1p endonuclease bound to a small nucleolar RNA substrate shows binding to a 13 bp stem capped by a tetraloop, with the protein again making contacts with both stem and loop (58). With respect to the longer SRSs, the mammalian Staufen1-binding site within ARF1 mRNA appears to be a 19 bp stem with no adjacent hairpin loop (i.e. the ‘loop’ enclosed by the stem contains multiple hairpin loop structures), and mutations that shorten this stem reduce Staufen binding (32). That different dsRBDS in the same protein may have different binding preferences is supported by data from mammalian ADAR2: one of its dsRBDS prefers hairpin loops, whereas the other prefers duplexes that contain mismatches within internal loops (83).

Staufen levels as a determinant of target mRNA selection

Our data suggest that modest changes in the levels of Staufen might have a significant effect on its compendium of bound mRNAs. Thus, mechanisms that regulate the levels of Staufen could have biologically relevant effects on the processes that Staufen regulates. This has important implications for RIP experiments that use over-expression of tagged Staufen proteins, and likely other dsRBPs, in cell lines or in vivo, as they may lead to identification of spurious target mRNAs. It may be argued that Staufen and other dsRBPs are particularly prone to artefactual target mRNA identification because of their propensity to bind a range of double-stranded motifs rather than to specific sequences. However, given the low complexity and the redundancy within the recognition sites of many sequence-specific RBPs, it is possible that target site selection for many RBPs is strongly influenced by RBP levels.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS

The authors thank Daniel St Johnston (Cambridge, UK) for providing the GFP-Staufen Drosophila stock and the rabbit polyclonal anti-Staufen antibody, Luc DesGroseillers (Montreal, Canada) for providing the raw data from the human Staufen1 and 2 microarray experiments and Anna Soltyk (Canadian Drosophila Microarray Centre, University of Toronto, Mississauga) for help with the microarray experiments. Extensive use was made during this study of the following resources: Fly-FISH, FlyBase, the Bloomington Drosophila Stock Center, and the Berkeley Drosophila Genome Project. H.D.L. and C.A.S. conceived of the project and designed its overall goals; synthetic antibody RIP-Chip was carried out by J.D.L.; anti-GFP RIP-Chip and western blots by K.A.; RT-qPCR validation experiments by J.D.L. and K.A.; GO term analysis by J.D.L. and K.A.; subcellular localization analysis by J.D.L.; 3’UTR length analysis, motif finding and structural analysis by X.L. under the guidance of Q.D.M.; J.D.L., K.A. and X.L. wrote the first draft of the manuscript, which was revised by H.D.L. and Q.D.M. with input from the other authors. J.D.L. is co-supervised by H.D.L. and C.A.S.; K.A. by J.T.W. and H.D.L.; X.L. by Q.D.M. and H.D.L.

FUNDING

Canadian Institutes of Health Research [MOP-14409 to H.D.L. and MOP-125894 to Q.D.M.]; University of Toronto Open Scholarships (to J.D.L., K.A. and X.L.). Funding for open access charge: Canadian Institutes of Health Research.

Conflict of interest statement. None declared.

REFERENCES

1. Lunde,B.M., Moore,C. and Varani,G. (2007) RNA-binding proteins: modular design for efficient function. Nat. Rev. Mol. Cell. Biol., 8, 479-490.
2. Bycroft,M., Gruenert,S., Murzin,A.G., Proctor,M. and St Johnston,D. (1995) NMR solution structure of a dsRNA binding domain from Drosophila staufen protein reveals homology to the N-terminal domain of ribosomal protein S5. EMBO J., 14, 3563–3571.
3. Kharrat,A., Macias,M.J., Gibson,T.J., Nilges,M. and Pastore,A. (1995) Structure of the dsRNA binding domain of E. coli RNase III. EMBO J., 14, 3572–3584.
4. Nanduri,S., Carpick,B.W., Yang,Y., Williams,B.R. and Qin,J. (1998) Structure of the double-stranded RNA-binding domain of the protein kinase PKR reveals the molecular basis of its dsRNA-mediated activation. *EMBO J.*, 17, 5458–5465.

5. St Johnston,D., Brown,N.H., Gall,J.G. and Jantsch,M. (1992) A conserved double-stranded RNA-binding domain. *Proc. Natl Acad. Sci. USA.*, 89, 10979–10983.

