Cavin1 intrinsically disordered domains are essential for fuzzy electrostatic interactions and caveola formation

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Caveolae are spherically shaped nanodomains of the plasma membrane, generated by cooperative assembly of caveolin and cavin proteins. Cavins are cytosolic peripheral membrane proteins with negatively charged intrinsically disordered regions that flank positively charged α-helical regions. Here, we show that the three disordered domains of Cavin1 are essential for caveola formation and dynamic trafficking of caveolae. Electrostatic interactions between disordered regions and α-helical regions promote liquid-liquid phase separation behaviour of Cavin1 in vitro, assembly of Cavin1 oligomers in solution, generation of membrane curvature, association with caveolin-1, and Cavin1 recruitment to caveolae in cells. Removal of the first disordered region causes irreversible gel formation in vitro and results in aberrant caveola trafficking through the endosomal system. We propose a model for caveola assembly whereby fuzzy electrostatic interactions between Cavin1 and caveolin-1 proteins, combined with membrane lipid interactions, are required to generate membrane curvature and a metastable caveola coat.

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CAV1 is able to generate a metastable caveola coat.

**Results**

**Cavin1 forms electrostatically driven oligomers that depend on DR1 and DR3 domains.** The cavin family proteins all share distinguishing structural similarities with each other, consisting of disordered N-terminal and C-terminal domains DR1 and DR3, a central disordered region DR2, and interspersed α-helical coiled-coil region HR1 and predicted α-helical region HR2 (Fig. 1a). The trimeric coiled-coil HR1 domain and C-terminal HR2 domains are both rich in basic amino acid residues, while the three DR domains instead possess a high proportion of acidic amino acid residues. This alternating electrostatic charge distribution is a distinctive and conserved feature of all family members (Supplementary Fig. 1A), indicating it is an essential characteristic of the proteins. We also used the D2P2 web server to analyse the sequence of Cavin1 for predicted regions of disorder, and confirmed that the DR1, DR2 and DR3 regions are predicted to be intrinsically disordered as suggested by previous secondary structure analyses (Supplementary Fig. 1B). Interestingly, sites of phosphorylation in Cavin1 are predominantly found in the DR1, DR2, and DR3 domains, while sites of ubiquitylation are concentrated in the HR1 and HR2 regions. In subsequent experiments the boundaries of the mouse Cavin1 domains are defined as: DR1 (1–44), HR1 (45–155), DR2 (156–209), HR2 (210–310) and DR3 (311–392) (Fig. 1a). Expression constructs used in this study are outlined in Supplementary Fig. 2.

We recently proposed that the predominantly negatively charged DR sequences of Cavin1 may associate with the positively charged HR domains to promote intramolecular and/or intermolecular electrostatic interactions required for coat assembly. To probe the role of electrostatic interactions, we used fluorescence correlation spectroscopy (FCS) to measure the diffusional purities of purified GFP-tagged Cavin1 in both 500 mM NaCl (high salt) and 150 mM NaCl (iso-osmotic salt concentration). According to polymer theory, the diffusivity of protein molecules in solution decreases with increasing intermolecular interactions due to molecular crowding limiting its molecular motion. GFP-Cavin1 (100 nM concentration) showed a remarkable decrease in its diffusivity with the reduction of ionic strength from 500 mM NaCl (12.01 ± 3.03 µm²/s) to 150 mM NaCl (4.02 ± 0.50 µm²/s) (Fig. 1b). This indicates that at physiological salt concentrations Cavin1 can form homomeric oligomers with an average hydrodynamic radius ~55 nm, similar to those observed previously, and that this self-association is dependent on electrostatic interactions. In contrast to full-length Cavin1, removal of either N-terminal or C-terminal DR1 or DR3 domains prevents this electrostatically driven self-assembly at lower physiological salt concentrations (Fig. 1b).

Next, we sought to understand the role of DR sequences in oligomeric assembly of Cavin1 in a more representative cellular milieu. For these experiments we used MCF7 cells, which lack caveola and do not express any caveolin or cavin proteins. GFP-tagged Cavin1 proteins were transiently expressed and FCS
analysis was used to measure the diffusivity of each protein in cell lysates (all at 150 mM NaCl). Full length GFP-Cavin1 in MCF7 cell lysates forms relatively heterogeneous large molecular weight species in solution with slow diffusive properties (6.35 ± 2.35 µm²/s) (Fig. 1c) similar to purified recombinant GFP-Cavin1. In contrast to purified recombinant GFP-Cavin1-ΔDR1, the N-terminal DR1 deletion in cell lysates showed a similar (although tending to faster) rate of diffusion to the full-length protein (7.65 ± 4.40 µm²/s) (Fig. 1c). Complete deletion of the C-terminal DR3 region of GFP-Cavin1-ΔDR3 however, significantly increased the diffusivity of Cavin1 in MCF7 lysates (18.54 ± 6.22 µm²/s), similar to the recombinant GFP-Cavin1-ΔDR3 (Fig. 1c). Overall these studies demonstrate a role for the DR sequences in electrostatically driven oligomerisation of Cavin1 in solution.

Cavin1 undergoes liquid–liquid phase separation (LLPS) influenced by the DR domains. There is an increasing awareness of the role of intrinsically disordered sequences in generating membraneless organelles via liquid–liquid phase separation (LLPS) or ‘demixing’ of proteins and associated molecules in solution. Demixing or LLPS can be driven by a variety of mechanisms, including cation–π and π–π stacking, interactions with polyanions such as RNA, and intermolecular electrostatic interactions. A number of recent studies have shown that membranes can be platforms for nucleating and transporting phase-separated assemblies, or in turn be regulated and organised via LLPS-mediated processes. It has also been proposed that the formation of phase-separated condensates can perform physical work on their surroundings, including at the membrane–cytosol interface to generate membrane curvature. Because of the demonstrated importance of disordered regions in Cavin1 for its assembly behaviour, we assessed whether purified Cavin1 is able to form supramolecular assemblies leading to LLPS in vitro.

Purified recombinant GFP-Cavin1 expressed in E. coli remains dispersed in solution at both physiological NaCl concentration (150 mM) and at high NaCl concentration (750 mM) within a protein concentration range of 1–10 µM (Supplementary Fig. 3A). However, when Dextran T-500 (1.25% w/v) was added as a macromolecular crowding agent, full-length GFP-Cavin1 rapidly formed spherical liquid droplets at 150 mM NaCl even at low protein concentrations (0.1 µM) (Fig. 2a and Supplementary Fig. 3B, C). This is well below the estimated cellular concentration of Cavin1 of 3 µM. These droplets increased in size with increasing protein concentration in the range 1–10 µM (Fig. 2a). Increasing the salt concentration strongly inhibited the ability of GFP-Cavin1 to undergo LLPS, consistent with a role for electrostatic intermolecular interactions. We also tested both purified full-length Cavin1-GFP isolated from mammalian HEK293 cells, and unpurified Cavin1-GFP in MCF7 cell lysates, and found that both preparations underwent similar salt-sensitive LLPS (Supplementary Fig. 3D, 3E) although this required a higher concentration of Dextran T-500 (3%), possibly due to the presence of other bound proteins, lipids, or post-translational modifications such as phosphorylation, ubiquitylation, and SUMOylation present in the mammalian cell expression system. Higher concentrations of Dextran T-500 (3%) did not significantly alter the LLPS behaviour of GFP-Cavin1 (Supplementary Fig. 3F). Lastly, we assessed if Cavin1 could undergo LLPS in cells. When overexpressed in MCF7 cells, which lack caveolae due to the absence of CAV1, we found that GFP-Cavin1 on its own remained diffuse and did not form droplets (Supplementary Fig. 4A, B). However, after testing several conditions we discovered that if cells were treated with cholesterol following serum starvation GFP-Cavin1 rapidly formed cytoplasmic condensates as well as membrane–associated tubules (Supplementary Fig. 4C, D). We then expressed GFP-Cavin1 in MCF7 cells together with an
mCherry-CAAX construct as a plasma membrane marker (Supplementary Fig. 4E). After cholesterol addition, we observed plasma membrane localisation of GFP-Cavin1 and formation of GFP-Cavin1 and mCherry-CAAX positive plasma membrane-associated tubules. Similar results were observed in CAV1\(^{-/-}\) mouse embryonic fibroblasts (MEFs) (Supplementary Fig. 4F). We speculate that cholesterol may alter the normal equilibrium of Cavin1’s interaction with phospholipid membranes, thus promoting self-association and condensation.

