Research Letter

Two Distantly Spaced Basic Patches in the Flexible Domain of Huntingtin-Interacting Protein 1 (HIP1) Are Essential for the Binding of Clathrin Light Chain

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The interaction between HIP family proteins (HIP1 and HIP12/1R) and clathrin is fundamental to endocytosis. We used circular dichroism (CD) to study the stability of an HIP1 subfragment (aa468-530) that is splayed open. CD thermal melts show HIP1 468-530 is only stable at low temperatures, but this HIP1 fragment contains a structural unit that does not melt out even at 83°C. We then created HIP1 mutants to probe our hypothesis that a short hydrophobic path in the opened region is the binding site for clathrin light chain. We found that the binding of hub/LCb was sensitive to mutating two distantly separated basic residues (K474 and K494). The basic patches marked by K474 and K494 are conserved in HIP12/1R. The lack of conservation in sla2p (S. cerevisiae), HIP1 from D. melanogaster, and HIP1 homolog ZK370.3 from C. elegans implies the binding of HIP1 and HIP1 homologs to clathrin light chain may be different in these organisms.

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1. Introduction

Huntingtin-interacting protein 1 (HIP1) and its relative, HIP12/1R, contribute to the budding of clathrin-coated vesicles (CCVs) [1, 2]. The yeast homolog of HIP1, sla2p, is required for the development of CCVs in yeast [3, 4]. There are shared regions in HIP1 and HIP12/1R that mediate the binding of clathrin, but there are also sites that are unique to each HIP protein. For example, HIP1, but not HIP12/1R, has a clathrin box motif (332LMDMD) which functions to bind the N-terminal beta propeller domain of clathrin [5]. HIP1 and HIP12/1R do not interact with adaptor protein 2 (AP2) in the same way. The AP2-binding FXDXF and DPF motifs (X denotes any amino acid) are only in HIP1 [5]. HIP12/1R apparently has a higher affinity for F-actin [6], suggesting that HIP1 and HIP12/1R play distinct roles in the formation of CCVs. A number of amino acids (L486 and R487 in the DLL486R487KN region) in the coiled-coil domain of HIP1 and HIP12/1R have been identified to impact the binding of clathrin light chain (CLC) [7], but we do not know if L486 and R487 constitute the entire CLC binding determinants in HIP proteins. We recently determined the crystal structures of two contiguous subfragments of the HIP1 coiled-coil domain (PDB files: 2NO2 and 2QA7) that span the clathrin light chain-binding region [8, 9]. The DLL486R487KN stretch was embedded in the splayed opened region that we first discovered in HIP1 482-586 (2NO2) [8] and later found in HIP1 371-481 (2QA7) [9]. In this research letter we define the inherent stability of a segment of HIP1 that contains the opened region and ask if electrostatic interactions help drive the binding of CLC. We generated an HIP1 subfragment (aa468-530) that spans the opened region for circular dichroism (CD) experiments to assess this domain’s intrinsic stability. We then studied if two distantly spaced positively charged patches in the opened region played any role in the binding of CLC. Here we report that the HIP1 468-530 construct is unstable, but paradoxically a heat-resistant structural unit is present within this subfragment. We found that the two basic patches in the flexible part of HIP1 (centered on K474 and K494) are crucial for the binding of CLC. These electrostatic determinants are part of a solvent exposed hydrophobic surface that we previously argued was suitable for CLC.
Figure 1: Structural features of the HIP1 clathrin light chain-binding site. Two distantly separated basic amino acids in the opened region of HIP1 mediate the binding of clathrin light chain. The assembled model was generated using PDB files 2NO2 [8] and 2QA7 [9]. For the sake of clarity, we only show a portion of the second HIP1 helix (dark grey) that stops before Y468. The yellow bar marks the position of the solvent exposed hydrophobic S3 path [8]. The new data in Figure 3 indicate that K474 is a strong determinant for binding and imply that S3 path begins before the DLL486R487KN region. The N- and C-termini of the HIP1 crystal structure are labeled N and C. The numbers 1–3 along the yellow bar mark the position of amino acids that control the binding of clathrin light chain (position 1: K474; position 2: L486 and R487) mark the position of amino acids that control the binding of clathrin light chain (position 1: K474; position 2: L486 and R487, reported by the McPherson group [7]; and position 3: K494). K474 and K494 are ~50 Å apart, R500, R508 or K511 (see arrow) do not participate in binding and therefore define the boundary of the light chain-binding site. The HIP1 468-530 subfragment in the CD studies in Figure 2 spans across an opened region of the HIP1 coiled coil in our 2NO2 and 2QA7 crystal structures. The HIP1 model was created using PyMol (http://www.pymol.org).