6. Bevilacqua,P.C. and Coulth,T.R. (1996) Minor-groove recognition of double-stranded RNA by the double-stranded RNA-binding domain from the RNA-activated protein kinase PKR. *Biochemistry*, 35, 9983–9994.

7. Schupbach,T. and Wieschaus,E. (1986) Germline autonomy of maternal-effect mutations altering the embryonic body pattern of *Drosophila*. *Dev. Biol.*, 113, 443–448.

8. St Johnston,D., Bruchle,D. and Nusslein-Volhard,C. (1991) Staufen, a gene required to localize maternal RNAs in the *Drosophila* egg. *Cell*, 66, 51–63.

9. Mickle,J., Dufau,M.L., Zatloukal,K. and Koopman,P. (1991) Distinct roles of two conserved Staufen domains in oskar mRNA localization and translation. *EMBO J.*, 19, 1366–1377.

10. Schuld,J.L., Smith,J.L. and Macdonald,P.M. (1991) Oskar mRNA is localized to the posterior pole of the *Drosophila* oocyte. *Cell*, 66, 23–35.

11. Ephrussi,A., Dickinson,L.K. and Lehmann,R. (1991) Oskar organizes the germ plasm and directs localization of the posterior determinant nanos. *Cell*, 67, 37–50.

12. St Johnston,D., Driever,W., Berletch,B., Richstein,S. and Nusslein-Volhard,C. (1990) Multiple steps in the localization of bicoid RNA to the anterior pole of the *Drosophila* oocyte. *Development*, 107, 13–19.

13. Ferrandon,D., Elphick,L., Nusslein-Volhard,C. and St Johnston,D. (1994) Staufen protein associates with the 3'UTR of bicoid mRNA to form particles that move in a microtubule-dependent manner. *Cell*, 79, 1221–1232.

14. Broadus,J. and Doe,C.Q. (1997) Extrinsic cues, intrinsic cues and microfilaments regulate asymmetric protein localization in *Drosophila* neuroblasts. *Curr. Biol.*, 7, 827–835.

15. Broadus,J., Fuerstenberg,S. and Doe,C.Q. (1998) Staufen-dependent localization of prospero mRNA contributes to neuroblast daughter-cell fate. *Nature*, 391, 792–795.

16. Schuldi,A., Adams,J.H., Davidson,C.M., Micklem,D.R., Wier,J., Kiebler,M. and Jantsch,M. (1992) Distinct roles of two conserved Staufen domains in oskar mRNA localization and translation. *Proc. Natl Acad. Sci. USA.*, 89, 10979–10983.

17. Li,P., Yang,X., Wasser,M., Cai,Y. and Chia,W. (1997) Insusceptible and Staufen mediate asymmetric localization and segregation of prospero RNA during *Drosophila* neuroblast cell divisions. *Cell*, 90, 437–447.

18. Kiebler,M.A., Hemraj,J., Verkade,P., Kohrmann,M., Fortes,P., Marion,R.M., Oriti,J. and Dotti,C.G. (1999) The mammalian staufen protein localizes to the somatodendritic domain of cultured hippocampal neurons: implications for its involvement in mRNA transport. J. Neurosci., 19, 288–297.

19. Vessey,J.P., Macchi,P., Stein,J.M., Mikl,M., Hawker,K.N., Vogelsang,P., Wieczorek,K., Vendra,G., Riefler,J. and Will,J.B. (2008) A loss of function allele for murine Staufen1 leads to impairment of dendritic Staufen1-RNP delivery and dendritic spine morphology. *Proc. Natl Acad. Sci. USA.*, 105, 16374–16379.

20. Goetz,B., Tung,Y., Xie,Y., Dorostkar,M., Thomas,S., Puhl,U., Boecker,S. and Kiebler,M.A. (2006) The brain-specific double-stranded RNA-binding protein Staufen2 is required for dendritic spine morphogenesis. J. Cell Biol., 172, 221–231.

21. Duchaine,T.F., Hemraj,J., Furic,L., Deitinghoff,A., Kiebler,M.A. and DesGrosseillers,L. (2002) Staufen2 isoforms localize to the somatodendritic domain of mammalian neurons and interact with different organelles. J. Cell. Sci., 115, 3285–3295.