Deletion of the C-terminal DR3 domain had a small but reproducible effect on the tendency of Cavin1 to undergo LLPS in vitro, with droplet formation showing greater sensitivity to increasing ionic strength and protein concentration (Fig. 2a and Supplementary Fig. 3B, C). Similar results were observed in CAV1\(^{-/-}\) mouse embryonic fibroblasts (MEFs) (Supplementary Fig. 4F). We speculate that cholesterol may alter the normal equilibrium of Cavin1’s interaction with phospholipid membranes, thus promoting self-association and condensation.

Deletion of the C-terminal DR3 domain had a small but reproducible effect on the tendency of Cavin1 to undergo LLPS in vitro, with droplet formation showing greater sensitivity to increasing ionic strength and protein concentration (Fig. 2a and Supplementary Fig. 3B, C). Deletion of the DR1 domain however had a dramatic effect, leading Cavin1 to transition into non-spherical coacervates at all protein and salt concentrations (Fig. 2a and Supplementary Fig. 3B, S3C). Fluorescence recovery after photobleaching (FRAP) was used to analyse the diffusion of proteins within the liquid droplets and the ability of GFP-Cavin1 to exchange with bulk solution. GFP-Cavin1 (\(\tau_{1/2} \sim 20\) s) and GFP-Cavin1-\(\Delta DR3\) (\(\tau_{1/2} \sim 10\) s) showed rapid fluorescence recovery after photobleaching, indicating there is ready exchange of protein molecules within the droplets as expected for liquid droplets (Fig. 2b, c). GFP-Cavin1-\(\Delta DR1\), however, showed virtually no recovery (Fig. 2b, c), suggesting that gel formation has occurred and the truncated protein is unable to diffuse within the condensates\(^{19,43}\). Overall, these analyses highlight the importance of electrostatic interactions in promoting self-association and subsequent LLPS behaviour by Cavin1, and points to distinct roles of DR1 and DR3 sequences in this process.

Cavin1 promotes co-phase separation with N-terminal regions of CAV1. Although Cavin1 and CAV1 are associated together in caveolae, it remains unclear whether they interact with each other via direct protein–protein interactions. CAV1 has a unique structural domain architecture shared with other caveolins, consisting of an N-terminal disordered region (DR) (1–60), followed by an oligomerization domain (OD) (61–80), scaffolding domain (CSD) (81–100), intramembrane domain (IMD) (101–133), and a C-terminal membrane associated \(\alpha\)-helical domain (134–179) (Fig. 3a and Supplementary Fig. 5A, B)\(^{2,5}\). We hypothesised that the N-terminal disordered sequence of CAV1 may enable CAV1–Cavin1 association through interactions involving liquid–liquid phase separation. To test this, we first purified full length CAV1 fused with maltose binding protein (MBP) and GFP-binding nanobody protein (GBP)\(^{44}\) in non-ionic detergent n-dodecyl \(\beta\)-\(d\)-maltoside (DDM) (1.2 mM). MBP–GBP–CAV1 was labelled bound with GFP for visualisation, and unlabelled purified Cavin1 was used in a co-phase separation assay (Fig. 3b, c).
**Fig. 3** **CAV1 N-terminus co-phase separates with Cavin1.**

**a** Schematic representation of CAV1 domain architecture and design of various domain/point mutations. LLPS assays with MBP-GBP-CAV1 and cavin1 independently (b) and in mixture (c). MBP–GBP–CAV1 does not undergo LLPS in isolation but co-phase separates with Cavin1. Among all truncation mutations tested, only mCherry-CAV1 (1-100) was able to co-phase separate with Cavin1. Among CAV1 DR point mutations (M1–M5), mutants M1, M2 and M4 were able to co-phase separate with GFP-Cavin1 while total charge inversion mutant M3 and mutant M5 failed to co-phase separate highlighting the importance of charged residues in CAV1–Cavin1 association. Scale bars = 10 μm; enlarged boxes scale bar = 5 μm.

| **N-Terminal DR** | **OD** | **GSD** | **IMD** | **C-terminal helical domain** |
|-------------------|-------|--------|--------|-----------------------------|
| 1                 |       |        |        |                             |
|                   | 30    | 80     | 100    | 133                         |
| mCherry CAV1 (1-30) |       |        |        |                             |
| mCherry CAV1 (30-80) |       |        |        |                             |
| mCherry CAV1 (1-80) |       |        |        |                             |
| mCherry CAV1 (1-100) |   100  |        |        |                             |

CAV1 (1-100)

WT: M5GGKDYSGLEHLYTPIVRQGN1YKPPNAEMSEKQYVDAHTKEILDVRDPKHLNDVVKIDFDIVIAEPEGTSFDGIKASAFTFTFTKTYWY
M1: M5GGKDYSGLEHLYTPIVRQGN1YKPPNAEMSEKQYVDAHTKEILDVRDPKHLNDVVKIDFDIVIAEPEGTSFDGIKASAFTFTFTKTYWY
M2: M5GGKDYSGLEHLYTPIVRQGN1YKPPNAEMSEKQYVDAHTKEILDVRDPKHLNDVVKIDFDIVIAEPEGTSFDGIKASAFTFTFTKTYWY
M3: M5GGKDYSGLEHLYTPIVRQGN1YKPPNAEMSEKQYVDAHTKEILDVRDPKHLNDVVKIDFDIVIAEPEGTSFDGIKASAFTFTFTKTYWY
M4: M5GGKDYSGLEHLYTPIVRQGN1YKPPNAEMSEKQYVDAHTKEILDVRDPKHLNDVVKIDFDIVIAEPEGTSFDGIKASAFTFTFTKTYWY
M5: M5GGKDYSGLEHLYTPIVRQGN1YKPPNAEMSEKQYVDAHTKEILDVRDPKHLNDVVKIDFDIVIAEPEGTSFDGIKASAFTFTFTKTYWY
Like GFP–Cavin1, unlabelled Cavin1 formed liquid droplets with addition of dextran T-500 (1.25% w/v) observed in bright field image as transparent liquid drops. In the absence of Cavin1 MBP–GBP–CAV1 did not undergo LLPS on its own (Fig. 3b). However, when GFP-labelled MBP–GBP–CAV1 was mixed with Cavin1 it was recruited to Cavin1 liquid droplets (Fig. 3c). Interestingly MBP–GBP–CAV1 appeared to form a shell around the Cavin1 droplets rather than complete co-mixing.

We next probed the mutual roles of Cavin1 and CAV1 disordered sequences in Cavin1–CAV1 co-phase separation. To this end we generated several mCherry-tagged truncation mutants in the region CAV1 (1–100) encompassing the disordered N-termus, oligomerization and scaffolding domains (Fig. 3a). Similar to previous reports we found that mCherry-CAV1(1–100) formed a higher molecular weight oligomer by gel-filtration, while any C-terminal truncations of this sequence resulted in monomeric proteins (Supplementary Fig. 5C).13 Like full-length MBP–GBP–CAV1, the mcherry-CAV1(1–100) sequence was able to undergo LLPS with GFP–Cavin1 droplets, again forming an outer shell around the GFP–Cavin1 droplets (Fig. 3d). In contrast truncated mCherry-CAV1 constructs (1–30), (30–80) and (1–80) were all unable to co-phase separate with GFP–Cavin1, and we also observed similar results using unlabelled Cavin1 (Supplementary Fig. 5D). Interestingly, while mcherry-CAV1(1–100) was able to co-phase separate with the Cavin1 N-terminal deletion GFP–CAV1-ADR1, it did not associate with droplets formed by the C-terminal deletion of GFP–CAV1-ΔDR3 (Fig. 3d) suggesting that the Cavin1 DR3 sequences are essential for CAV1–Cavin1 association.

