

1 Superresolution microscopy reveals partial preassembly and subsequent bending of 2 the clathrin coat during endocytosis

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23

24 Abstract

25 Eukaryotic cells use clathrin-mediated endocytosis to take up a large range of
26 extracellular cargos. During endocytosis, a clathrin coat forms on the plasma membrane, but
27 it remains controversial when and how it is remodeled into a spherical vesicle.

28 Here, we use 3D superresolution microscopy to determine the precise geometry of the
29 clathrin coat at large numbers of endocytic sites. Through pseudo-temporal sorting, we
30 determine the average trajectory of clathrin remodeling during endocytosis. We find that
31 clathrin coats assemble first on flat membranes to 50% of the coat area, before they become
32 rapidly and continuously bent, and confirm this mechanism in three cell lines. We introduce
33 the *cooperative curvature model*, which is based on positive feedback for curvature
34 generation. It accurately describes the measured shapes and dynamics of the clathrin coat
35 and could represent a general mechanism for clathrin coat remodeling on the plasma
36 membrane.

37

38 Introduction

39 Endocytosis is an essential function of eukaryotic cells to internalize molecules from their
40 surface. The major endocytic pathway is clathrin-mediated endocytosis (CME), which
41 supports the uptake of many diverse cargos including nutrients, signaling molecules,
42 membrane proteins, and pathogens. During CME, a dense coat of proteins self-assembles on
43 the inner leaflet of the plasma membrane. The membrane itself is bent into an Ω -shaped
44 invagination that is eventually pinched off to form a coated vesicle, completing endocytosis
45 (Kaksonen and Roux, 2018).

46 The major component of the coat are clathrin triskelia, which comprise three heavy and
47 three light chains (Fotin et al., 2004). When triskelia are bound to the plasma membrane by
48 adaptor proteins, they form ordered clathrin lattices. The structural flexibility of triskelia
49 allows these lattices to adapt variable ratios of predominantly pentagonal and hexagonal
50 faces, forming both flat and curved geometries.

51 Both geometries have been observed *in vivo* and *in vitro*, and their structure has been
52 well characterized *in vitro* from a structural biology perspective (Cheng et al., 2007;
53 Dannhauser and Ungewickell, 2012; Fotin et al., 2004; Heuser and Kirchhausen, 1985; Morris
54 et al., 2019; Pearse, 1976; Smith et al., 1998; Takei et al., 1998; Ungewickell and Branton,
55 1981). However, it remains elusive how clathrin coat formation and membrane bending are
56 temporally and causally related during endocytosis in cells.

57 In EM micrographs, it was observed early on that clathrin lattices can assume many
58 different curvatures in cells (Heuser, 1980). Since then, two main models of clathrin coat
59 formation during endocytosis have been put forward. In the *constant area model (CAM)*,
60 clathrin grows to its final surface area as a flat coat, which then becomes continuously more
61 curved until vesicle formation is complete. This model assumes that all the differently curved
62 clathrin structures are endocytic intermediates. Early observations suggested the presence of
63 pentagonal and hexagonal faces in isolated coated vesicles (Kanaseki and Kadota, 1969).
64 Combined with the observation of flat lattices being enriched in hexagons, it was suggested
65 that the integration of at least 12 pentagonal faces is a prerequisite for the formation of a
66 spherical structure (Heuser, 1980). However, this would require extensive lattice remodeling
67 which was deemed thermodynamically and structurally unfavorable and thus unlikely to
68 occur (Kirchhausen, 1993). The *constant curvature model (CCM)* was therefore formulated,
69 which assumes that flat clathrin structures are not endocytic precursors. Instead, it was
70 proposed that the endocytic clathrin coat assumes its final curvature, i.e. the curvature of the
71 vesicle, directly from the start, while continuously growing in surface area over time.

72 The constant curvature model had been prevalent in the field, but was recently challenged by
73 reports that flat clathrin coats can indeed change curvature during endocytosis (Avinoam et
74 al., 2015; Bucher et al., 2018; Scott et al., 2018). This has once again boosted interest in
75 clathrin remodeling (Chen and Schmid, 2020; Kaksonen and Roux, 2018; Sochacki and
76 Taraska, 2018). Recent years have seen numerous studies based on diverse methods that did
77 not converge on a common result. Reports supported either the constant area model

78 (Avinoam et al., 2015; Sochacki et al., 2021), the constant curvature model (Willy et al., 2021),
79 a combination of both models (Bucher et al., 2018; Tagiltsev et al., 2021; Yoshida et al., 2018)
80 or the simultaneous existence of both models within cells (Scott et al., 2018). To conclusively
81 understand this complex process, it would be desirable to directly visualize the three-
82 dimensional nanoscale structure of the endocytic clathrin coat in living cells. Unfortunately,
83 however, to date no method offers the necessary spatial and temporal resolution to do that.
84 Here, we aim to circumvent this limitation by densely sampling the entire endocytic process
85 in fixed cells, and subsequently reconstructing the dynamic information.

86 We developed a superresolution microscopy approach to quantitatively study the three-
87 dimensional (3D) clathrin coat architecture at endocytic sites. We used a novel model fitting
88 framework to extract geometric parameters that allowed us to sort static images of clathrin
89 lattices according to their progression along the endocytic timeline. The inferred endocytic
90 dynamics allowed us to reconstruct the stereotypic remodeling of clathrin during endocytosis
91 at the nanoscale.

92 In summary, we found that a clathrin coat first partially assembles on a flat membrane,
93 and then simultaneously grows in surface area and coat curvature. While initial bending
94 occurs rapidly, it later slows down and endocytic sites are eventually paused in a state of high
95 curvature before vesicle scission. This trend is conserved across cell lines, suggesting it is a
96 common attribute that is not affected by cell type. Based on this data, we developed a new
97 kinetic growth model, the *cooperative curvature model (CoopCM)*. It describes coat area
98 growth as addition of clathrin to the lattice edge with a constant rate and assumes that the
99 curvature of the coat increases towards a preferred curvature, driven by the cooperative
100 interplay of clathrin triskelia within the lattice. The CoopCM predicts a fast initial curvature
101 increase that is slowed down progressively as the coat becomes spherical, and shows
102 excellent agreement with experimental data.

103

104 Results

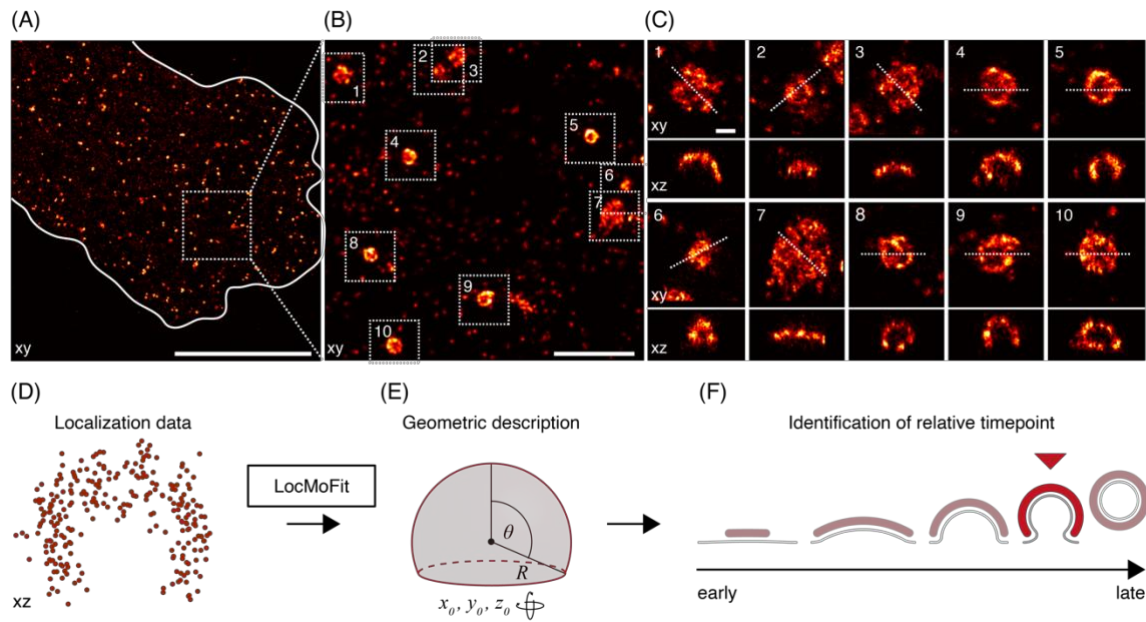
105 *Quantitative 3D superresolution imaging of clathrin structures*

106 Here, we used 3D single-molecule localization microscopy (SMLM) to systematically
107 determine the precise geometry of individual clathrin-coated structures at the plasma
108 membrane. For this, we optimized the sample preparation to label clathrin at endocytic sites
109 as densely as possible using indirect immunofluorescence with polyclonal antibodies against
110 clathrin light and heavy chains, which was crucial to achieve high quality SMLM (Mund and
111 Ries, 2020). We then localized sparsely activated single fluorophores by fitting an
112 experimentally derived model of the astigmatic point-spread function using a method we
113 developed recently (Li et al., 2018). This improved the resolution to about 10 nm in x/y and
114 30 nm in z (based on modal values of the localization precision at 3.9 nm in x/y and 12.5 nm
115 in z, see Methods), and reduced typically observed image distortions. The achieved image
116 quality allowed us to clearly visualize and quantify the 3D clathrin coat shape at different
117 stages of endocytic site maturation (Figure 1A-C).

118 The large majority of sites were single structures that were well-isolated from each other,
119 and exhibited great structural diversity. In addition, we noted several clusters of closely
120 juxtaposed endocytic sites (Supplementary Figure 1A). In the isolated sites, we observe a
121 variety of shapes including flat, curved, dome-shaped and spherical structures of different
122 sizes (Figure 1C), indicating that endocytosis has been arrested by fixation at different time
123 points during endocytic progression.

124 To quantify size and shape of individual clathrin coats, we used LocMoFit, a
125 computational pipeline based on a maximum-likelihood model fitting framework that we
126 developed previously (Wu et al., 2021). This framework directly fits 3D geometric models to
127 the 3D point cloud of localizations (Figure 1D-E) instead of rendered images, and thereby
128 harnesses the full information contained in the SMLM data, including for instance the
129 localization precision, rather than just spatial coordinates.

130 We describe the clathrin coat as a spherical cap that is defined by a radius r and a closing
131 angle θ (Figure 1E). Our model also accounted for coat thickness, antibody size and blurring
132 due to the localization precision (Methods). Hence, it describes both shallow and highly
133 curved structures equally well. Moreover, because θ increases during endocytosis from 0° at
134 flat sites to 180° at complete vesicles, we could use this parameter to sort the individual
135 images according to endocytic progress (Figure 1D).



136

137 **Figure 1 – 3D SMLM of clathrin-coated structures.** (A) 3D SMLM image of clathrin immunolabeled with AF647
 138 at the bottom membrane of a SK-MEL-2 cell. (B) Enlarged view of the region indicated in (A). (C) All structures
 139 found in (B) are shown in top view (xy) and as 50 nm-thick z-slices (xz), orientation of slice indicated by dotted
 140 line). Scale bars are 10 μm (A); 1 μm (B); and 100 nm (C). (D) Geometric analysis pipeline. All clathrin coats are
 141 segmented, and their localizations are directly fitted with a spherical cap model using the fitting framework
 142 LocMoFit (Wu et al., 2021). (E) In LocMoFit, individual clathrin coats are parametrized by their size (radius R),
 143 closing angle θ , position (x_0, y_0, z_0) and 3D orientation. (E) Using θ as a proxy for endocytic progression, the
 144 relative endocytic time point for each structure can be determined.