22. Kusek,G., Campell,B., Doyle,F., Tenenbaum,S.A., Kiebler,M. and Temple,S. (2012) Asymmetric segregation of the double-stranded RNA-binding protein Staufen2 during mammalian neural stem cell divisions promotes lineage progression. *Cell Stem Cell*, 11, 505–516.

23. Vessey,J.P., Macchi,P., Burns,S.E., Kiebler,M.A., Kaplan,D.R. and Miller,F.D. (2012) An asymmetrically localized Staufen2-dependent RNA complex regulates maintenance of mammalian neural stem cells. *Cell Stem Cell*, 11, 517–528.

24. Kim,Y.K., Furic,L., DesGrosseillers,L. and Maquat,L.E. (2005) Mammalian Staufen1 recruits Upf1 to specific mRNA 3'UTRs so as to elicit mRNA decay. *Cell*, 120, 195–208.

25. Park,E., Gleghorn,M.L. and Maquat,L.E. (2013) Staufen2 functions in Staufen1-mediated mRNA decay by binding to itself and its partner Upf1 helicase, but not ATPase activity. *Proc. Natl Acad. Sci. USA.*, 110, 405–412.

26. Dugrè-Brisson,S., Elvira,G., Boulay,K., Chatel-Chaix,L., Mouland,A.J. and DesGrosseillers,L. (2005) The RNA-binding protein Staufen1 is increased in DM1 skeletal muscle and promotes alternative pre-mRNA splicing. *J. Cell. Biol.*, 169, 699–712.

27. Elbarbary,R.A., Li,W., Tian,B. and Maquat,L.E. (2013) STAUF binding 3' UTR IRAKus complements nuclear retention to protect cells from PKR-mediated translational shutdown. *Genes Dev.*, 27, 1495–1510.

28. Legendre,J.B., Campbell,Z.T., Kroll-Conner,P., Anderson,P., Kimble,J. and Watkins,M. (2012) RNA targets and specificity of Staufen, a double-stranded RNA-binding protein in *C. elegans*. *J. Biol. Chem.*, 288, 2532–2545.

29. Ramos,A., Grunert,S., Adams,J., Mickle,D.R., Proctor,M.R., Freud,S., Bycroft,M., St Johnston,D. and Varani,G. (2000) RNA recognition by a Staufen double-stranded RNA-binding domain. *EMBO J.*, 19, 997–1009.

30. Ferrandon,D., Koh,I., Westhof,E. and Nusslein-Volhard,C. (1997) RNA-RNA interaction is required for the formation of specific bicoid mRNA 3' UTR-STAUFEN ribonucleoprotein particles. *EMBO J.*, 16, 1751–1758.

31. Kim,Y.K., Furic,L., Parisien,M., Major,F., DesGrosseillers,L. and Maquat,L.E. (2007) Staufen1 regulates diverse classes of mammalian transcripts. *EMBO J.*, 26, 2670–2681.

32. Furic,L., Maher-Laporte,M. and DesGrosseillers,L. (2008) A genome-wide approach identifies distinct but overlapping subsets of cellular mRNAs associated with Staufen1- and Staufen2-containing ribonucleoprotein complexes. *RNA*, 14, 324–335.

33. Gong,C. and Maquat,L.E. (2011) RNA-RNAs transactivate STAUF1-mediated mRNA decay by duplexing with 3' UTRs via AUp elements. *Nature*, 470, 284–288.

34. Wang,J., Gong,C. and Maquat,L.E. (2013) Control of myogenesis by rodent sRISC-containing IncRNAs. *Genes Dev.*, 27, 793–804.

35. Javer,J.D., Anciuciu,C., Kollazzo,P., Westwood,J.T., Sidhu,S.S., Liphschitz,H.D. and Smibert,C.A. (2012) Synthetic antibodies as tools to probe RNA-binding protein function. *Mol. Biosyst.*, 8, 1650–1657.

36. Lehmann,R. and Nusslein-Volhard,C. (1991) The maternal gene nanos has a central role in posterior pattern formation of the *Drosophila* embryo. *Development*, 112, 679–691.