Sequence alignment of CAV1, CAV2 and CAV3 highlighted several interesting features including an overall conserved but disordered region (30–80) containing two identical motifs, 54RDP56 and 68FEDVIAEP75 (Supplementary Fig. 5A, B). The N-terminal CAV1 disordered region (1–30) however, was not conserved in CAV2 or CAV3. To further pinpoint the sequence requirements of CAV1 and Cavin1 interaction, we made five mutations in mCherry-CAV1(1–100) (Fig. 3a). The two highly conserved motifs 54RDP56 and 68FEDVIAEP75 were mutated to alanine, or random glycine and serine (mutants M1 and M2, respectively). The last three mutants (M3, M4 and M5) replaced charged residues (Glu, Asp, Arg, Lys) with alanine in the entire region CAV1 (1–100) M1, M2 and M4 underwent co-phase separation with GFP-Cavin1 similar to the wild-type CAV1 (1–100), while mutants M3 and M5 failed to associate with Cavin1 droplets (Fig. 3d). These results confirm that the mcherry-CAV1(1–100) interaction with GFP-Cavin1 is highly specific and depends on charged residues within the CAV1 (30–60) region. Overall, these studies indicate that the association between CAV1 and Cavin1 may be driven at least in part by interactions involving liquid phase condensation, with co-mixing mediated by their respective disordered sequences.

Studies of caveolin mutants in cells are typically challenging due to their disrupted trafficking and mis-localisation.24,45–51 Nevertheless, we assessed the localisation of the N-terminal GFP tagged CAV1 mutants (M1–M5) in the context of the full-length protein and in the presence of Cavin1–mCherry in MCF7 cells (Supplementary Fig. 6A). GFP-CAV1-WT showed the familiar punctate distribution in MCF7 cells that co-localised with Cavin1–mCherry, as did the mutant M4. In contrast the GFP-CAV1 mutants M1 and M5 were not associated with mCherry-Cavin1 at the plasma membrane, and the M2 and M3 mutants were either not expressed or rapidly degraded and could not be detected. Comparison with several organelle markers indicated that mutant GFP-CAV1-M1 was mis-trafficked and accumulated in the Golgi, similar to what was seen with the analogous CAV3(R26Q) dystrophic mutation55 (Fig. S6B–E). GFP-CAV1-M5 was mostly mis-localised to lipid droplets, with some diffuse plasma membrane localisation and overlap with endosomes. This phenotype was similar to that observed previously when a putative COPII-binding sequence in the CAV1 N-terminus was mutated (D67G).5 Overall these experiments generally correlate with in vitro studies, where mutations impacting co-phase separation with Cavin1 do not associate with Cavin1 at the cell surface, are unable to form caveolae and are either degraded or mis-localised in cells.

The Cavin1 DR sequences are essential for membrane remodelling in vitro. We previously showed that Cavin1 and Cavin2 possess an intrinsic ability to tubulate artificial lipid membranes using negative stain electron microscopy.11 To examine this membrane remodelling by Cavin1 at higher resolution, we first performed cryoelectron microscopy (cryoEM) analysis of samples after mixing purified Cavin1 with small unilamellar vesicles (SUVs) composed of Folch lipid extracts. We observed formation of an extensive network of membrane tubules (34 ± 5 nm diameter, 12 tubules, and 2 independent experiments) possessing a Cavin1 protein coat using both negative stain electron microscopy and cryoEM (Fig. 4a). Although tubulation of Folch membranes was most efficient, Cavin1 could also tubulate liposomes consisting of PC/PE/PI(4,5)P2 (Supplementary Fig. 7A). In addition, Cavin1-GFP expressed and purified from HEK293 cells could also tubulate Folch membranes similarly to the bacterially expressed protein (Supplementary Fig. 7B). Examination of the Cavin1-coated tubules by cryoelectron tomography (cryoET) revealed a striated but relatively heterogeneous pattern of protein densities around the tubules (Fig. 4b and Supplementary movie 1). These are similar to structures previously observed on the cytosolic face of caveolae using fast-freeze deep-etch52,53 and conventional EM methods54–56, and with the elongated rod-like structures of isolated Cavin1 observed by negative staining EM11. These experiments indicate that Cavin1 possesses an inherent membrane remodelling activity, driven by large scale oligomeric assembly on the membrane surface.

The importance of Cavin1 DR domains in self-association and LLPS raised the question as to whether they play a role in its ability to physically remodel membranes. To this end, we used the in vitro membrane remodelling assay to investigate their importance in generating membrane curvature. We expressed and purified a range of Cavin1 DR domain truncations with an N-terminal His-ubiquitin (HisUb) tag (Fig. 4c and Supplementary Fig. 2), and used the membrane tubulation assay combined with negative stain EM to analyse their ability to remodel mammalian (Folch) synthetic phospholipid membranes (Fig. 4d). Complete removal of either the N-terminal DR1 or C-terminal DR3 domains abolished the ability of Cavin1 to tubulate liposomes in vitro. Shorter truncations showed that while the N-terminal DR1 deletion Cavin1 (10–392) still formed membrane tubules, these were relatively infrequent and of a smaller diameter (~10 nm), whereas further deletion of N-terminal DR1 sequences in Cavin1 (30–392) prevented the formation of membrane tubules altogether. The C-terminal DR3 deletion mutant Cavin1 (1–345) formed membrane tubules similar to full length Cavin1. However, the deletion of further amino acids from the
C-terminus in Cavin1 (1–330) completely inhibited membrane tubulation. For those DR truncation mutants that lacked membrane remodelling activity we observed instead a propensity to cause liposome clustering. This likely occurs because these Cavin1 constructs can now bind adjacent phospholipid vesicles via multiple positively charged surfaces of the HR1 and HR2 domains, unrestrained by compensating negatively-charged DR1 and DR3 sequences. Overall, these studies define a core Cavin1 sequence (10–345) required for Cavin1 to efficiently promote membrane curvature.

We next examined the ability of purified GFP-Cavin1 to modulate Folch lipid giant multilamellar vesicles (GMVs) doped with 0.1 mol% fluorescent Bodipy-TMR PI(4,5)P2 analogue. GFP-Cavin1 showed strong localised clustering at the membrane surface compared to other membrane remodelling proteins (Fig. 5a) and possessed a remarkable membrane sculpting activity as indicated by the rapid collapse of GMVs over a period of several minutes (Fig. 5b). We performed similar experiments with Rhodamine B-PE as a fluorescent marker and observed the same protein clustering and membrane sculpting activity (Supplementary Fig. 7C). In contrast, although both the N-terminal and C-terminal DR deletions of Cavin1 still bound efficiently to GMVs, they did not display any significant membrane sculpting activity (Fig. 5c, d). With Cavin1-ΔDR1 we also often observed a characteristic accumulation of the protein at the interface between adjoining vesicles leading to the clustering of the GMVs (Fig. 5e). Overall, these studies using GMVs and SUVs show that while the DR1 and DR3 domains are dispensable for membrane binding, they have an essential role in the ability of Cavin1 to sculpt the curvature of phospholipid membranes.