2. Results and Discussion

2.1. Coiled-Coil Segment of HIP1 that Includes Determinants for Clathrin Light Chain Is Intrinsically Unstable. The binding of CLC to HIP1 requires L486 or R487 [7] (human HIP1 numbering (accession number NP 005329), conserved in HIP12/1R). We recently located L486 and R487 to a flexible region of HIP1 that mediates the binding of Huntingtin interacting protein 1 interactor (HIPPI) [9]. To probe if those in S3 path contributed, we made 5 GST-HIP1h (HIP1h is aa370-644) 370-644 mutants (K494A, K494E, R500E, R508E, and K511E). We performed GST pulldowns to evaluate the binding of 6HisHub/6HisLCb (hub is central third of clathrin) to closely mimic how HIP1 interacts with clathrin baskets in cells. Every GST pulldown was done at least three times, using freshly isolated proteins and charged GST beads each time. Clathrin hub (N-terminally histidine-tagged bovine clathrin light chain b with the neuronal insert) was detected by western blotting with a commercial histidine tag antibody. LCb was blotted with CON.1 monoclonal antibody and GST-HIP1h constructs were visualized with a commercial GST antibody. The anti-GST bands in Figures 3(a) and 3(b) showed that the GST-HIP1h levels were balanced (loading control). The negative controls in lanes 1–3 in Figure 3(a) and lanes 1–3 in Figure 3(b) show that the GST signals from each binding experiment were not random interactions, but reflected true binding events. The level of 6HisHub/6HisLCb captured by GST-HIP1h is shown in lane 4 in Figures 3(a) and 3(b), and as expected, required bound LCb (compare lanes 3 and 4 in Figures 3(a) and 3(b)). We did not remove the histidine tag participate in binding CLC. K494 (see position 3, Figure 1) is in S3 path (yellow bar, Figure 1) previously described [8] and is followed by a cluster of basic residues (indicated by the arrow in Figure 1). K474 (position 1) is located before the DLL486R487KN region, close to the part of HIP1 that mediates the binding of Huntingtin interacting protein 1 interactor (HIPPI) [9]. The position of the 222 nm signal did not change with temperature, but the 208 nm signal shifted as the sample was heated. The 222/208 nm ratios were calculated from the raw CD data. Because the signal below 200 nm was noisy, we did not attempt to get the helix content from the CD data.

Figure 2: HIP1 468-530 subfragment in the opened region is stable only at low temperature. Equilibrium far UV CD scans were performed at 4°C (open triangle), 10°C (open diamond), 37°C (cross), and 83°C (open circle). The position of the 222 nm signal did not change with temperature, but the 208 nm signal shifted as the sample was heated. The 222/208 nm ratios were calculated from the raw CD data. Because the signal below 200 nm was noisy, we did not attempt to get the helix content from the CD data.

2.2. Basic Patches Centered on K474 and K494 in HIP1 Are Essential for Binding Clathrin Light Chain. We investigated if a series of basic patches in the opened region could
on LCb used for purification because control experiments showed that the tag did not interfere with the GST pulldowns (data not shown). The binding of 6Hishub/6HisLCb was significantly blocked when HIP1h K494 was replaced with glutamic acid (K494E mutant, lane 5 in Figure 3(a)) or with alanine (lane 5, Figure 3(b)). In contrast, we saw no detectable impact when R500, directly above K494 (see Figure 1 for location), was changed to glutamic acid (see lane 6, Figure 3(a)) or alanine (data not shown). Consistent with data in lanes 5 and 6 in Figure 3(a), the K494E/R500E double mutant did not bind 6Hishub/6HisLCb (lane 7, Figure 3(a)). Next we evaluated a group of basic residues close to the C-terminal end of S3 path (indicated by an arrow in Figure 1). Lanes 8 and 9 in Figure 3(a) show that the R508E and K511E charge flip mutants bound 6Hishub/6HisLCb similar to the wild type control (compare lane 4 with lanes 8-9 in Figure 3(a)). We conclude from these data that K494 located in the hydrophobic S3 path is required to bind CLC, but R500, R508, and K511 around this path do not participate. This suggests that the CLC binding site does not go beyond the boundaries of S3 path that is defined by R500, R508, and K511.