37. Ren,B., Robert,F., Wyrick,J.J., Aparicio,O., Jennings,E.G., Simon,L., Zeitlinger,J., Schreiber,J., Hannett,N., Kanin,E. and Green,M. (2000) Genome-wide location and function of DNA binding proteins. *Science*, 290, 2306–2309.

38. Tusher,V.G., Tibshirani,R. and Chu,G. (2001) Significance analysis of microarrays applied to the ionizing radiation response. *Proc. Natl Acad. Sci. USA.*, 98, 5116–5121.

39. Saeed,A.I., Bhagabati,N.K., Brasted,J.C., Liang,W., Sharov,V., Howe,E.A., Li,J., Thiagarajan,M., White,J.A. and Quackenbush,J. (2006) TM4 microarray software suite. *Methods Enzymol.*, 411, 134–193.

40. Saeed,A.I., Brasted,J.C., Liang,W., Sharov,V., Howe,E.A., Li,J., Thiagarajan,M., White,J.A. and Quackenbush,J. (2003) TM4: a free, open-source system for microarray data management and analysis. *Biotechniques*, 34, 374–378.

41. Edgar,R., Domrachev,M. and Lash,A.E. (2002) Gene expression omnibus: NCBI gene expression and hybridization array data repository. *Nucleic Acids Res.*, 30, 207–210.

42. Huang-da,W., Sherman,B.T. and Lempicki,R.A. (2009) Systematic and integrative analysis of large gene lists using DAVID bioinformatics resources. *Nat. Protoc.*, 4, 44–57.
44. Huang da., W., Sherman, B.T. and Lempicki, R.A. (2009) Bioinformatics enrichment tools: paths toward the comprehensive functional analysis of large gene lists. *Nucleic Acids Res.*, 37, 1–13.

45. Bernhart, S.H., Hofacker, I.L. and Stadler, P.F. (2006) Local RNA base pairing probabilities in large sequences. *Bioinformatics*, 22, 614–615.

46. Schliep, S., Maticzka, D., Mohl, M., Gagnon, J.N., Brown, C.M. and Backofen, R. (2012) Global or local? predicting secondary structure and accessibility in mRNAs. *Nucleic Acids Res.*, 40, 5215–5226.

47. Ding, Y. and Lawrence, C.E. (2003) A statistical sampling algorithm for RNA secondary structure prediction. *Nucleic Acids Res.*, 31, 7290–7301.

48. Hulsken, T., de Vlieg, J. and Alkema, W. (2008) BioVenn - a web application for the comparison and visualization of biological lists using area-proportional venn diagrams. *BMC Genomics*, 9, 488.

49. Lécuyer, E., Yoshida, H., Parthasarathy, N., Alm, C., Babuk, T., Cerovina, T., Hughes, T.R., Tomancak, P. and Krause, H.M. (2007) Global analysis of mRNA localization reveals a prominent role in organizing cellular architecture and function. *Cell*, 131, 174–187.

50. Tomancak, P., Beaton, A., Weiszmann, R., Kwan, E., Shu, S., Lewis, S.E., Richards, S., Ashburner, M., Hartenstein, V., Celniker, S.E. et al. (2002) Systematic determination of patterns of gene expression during *Drosophila* embryogenesis. *Genome Biol.*, 3, RESEARCH0088.

51. Tomancak, P., Berman, B.P., Beaton, A., Weiszmann, R., Kwan, E., Hartenstein, V., Celniker, S.E. and Rubin, G.M. (2007) Global analysis of patterns of gene expression during *Drosophila* embryogenesis. *Genome Biol.*, 8, R145.

52. Graveley, B.R., Brooks, A.N., Carlson, J.W., Duff, M.O., Landolin, J.M., Yang, L., Artieri, C.G., van Baren, M.J., Boley, N., Booth, B.W. et al. (2011) The developmental transcriptome of *Drosophila melanogaster*. *Nature*, 471, 473–479.

53. Gerber, A.P., Luschnig, S., Krasnow, M.A., Brown, P.O. and Herschlag, D. (2006) Genome-wide identification of mRNAs associated with the translational regulator PUMILIO in *Drosophila* melanogaster. *Proc. Natl Acad. Sci. USA*, 103, 4487–4492.