Cavin1 disordered sequences are essential for interacting with CAV1 and forming caveolae. Our studies in vitro highlight several properties of Cavin1 that are dependent on its disordered sequences. Firstly, DR1 and DR3 of Cavin1 are important for the formation of a large scale associated state and LLPS promoted by electrostatic interactions, and the DR1 domain is required for the dynamic properties of Cavin1 in LLPS; removal of the DR1 domain results in gel formation and prevents its free diffusion within the condensates. The Cavin1-ΔDR1 construct also displays a capacity to bind and cluster membrane vesicles in vitro. Secondly, minimal sequences of Cavin1 DR1 and DR3 are required for membrane remodelling. Lastly, the C-terminal DR3 domain of Cavin1 is required for the association with CAV1 in co-mixed liquid droplets in vitro.

To examine the importance of Cavin1 disordered N-terminal and C-terminal domains to functional caveola formation, we next analysed the localisation of the DR1 and DR3 truncation mutants in cells using either standard confocal microscopy (Supplementary Fig. 8A) or confocal fluorescence with Airyscan super-resolution imaging (Fig. 6a). The prostate cancer PC3 cell line was used, which expresses CAV1 but does not express any members of the Cavin family so that CAV1 is diffusely localised at the plasma membrane (Supplementary Fig. 8A). Expression of full-length GFP-Cavin1 in PC3 cells fully restores the formation of caveolae with CAV1 (in the absence of other cavins), providing a functional readout for Cavin1 activity (Fig. 6a). Full length GFP-Cavin1 showed a characteristic punctate distribution and co-localised with CAV1 at the plasma membrane (Supplementary Figs. 6A and 8A). In contrast, after removal of the C-terminal domain GFP-Cavin1-ΔDR3 is unable to promote caveola formation, does not co-localise with CAV1, and now...
associates extensively with microtubules. This is consistent with a previous report of a similar C-terminal truncated Cavin1 (residues 1–322) in CHO cells.60 FRAP analysis of GFP-Cavin1-ΔDR3 on microtubules showed a fast fluorescence recovery, indicating a dynamic exchange with the cytoplasm or diffusion along the microtubules (Fig. 6b and Supplementary movie 2). Interestingly, when microtubules were depolymerised with nocodazole this resulted in redistribution and condensation.

Fig. 5 Removing the Cavin1 DR domains prevents deformation of GMV membranes. a Purified Ub-tagged and GFP-tagged Cavin1 shows strong localised clustering on the surface of Folch giant multilamellar vesicles (GMV) containing Bodipy-TMR-labelled PI(4,5)P2 (0.1 mol%). Scale bar = 5 μm Cavin1 (b), Cavin1-ΔDR3 (c) or Cavin1-ΔDR1 (d) were incubated with Folch GMVs containing Bodipy-TMR-labelled PI(4,5)P2 (0.1 mol%), allowed to settle on glass coverslips and images were acquired one frame per second. Frame numbers are indicated in PI(4,5)P2 channel (red). e GMVs incubated with Cavin1-ΔDR1 were often observed to be tethered to each other with Cavin1-ΔDR1 concentrated at the contact sites. Scale bar = 10 μm.
Fig. 6 The Cavin1 DR domains are essential for caveola formation with CAV1. a GFP-tagged Cavin1 and truncations (green) were expressed for 24 h in PC3 cells, fixed and immunolabelled for Caveolin1 (CAV1) (red). Full length Cavin1 forms typical caveola puncta, colocalising with CAV1 at the cell surface. Cavin1-ΔDR1 mutant expression leads to formation of tethered intracellular CAV1-positive clusters. Cavin1-ΔDR3 shows cytoplasmic and microtubule localisation. Images were collected using a Zeiss fast Airyscan microscope. Scale bar = 10 μm; enlarged boxes scale bar — 5 μm. Fluorescence recovery after photobleaching (FRAP) analysis of GFP-Cavin1-ΔDR3 before (b) and after (c) nocodazole (10 μM) addition. Scale bar — 5 μm. d In PC3 cells GFP-tagged Cavin1-ΔDR1 truncation shows colocalization with the early endosomal marker (EEA1) (red) and CAV1 (blue). Inset shows merge images of GFP-Cavin1-ΔDR1/EEA1 and GFP-Cavin1-ΔDR1/CAV1. Scale bar = 10 μm; enlarged boxes scale bar — 5 μm. e GFP-tagged Cavin1 and Cavin1-ΔDR1 were visualised in PC3 cells by electron microscopy and labelling of GFP tagged proteins using APEX-GBP staining. Scale bar = 1 and 2 μm. Enlarged boxes scale bar — 500 nm and 1 μm. f Live imaging of PC3 cells expressing Rab5a-mCherry with either GFP-Cavin1 or GFP-Cavin1-ΔDR1. Images were acquired one frame per 4 s and frame numbers are indicated in boxes. Arrows indicate mCherry/GFP signal co-localisation or separation event. Scale bar — 2 μm.
of GFP-Cavin1-ΔDR3 to form spherical droplets in the cytosol (Fig. 6c and Supplementary Fig. 8B and Supplementary movie 3). These also showed fast exchange of protein molecules with the bulk cytoplasm suggestive of liquid-droplet behaviour, and consistent with droplet formation by Cavin1-ΔDR3 in vitro. This indicates a dynamic equilibrium exists between cytosolic, liquid droplet and microtubule-associated states of the GFP-Cavin1-ΔDR3 truncation.

Strikingly, expression of GFP-Cavin1-ΔDR1 resulted in the formation of large intracellular structures that also contained endogenous CAV1 (Fig. 6a and Supplementary Fig. 8A). A C-terminal tagged Cavin1-ΔDR1-GFP construct showed similar clusters co-localised with CAV1, confirming this phenotype is not influenced by the location of the GFP tag (Supplementary Fig. 8C). To analyse these structures in more detail, we performed co-localisation experiments of GFP-Cavin1-ΔDR1 with various cellular markers. While no overlap was seen with the Golgi complex, lysosomal or recycling endosomal membrane markers, a significant proportion of GFP-Cavin1-ΔDR1 and endogenous CAV1 were found to co-localise with the early endosomal marker EEA1 (Supplementary Fig. 9A). Airyscan microscopy revealed that clusters of EEA1-positive endosomes surrounded the GFP-Cavin1-ΔDR1 and CAV1-positive structures (Fig. 6d and Supplementary Fig. 9B). We then performed transferrin uptake assays in PC3 cells using transferrin labelled with Alex-488 fluorescent dye. Transferrin positive endosomes showed little overlap with full-length mCherry-Cavin1-positive spots on the cell surface (Supplementary Fig. 10). However, the mCherry-Cavin1-ΔDR1 construct formed intracellular clusters with transferrin positive endosomes surrounding them similar to EEA1. These large intracellular structures were visualised by APEX labelling and electron microscopy imaging61 of GFP-Cavin1-ΔDR1 in PC3 cells, revealing intracellular assemblies consisting of large clusters of vesicles with a surrounding halo of GFP-Cavin1-ΔDR1 labelling (Fig. 6e). In contrast, GFP-Cavin1 expression resulted in formation of the characteristic single caveolae and rosettes of caveolae at the plasma membrane as expected (Fig. 6e).

We lastly performed live imaging of PC3 cells expressing either GFP-Cavin1 or GFP-Cavin1-ΔDR1 with Rab5a-mCherry as a marker of early endosomes. Caveolae are consistently localised to the trailing edge of migrating cells, where constant membrane remodelling events are occurring62. In migrating PC3 cells we observe dynamic GFP-Cavin1 positive caveola puncta undergoing transient fission and fusion events and kiss-and-run interactions with Rab5a-mCherry positive endosomes similar to previous observations63 (Fig. 6f and Supplementary movie 4). In contrast, GFP-Cavin1-ΔDR1 initially showed plasma membrane puncta fusion events similar to GFP-Cavin1 (imaged at an early 12 h time point following transfection before larger immobole condensates are formed), but over time resulted in formation of the larger structures that stably associated with Rab5a positive endosomes (Fig. 6f and Supplementary movie 5). This suggests that the DR1 domain is important for the dynamics of intracellular trafficking and recycling of caveolae at endosomes. Overall, our results show that disordered sequences of Cavin1 are essential for generating caveolae, but that each DR domain has a distinct function. Removing the C-terminal DR3 domain prevents interaction with CAV1 and results in mis-localisation to the cytoplasm and abnormal association with microtubules. Removing the N-terminal DR1, which results in gel formation and membrane clustering in vitro, allows initial caveola formation with CAV1 at the plasma membrane, but then causes subsequent accumulation of aberrant intracellular protein and membrane assemblies with a subset of early endosomes unable to recycle to the plasma membrane.