The assembled HIP1 model (2NO2 joined to 2QA7) in Figure 1 shows how the positive patch centered on K474 is oriented relative to K494 (separated by ∼50 Å). Our data in Figure 3(b) show a dramatic drop in the binding of 6Hishub/6HisLCb when K474 was mutated to alanine (see lanes 4 and 6 (K494A in lane 5 for comparison)). This result demonstrates for the first time that K474 upstream the DLL486R487KN region is necessary for binding 6Hishub/6HisLCb and suggests that the CLC binding site may be more extensive than previously thought. We predict that the two basic patches in HIP1 we have defined here are also present in HIP12/R because K474 and K494 (HIP1 numbering) are both conserved in this protein. Finally, we looked if K474, R487, and K494 were conserved in HIP1 from vertebrates Mus musculus, Rattus norvegicus, Xenopus laevis, Drosophila melanogaster, and in sla2p (HIP1 homolog from Saccharomyces cerevisiae) and ZK370.3 (HIP1 homolog from Caenorhabditis elegans). This analysis showed that K474 was conserved in all the vertebrates but was V in C. elegans, L in D. melanogaster, and M in sla2p. R487 was conserved in all the vertebrates but was E in both D. melanogaster and sla2p and T in C. elegans. K494 was conserved in all vertebrates and invertebrate HIP1 proteins, except sla2p (K474 (Q in S. cerevisiae), R487 (T), and K494 (D)). The natural mutations in sla2p, HIP1 from D. melanogaster, and ZK370.3 from C. elegans could mean that the binding of HIP1 and HIP1 homologs to clathrin light chain in these organisms is different.

3. Materials and Methods

3.1. Materials. Triton X-100, Tween-20, beta-mercaptoethanol (βME), TRIZMA base, and BIS-TRIS were from Sigma-Aldrich (St. Louis, Mo, USA). Sodium phosphate dibasic (Na2HPO4) was from EMD Chemicals (Gibbstown, NJ, USA); Luria broth was from EMD Biosciences (Sparks, Md, USA); tris(2-carboxyethyl)-phosphine was from Sigma-Aldrich. Pfu turbo was from Stratagene (La Jolla, Calif, USA) and primers were from Integrated DNA Technologies (Coralville, Iowa, USA). The Pierce Coomassie Plus Bradford reagent kit was purchased from Fisher Scientific (Hanover Park, Ill, USA). Chromatography resins, columns, and standards were purchased from GE Healthcare (Piscataway, NJ, USA). CON.1 antibody was bought from Covance (Cumberland, Va, USA); restriction grade thrombin and the anti-His antibody were obtained from Novagen (La Jolla, Calif, USA). Coomassie G-250 stain and Immun-Star chemiluminescent kit were from Bio-Rad Laboratories (Hercules, Calif, USA).

3.2. Construction of GST-HIP1h Mutants and 6HisLCb. The plasmid encoding the original N-terminal GST tagged HIP1h 370-664 was a gift from the McPherson group. The various GST-HIP1h mutants used in this work were created using the QuikChange mutagenesis protocol (Stratagene). The sequence was confirmed by DNA sequencing (IMBI, Indiana University) and then transformed into Rosetta 2 (DE3) pLysS cells (Novagen). Standard cloning was used to insert neuronal LCb DNA in pET15b to generate 6HisLCb. The recombinant 6HisLCb plasmid was transformed into BL21 (DE3) pLysS cells for overexpression.