145 **The clathrin coat grows in area and becomes more curved during endocytosis**

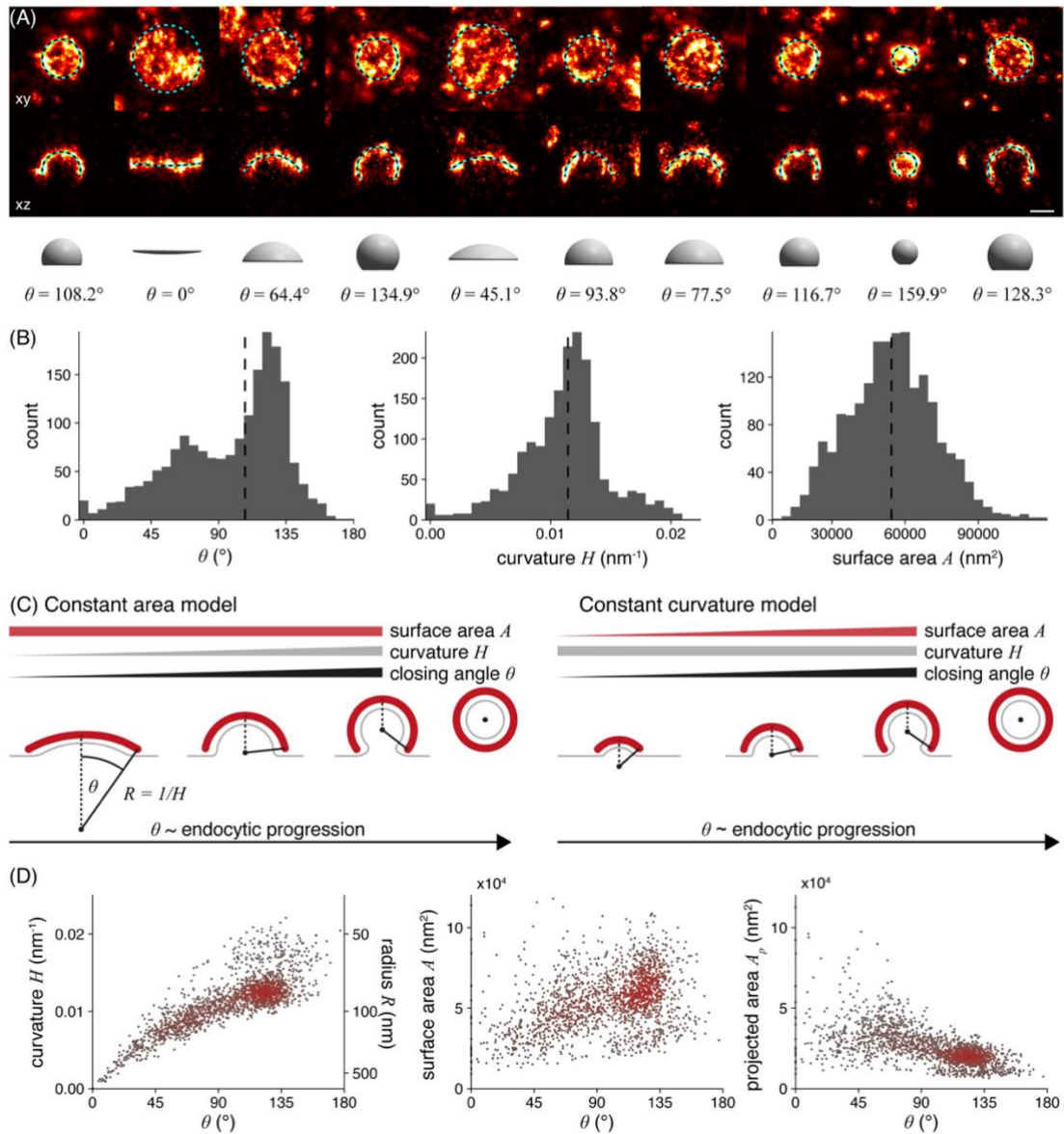
146 We first imaged immunostained clathrin in chemically fixed SK-MEL-2 cells, where we
 147 focused on the bottom plasma membrane that was densely covered by endocytic sites. These
 148 cells have been extensively studied and are established model cells for clathrin-mediated
 149 endocytosis with well-characterized endocytic dynamics (Aguet et al., 2013; Avinoam et al.,
 150 2015; Doyon et al., 2011; Kaplan et al., 2020). Using the 3D model fitting pipeline, we
 151 determined radius R , closing angle θ , position and rotation of 1798 endocytic sites from 13
 152 cells (Figure 2A) with high accuracy (Supplementary Figure 2). We found that two structural
 153 intermediates are enriched, while others are rare and only a small fraction of sites were
 154 completely flat (Figure 2B). Slightly curved sites with $\theta \approx 70^\circ$ and strongly curved sites with
 155 $\theta \approx 130^\circ$ were enriched, indicating that those structural states are more frequently found in
 156 cells. We only rarely obtained sites with $\theta \approx 180^\circ$, which would be expected for complete
 157 spherical coats, even though fully formed vesicles are found in our data (Supplementary
 158 Figure 1B). This indicates that at the time point of scission, the clathrin coat of nascent vesicles
 159 is still incomplete at the neck, or that the kinetics of scission are too transient for detection.
 160 Deeply invaginated clathrin coated pits could further be sheared off during sample
 161 preparation, thus evading our detection. Curvatures $H = 1/R$ ranged from 0 to 0.022 nm^{-1}

162 with a median of 0.011 nm^{-1} , corresponding to a median radius of 87 nm, and surface areas
163 ranged from $9,000 \text{ nm}^2$ to $140,000 \text{ nm}^2$ with a median of $54,000 \text{ nm}^2$. These values agree well
164 with previous measurements of the vesicular coat using EM (Avinoam et al., 2015), median
165 curvature 0.015 nm^{-1} , median surface $54,500 \text{ nm}^2$).

166 We then wanted to understand how the clathrin coat geometry changes during
167 endocytosis. To this end, we used the closing angle parameter θ to sort endocytic sites
168 relative to each other in time. Irrespective of whether endocytosis follows a constant
169 curvature or constant area model, θ monotonically increases from early to late endocytic
170 time points (Figure 2C), and can thus be used as unbiased proxy for endocytic progression
171 (Avinoam et al., 2015).

172 The curvature H was strongly correlated with θ , indicating that coat curvature increases
173 continuously during endocytosis (Figure 2D). Similarly, the surface area increased from $32,000$
174 nm^2 (median of 5% of sites with smallest theta) to $50,000 \text{ nm}^2$ (median of 5% of sites with
175 highest theta). The projected area decreased from $31,000 \text{ nm}^2$ to $13,000 \text{ nm}^2$ (median of 5%
176 of sites with lowest and highest theta respectively), which is in close agreement with previous
177 EM measurements (Bucher et al., 2018). It is readily obvious that our data are incompatible
178 with the constant curvature model, as the curvature is not constant, but increases
179 monotonically with θ . Just as clearly, our data do not support the constant area model,
180 because the coat surface also increases during endocytosis.

181 Almost all data points are part of one continuous point cloud (Figure 2D), indicating a
182 continuous transition between structural states during endocytosis. We noticed an additional
183 small, disconnected set of data points representing 8.5% of all sites that correspond to
184 endocytic sites with curvatures above 0.016 nm^{-1} and θ of 80° - 180° (Figure 2D, and example
185 structures in Supplementary Figure 3). In a control experiment to check whether these are
186 endocytic structures, we selectively analyzed clathrin structures that colocalized with AP-2, a
187 bona fide marker for CME. We did not observe AP-2 in any of the disconnected sites, from
188 which we conclude that they indeed do not belong to CME. These small structures could
189 represent coated vesicles from the *trans* Golgi (Supplementary Figure 4).



190

191 **Figure 2 – Quantitative analysis of clathrin-coated structures in SK-MEL-2 cells.** (A) Clathrin coat geometry is
 192 quantified using LocMoFit. The fitted spherical cap model is drawn as circle with its outer radius (top row, xy), as
 193 cross-sectional profile (50 nm xz slices, middle row), and as surface with the corresponding closing angle
 194 θ (bottom row). Scale bar 100 nm. (B) Distributions of closing angle θ (median = 108°), curvature H (median =
 195 0.011 nm^{-1}) and surface area A (median = $54,000 \text{ nm}^2$) of endocytic sites in SK-MEL-2 cells ($n = 1798$ sites, $N = 13$
 196 cells) as determined from the model fit. (C) Two previously proposed mechanistic models for clathrin coat
 197 assembly during endocytosis (for details see text). In both scenarios θ increases monotonically and is thus a
 198 proxy for endocytic progression. (D) Development of curvature, surface area and projected area of the clathrin
 199 coat during endocytosis. Color indicates point density.

200

201 ***The cooperative curvature model of clathrin coat remodeling***

202 Our data clearly showed that clathrin coats grow and become more curved as the plasma
203 membrane gets bent to produce a vesicle. Because the data quantitatively describes all 3D
204 geometries that the endocytic clathrin coat assumes, it allowed us to move towards a
205 mathematical model of clathrin coat formation during endocytosis.

206 Here, we introduce the *Cooperative Curvature Model (CoopCM)*, which describes coat
207 growth based on known structural and dynamical properties of clathrin coats (Figure 3A).
208 Firstly, we assume that the clathrin coat starts growing on a flat membrane. While triskelia
209 have been shown to exchange dynamically during endocytosis (Avinoam et al., 2015), net
210 growth of the area A occurs via the addition of triskelia at the lattice edge \mathcal{E} with a constant
211 growth rate k_{on} (Equation 1).

$$212 \quad \frac{d}{dt}A = \dot{A} = k_{\text{on}}\mathcal{E} \quad (1)$$

213 Moreover, we assume that the intrinsic pucker angle of individual triskelia and their
214 interactions in a lattice together translate into an overall preferred curvature H_0 of the
215 clathrin coat as a whole. However, the initially flat assembly suggests that this preferred
216 curvature cannot be realized in an immature lattice. We hypothesize that cooperative
217 processes in the coat are required to increase curvature. Coat curvature H then increases
218 asymptotically towards H_0 at an initial rate γ , slowing down towards zero when the preferred
219 curvature is reached (Equation 2).

$$220 \quad \frac{d}{d\theta}H = \gamma \left(1 - \frac{H^2}{H_0^2}\right) \quad (2)$$

221 Choosing a quadratic dependence of the rate of curvature change on curvature is a simple
222 way to represent cooperativity in the lattice, which recently has been demonstrated in
223 experiments (Sochacki et al., 2021; Zeno et al., 2021) and fits our data more accurately than
224 the linear relationship, which would correspond to a less cooperative process (Supplementary
225 Note). Equation 2 can be solved analytically to yield an expression of the curvature H
226 depending on θ (Equation 3), which can then be fitted to our data.

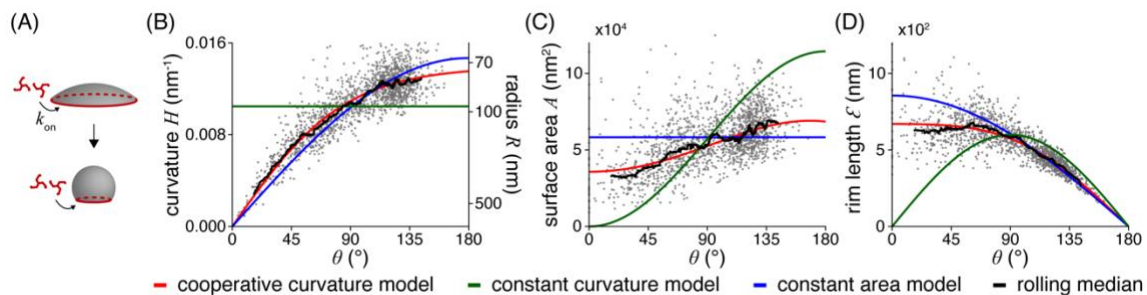
$$227 \quad H(\theta) = H_0 \tanh\left(\frac{\gamma\theta}{H_0}\right) \quad (3)$$

228 Analogous expressions for other geometric parameters like surface area can be derived
229 straightforwardly.

230 The fit of this CoopCM to the data of curvature H in relation to closing angle θ shows
231 excellent agreement (Figure 3B). In comparison, the CAM fitted the curvature data slightly
232 worse than the CoopCM, and the constant curvature model agreed poorly with the data. The
233 improved fit of the CoopCM compared to the CAM or CCM is not a result of the additional
234 fitting parameter, as we showed using the Bayesian Information Criterion (see Supplementary
235 Note).

236 From the curvature fits, we calculated the corresponding curves for surface area (Figure
 237 3C) and the edge length of the clathrin coat (Figure 3D). The edge length decreases
 238 monotonically and approaches zero at $\theta = 180^\circ$, thereby stopping growth according to
 239 Equation 1. Both graphs again highlight the very close agreement of the model prediction
 240 with the experimental data. From the fit, we determined that invagination occurs when about
 241 half of the total coat area has grown ($A_0 = 0.51$), and that the preferred curvature of a
 242 clathrin coat is $H_0 = 0.014 \text{ nm}^{-1}$, corresponding to a radius of $R_0 = 72 \text{ nm}$. The model yields
 243 nearly identical parameters when surface area or rim length are fitted instead of curvature
 244 (Table 1), highlighting its robustness.