Minimal Cavin1 DR sequences needed for membrane remodelling are also essential for caveola formation. Using the series of truncations tested in vitro for membrane remodelling activity, we next asked if the same minimal sequences are sufficient for caveola formation in cells. GFP-Cavin1(10–392) showed a relatively normal localisation with CAV1 puncta at the cell surface. However, GFP-Cavin1(30–392) formed large intracellular puncta and clusters that co-localised with CAV1 (Fig. 7a), and also showed a partial co-localisation with EEA1 (Supplementary Fig. 9C), similar to Cavin1 with the complete DR1 domain removed. Thus, deletion of the N-terminal DR1 sequence of Cavin1 has a progressive effect on the re-distribution of caveolae from the plasma membrane to intracellular endocytic compartments.

The C-terminal DR3 truncation GFP-Cavin1(1–345) retained a normal ability to generate plasma membrane puncta that co-localised with CAV1 (Fig. 7a), and APEX labelling and electron microscopy of GFP-Cavin1(1–345) showed its typical localisation to caveolae at the plasma membrane (Fig. 7b). Further deletion of C-terminal DR3 sequences in GFP-Cavin1(1–330), however, resulted in a total cytosolic redistribution. The C-terminal truncations show that amino acids (346–392) are dispensable for generating caveolae in PC3 cells, while residues 330–345 are essential. Finally, we used a proximity ligation assay (PLA)10 to assess the interactions of Cavin1 C-terminal truncations with CAV1 at the plasma membrane. PLA analyses correlated with the cellular imaging of the GFP constructs, showing that the mutant Cavin1(1–345) can interact with (or is at least in close proximity to) CAV1, while the shorter truncations Cavin1(1–330) and Cavin1(1–310) do not (Fig. 7c and Supplementary Fig. 11).

Specific DR sequences are essential for the ability of Cavin1 to form caveolae. The disordered sequences 1–30 and 310–345 in DR1 and DR3 are required for Cavin1 to efficiently self-associate, remodel synthetic phospholipid membranes in vitro, and promote caveola formation with CAV1 in cells. To examine these sequences in more detail we generated a series of specific mutations in the DR1, DR2 and DR3 domains in the context of the minimal functional construct Cavin1(1–345) (Fig. 8a). Beginning with DR1 (residues 1–30), we first tested whether the acidic amino acids were important by mutating the Glu/Asp residues to alanine (DR1mut1). When GFP-tagged Cavin1(1–345) DR1mut1 was expressed in PC3 cells it formed large intracellular puncta that co-localised with CAV1 (Fig. 8b), and also colocalised with a subpopulation of EEA1-positive endosomes, but not LAMP1 or GM130 (Supplementary Fig. 12A). APEX labelling and imaging by EM showed clusters of GFP-Cavin1(1–345) DR1mut1 that appeared identical to those formed by either GFP-Cavin1-ΔDR1 or GFP-Cavin1(30–392) (Supplementary Fig. 12B). By FCS, this variant showed a significant increase in diffusivity with respect to wild-type Cavin1(1–345), indicating that its net negative charge is important for self-association (Supplementary Fig. 12C). More precise mutation of Asp/Glu residues in the first ten amino acids of DR1 (DR1mut2) had no qualitative effect on the ability of GFP-Cavin1(1–345) to form caveolae, while altering the Asp/Glu residues in amino acids 10–30 of DR1 (DR1mut3) resulted in the same phenotype as mutant DR1mut1 (or complete deletion of DR1), forming large intracellular clusters with CAV1 (Fig. 8a, b). We next substituted DR1(1–30) with random Gly/Ser sequences, while maintaining the relative positions of acidic Asp/Glu residues and prolines (DR1mut4). The objective was to determine if any other sequences apart from the acidic side chains contributed to the activity of the domain. In MCF7 cell lysates the DR1mut4 mutant did not show a major difference in diffusivity by FCS compared to wild-type Cavin1(1–345) indicating that only the
acidic side-chains in the DR1 region are necessary for self-association (Supplementary Fig. 12C). The subcellular localisation of GFP-Cavin1(1–345) mutant DR1mut4 in PC3 cells also showed co-localisation with CAV1 at the plasma membrane (Fig. 8b), indicating that it is the electrostatic properties of the DR1 sequence that are most important for its function and not the specific sequence itself. However, the spacing of acidic residues in DR1 is critical, as complete removal of surrounding sequences (DR1mut5) also results in GFP-Cavin1(1–345) mislocalisation. An analogous result was observed for the central DR2 domain of Cavin1, where mutation of the acidic residues (DR2mut6) abolished caveola recruitment in PC3 cells and prevented self-association in MCF7 cell lysates, but altering the surrounding sequences while maintaining negative charges had no effect on caveola formation (DR2mut7) (Fig. 8c and Supplementary Fig. 12C). Thus, the presence and the spacing of acidic sequences in DR1 and DR2 are essential for normal caveola formation, but their specific surrounding sequences are not.

Lastly, we assessed the roles of specific sequences in the essential DR3 region (residues 311–345). The mutation of acidic Asp/Glu residues in GFP-tagged Cavin1(1–345) (DR3mut8) resulted in a diffuse cytosolic localisation in PC3 cells (Fig. 8d) and prevented self-association in FCS measurements (Supplementary Fig. 12C). These results show that the acidic Glu/Asp residues in the Cavin1 DR1, DR2 and DR3 domains are all essential for oligomeric interactions and forming caveolae with CAV1 at the cell surface. The acidic side-chain mutations result in identical phenotypes to the complete truncation of the DR1 and DR3 domains. In contrast to the DR1 and DR2 domains however, we found that altering everything in DR3 other than Asp/Glu residues (DR3mut9) resulted in a protein with a normal ability to self-associate (Supplementary Fig. 12C), but that was unable to restore caveola formation with CAV1 in PC3 cells (Fig. 8d). This protein was generally cytosolic, but in some cells we observed the formation of numerous spherical cytoplasmic structures, that dynamically exchange with the cytosol as shown by FRAP analysis and regularly undergo fusion, suggesting the protein has undergone LLPS and droplet formation (Fig. 8e and Supplementary Fig. 12D and Supplementary movie 6). Remarkably however, unlike the complete DR3 deletion purified Cavin1(1–345) DR3mut9 is still able to remodel and tubulate synthetic liposomes in vitro (Supplementary Fig. 12E). This shows that while specific sequences in the Cavin1 region 310–345 are dispensable for large scale oligomer formation, LLPS and membrane remodelling, they are still essential for recruitment to caveolae with CAV1 in cells. The acidic side chains in this region, however, are required for all of these functional Cavin1 activities (mutant DR3mut8). To refine this further, we designed three shorter variants of the DR3mut9 mutation, DR3mut10 (311–320), DR3mut11 (321–331), and DR3mut12 (332–345). While mutant DR3mut12 behaved like wild-type Cavin1 and formed normal caveolae, both mutant DR3mut10 and DR3mut11 showed a cytosolic distribution similar to DR3mut9 (Fig. 8d). Therefore, specific sequences in the Cavin1 DR3 region