3.3. Protein Overexpression and Purification. The recombinant GST-HIP1h constructs were grown at 37°C in 1 L Luria broth (LB) to an O.D. 600 of 0.5–0.8 units. The incubation temperature was dropped to 30°C cells and protein expression was induced with IPTG (100 µg/mL final concentration). Cells were harvested after 3 hours at 30°C and bacterial pellets were frozen at −80°C before use. Bacterial pellets were resuspended in 50 mL of 1X PBS (10 mM Na2HPO4, 1.8 mM KH2PO4 (pH 7.3), 140 mM NaCl, 2.7 mM KCl, supplemented with 0.25 mL of 1 M DTT, 0.25 mL of protease inhibitor cocktail (Sigma), and 2 mL of PMSF (17.4 mg/mL in 2-propanol)). After sonication, 2.5 mL of 20% (v/v) Triton X-100 was added and the lysate was rotated at room temperature for ~30 minutes. The crude bacterial lysate was spun at 12000 g (4°C) for 10 minutes. The supernatant was mixed with ~5 mL of glutathione Sepharose 4B (Amersham) resin suspended in PBS. The GST-HIP1h constructs were eluted from the column with 50 mL of 3 mg/mL L-glutathione (sigma) in (PBS) at pH 8.0 and dialyzed overnight against the same buffer. For CD experiments the purification protocol was modified so that we can cleave the GST tag in the GST column. After bacterial lysate was added to GST beads, the mixture was rocked at room temperature for 1-2 hours. The beads were spun down at 500 g at 4°C with 5 minutes and the supernatant was poured off. The wet beads were transferred to a column and washed slowly with 110 mL 1X PBS and then 1 unit thrombin per mg of protein was diluted into 3.5 mL of 1X PBS and added to the column. After digestion, the HIP1h constructs were further purified on Superdex 75 column (GE Healthcare) equilibrated with 1X PBS (at room temperature). Column
fractions were pooled and dialyzed at 4°C against 10 mM potassium phosphate buffer at pH 7.9.

The clathrin hub construct was purified as previously described [13] and the 6HisLCb was purified in a single step using the nickel affinity resin. After the crude lysate was added to charged Sepharose nickel resin and incubated at 4°C for 10 minutes, the beads were gently spun down and washed with 25 mL of 10 mM Na2HPO4, 10 mM imidazole, 0.5 M NaCl, pH 7.4 buffer (buffer A). The beads were then washed with 50 mL of buffer A that contained 0.5 M imidazole. The 6HisLCb was eluted off the column with buffer A that contained 205 mM imidazole. EDTA was added to the sample (1 mL 0.5 M EDTA per 10 mL of protein) and dialyzed against 10 mM tris, pH 7.9 overnight at 4°C (dialysis buffer includes BME, and a cocktail of protease inhibitor).

3.4. CD Measurements. Purified HIP1 468-530 was diluted with 10 mM potassium phosphate buffer at pH 7.9 to 0.5 mg/mL for CD measurements at different temperatures. CD data was collected using a Jasco J-175 circular dichroism spectropolarimeter with thermally controlled sample cells. The first CD scan was taken at 4°C and then the temperature was changed to the indicated temperatures in Figure 2. The sample was allowed to sit at each indicated temperature for several minutes before taking the CD scan.

3.5. GST Pull Down Assays. Glutathione Sepharose 4B resin (GE Healthcare 17-0756-01) was washed three times with 1 mL of PBS. Protein concentrations were determined by Bradford assays (Pierce 23236). Equal molar amounts of GST and GST-Hip1h proteins (1 μM) and 1 mL of PBS were added to resin and incubated on a rotating platform at 4°C for 1 hour. Unbound proteins were removed by washing three times with binding buffer (50 mM Tris, 200 mM KCl, 1 mM EDTA, 1% Triton X-100, 50 mM imidazole, 0.5 mg/mL ovalbumin, pH 8.0). 6HisLCb and Hub alone were combined in a 3:1 ratio and allowed to incubate at 4°C for at least 30 minutes. 25 μL of GST or GST-Hip1h bound beads, 0.4 nmol of Hub Alone or 1.9 nmol of Hub 6HisLCb complex, 375 μL of room temperature binding buffer were added to MicroSpin columns (GE Healthcare 27-3565-01) and incubated on rotator at 4°C for 1 hour. Beads were washed six times with 0.5 mL of binding buffer + 16 mM imidazole and supernatant removed by centrifugation. After final wash, 55 μL of 2x SDS gel loading buffer was added to beads. Spin columns were closed placed in 1.7 mL Eppendorf tubes, heated for 10 minutes at 80–90°C, and spun down to collect samples.

3.6. Western Blots. The bound proteins were resolved by SDS-PAGE and analyzed by standard western blotting. After transfer the nitrocellulose membrane was stained with Ponceau stain and cut to separate proteins for blotting. Hub was detected with anti-His monoclonal antibody (Novagen 70796-3), GST and GST-Hip1h were detected with anti-GST monoclonal antibody (Covance MMS-112P), and 6HisLCb was detected with clathrin light chain monoclonal antibody (CON.1) (Covance MMS-423P). Binding was detected with
Immun-Star chemiluminescent protein detection system (BioRad 170-5010).

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References


