Systematic discovery of new recognition peptides mediating protein interaction networks

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Supplementary Information

Interaction data
We searched for new motifs among sets of proteins sharing an interaction partner. We used interactions determined by genome-scale yeast two-hybrid screens or mined from the literature to create sets of four or more proteins sharing an interaction partner. For Yeast (S. cerevisiae) we used all data from Uetz et al. [1] and core-data from Ito et al. [2]. For Fly (D. melanogaster) we used those with a confidence ≥0.5 from Giot et al. [3], and for Nematode (C. elegans) we used the core 1, core 2 and literature data from Li et al. [4]. For Human (H. sapiens) we used interaction data extracted from the literature in the Human Proteome Resource Database (HPRD [5]). We did not use data from affinity purifications as they capture fewer direct, transient interactions [6] typical of those involving linear motifs. To find motifs of a more general nature (i.e. core motifs typical for a domain family), we pooled proteins into ‘domain’ sets (as distinct from ‘protein’ sets) if their interaction partners shared a common domain (Fig S1).

Domain versus protein sets
We constructed domain sets by pooling the all interaction partners for proteins containing a particular domain (Fig S1). As shown in the Figure S1, this pooling can
sometimes lead to the detection of motifs that might otherwise be missed because of too few interaction partners when considering single proteins. However, this can also lead to motifs being assigned to the wrong domain (e.g. SH3 PxxP motifs can be assigned to SH2 domains since SH3 & SH2 domains often co-occur).

Figure S1
Interaction maps are probed for protein interaction sets (A): partners of proteins with multiple interactions are clustered together where there are no known sequence feature present (B). Domains and homologous regions are then identified (B) and removed prior to running exhaustive pattern discovery (C) to produce list of motifs ranked by their probabilities P (D). Hypothetical motifs are shown as coloured squares in C & D. 'Proteins' in D gives the set of proteins containing at least one copy of the motif.

I. Shows two separate cases of protein sets containing interactions partners for proteins X and Y that are pooled owing to the presence of a common domain to give a domain set shown in (II).
Recombinant DNA constructs
We cloned Dynein light chain (Cdlc2, 10 kDa) and Translin (Translin, 27 kDa) from Drosophila melanogaster directly into pET41b+ vector (Novagen); PP1 from H. sapiens (PP1, 37 kDa; N.B. the human and yeast proteins are 93% identical), kindly provided by Mathieu Böllen (Katholieke Universiteit Leuven), in a pTacTac vector [7]; and Elongin C from H. sapiens (EloC, 15 kDa; N.B. the human and fly proteins are 94% identical), kindly provided by Joan Conaway (Stowers Institute, Kansas City), in a pRSETB vector (Invitrogen).

Protein Expression
We used positive clones to transform E. coli BL21(DE3) (Stratagene), and grew cells at 37 °C in 6 x 1l of Luria-Bertani broth supplemented with either 50 µg/ml Ampicillin (PP1 and EloC) or 30 µg/ml (Cdlc2 and Translin) until absorbance at 600 nm reached 0.4. We then induced over-expression by the addition of 40 µg/ml Isopropyl-ß-D-thiogalactopyranosid (IPTG) and grew cells for 16h at 16°C.

We harvested the cells and re-suspended them in 80 ml PBS together with one Complete™ protease inhibitor cocktail tablet (Roche) and DNaseA. We then lysed them using a French Press (Emulsiflex C5) at 1000-1500psi. We cleared the lysate by centrifugation at 40000 rpm for 50 min, and loaded the soluble fraction onto various purification columns depending on the construct.

Protein Purification
We purified all proteins using a Biologic HR workstation (Bio-Rad). For Cdlc2 and Translin we used a 5ml GST-Trap column (Pharmacia) equilibrated in PBS and eluted in 50 mM Tris-HCl pH 8.0, 10 mM Glutathione; we purified PP1 as described previously [8]; and we used a MonoQ HR 16/10 (Pharmacia) to purify EloC. We also used a previous preparation of human abl-SH3 domain [9]. We were unable to purify Elongin C to the concentration required for fluorescence polarization measurements.
Fluorescence Polarization

We obtained FITC N-terminal labeled micro-scale peptides (JBT Peptide Technologies, Berlin) for fluorescence polarization measurements. A preliminary test with abl-SH3 binding peptide (GAPTYSPPPPP) demonstrated that we could both observe binding and compute a $K_D$ of 3.0±0.4 μM roughly comparable to that obtained previously by measuring fluorescence of tryptophans in the abl-SH3 domain (0.4μM for a peptide lacking the N-terminal Glycine) [10].

For each protein/motif pair we selected peptides from two proteins containing the predicted motif, with an additional two amino acids from the N- and C- termini. For one of these we also obtained a mutant peptide where the positions specifying the motif had been mutated to Alanine. The peptides were as follows: Cdcl2 native ERAIQTENA (CG33213/104-112), mutant ERAAAAENA; PP1 native DPDAEDEDQDS (Scd5/302-312) mutant DPAAEAEDQAS; Translin VSVPARVYSPV (CG8965/341-352), mutant VSASPAAVAAPV. When binding was observed (PP1, Translin), we tested whether the binding was non-specific by assaying a selection of other peptides arbitrarily chosen from other protein/motif pairs (an example is shown in Fig 3B & Fig 4B; none of the arbitrary peptides bound specifically). For this protein we also tried an additional peptide containing the canonical TQT motif (PFGTQTSPPP, from CG32130/215-223). No binding was observed with any of the peptides for Cdcl2, even at protein concentrations as high as 400 μM.

We suspended peptides to working concentrations (10 nM final concentration) in PBS, pH 7.4. We then added increasing concentrations of the recombinant protein to PBS solution and mixed with the FITC-labeled peptides on 384-well Proxiplate (PerkinElmer, USA), and incubated each sample for 5 min at room temperature. We then measured Fluorescence polarization on an EnVision HTS (PerkinElmer, USA) according to the manufacturers’ instructions. No increase in the fluorescence polarization was observed upon longer incubations. We plotted saturation binding curves with the PRISM software (Graph-Pad, San Diego, USA), and used it to compute dissociation constants ($K_D$) via non-linear regression using the results from three individual experiments.
References