Fig. 7 Definition of the minimal DR sequences required for Cavin1 function. a GFP-tagged Cavin1 DR domain truncation mutants (green) (Fig. 4c) were expressed in PC3 cells and immunolabelled with CAV1 (red). Residues 1–30 in DR1 and 330–345 are required for caveola formation. Scale bar = 10 μm; enlarged proximity ligation assay (PLA) analyses show that truncation of Cavin1 from the C-terminus beyond residue 345 results in loss of association with CAV1. PLA signal was quantified as dots per cell for specific interaction between GFP-tagged proteins and CAV1, n = 33 over two independent experiments. Error bars indicate mean ± SD, ***P < 0.001. Source data for Fig. 7c are provided as a Source Data file.
311–331 are required for CAV1 association and caveola formation, while acidic residues within DR3 region (332–345) are essential for promoting electrostatic oligomeric Cavin1 assembly. Lastly, we quantified the co-localisation of those Cavin1 DR mutants that still retained prominent association with CAV1 (Fig. S12F). While the sequences altered in these constructs are not strictly required for caveola formation (e.g., DR1mut2 or DR3mut12) or CAV1 interaction (e.g., DR1mut1, DR2mut1, DR2mut2, DR3mut1, DR3mut2, DR3mut3, DR3mut4, DR3mut5, DR3mut6, DR3mut7, DR3mut8, DR3mut9, DR3mut10, DR3mut11, DR3mut12).

Fig. 8 Sequence requirements of the Cavin1 DR domains in caveola assembly. a Schematic diagram of Cavin1(1–345) with the sequences of the various point mutants indicated. DR1, DR2 and DR3 mutations occur in the regions 1–30, 161–197 and 311–345 respectively. b GFP-tagged Cavin1(1–345) DR1 domain mutants (green), c DR2 domain mutants and d DR3 domain mutants expressed in PC3 cells and immunolabelled with endogenous CAV1 (red). Images in b–d were by Airyscan confocal microscopy. Scale bar = 10 μm; enlarged boxes scale bar = 5 μm. e FRAP analysis of Cavin1 (1–345) DR3mut9 mutant showing fast recovery of fluorescence in cytosolic droplets and also droplet fusion events (marked by arrow). Scale bar = 10 μm; enlarged boxes scale bar = 5 μm.
DR1mut3 or DR1mut5), they all showed a marginal reduction in co-localisation suggesting they make a minor contribution to Cavin1-CAV1 interactions. Altogether, these studies demonstrate the critical importance of acidic residues in all three DR domains for promoting electrostatic intermolecular interactions and caveola formation; while specific sequences in Cavin1 DR3 region (311–331) are necessary for Cavin1 and CAV1 association for caveola recruitment.

Discussion

Despite the fact that intrinsically disordered sequences are a prominent and highly conserved feature of all cavins, no previous studies have explicitly addressed their functional importance. We now show that they are essential for caveola formation. In addition, they also regulate the ability of Cavin1 to self-associate and undergo LLPS in vitro, where Cavin1 shows the classical properties of LLPS as demonstrated by phase separation that is sensitive to protein concentration, ionic strength, molecular crowding agents, and by the rapid exchange of protein in Cavin1 droplets as shown by FRAP. The sensitivity of this LLPS to salt concentration indicates an electrostatically driven Cavin1 condensation. We demonstrate the distinct roles of the disordered DR domains of Cavin1 in LLPS behaviour, including a mutant protein lacking the DR1 domain that still self-associates but no longer shows the dynamic exchange properties of the full-length protein. In addition, CAV1 was also able to associate with Cavin1 generated liquid droplets, an interaction that is dependent on their mutual disordered sequences. Our cellular studies show that acidic residues in all three Cavin1 disordered sequences (DR1, DR2 and DR3) are essential for generating caveolae with CAV1 at the plasma membrane. Deletion or mutation of these regions in Cavin1 result in mislocalisation and an inability to form plasma membrane caveola invaginations. Interestingly the N-terminal and C-terminal sequences play divergent roles in this process. Deletion of the N-terminal DR1 domain affects caveola dynamics and leads to the formation of large intracellular clusters of Cavin1, CAV1, and endosomal membrane vesicles. In contrast, deletion of the DR3 domain prevents CAV1 association in vitro and in vivo and results in dynamic microtubule association or cytoplasmic droplet formation. We speculate that Cavin1-ΔDR3 association with microtubules may share mechanistic similarities with the condensation of Tau on microtubules, or interactions of multivalent positively-charged peptides with the C-terminal acidic tails of tubulin subunits, but this will require further study.

To better appreciate and visualise the role of the disordered DR domains in Cavin1 activity, we constructed a theoretical structural model of the protein, building on the assumption that the functional Cavin1 unit is a homotrimer based on the coiled-coil structure of its N-terminal HR1 domain (see “Materials and methods” section) (Fig. 9a). This model points to several interesting features of the Cavin1 protein. Firstly, the combined DR1, DR2 and DR3 domains account for more than 50% of the total Cavin1 sequence. In other words, Cavin1 is not a typical globular protein but rather consists of large random-coil elements tethered by α-helical structural cores. Secondly, as suggested by sequence analyses (Supplementary Fig. 1), there is a distinctive electrostatic pattern to the structure, with the α-helical domains providing positively charged surfaces for membrane association, and the disordered regions having a generally negatively charged nature. A likely consequence of this is that electrostatic repulsion will cause these DR domains to orient outwards when Cavin1 is in contact with membranes, and we propose they will also form transient electrostatic interactions with the HR domains of neighbouring Cavin1 molecules (Fig. 9b). Notably, multiple theoretical and experimental studies have shown that the sequence-specific electrostatically driven interactions between disordered proteins can lead to LLPS and high affinity protein complex formation under physiological conditions, with the tendency to phase separate (or undergo ‘complex coacervation’) increasing as the ‘blockiness’ of the charge distribution increases.

Our studies of the DR domains of Cavin1 confirm that the acidic residues within these domains are essential for the formation of caveolae in cells and promotion of membrane remodelling in vitro. By what mechanism might the DR domains contribute to these membrane sculpting activities? Several recent studies have demonstrated the ability of intrinsically disordered sequences to generate membrane curvature when coupled to membrane binding domains. This is caused by molecular crowding of the disordered sequences leading the proteins to partition with curved or convex membranes so as to increase their conformational entropy; and this can also be enhanced by electrostatic repulsive forces both between the disordered domains and with the membrane itself (Fig. 8). One possible mechanism we can propose for Cavin1-driven membrane curvature is that negatively charged DR sequences and positively charged HR regions of Cavin1 combine to promote self-association, membrane interaction and protein crowding at the membrane surface leading to subsequent membrane bending. In the absence of CAV1 and at high protein concentrations in vitro, or under certain conditions in cells, Cavin1 can generate arrays of protein oligomers to form membrane tubules. Under normal conditions however, the process of generating membrane curvature is tightly regulated by CAV1, EHD2 and Pacsin2, and also specific membrane lipids, to restrict Cavin1 remodelling activity only to caveolae. We see an almost complete correlation between the ability of different Cavin1 truncations and mutants to tubulate membranes in vitro and the ability to form caveolae in situ. The notable exception to this is that alteration of sequences in the DR3 region 310–331 does not affect the ability of Cavin1 to assemble into oligomers and efficiently tubulate synthetic membranes, but still results in a failure to generate caveolae in cells. This implies these specific sequences in the Cavin1 DR3 region are additionally required for Cavin1 recruitment to CAV1-positive membrane domains through interactions with the disordered CAV1 N-terminus.