245 We then decided to test if the observed mode of clathrin remodeling is specific for SK-
 246 MEL-2 cells. For this, we analyzed the endocytic clathrin ultrastructure also in U2OS cells and
 247 3T3 mouse fibroblasts (Supplementary Figure 5 and Table 2-3). In these cell lines, just like in
 248 SK-MEL-2 cells, the curvature as well as the surface area continuously increase throughout
 249 endocytosis. We observe that the preferred curvature of the clathrin coat is smaller in U2OS
 250 ($H_0 = 0.012 \text{ nm}^{-1}$, $R_0 = 85 \text{ nm}$) and 3T3 cells ($H_0 = 0.011 \text{ nm}^{-1}$, $R_0 = 89.7 \text{ nm}$) compared
 251 to SK-MEL-2 ($H_0 = 0.014 \text{ nm}^{-1}$, $R_0 = 72 \text{ nm}$). This suggests that the average size of vesicles
 252 formed in CME is cell line specific. The fraction of the surface area acquired on the flat
 253 membrane is very similar for all three cell lines, with U2OS derived sites initiating curvature
 254 at $A_0 = 0.52$ and 3T3 sites at $A_0 = 0.45$ (Table 2-3). Taken together, we have shown in
 255 several cell lines that clathrin coats neither grow via constant curvature or constant area
 256 pathways, but rather first grow flat and then acquire curvature and more surface area
 257 simultaneously, with a non-linear mode of curvature generation.



258

259 **Figure 3 – Model for clathrin coat growth.** (A) Schematic of the cooperative curvature model, where clathrin
 260 lattices grow by addition of triskelia to the rim at a constant growth rate k_{on} . Curvature generation is driven
 261 towards a preferred curvature, ultimately creating a spherical vesicle. (B) Distinct clathrin growth models and
 262 rolling median (window width = 82 sites) fitted to curvature over θ . The resulting fitting parameters are then
 263 used to map the same models also over (C) surface area and (D) rim length. ($n = 1645$ sites, $N = 13$ cells).

264

265

266 **Temporal reconstruction of structural dynamics in endocytosis**

267 We systematically segment all endocytic sites in the superresolution images, and thereby
268 obtain a large dataset where all endocytic time points are sampled homogenously. The
269 distribution of structural states within the dataset is thus representative of their lifetime, with
270 long-lived, stable curvature states being overrepresented in the data compared to transient
271 states. This opens up the possibility to reconstruct the temporal progression of clathrin
272 remodeling during endocytosis, and to ask whether clathrin coats acquire their curvatures
273 with a constant rate, or if there are certain curvature transitions that occur more rapidly than
274 others.

275 For this, we sorted all endocytic sites by θ . The rank of an endocytic site thus corresponds
276 to its pseudotime, which describes its relative time point between 0 and 1 along the endocytic
277 trajectory (Figure 4A).

278 As our model (Equation 1) describes the dynamic growth of the clathrin coat, we can solve
279 it to derive an expression for θ over time t (Equation 4), where the coat starts out flat and
280 then starts to generate curvature, increasing θ over time.

$$281 \quad \theta(t) = \sqrt{\frac{24\gamma k_{\text{on}}}{8\gamma^2 H_0^{-2} - 1} t} \quad (4)$$

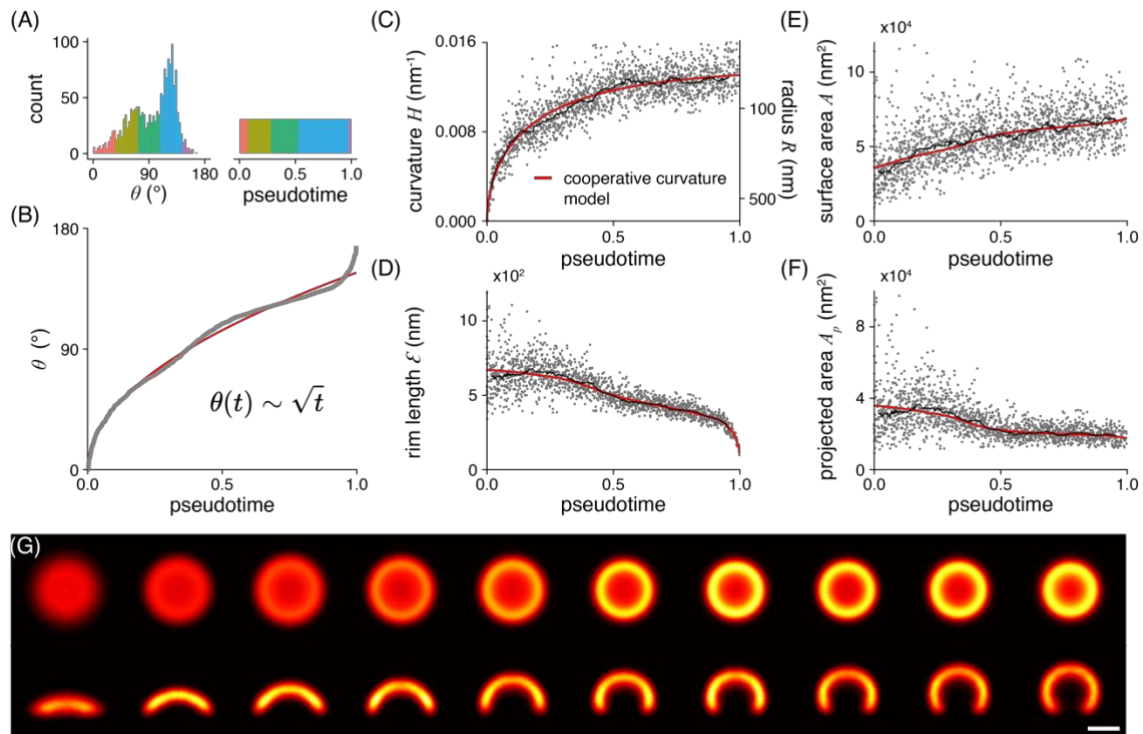
282 The square root dependence of θ on time t reflects the slowing down of curvature
283 generation as the clathrin coat approaches its preferred curvature. This expression fits the
284 pseudotime-resolved data remarkably well (Figure 4B). Consistent with our previous
285 reasoning, a linear model did not agree well with the data (Supplementary Note), since it gives
286 a linear invagination speed for small t , emphasizing the validity of our cooperative curvature
287 model, which leads to the characteristic square root dependence. The data slightly deviates
288 from the model in the early phase, and more notably in the very end for pseudotimes close
289 to 1. This potentially indicates that clathrin geometry is influenced by other factors besides
290 the coat itself close to vesicle scission.

291 Because this pseudotime resolved dataset was generated from a large number of
292 endocytic sites in many cells, we effectively generated the average trajectories of how
293 curvature, surface area, projected area and lattice edge change during endocytosis in SK-MEL-
294 2 cells (Figure 4C-F). We observe comparatively few flat clathrin coats. This shows that clathrin
295 lattices are only transiently flat in the beginning of endocytosis, and might represent an
296 energetically unfavorable conformation (Figure 4B). A fast transitions from a flat to a curved
297 structure could further be mediated independently of clathrin coat (Zhao et al., 2017), e.g. by
298 the action of BAR domain proteins (Henne et al., 2010), or at the base of filopodial projections.
299 We further find comparatively many structures with a curvature of $\theta \approx 70^\circ$ and $\theta \approx 130^\circ$,
300 indicating more long-lived endocytic stages. As the surface area is constantly increasing over
301 pseudotime, we assume that the enrichment at $\theta \approx 130^\circ$ represents a stalling point, where
302 vesicles are formed by addition of final clathrin triskelia and potentially other factors that
303 enable the recruitment and mechanic function of dynamin during vesicle scission. Similarly,

304 the enrichment at $\theta \approx 70^\circ$ could be indicative of further recruitment of regulatory
305 components, potentially supporting a previously suggested endocytic checkpoint (Loerke et
306 al., 2009). This functional interpretation is supported by the observation of similar peaks in
307 U2OS (at $\theta \approx 60^\circ$ and $\approx 130^\circ$) as well as 3T3 cells (at $\theta \approx 50^\circ$ and $\approx 130^\circ$).

308 Within the first 10% of pseudotemporal progression, the clathrin coat rapidly acquires
309 shallow curvature of $H = 0.007 \text{ nm}^{-1}$ ($R = 134 \text{ nm}$). It then becomes gradually more bent up
310 to $\approx 60\%$ of the endocytic timeline, when the sites reach an average curvature of 0.012 nm^{-1}
311 ($R = 83 \text{ nm}$). During the last $\approx 40\%$ of progression, the curvature increase almost stops, until
312 vesicle scission occurs. Interestingly, the earliest sites in our dataset already contain $\approx 50\%$ of
313 the surface area of the final vesicles, which is also reflected in the fitting results ($A_0 = 0.51$)
314 (Figure 4E). This indicates that the initial assembly of the clathrin coat occurs very rapidly, or
315 is in part below our detection limit.

316 We performed identical analyses for U2OS and 3T3 derived clathrin sites and obtained
317 highly similar results as for SK-MEL-2 cells, indicating that this trajectory represents a general
318 pathway for clathrin coat remodeling during endocytosis (Supplementary Figure 5 and Tables
319 2-3). Finally, we made a 3D nanoscale movie, which vividly illustrates assembly and
320 remodeling of the clathrin coat during endocytosis, from pseudotime resolved averages of
321 our data (Figure 4G). This yielded a nanoscale pseudo-temporal movie of endocytosis in SK-
322 MEL-2 cells (Supplementary Movie).



323

324 **Figure 4 – Temporal reconstruction of clathrin coat remodeling.** (A) Endocytic sites are sorted by θ to
 325 reconstruct pseudotime. Enriched θ states, for example the peak at 135° , represent long-lived states that remain
 326 for extended periods in pseudotime. Color code represents the same number of sites in both histograms. (B) The
 327 square-root dependence between θ and pseudotime approximated by the cooperative curvature model (red
 328 line). (C) Curvature generation over pseudotime is approximated by the cooperative curvature model. Fit results
 329 in (C) were used to describe (D) rim length, (E) surface area and (F) projected area change over pseudotime. A
 330 rolling median (window of 82 sites) is plotted alongside (black line). (G) Superresolution averages for distinct
 331 endocytic stages, resulting from all collected snapshots. Each bin contains the same number of snapshots of
 332 clathrin-coated structures sorted along their pseudotime ($n = 163$ per bin), so that all bins represent equally long
 333 pseudotime intervals. Individual sites were rescaled to the median bin radius and aligned by their center
 334 coordinates as well as rotation angles. Scale bar is 100 nm. ($n = 1645$ sites, $N = 13$ cells).

335 Discussion

336 *Quantitative description of clathrin coat ultrastructure*

337 The nature of clathrin assembly during endocytosis has become a classical question in cell
338 biology that remains unresolved (reviewed in (Chen and Schmid, 2020; Sochacki and Taraska,
339 2018)). There have been two main competing mechanistic models of how triskelia form a
340 coat: the constant curvature model predicts that the clathrin coat assumes its final curvature
341 directly from the start, while continuously increasing in surface area over time. In contrast,
342 the constant area model predicts that clathrin grows to its final surface area as a flat coat,
343 which then becomes continuously more curved until vesicle formation is complete.

344 Each model is supported by a number of studies, which mostly relied on electron
345 microscopy techniques. Among them, 3D correlative electron tomography yields exquisite 3D
346 endocytic membrane shapes (Avinoam et al., 2015), but can be tedious to extend towards
347 large numbers of cells. Platinum replica electron microscopy offers large fields of view and
348 thus superb throughput, but gives limited 3D resolution (Bucher et al., 2018; Sochacki et al.,
349 2017).

350 We reason that the trajectory of clathrin coat assembly during endocytosis could best be
351 determined by systematically imaging large numbers of endocytic sites with high 3D
352 resolution. Therefore, we used SMLM that combines high 3D resolution with high throughput.
353 Although lower than in electron microscopy, the resolution we achieved allowed us to resolve
354 the precise 3D coat ultrastructure of clathrin-coated pits. Importantly, due to the molecular
355 specificity for clathrin, we were able to segment all endocytic sites at the bottom surface of
356 the cells in our images in an unbiased way, thus ensuring homogenous sampling of the
357 structural variety of clathrin coats.

358 We applied a novel maximum-likelihood based fitting framework that we developed
359 recently (Wu et al., 2021), which allows fitting complex geometric models to localization point
360 clouds. Because the fit is applied directly to the localizations, it considers all quantitative
361 parameters of all localizations, most notably localization uncertainty. This results in the
362 precise and reliable quantification of the underlying structures (Wu, Tschanz & Krupnik et al.,
363 2020), even when taking linkage error through indirect immunolabeling into account
364 (Supplementary Figure 6).