54. Lopez de Silanes, I., Zhan, M., Lal, A., Yang, X. and Gorospe, M. (2008) Identification of a target RNA motif for RNA-binding protein HuR. *Science*, 320, 753–757.

55. Gama-Carvalho, M., Barbosa-Morais, N.L., Brodsky, A.S., Lopez de Silanes, I., Zhan, M., Lal, A., Yang, X. and Gorospe, M. (2009) Genome-wide identification of mRNAs associated with the translational regulator PUMILIO in *Drosophila* melanogaster. *Nature*, 471, 473–479.

56. Tomancak, P., Beaton, A., Weiszmann, R., Kwan, E., Shu, S., Lewis, S.E., Richards, S., Ashburner, M., Hartenstein, V., Celniker, S.E. et al. (2002) Systematic determination of patterns of gene expression during *Drosophila* embryogenesis. *Genome Biol.*, 3, RESEARCH0088.

57. Tomancak, P., Berman, B.P., Beaton, A., Weiszmann, R., Kwan, E., Hartenstein, V., Celniker, S.E. and Rubin, G.M. (2007) Global analysis of patterns of gene expression during *Drosophila* embryogenesis. *Genome Biol.*, 8, R145.

58. Lopez de Silanes, I., Zhan, M., Lal, A., Yang, X. and Gorospe, M. (2008) Identification of a target RNA motif for RNA-binding protein HuR. *Science*, 320, 753–757.

59. Wu, H., Henras, A., Chanfreau, G. and Feigon, J. (2004) Structural protein HuR. *Science*, 320, 753–757.

60. Harrison, M.M., Botchan, M.R. and Cline, T.W. (2010) The zinc-finger protein Zelda is a key activator of the early zygotic genome in *Drosophila melanogaster*. *Dev. Biol.*, 338, 15864–15869.

61. Harrison, M.M., Li, X.Y., Kaplan, T., Botchan, M.R. and Cline, T.W. (2010) Genome-wide identification of alternative splice mRNA species despite evolution of diverse mRNA target sets. *Mol. Cell. Biol.*, 28, 4093–4103.

62. Li, X., Quon, G., Lipshitz, H.D. and Morris, Q. (2010) Predicting in vivo binding sites of RNA-binding proteins using mRNA secondary structure. *RNA*, 16, 1096–1107.

63. Wu, H., Henras, A., Chanfreau, G. and Feigon, J. (2004) Structural basis for recognition of the AGN7 tetraloop RNA fold by the double-stranded RNA-binding domain of Rnt1p RNAIII. *Proc. Natl Acad. Sci. USA*, 101, 8307–8312.

64. Liang, H.L., Nien, C.Y., Liu, H.Y., Metzstein, M.M., Kirov, N. and Rushlow, C. (2008) The zinc-finger protein Zelda is a key activator of the early zygotic genome in *Drosophila melanogaster*. *Nature*, 456, 400–403.

65. Harrison, M.M., Botchan, M.R. and Cline, T.W. (2010) Grainyhead localization signal prediction. *BMC Bioinformatics*, 10, 202.
Emeson, R.B. et al. (2010) The solution structure of the ADAR2 dsRBM-RNA complex reveals a sequence-specific readout of the minor groove. Cell, 143, 225–237.

Katze, M.G., DeCorato, D., Safer, B., Galabru, J. and Hovanessian, A.G. (1987) Adenovirus VAI RNA complexes with the 68 000 Mr protein kinase to regulate its autophosphorylation and activity. EMBO J., 6, 689–697.

Davis, S. and Watson, J.C. (1996) In vitro activation of the interferon-induced, double-stranded RNA-dependent protein kinase PKR by RNA from the 3’ untranslated regions of human alpha-tropomyosin. Proc. Natl Acad. Sci. USA, 93, 508–513.

Robertson, H.D., Manche, L. and Mathews, M.B. (1996) Paradoxical interactions between human delta hepatitis agent RNA and the cellular protein kinase PKR. J. Virol., 70, 5611–5617.

Gleghorn, M.L., Gong, C., Kielkopf, C.L. and Maquat, L.E. (2013) Staufen1 dimerizes through a conserved motif and a degenerate dsRNA-binding domain to promote mRNA decay. Nat. Struct. Mol. Biol., 20, 515–524.