A second mechanism for membrane curvature suggested by our results (and not mutually exclusive with a role for molecular crowding) is the formation of phase-separated Cavin1 domains that incorporate membrane-embedded CAV1. Intrinsically disordered regions of proteins have gained significant attention for their ability to promote LLPS, or biomolecular condensation, with important biological functions such as stress granule formation, assembly of nuclear substructures and sensing changes in cellular homeostasis. The plasma membrane and surfaces of intracellular compartments including the ER and lysosomes have been found to play a role in LLPS, acting as sites of droplet nucleation or as platforms for transport of phase-separated assemblies for example. It has also recently been proposed that biomolecular condensates associated with phospholipid membranes might possess emergent mechanical properties that can result in membrane curvature generation. This is depicted in schematic form in Fig. 9d. Here, we have shown for the first time that purified Cavin1 can readily undergo LLPS under near physiological conditions and is able to recruit CAV1 through interactions involving LLPS. The DR1 and DR3 domains contribute to this process, although neither domain is strictly essential. Indeed, mutations in DR3 that maintain its negative charge but prevent CAV1 interaction at the plasma membrane actually promote GFP-Cavin1 liquid droplet formation in cells. Notably, removal of
the DR1 domain results in apparent gel formation rather than liquid droplet assembly in vitro, and within cells results in a striking accumulation of large intracellular structures that also contain CAV1. These are formed by endocytic redistribution of caveola structures from the cell surface and accumulation with early endosomal membranes. Caveolae, positive for both CAV1 and Cavin1, have been shown to bud from the plasma membrane and fuse with early endosomal compartments, and this would almost certainly require dynamic remodelling of the protein coat to allow the fusion process to occur. We postulate that the intracellular structures we observe with Cavin1-ΔDR1 are formed by internalised caveolae, which have become trapped during the stage of early endosomal fusion. This may be due to the DR1-truncated Cavin1 being unable to undergo normal dynamic exchange, as suggested by its gel-forming properties and its propensity to cluster membrane vesicles, causing inhibition or slowing of the docking and fusion with the early endosome in a process involving EEA1 and Rab5α.

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**Fig. 9 Model for the role of Cavin1 DR domains in LLPS and caveola formation.**

*a* Structural model of a Cavin1 homotrimeric assembly. The trimeric HR1 coiled-coil domain is derived from the crystal structure of the mouse Cavin1 HR1 domain, the UC1 and HR2 domains are modelled as described previously, and the DR domains are modelled as random coil structures (see “Methods” section for further details). The structure is shown in ribbon diagram (top) and with an electrostatic surface representation (bottom). 

*b* Proposed orientation of Cavin1 proteins on the membrane surface, with membrane-binding HR1 and HR2 domains associated with the phospholipid bilayer and negatively charged DR sequences directed outwards due to electrostatic repulsion.

*c* Potential role of Cavin1 disordered sequences in membrane curvature generation due to steric crowding. This concept is largely derived from previous studies of other membrane-associated proteins.

*d* Potential role of Cavin1 fuzzy interactions and LLPS in membrane curvature generation, CAV1 interaction and caveola formation.
Our data indicates that the assembly of caveolae by Cavin1 strictly depends on a ‘fuzzy’ network of interactions promoted by electrostatic associations, with an essential role for the intrinsically disordered DR domains of Cavin1 in self-association, CAV1 interaction, membrane remodelling and ultimately caveola formation. Fuzzy interactions are defined broadly as those that involve dynamic, exchanging, multivalent interactions with varying degrees of protein disorder or structural ambiguity83–86. This provides versatility and reversibility in protein–protein interactions, and such fuzzy interactions are also proposed to be a driver of protein phase transitions19. One of the historically consistent observations regarding caveolae is that they do not possess an obvious or highly ordered coat morphology akin to clathrin or COP-coated vesicles. In previous studies of caveola architecture it is notable that while some recurring structures are observed, the general appearance of the caveola surface is highly heterogeneous83–86. Our model for caveola assembly and structure differs markedly from other classical membrane coats such as clathrin or COP1 and COP2, which are built from symmetrical arrays of structured protein interactions. While structural elements of cavins and caveolins will likely produce semi-segmental spacings between the building blocks, the flexible nature of the disordered domains that provide the ‘glue’ for caveola assembly mean that the overall organisation of the coat will be highly dynamic. Caveola formation is the result of multiple low affinity fuzzy interactions between Cavin1, CAV1 and membrane lipids, and we propose that this leads to a metastability in caveola structure that is important in both the dynamic cycling of caveolae through the endocytic pathway and also for their ability to respond to stresses by rapid disassembly.

Fluorescence correlation spectroscopy (FCS) analysis. FCS analysis was performed on a Zeiss LSM 710 confocal microscope equipped with 405/1.2W C-Apochromat and twin Big GaAsP detectors capable of single molecule detection. Briefly, MCF7 cells were lysed in a buffer (200 μl) containing 25 mM Tris (pH 7.4), 150 mM NaCl while passing through a 27G needle six times. Lysates were then used for FCS analysis. Purified tagged cavin proteins were prepared for FCS by dilution of respective stock solutions of cavin truncates in either 500GF or 150GF buffer to achieve 0.1 μM protein concentration with subsequent centrifugation at 17,000 × g for 10 min. At the beginning of each FCS session on a Zeiss LSM 710, pinhole size was optimised to BODIPY-FL maleimide DIBO (~37 kDa) and TAMRA DIBO (~37 kDa) were measured for each measurement was exported from the Zeiss analysis programme and plotted for all constructs in Graph pad Prism software. Hydrodynamic radius calculations were done using Stokes–Einstein equation with basic assumption of perfect spherical object diffusion. Stokes–Einstein equation: \(D = \frac{K_BT}{6\pi
u r} \), where \(K_B\) is the Boltzmann constant, \(T\) is the temperature (298 K), \(\nu\) is the fluid viscosity (Pa·s) and \(r\) is the hydrodynamic (Stokes) radius of spherical particle.

In vitro phase separation assays. Purified GFP-tagged Cavin1 proteins, or mixtures of mCherry-Cavin1 and GFP-Cavin1 proteins, were diluted to respective protein and/or salt concentrations prior to addition of dextran T-500 (Pharmacia). Dextran solution was added on the top of protein solution without any mixing to allow natural diffusion of dextran. Image acquisition and fluorescence recovery after photobleaching (FRAP) assays were performed after 2 min wait period to allow phase separated droplets to settle. Phase separation analysis was done within 10 min post addition of dextran. Non-bleaching image acquisition conditions were established before performing FRAP assay. FRAP analysis was done by bleaching rectangular area (2 μm × 1 μm approximately) within protein droplet using 488 nm Argon laser and subsequent image acquisition was done one frame per second. Recovering curves were normalized to the maximum fluorescence bleaching correction using formula \(F(t) = \frac{F_{immediate}}{F_{postbleach}}\). Normalised data points were used to perform non-linear

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exponential recovery fit using equations within ImageJ 1.50 g or Prism version 8 to obtain half-life value for fluorescence recovery of respective protein.

Liposome preparation and in vitro membrane tubulation assay. Liposomes were prepared by mixing of 10 μ 1 mL mm stock solution of Folch lipids (bovine brain extract lipid—Folch I fraction Sigma Aldrich B1502) with 50 μ 1 mL chloroform in a round-bottom flask. The mixtures were dried gently by a stream of nitrogen first and under vacuum overnight thereafter. Liposomes were rehydrated in 500 μL 150GFP buffer, followed by repetitive freeze-thaw cycles for 3–5 times, and then spread on a mixture of dry ice and acetone followed with 60 °C water. The liposomes were then extruded through a 400-nm polycarbonate membrane 21 times using an Avanti mini-extru-grator to generate large unilamellar lipid vesicles (LUVs).

A 5-μl volume of purified Cav1 variants [0.1 mg/mL (1.5–2 μM)] was mixed with 5 μl 200 μM liposomes for 1–3 min at room temperature. Samples were then quickly spotted onto formvar-carbon coated electron microscopy grids (Cu/Pd grids 200 mesh hexagonal – PreciSciTech - GPC-PD200H) for 10 s and excess samples were removed by blotting at corner using Whatman filter paper. This is followed by 2–3 distilled water washes in similar fashion and subsequent application of 1% uranyl acetate stain. The excess of stain was removed by blotting and grids were allowed to air dry for a while before viewing under the electron microscope. Final images were acquired on JEOL 1011 electron microscope at 80 kV.