365 We described the clathrin coat as a spherical cap, which matched the large majority of
366 sites very well. Additionally, in our data, we observed that some sites are asymmetric,
367 ellipsoidal, or deformed more irregularly (Supplementary Figure 1C). We do not currently
368 evaluate this asymmetry. Rather, we introduced rotational symmetry during the averaging
369 (Figure 4G), where we aligned each site based on their model parameters and thus averaged
370 out many irregularities. A deviation of the cross-sectional profile from a circle is nevertheless
371 preserved in the averaging (Supplementary Figure 8D - F), and in future studies more complex
372 geometric models will enable analyzing the structure and asymmetries of the coat in more
373 detail.

374

375 ***Flat clathrin-coated structures***

376 In the first step of our analysis pipeline, we segmented all clathrin-coated structures in
377 the superresolution images. We thereby aimed to achieve homogenous sampling to capture
378 all endocytic intermediates with the same probability irrespective of time point, size, and
379 orientation. Interestingly, the surfaces of the earliest detectable endocytic sites already
380 contained half the surface area of sites in the latest CME stages. How do these earliest sites
381 assemble? We cannot sort the very flat sites, which are comparatively rare in our dataset, in
382 time using θ because their θ is close to zero. However, their relative fraction among all sites
383 is still informative. As short-lived states are under-represented, the relative absence of small
384 flat sites in our data indicates that the initial assembly occurs rapidly.

385 In addition to the short lifetimes of small flat sites, two technical reasons could potentially
386 contribute to their rare occurrence in our data. Due to their small size and potentially sub-
387 stoichiometric labeling, which is also noticeable as holes in larger structures (Supplementary
388 Figure 1D), they might not have sufficient immunostaining signal and thus they might be hard
389 to be differentiated from small clusters of primary and secondary antibodies that are
390 routinely observed in SMLM images. Additionally, the very first small clathrin assemblies
391 might be meta-stable, with mostly loosely bound triskelia, and thus might not be stabilized
392 well by chemical fixation. However, it is important to note that very small sites are also rarely
393 found in previously published electron microscopy datasets (Avinoam et al., 2015; Bucher et
394 al., 2018; Sochacki et al., 2021). Since EM should not suffer from the same technical
395 limitations, it seems likely that these states are indeed short-lived.

396 Generally, completely flat clathrin coats are rare in our data. Instead, we observed that
397 clathrin coats quickly acquire curvature already early in endocytosis. This agrees with the
398 early observation made using EM (Heuser, 1980) that even flat structures already had some
399 small degree of curvature. Interestingly, an enrichment of pentagons was observed towards
400 the edges of flat lattices, leading to the hypothesis that even without additional formation of
401 pentagons these flat lattices have the built-in capacity to become curved, and are not very
402 stable. This notion is in agreement with recent demonstrations that flat clathrin coats store
403 energy, which can drive coat bending without additional factors (Sochacki et al., 2021;
404 Tagiltsev et al., 2021).

405 The importance of curvature generation for clathrin coat maturation is supported by
406 previous studies that suggest failure to initiate significant curvature as a hallmark of abortive
407 events (Loerke et al., 2009; Wang et al., 2020). Observed in live-cell studies, these dim and
408 short-lived events fail to significantly increase in intensity and do not undergo scission (Loerke
409 et al., 2009; Mettlen et al., 2009). As structures in our data set mostly contain at least shallow
410 curvatures at minimally half the final coat area, we are likely not capturing these abortive
411 events, and they have negligible impact on our analysis.

412

413 ***The cooperative curvature model***

414 Here, we visualized the shapes of clathrin coats with high 3D resolution. This was crucial
415 to precisely measure the clathrin curvature throughout the endocytic timeline ranging from
416 flat to highly curved structures. Especially shallow curvatures are hard to accurately assess
417 with 2D imaging. Thus, our data enabled us to robustly test different growth models for
418 clathrin during endocytosis.

419 The constant curvature model predicts a continuous increase in surface area, constant
420 coat curvature, as well as a rim length that increases until a half spherical geometry is reached,
421 and then decreases again. Our experimental data, most notably the increase in curvature and
422 monotonical decrease in rim length, are incompatible with these predictions, thus ruling out
423 the constant curvature model (Figure 3, green lines).

424 The constant area model, in a strict interpretation, on the other hand predicts the coat
425 to assemble completely on a flat membrane, after which its area stays constant and only
426 curvature increases to form a spherical vesicle. These predictions agree reasonably well with
427 the data for curvature propagation during CME, but fail to explain the monotonic increase in
428 coat surface over time. Thus, the continuous increase of surface area that we observed here
429 rule out the constant area model as well (Figure 3, blue lines). Interestingly, earlier work
430 compatible with the constant area model already suggested the presence of shallow
431 curvatures in even the flattest clathrin coated structures (Avinoam et al., 2015; Bucher et al.,
432 2018; Heuser, 1980)

433 We found that around half of the clathrin coat area has preassembled when plasma
434 membrane invagination begins (Figure 2D, 4C), agreeing well with previous reports (Bucher
435 et al., 2018; Scott et al., 2018), after which the coat keeps growing gradually. Based on this
436 observation, we developed a mathematical model that for the first time considers that
437 curvature is generated in a positive (non-linear) feedback loop. Our cooperative curvature
438 model assumes that net area growth of the clathrin coat occurs via triskelia addition to the
439 many available binding sites at the coat edge. We proposed that this process depends on the
440 number of free binding sites, which scale with the rim length, and can be described by a single
441 kinetic constant k_{on} . Triskelia addition within the coat is still likely to occur at a lower rate, as
442 clathrin lattices can be unsaturated and have defects (Frey et al., 2020; Sochacki et al., 2021)
443 and triskelia can exchange (Avinoam et al., 2015), which is energetically favorable for
444 curvature changes within these lattices (Frey and Schwarz, 2020; Sochacki et al., 2021;
445 Tagiltsev et al., 2021). While growth at the rim accounts for increase in surface area, curvature
446 is most likely generated in the bulk of the coat. Here we assumed a non-linear relation
447 between the rate of curvature increase and curvature, which reflects cooperativity in the
448 lattice, e.g. due to rearrangements of neighboring triskelia or larger regions thereof. This
449 assumption of cooperativity is supported by recent experiments, which suggest clathrin to
450 exhibit curvature sensing properties, preferentially assembling at pre-bent membranes
451 (Sochacki et al., 2021; Zeno et al., 2021). The model described above shows an excellent fit
452 between theory and experiment, and further predicts a square root dependence of theta over

453 time. This describes curvature generation during clathrin coat maturation as a non-linear
454 mechanism driven by a positive feedback of multiple triskelia, slowing down once the coat
455 approaches a preferred degree of curvature.

456 Even though our model agrees very well with the data, it still exhibits certain limitations worth
457 mentioning:

458 Firstly, it only considers the coat itself and ignores the plethora of other proteins within
459 the endocytic machinery. These include numerous BAR domain proteins that are dynamically
460 recruited and disassembled during endocytosis and thus are bound to influence membrane
461 curvature at endocytic sites, as well as a variety of clathrin adaptor proteins, whose presence
462 or absence could explain the cell-type specific differences in average vesicle sizes that we
463 observed (Supplementary Figure 5). Taken together, these factors could explain the
464 imperfection of our model in the very beginning, and the final part of the timeline (Figure 4A),
465 where vesicle scission is driven by the fast-acting mechanoenzyme dynamin. Additionally, we
466 only modeled clathrin recruitment and ignore clathrin disassembly, which could be mediated
467 by adaptor unbinding (Taylor et al., 2011) and uncoating factors including auxilin (He et al.,
468 2020) that are recruited before the end of vesicle scission. We also assumed that the clathrin
469 coat has constant properties, most notably that the intrinsic coat-driven curvature generation
470 towards its preferred curvature occurs unidirectionally and remains the same throughout
471 endocytosis (equation 3). It is however likely that the properties of the clathrin coat change
472 during endocytosis, e.g. by coat stiffening or increasingly tight interactions between triskelia
473 (Frey and Schwarz, 2020; Frey et al., 2020; Sochacki et al., 2021).

474 Secondly, we reconstructed an average trajectory of clathrin coat remodeling, generated
475 from many individual static snapshots, thereby averaging conceivable different pathways of
476 CME. However, different pathways with substantially changed relationships between the
477 parameters like curvature and theta would be visible in the corresponding plots as separate
478 point clouds. This is not the case, rather we observed a continuous correlation between
479 curvature and theta following a single trajectory, indicating that CME follows a single,
480 stereotypic pathway. We did identify a small, disconnected population of sites in our data set
481 that most likely originate from a distinct cellular mechanism (Supplementary Figure 3 and 4).
482 While this indicates that our approach should capture potentially different pathways of
483 clathrin coated vesicle formation, we cannot exclude that minor mechanistic variations are
484 included into the average trajectory.

485 Clathrin recruitment has been quantified extensively using live-cell microscopy, resolving
486 the sequential recruitment and dynamic characteristics of many important endocytic
487 components (Aguet et al., 2013; Cocucci et al., 2012; Doyon et al., 2011; He et al., 2017; Jin
488 et al., 2021; Loerke et al., 2009; Mettlen et al., 2009; Saffarian et al., 2009; Schöneberg et al.,
489 2018; Taylor et al., 2011). We wondered if it is possible to correlate our pseudotime
490 reconstruction with previously reported real-time dynamics of clathrin, which shows an initial
491 fluorescence intensity increase, a plateau, and finally a sharp intensity decrease after vesicle
492 scission. While we observed a correlation between the number of clathrin localizations and

493 surface area (Supplementary Figure 7), we note that indirect immunolabeling is not always
494 quantitative, which complicates a direct comparison of previously reported live-cell
495 fluorescence intensity over time and our pseudotime trajectory data. Nevertheless, we
496 speculate that the initial fast intensity increase in live-cell studies likely corresponds to the
497 initial growth of the flat clathrin coat that escapes our detection due to the fast dynamics and
498 small size. The slower fluorescence increase and subsequent plateauing then coincides with
499 curvature generation and final addition of triskelia at the coat rim as resolved in detail in our
500 pseudotime data. To better understand the correlation between changes in the nanoscale
501 architecture of clathrin coats and dynamic consequences, we would ultimately require a
502 method that combines high structural with temporal resolution. In a recent publication, this
503 was attempted using a live-cell 2D superresolution microscopy approach (Willy et al., 2021).
504 This study reported an increasing projected area of clathrin over time, suggested that
505 curvature is present in the earliest detectable clathrin structures, and concluded that CME
506 follows the constant curvature model. Although we also find that completely flat structures
507 are rare and curvature is initiated before the final surface is acquired, our data is entirely
508 incompatible with the CCM. This is especially true in the first half of endocytosis with shallow
509 but progressively increasing curvatures (Figure 3), which is challenging to measure using 2D
510 imaging. This shows that it remains highly desirable to ultimately image the 3D nanoscale
511 architecture of the clathrin coat in living cells in real time.

512 In summary, we characterized the dynamics and geometries of the clathrin coat in
513 endocytosis by combining 3D superresolution microscopy, quantitative analysis, and
514 mathematical modeling. We found that clathrin bending and assembly occur simultaneously
515 along a precisely defined trajectory. We anticipate that this work will be foundational to
516 further study the structural mechanism of endocytosis, both under physiological conditions,
517 and in disease, where changes in CME likely occur very frequently and have just recently been
518 shown to have profound functional consequences (Moulay et al., 2020; Xiao Guan-Yu et al.,
519 2018).

520

521 **Acknowledgements**

522 We thank the entire Ries and Kaksonen labs for fruitful discussions and support. This work
523 was supported by the European Research Council (ERC CoG-724489 to J.R.), the National
524 Institutes of Health Common Fund 4D Nucleome Program (Grant U01 to J.R.), the Human
525 Frontier Science Program (RGY0065/2017 to J.R.), the EMBL Interdisciplinary Postdoc
526 Programme (EIPOD) under Marie Curie Actions COFUND (Grant 229597 to O.A.), the
527 European Molecular Biology Laboratory (M.M., A.T., Y.-L. W. and J.R.) and the Swiss National
528 Science Foundation (grant 310030B_182825 and NCCR Chemical Biology to MK). O.A is an
529 incumbent of the Miriam Berman Presidential Development Chair.

530 **Author contributions**

531 M.M., O.A., M.K. and J.R. conceived the study, M.M., A.T., J.M. and O.A. performed
532 experiments, M.M., A.T., Y.-L.W., J.M., O.A. and J.R. analyzed superresolution data; F.F. and
533 U.S. developed the cooperative curvature model; J.R. supervised the study. M.M., A.T., F.F.,
534 J.R. wrote the manuscript with input from all authors.