Giant multilamellar vesicle (GMV) experiments. GMV were prepared using electroformation method described before90. Briefly, lipids mixture dissolved in chloroform/methanol solution was gently applied to indium-tin-oxide coated glass slide (Sigma-Aldrich Cat. No. D63908) as multiple layers. This solution was then dried under constant stream of nitrogen to remove organic solvent and further dried under vacuum overnight. Next day, electroformation was performed at 50 °C in 150GF buffer for 1 h. Vesicles were used immediately for experiments.

Cryoelectron microscopy/tomography of Cav1-coated membrane tubules. Liposome tubulation reaction was assembled as described in the previous section and subsequent to centrifugation after a 1.3-min incubation period, for vitrification the sample was applied to Lacey carbon grids (EMS, Hatfield, PA, USA) using a Vitrobot Mark II (FEI, Eindhoven, NL) plunge freezer with 4 μl of sample, 6 s blotting time and a −3 mm offset at 24 °C and 100% humidity. Images were collected on a Tecnai G2 F30 TEM (FEI, Eindhoven, NL) operated at 300 kV at a magnification of ×85000, with 5 μm defocus on a Gatan K2 summit camera in counting mode for a final pixel size of 3.5 Å per pixel. Images were processed in either IMOD (version 4.9) or ImageJ. Tilt-series were acquired on a Talos Arctica TEM (Thermo Fisher Scientific- FEI, Eindhoven, NL) operated at 200 kV and at a magnification of ×45000 (final pixel size 3.1 Å per pixel). Images were recorded using the microscope software Tomography (Thermo Fisher Scientific- FEI, NL) on Falcon 3 (Thermo Fisher Scientific- FEI, NL) camera operated in counting mode at an angular range of −60° to 60° in a bidirectional fashion and at an angular increment of 2°. The defocus was set to −5 μm. Unbinned movies of eight frames with a set dose rate of 1.7 e Å−2 frame−1 were acquired and then subjected to DPPE-cholesterol (Sigma-Aldrich Cat. No. D3405) reaction was then performed as follows. Briefly, cells were washed with DAB/ cadaclyte mixture (DAB final concentration—1 mg/ml for 2 min, then treated with DAB/cadaclyte + 5.88 mM H2O2 (hydrogen peroxide, Sigma-Aldrich Cat. No. H31009) for 20 min. Cells were then washed with 0.1 M sodium cadaclyte buffer and contrasted with 1% osmium tetroxide for 2 min. Cells were then embedded in LX112 resin and thin sections were cut as described previously81. Images were acquired on JEOL 1011 electron microscope fitted with a Morada CCD camera (Olympus) under the control of ITEM software and operated at 80 kV.

Immunofluorescence analysis, live cell imaging and proximity ligation assay (PLA). PC3 cells were grown at about ~50% confluence in RPMI 1640 medium supplemented with 10% FBS. Cells were then transfected with respective Cav1 variant mutants using Lipofectamine 3000 (Invitrogen) as per manufacturer’s instructions. Cells were fixed 24 h after transfection with 4% paraformaldehyde in phosphate-buffered saline (PBS) at 4 °C and subsequently permeabilised with 0.1% Triton X-100 in PBS for 7 min. Cells were probed with CAV1 antibody (1:500 dilution) and anti-Rabbit secondary antibody Alexa Fluor® 561 conjugate (Dilution 1:400). For Transferrin uptake assays, PC3 cells expressing either GFP-Cavin1 or GFP-Cavin1–ΔDR1 were incubated with transferrin labelled with Alexa-488 (5 μg/ml) for 1 h at 37 °C. Cells were then washed three times with ice cold PBS and were subsequently fixed with 4% paraformaldehyde in PBS for all experiments except live imaging. Cholesterol addition experiments were performed in MCF7 cells expressing GFP tagged Cavin1 with serum starvation (Serum free DMEM + 1% BSA, 1 h) prior to the addition of water soluble analogue of Cholesterol (Sigma-Aldrich Cat. No. C4951). Cells were incubated in DMEM media containing Cholesterol for 40 min at 37 °C with immediate fixing using 4% paraformaldehyde in phosphate-buffered saline (PBS) at room temperature. Confocal images (1024 × 1024) were acquired on Zeiss inverted LSM 880 coupled with fast airyscan detector (Carl Zeiss, Inc) equipped with 63x oil immersion objective, NA 1.4. Images were acquired at different laser power for GFP tagged truncation mutants and detector gain settings in order to avoid oversaturation of pixels. All images were processed for brightness/contrast (histogram) adjustment for visualisation using ImageJ. For live cell imaging and FRAP analysis, cells were plated on glass bottom-well (1.5 petri dishes (ibidi) and allowed to grow at about ~70% confluence and transfected with respective Cavin1 mutants. For bleaching, 488 nm laser at 100% attenuation power was used for 20 iterations and subsequent imaging was done at one frame per second. Airyscan processing was done automatically in Zeiss software (ZEN 2.3). For PLA, PC3 cells were processed as described previously90. Images were then acquired on Zeiss LSM 710 and LSM 880 confocal microscope (Carl Zeiss, Inc) equipped with 63x oil immersion objective and quantification of PLA dots per cell was performed using find maxima function in ImageJ with offset of 25. For quantitative co-localisation, images (1024 × 1024) were acquired on Zeiss inverted LSM 880 in confocal mode and Pearson’s coefficient calculation was done using colo2 macro using ImageJ (https://imagej.net/ Colo2).

Constructing a structural model of mouse Cav1. A structural model of mouse Cav1 was built manually based both on known structures of the mouse Cav1 HR1 domain91 (PDB ID 4QKV), the previous model of the Cav1 unc12 UC1 region10, and secondary structure prediction of the Cav1 protein carefully cross-referenced to several Cav1 homologues and other Cavin family members11. Based on the homotrimeric coiled-coil structure of the HR1 domain we constructed our model under the assumption that a single Cav1 complex would consist of three separate Cav1 chains. The secondary structure predictions and previous crystal structure led us to define the following regions of Cav1 as either α-helical or random-coil: DR1, residues 1–48; random-coil; HR1, residues 49–147, α-helical (based on PDB 4QKV of mouse Cav1 HR1); DR2, residues 148–218, random-coil; HR2, α-helical for residues 219–242, random-coil for residues 243–244, α-helical for residues 245–278 (model from ref. 10); random-coil for residues 279–286, α-helical for residues 287–297; DR3, residues 298–392, random coil. Stretches of random-coil were built and added to α-helical domains manually in COOT Version 0.8.291 and the final model was subjected to simple geometry regulation in PHENIX Version 1.142. Structural images and electrostatic surface representations were rendered with PYMOL Version 2.3.1.

Statistics and reproducibility. Statistical analysis and P value calculations were performed by one-way ANOVA coupled with multiple comparisons at 95% confidence interval using graph pad Prism software. For the non-quantitative data in Figs. 3b–d, 4a, b, 5a–e, 6a–f, 7a, 8b–e and Supplementary Figs. 3A–F, 4A–F, 5C, 6A–E, 7A–C, 8A–C, 9A–C; 10, 11, pilot studies were carried out to assess the extent of variation between samples and qualitative observations of subsequent samples were consistent with these initial pilot studies. As such, these experiments were generally performed independently at least twice.

Reporting summary. Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability
All relevant data supporting the key findings of this study are available within the article and its Supplementary Information files or from the corresponding author upon reasonable request. The PDB file 4QKV is available at https://www.rcsb.org/structure/ 4QKV. A reporting summary for this Article is available as a Supplementary Information file. Source data are provided with this paper.

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