535 **Declaration of interests**

536 The authors declare no competing interests.

537 **Material and methods**

538 ***Cell culture***

539 SK-MEL-2 cells (gift from David Drubin, UC Berkeley, described in (Doyon et al., 2011))
540 were cultured adherently as described previously (Li et al., 2018) in DMEM F12 (Dulbecco's
541 modified Eagle's medium with Nutrient Mixture F-12) with GlutaMAX, phenol red (Thermo
542 Fisher; 10565018), 10% (v/v) FBS, ZellShield (Biochrom AG, Berlin, Germany), at 37 °C, under
543 an atmosphere with 5% CO₂ and 100% humidity.

544 U2OS cells (Cell Line Services; #300174) were cultured adherently as described previously
545 (Thevathasan et al., 2019) in high-glucose DMEM supplemented with 10% FBS, 2 mM l-
546 glutamine, non-essential amino acids, ZellShield at 37 °C under an atmosphere with 5% CO₂
547 and 100% humidity.

548 3T3 mouse fibroblasts (gift from Alba Diz-Muñoz, EMBL Heidelberg) were cultured
549 adherently in DMEM (4.5 g/L D-Glucose) supplemented with 1× MEM NEAA (catalog no.
550 11140-035, Gibco), 1× GlutaMAX (catalog no. 35050-038, Gibco) and 10% (v/v) fetal bovine
551 serum (catalog no. 10270-106, Gibco) at 37°C under an atmosphere with 5% CO₂, 100%
552 humidity.

553 ***Sample preparation for superresolution imaging of clathrin-coated pits***

554 Cells were fixed as described previously (Li et al., 2018) using 3% (w/v) formaldehyde 10
555 mM MES, pH 6.1, 150 mM NaCl, 5 mM EGTA, 5 mM d-glucose, 5 mM MgCl₂ for 20 min.
556 Fixation was quenched in 0.1% (w/v) NaBH₄ for 7 min. The sample was washed three times
557 with PBS and permeabilized for 15 min with 0.01% (w/v) digitonin (Sigma-Aldrich, St. Louis,
558 MO, USA) in PBS. The sample was then washed twice with PBS and blocked for 1h with 2%
559 (w/v) BSA/PBS, washed with PBS, and stained for 3–12 h with anti-clathrin light chain (sc-
560 28276; Santa Cruz Biotechnology, Dallas, TX, USA) and anti-clathrin heavy chain rabbit
561 polyclonal antibodies (ab21679; Abcam, Cambridge, UK) in 1% (w/v) BSA/PBS. After three
562 washes with PBS the sample was incubated for 3-4h with a secondary donkey anti-rabbit
563 antibody (711-005-152; Jackson ImmunoResearch, West Grove, PA, USA) that was conjugated
564 to Alexa Fluor 647–NHS at an average degree of labeling of 1.5. The sample was then washed
565 three times and mounted for imaging in blinking buffer (50 mM Tris/HCl pH 8, 10 mM NaCl,
566 10% (w/v) D-glucose, 500 µg ml⁻¹ glucose oxidase, 40 µg ml⁻¹ glucose catalase, 35 mM MEA in
567 H₂O).

568 For the analysis of the disconnected population of sites, 3T3 cells were transfected with
569 a plasmid encoding the sigma 2 subunit, fused to GFP (a kind gift from Steeve Boulant,
570 University of Florida), to obtain cells transiently expressing AP2-GFP. The transfection was
571 performed using a Lipofectamine™ 2000 reagent (Life Technologies) according to the
572 manufacturer's recommendations: 1 µg DNA was mixed with 50 µL OptiMEM I
573 (ThermoFisher). The same was done for 3 µL Lipofectamin in 50 µL OptiMEM I. Both solutions
574 were incubated for 5 min and then mixed together and incubated for additional 10 min, at

575 room temperature. The media of previously seeded cells was exchanged to pre-warmed
576 OptiMEM I, to which the DNA-Lipofectamin solution (100 μ L) was dropwise added. After
577 approximately 24 hours of incubation (at 5% CO₂, 37 °C), the medium was exchanged with
578 fresh growth medium. After additional incubation for approximately 16 hours, cells were fixed
579 according to the same protocol described above.

580 ***Superresolution microscopy***

581 SMLM images were acquired at room temperature (24 °C) on a customized microscope
582 (Mund et al., 2018) with a 160x NA 1.43 oil-immersion objective (Leica, Wetzlar, Germany).
583 Illumination was done using a a LightHub laser combiner (Omicron-Laserage Laserprodukte,
584 Dudenhofen, Germany) with Luxx 405, 488, and 638 Cobolt 561 lasers, which were triggered
585 using a Mojo FPGA (Embedded Micro, Denver, CO, USA) for microsecond pulsing control of
586 lasers. The lasers were guided through a speckle reducer (LSR-3005-17S-VIS; Optotune,
587 Dietikon, Switzerland) and coupled into a multimode fiber (M105L02S-A; Thorlabs, Newton,
588 NJ, USA). The output of the fiber is magnified and imaged into the sample. Fiber generated
589 fluorescence was removed using a laser clean-up filter (390/482/563/640 HC Quad; AHF,
590 Tübingen, Germany). The focus was stabilized using a closed-loop system, based on reflecting
591 the signal of a near-infrared laser by the coverslip and detecting the resulting signal on a
592 quadrant photodiode, resulting in focus stabilities of ± 10 nm over several hours. Fluorescence
593 emission was filtered using a 676/37 or a 700/100 bandpass filter (AHF) and recorded by an
594 EMCCD camera (Evolve512D; Photometrics, Tucson, AZ, USA). Typically, 100,000–300,000
595 frames were acquired using 15-ms or 30 ms exposure times and laser power densities of ~ 15
596 kW/cm². 405-nm laser intensity was adjusted automatically by changing pulse duration in
597 order to keep the number of localizations per frame constant during the acquisition.

598 For the analysis of the disconnected population of sites, one diffraction limited image was
599 additionally acquired before SMLM imaging. For this, a 488 laser at 1.4 kW/cm² was used to
600 take a single frame at 30 ms exposure time. Emission was filtered using a 525/50 bandpass
601 filter (AHF).

602 ***Data analysis***

603 All data analysis was conducted in SMAP ((Ries, 2020) based on MATLAB and available as
604 open source under github.com/jries/SMAP).

605 ***Superresolution image reconstruction***

606 For fitting the localizations, peaks were detected as maxima in raw camera images after
607 applying a difference-of-Gaussian filter. At those positions, cropped images of 13x13 pixels
608 were fitted with an experimentally derived PSF model (free fitting parameters: x , y , z , photons
609 per localization, background per pixel), using an MLE (Maximum likelihood estimation) fitter
610 (Li et al., 2018). The x , y and z positions were corrected for residual drift by a custom algorithm
611 based on redundant cross-correlation. In short, the data was distributed into 10 time bins of
612 equal length. For each bin a superresolution image was reconstructed. We then calculated

613 the image cross-correlations among all superresolution images and extracted the relative
614 displacements in x and y were from the position of the maximum in the cross-correlation
615 images. We then calculated the drift trajectory that best describes the relative displacements.
616 In a second step, the z-drift was measured in an analogous way, using intensity profiles in z
617 instead of images. Localizations persistent over consecutive frames (detected within 35 nm
618 from one another and with a maximum gap of one dark frame) were merged into one
619 localization by calculating the weighted average of x, y and z positions and the sums of
620 photons per localization and background. Localizations were filtered by the localization
621 precision in x,y (0–20 nm) and z (0–30 nm) to exclude dim localizations. The modal value for
622 the localization precision σ was 3.9 nm in x/y and 12.5 nm in z, leading to a resolution
623 estimate (calculated using the Full Width Half Maximum (FWHM) using $\text{FWHM} = 2\sqrt{(2\ln 2)\sigma}$)
624 of 9.2 nm in x/y and 29.4 nm in z (typical values based on representative image).
625 Superresolution images were constructed with every localization rendered as a two-
626 dimensional spherical Gaussian with a width of 3 nm. The red hot color map used represents
627 the density of localizations, and is scaled in a way that 0.03% of the pixel values are saturated.

628 *Quantitative geometric analysis of clathrin-coated structures*

629 Clathrin-coated structures were segmented semi-automatically. First, we manually
630 defined a region of interest excluding the edges of the cell. Then, the image was blurred using
631 a Gaussian filter with a sigma of 100 nm, and peaks were detected using a manually set
632 threshold. This typically yielded several hundreds of sites in a region of 30x30 μm . These
633 candidate sites were curated manually, and only single, well-isolated clathrin-coated
634 structures were retained in the dataset.

635 Next, these structures were analyzed using LocMoFit, an MLE-based model fitting
636 framework that we developed recently (Wu et al., 2021). LocMoFit directly fits localization
637 coordinates with the probability density function (PDF) of a parametrized geometric model.
638 In this study, we modeled clathrin-coated structures with a hollow spherical cap
639 parameterized by the surface area A and the closing angle θ . θ is defined as the angle
640 between the two vectors that point to the pole and to the rim, respectively, from the center
641 of the sphere. The position of the model is defined as the center of mass of the cap. In
642 practice, we discretized the cap by placing spiral points (Saff and Kuijlaars, 1997), or the
643 spherical Fibonacci lattice, on the surface of the cap to approximate an even distribution. In
644 LocMoFit, these points were treated as discrete fluorophore coordinates when constructing
645 the PDF of the model. During fitting, additional parameters including the center position and
646 the orientation of the model were determined with respect to the fluorophore coordinates,
647 and an extra uncertainty and the background weight were applied to the PDF. After fitting,
648 the sphere radius is derived as $r = \sqrt{A/2\pi(1 - \cos \theta)}$, projected area as $A_p = \pi \sin^2 \theta$ and
649 edge length as $\varepsilon = 2\pi \sin \theta$. For some flat sites where the fit produced slightly negative
650 curvature values, curvature H and θ were set to 0 nm^{-1} and 0° respectively in order to
651 approximate them as completely flat.

652 After model fitting, a second curation step was performed. With this we ensured that only
653 well fitted sites are included in the final data set. Sites were excluded if they were upside
654 down (clearly originating from an upper membrane), double-sites with two sites clearly
655 connected to each other or an adjacent flat structure, large plaques or structures non-
656 distinguishable from an antibody cluster.

657 *Pseudo-temporal reconstruction of clathrin remodeling during endocytosis*

658 To sort endocytic sites in pseudotime, they were sorted by the closing angle θ , assigning
659 each site a rank index. Flat sites with a manually assigned $H = 0 \text{ nm}^{-1}$ and $\theta = 0$ were all
660 assigned an index of 0. As pseudo-temporal sorting assumes that all sites are part of the same
661 endocytic trajectory, endocytic sites with curvatures above a cell line specific threshold ($H >$
662 0.016 nm^{-1} for SK-MEL-2; $H > 0.013 \text{ nm}^{-1}$ for U2OS; $H > 0.014 \text{ nm}^{-1}$ for 3T3) that form a visibly
663 disconnected point cloud (Supplementary Figures 3 and 4) were excluded for this analysis
664 (Supplementary Figure 5). To compute pseudo-temporal averages, the sites were spatially
665 and rotationally aligned, and rescaled by the average radius of all sites within the respective
666 bin. Because the geometric model is rotationally symmetric, we then performed rotational
667 averaging by generating 72 duplicates of each structure, rotating them by 5 degrees with
668 respect to each other, and averaging them. The supplementary movie was computed using a
669 sliding window approach, where each frame shows the average of 30 sites, and the frame-to-
670 frame increment is 20 sites. The median pseudotime of those 30 sites is indicated.

671 *Further data analysis*

672 All data that resulted from the quantitative geometric description of clathrin-coated
673 structures were further analyzed in R (R Core Team, 2020). Fitting of the growth models
674 (Tables 1-3) was performed on a filtered data set, excluding disconnected sites above a cell
675 line specific threshold, and sites of negative curvature. During fitting, for sites with $\theta = 0^\circ$ we
676 set $\theta = 0.0001^\circ$ to avoid division by 0.

677 For depicting the growth models in Figure 3 and 4, as well as Supplementary Figure 5,
678 parameters resulting from the $H(\theta)$ fit were used and mapped to the A , ε and A_p data.

679 *Simulations*

680 Simulations were performed using the simulation engine for SMLM data implemented in
681 SMAP and LocMoFit as described in (Thevathasan et al., 2019). The realistic simulations were
682 based on a two-state (bright and dark) fluorophore model plus bleaching (Sage et al., 2019)
683 and parameters (number of photons, background photons, and fluorophore on-time t_l)
684 extracted from our experiment. (1) First, we defined an equally distributed closing angle θ
685 from 0 to 180° and calculated the surface area $A(\theta) = 2\pi(1 - \cos\theta)/H(\theta)^2$ (see
686 Supplementary Note for details). Here $H(\theta)$ is defined as in Equation 3, with fitting
687 parameters determined in SK-MEL-2 (see Table 1). (2) With the defined model parameters,
688 we generated protein positions for each simulated structure by taking randomly drawn $N \propto$
689 A samples from the PDF of the hollow spherical cap with no uncertainty. (3) With a probability

690 $p_{\text{label}} = 0.6$, a fluorescent label was created at a protein position. (4) Linkage displacements
691 in x , y and z were added to a label and were determined as normally distributed random
692 variables with a variance corresponding to the linkage error of 5 nm. The fluorophore is
693 assumed to be freely rotating three-dimensionally between different blinks. (5) Each
694 fluorophore appeared at a random time and lived for a time t_l , determined as a random
695 variable from an exponential distribution with a mean of 1.6 frames. (6) A label had a
696 probability $p_{\text{react}} = .5$ to be reactivated and then appeared at a random later time point,
697 otherwise it was bleached. (7) When it was on, a fluorophore had a constant brightness. Thus,
698 the brightness in each frame was proportional to the fraction of the time the fluorophore was
699 on in each frame. (8) The emitted photons in each frame were determined as a random
700 Poisson variable with a mean value corresponding to the average brightness in the frame. (9)
701 For each frame, we calculated the CRLB (Cramér-Rao lower bound) in x , y and z from the
702 number of photons (with a mean of 11,000) and the background photons (130 per pixel)
703 based on the theoretical Gaussian PSF (Mortensen et al., 2010) or a 3D cspline PSF model
704 derived from beads calibrations (Li et al., 2018). This error was added to the true x , y and z
705 positions of the fluorophores as normally distributed random values with a variance
706 corresponding to the respective calculated CRLB.

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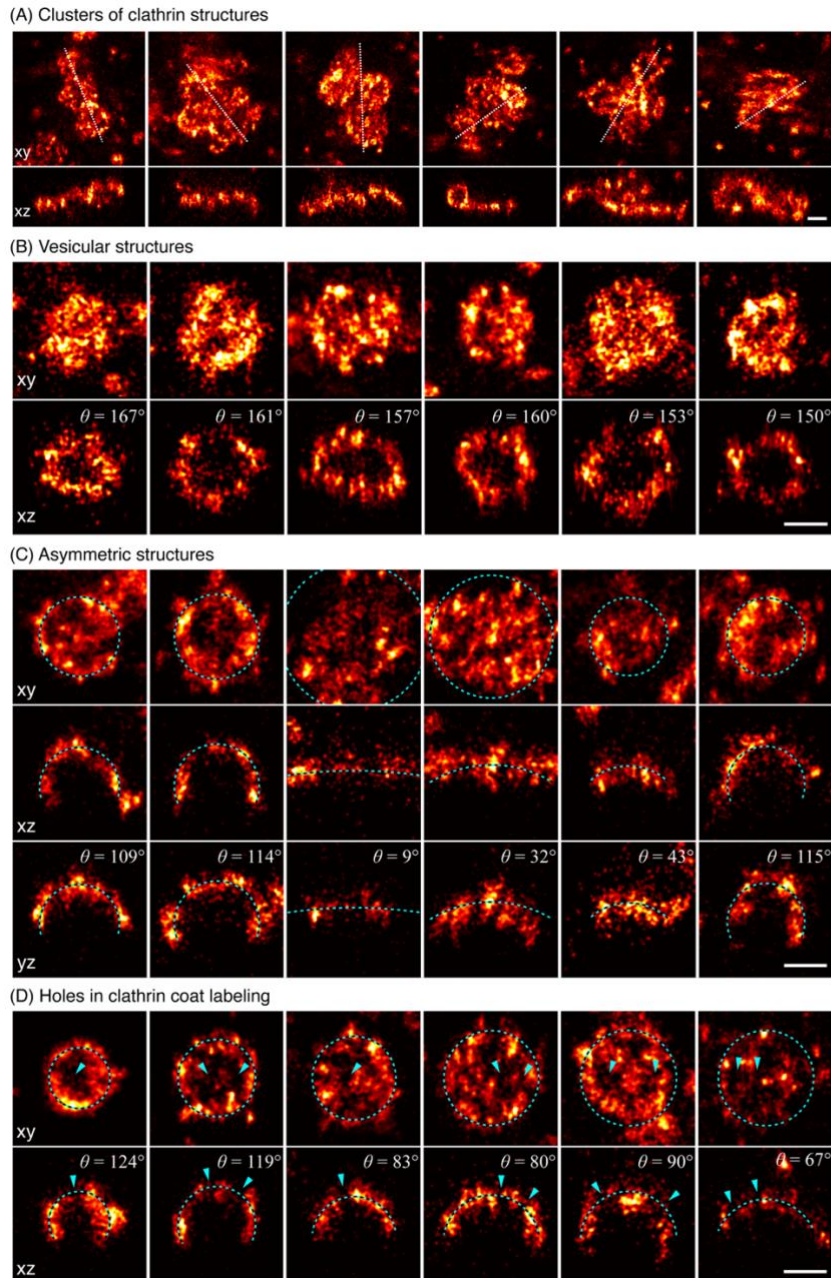
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850 **Supplementary Information**

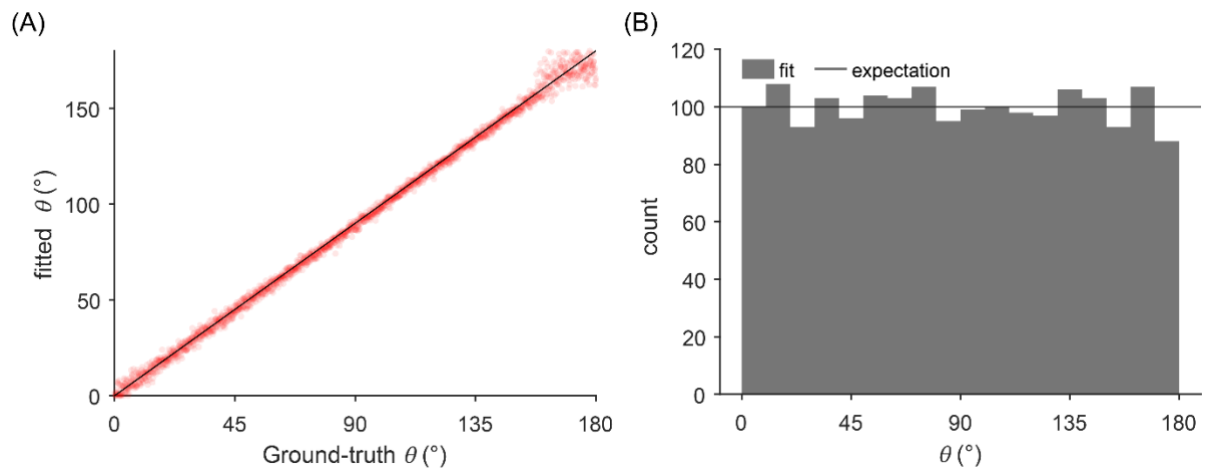


851

852 **Supplementary Figure 1 – Examples of diverse clathrin coat structures.** (A) Large clusters of clathrin molecules
853 excluded from further analysis. Shown in top view (xy) with dotted line indicating a 50 nm-thick z-slice (xz) shown
854 below. (B) Vesicular structures are sometimes fitted with a lower θ as expected. (C) While a spherical model
855 describes structure of most endocytic clathrin coats faithfully, there are few cases, as exemplified here, where
856 the elliptical and irregular shape of an assembling coat is difficult to approximate with a simple geometric model.
857 Two orthogonal 50 nm-thick z-slices are shown here in xz and yz, and the respective spherical model fit is plotted
858 as a dotted line. (D) Non-continuous labeling of clathrin manifests itself as holes in the coat, indicated with a blue
859 arrow. All scale bars are 100 nm.

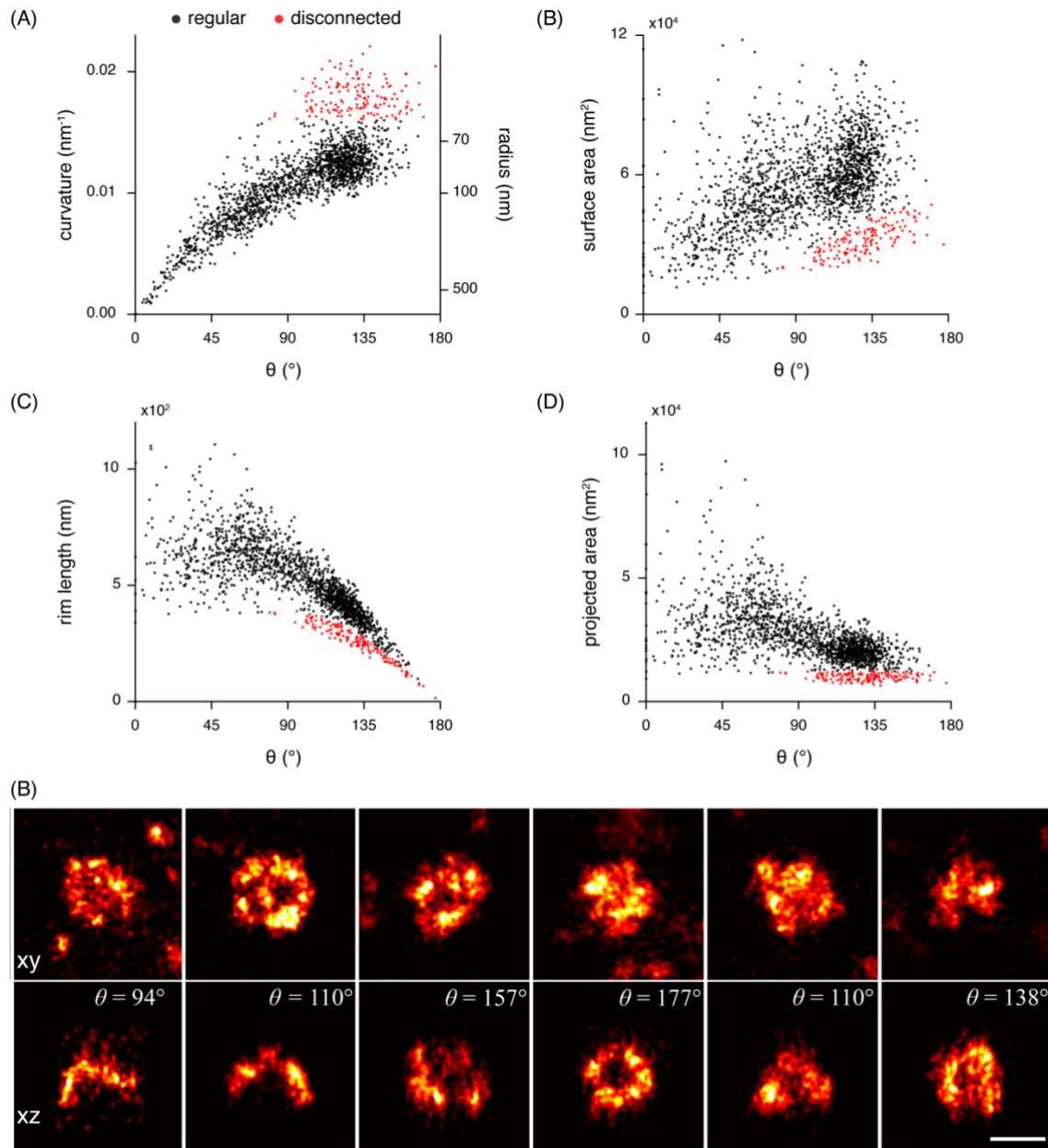
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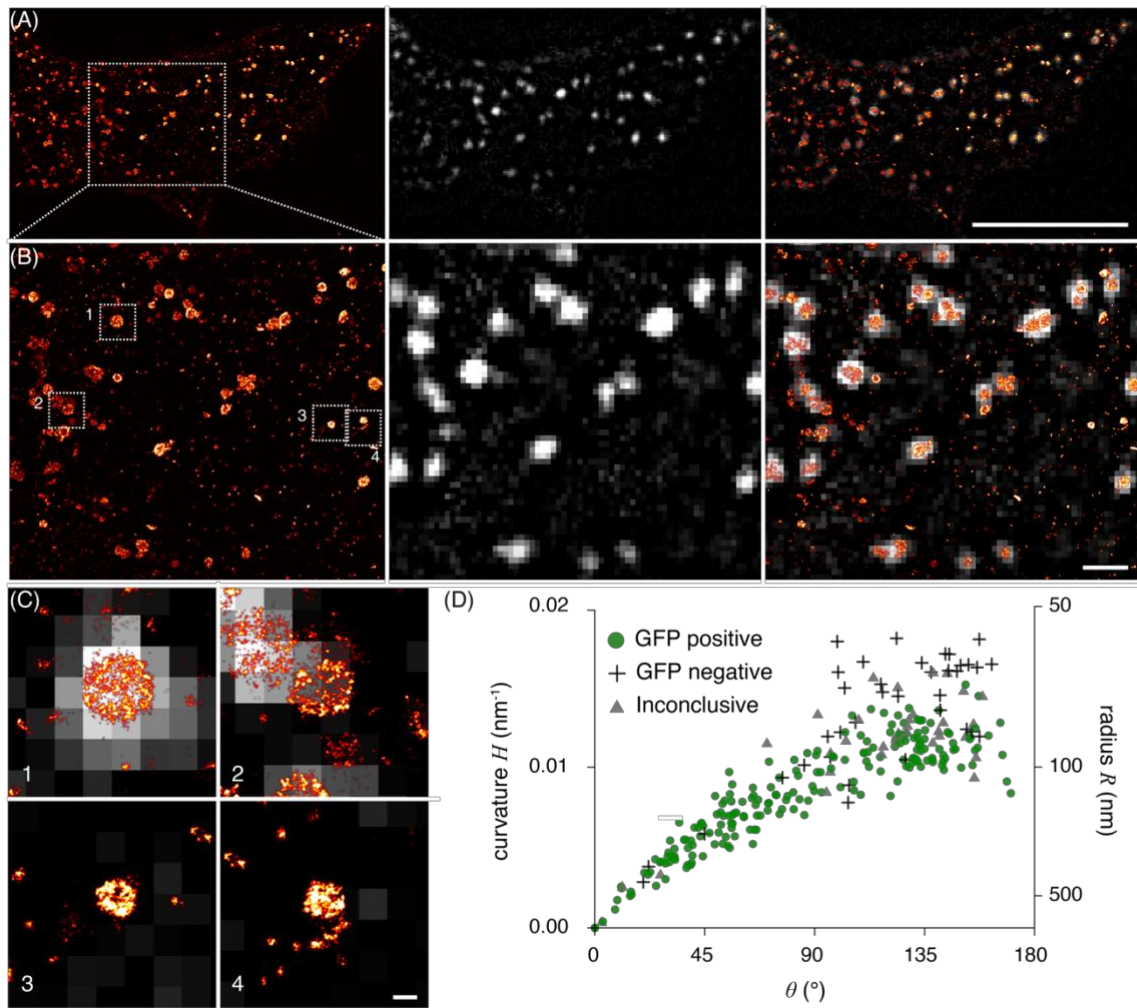
862

863 **Supplementary Figure 2 – Estimation error of the closing angle θ .** (A) The comparison of fitted θ of simulated
864 clathrin coat structures to corresponding ground-truth θ shows no systematic bias and a narrow spread of the
865 error across the ground-truth although the spread increases when $\theta < 20^\circ$ or $\theta > 160^\circ$. We reasoned that in
866 the earlier range, where the structures are flat, the slightly increased error corresponds to the insensitivity of θ
867 to flat structures. In the later range, where the vesicles are almost closed, the error was caused by undisguisable
868 tiny holes corresponding to the real vesicle openings and unlabeled clathrin in the coat. The fitted structures were
869 simulated to have similar quality as the experimental data and to distribute evenly across θ . (B) The distribution
870 of θ shows no significant error compared to the expectation corresponding to the evenly distributed θ , except
871 for the small potential underestimation of entirely closed coats.



872

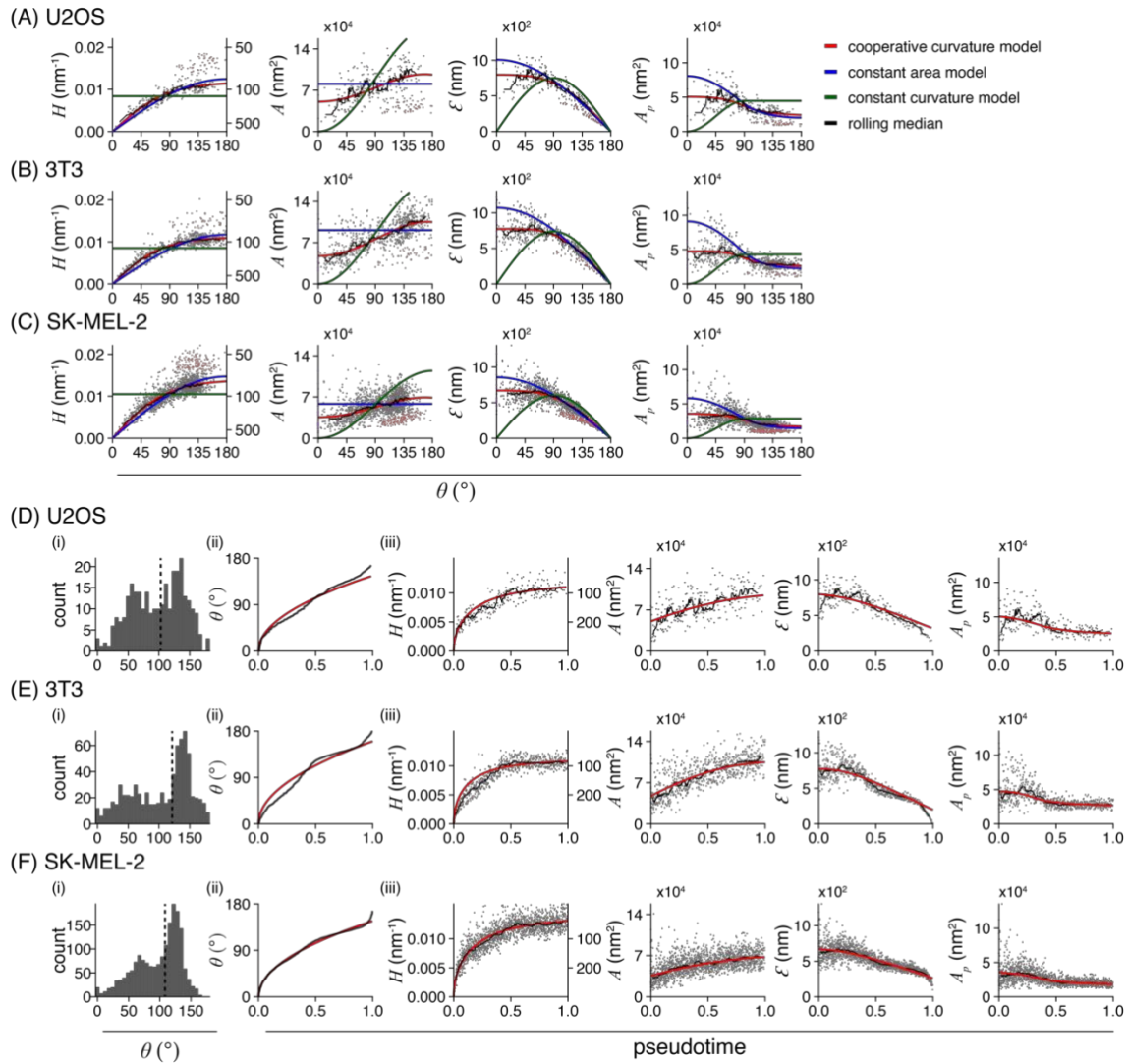
873 **Supplementary Figure 3 – Non-endocytic clathrin structures.** Results of LocMoFit analysis of clathrin structures
 874 in SK-MEL-2 cells ($N = 13$ cells, $n = 1798$ sites). (A) A strong correlation between curvature and θ can be observed
 875 for most structures ($n = 1645$ sites, black). A disconnected point cloud ($n = 153$ sites, 8.5%, red) indicates the
 876 presence of endocytosis-unrelated clathrin structures. The same distinct population of data points can be
 877 observed for the (B) surface area, (C) rim length, and (D) projected area. (B) Example structures from the
 878 disconnected population of sites in top view (xy) and 50 nm-thick z-slices (xz) and their respective θ values. Scale
 879 bar is 100 nm.



880

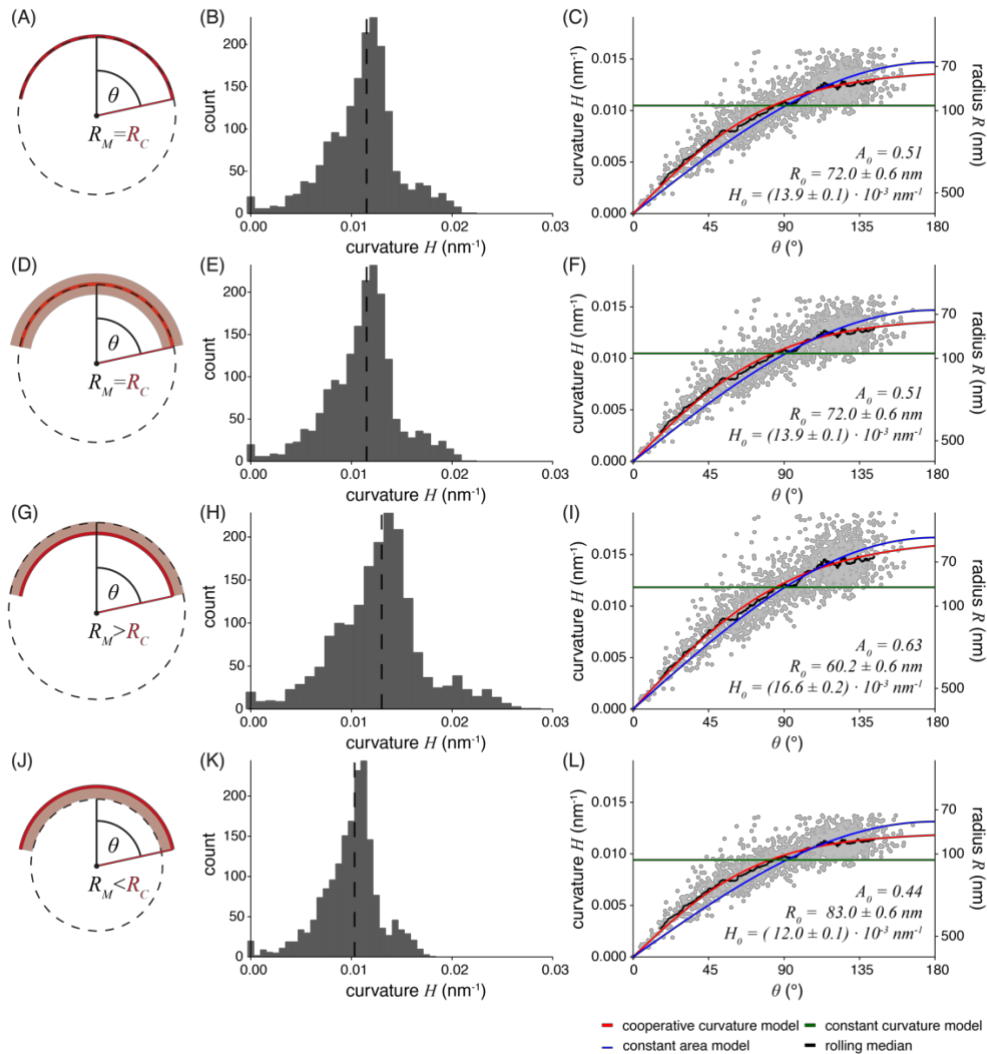
881 **Supplementary Figure 4 – Analysis of clathrin coats not following general trajectory of curvature generation.**

882 (A) 3T3 cell transiently overexpressing AP2-GFP. (left) Single-molecule localization microscopy image of
883 immunolabeled clathrin. (middle) Diffraction limited image of the AP2-GFP signal. (right) Overlay of the two
884 targets (scale bar: 10 μm). (B) Enlarged image of the section indicated in (A) (scale bar: 1 μm). (C) Example sites
885 indicated in (B) (scale bar: 100 nm). (1) Example for a structure annotated as “GFP positive”. (2) Example for an
886 “inconclusive” GFP signal. (3 and 4) Example of “GFP negative” structures”. (D) Analysis results when estimating
887 theta and curvature from clathrin structures, and annotating them depending on their AP2 signal. ($N = 3$ cells
888 and $n = 277$ sites). No AP2-GFP positive structures are found in the disconnected population of sites, suggesting
889 that they are most likely not generated via clathrin mediated endocytosis.



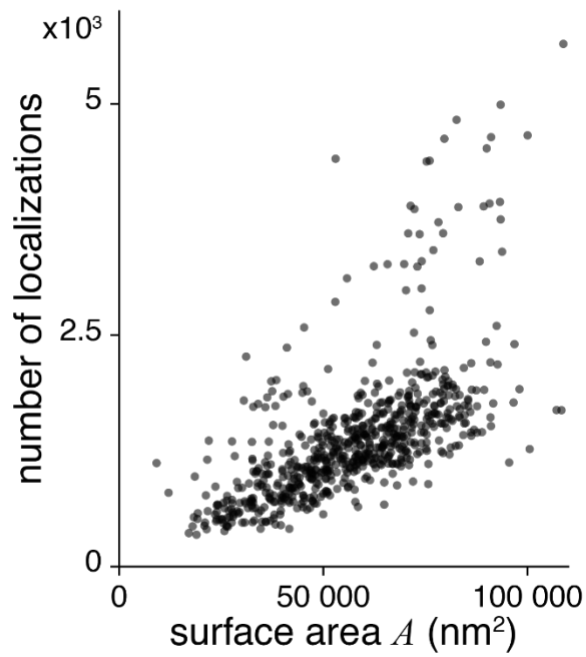
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891 **Supplementary Figure 5 – Clathrin coat remodeling in three different cell lines.** (A-C) Results of LocMoFit
 892 analysis for clathrin structures. Different growth models are fitted to curvature H over θ . The resulting fitting
 893 parameters are then used to map the same models also over surface area, rim length, and projected area (left
 894 to right). Purple: completely flat sites with $H = 0 \text{ nm}^{-1}$ and $\theta = 0$. Red: disconnected sites that were excluded from
 895 the fitting. Black line: rolling median (window width = 5% of total number of sites) (A) U2OS ($N = 3$ cell, $n_{\text{grey}} =$
 896 241 sites, $n_{\text{red}} = 53$ disconnected sites, $n_{\text{purple}} = 1$ completely flat sites), (B) 3T3 mouse fibroblasts ($N = 7$ cells, $n_{\text{grey}} =$
 897 688 sites, $n_{\text{red}} = 51$ disconnected sites, $n_{\text{purple}} = 8$ completely flat sites), and (C) SK-MEL-2 cells ($N = 13$ cells, $n_{\text{grey}} =$
 898 1631 sites, $n_{\text{red}} = 153$ disconnected sites, $n_{\text{purple}} = 14$ completely flat sites). (D-F) Temporal reconstruction of
 899 clathrin coat remodeling. (i) Distribution of θ slightly differs between cell lines, especially in the earlier states.
 900 Median θ shown as dotted lines correspond to 99.6° for U2OS; 121.4° for 3T3; and 108.5° for SK-MEL-2 cells. (ii)
 901 The cooperative curvature model (red line) highlights the square-root dependence between θ and pseudotime.
 902 (iii) The cooperative curvature model is used to describe the curvature H propagation over pseudotime. Resulting
 903 fitting parameters are then used to map the same model to surface area A , rim length ϵ and projected area A_p .
 904 A rolling median is plotted for in black (window width = 5% of total number of sites).



905

906 **Supplementary Figure 6 – Impact of linkage error on geometric model fit.** Indirect immunolabeling displaces
 907 the label from the target molecule by two antibodies. This generates a so-called linkage error of on average
 908 ± 10 nm, and resulting localizations might not accurately represent the underlying structure of interest (Früh et
 909 al., 2021). (A) Ideal case of no linkage error, where the model (dotted black line) with radius R_M accurately
 910 represents the underlying clathrin coat (red) with a radius R_C . (B) Histogram of quantified curvature with a
 911 median of 0.011 nm^{-1} . (C) Models fitted to curvature propagation over theta θ . (D) Uniform displacement of
 912 localizations (light red, ± 10 nm) due to unbiased labelling by the antibodies. The radius R_M still accurately
 913 represents the true radius R_C . (E) Histogram of the curvature, assuming no needed correction of R_M . Median:
 914 0.011 nm^{-1} . (F) Models fitted to curvature propagation over theta θ remain the same as for (C). (G) Biased labeling
 915 of the antibodies (light red, $+ 10$ nm) could result in an overestimation of R_M by 10 nm. (H) Histogram of the
 916 curvature corrected by subtracting 10 nm from quantified R_M . Median: 0.013 nm^{-1} . (I) Models fitted to corrected
 917 curvature propagation over theta θ . (J) Biased labeling could result in an underestimation of R_M by 10 nm. (K)
 918 Histogram of the curvature corrected for the overestimation in radius by adding 10 nm from quantified R_M .
 919 Median: 0.010 nm^{-1} . (L) Models fitted to corrected curvature propagation over theta. (C; F; I; L) Fitting parameters
 920 are A_0 : Fraction of surface area growing as a flat lattice before curvature initiation, defined as $A_0 = A(\theta =$
 921 $0.01)/A(\theta = \pi)$; R_0 : preferred radius of the clathrin coat fitted with CoopCM; H_0 : Preferred curvature of the
 922 clathrin coat fitted with CoopCM. While the fitting parameters scale with the error in radius estimation, the
 923 relationships among the parameters and thus our mechanistic interpretation by the cooperative curvature model
 924 still holds true.

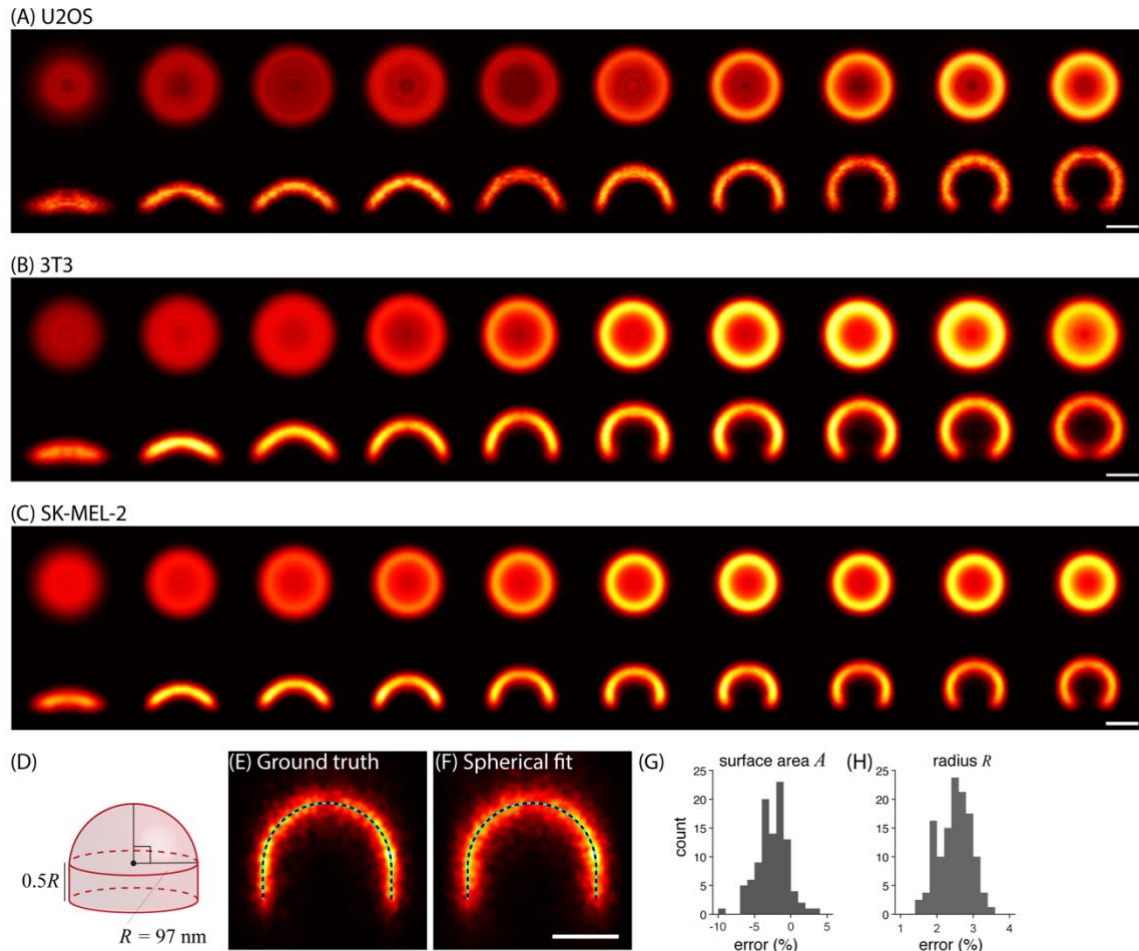


925

926 **Supplementary Figure 7 – Number of localizations versus surface area.** For $N = 6$ SK-MEL-2 cells, $n = 700$ sites

927 the number of localizations found in one clathrin coated structure was extracted. This is plotted against the

928 quantified surface area determined for each coat.



929

930 **Supplementary Figure 8 – Averaging of clathrin coated structures.** (A-C) Averages for distinct endocytic stages,
 931 resulting from all collected snapshots for (A) U2OS ($n = 241$ sites, $N = 3$ cells), (B) 3T3 ($n = 688$ sites, $N = 7$ cells),
 932 and (C) SK-MEL-2 cells ($n = 1645$ sites, $N = 13$ cells). Each bin contains the same number of snapshots of clathrin-
 933 coated structures sorted along their pseudotime ($n_{U2OS} = 24$ per bin, $n_{3T3} = 68$, $n_{SK-MEL-2} = 164$), so that all bins
 934 represent equally long pseudotime intervals. (D-H) Averaging preserves U-shapes in simulated endocytic sites.
 935 (D) A ground-truth structure was used for simulating U-shaped clathrin coats. The model for simulation was built
 936 by combining a hemisphere with a radius of $R = 97 \text{ nm}$ and a cylinder with the height $D = 0.5 R$. We choose the
 937 radius value according to the median radius of bin 6 in Figure 4G. This bin has a closing angle slightly larger than
 938 90° . 20-nm thick cross-sections of the averages ($N = 100$ sites) registered based on (E) the ground truth and (F)
 939 the spherical fit are shown with the ground-truth U-shaped model (dotted line). Histograms of normalized
 940 estimation errors are shown for the parameters (G) surface area and (H) radius, with mean values of -2.6% and
 941 2.5% respectively. The scale bar is 100 nm .

942

943

944 **Table 1 – Summary of clathrin coat growth model fits in SK-MEL-2.** Fitted parameter values for the constant
 945 area model (CAM), the constant curvature model (CCM) and the cooperative curvature model (CoopCM) when
 946 fitting curvature $H(\theta)$, surface area $A(\theta)$, rim length $\mathcal{E}(\theta)$ and projected area $A_p(\theta)$. A: Surface area fitted with
 947 CAM; R: Radius fitted with CCM; γ : Constant rate of curvature increase fitted with CoopCM; H_0 : Preferred
 948 curvature of the clathrin coat fitted with CoopCM; R_0 : preferred radius of the clathrin coat fitted with CoopCM;
 949 A_0 : Fraction of surface area growing as a flat lattice before curvature initiation, defined as $A_0 = A(\theta =$
 950 $0.01)/A(\theta = \pi)$; k_{on} : local growth rate obtained from $\theta(t)$ fitted with CoopCM and measured in nm per
 951 pseudotime units \tilde{s}^{-1} . $N = 13$ cells; $n = 1645$ sites.

		Curvature $H(\theta)$	Surface area $A(\theta)$	Rim length $\mathcal{E}(\theta)$	Projected area $A_p(\theta)$
CAM	A	$58300 \pm 400 \text{ nm}^2$	$56600 \pm 400 \text{ nm}^2$	$49400 \pm 500 \text{ nm}^2$	$48800 \pm 500 \text{ nm}^2$
CCM	R	$94.5 \pm 0.7 \text{ nm}$	$84.8 \pm 0.4 \text{ nm}$	$97.3 \pm 0.8 \text{ nm}$	$92.8 \pm 0.7 \text{ nm}$
CoopCM	γ	$(9.4 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(9.0 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(9.4 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(9.1 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$
	H_0	$(13.9 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(13.8 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(13.4 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(13.6 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$
	R_0	$72.0 \pm 0.6 \text{ nm}$	$72.2 \pm 0.6 \text{ nm}$	$74 \pm 0.8 \text{ nm}$	$73.4 \pm 0.9 \text{ nm}$
	A_0	0.51	0.55	0.49	0.52
	k_{on}	$78.1 \text{ nm}\tilde{s}^{-1}$	$72.2 \text{ nm}\tilde{s}^{-1}$	$83.4 \text{ nm}\tilde{s}^{-1}$	$78 \text{ nm}\tilde{s}^{-1}$

952 **Table 2 – Summary of clathrin coat growth model fits in U2OS.** Fitted parameter values for the constant area
 953 model (CAM), the constant curvature model (CCM) and the cooperative curvature model (CoopCM) when fitting
 954 curvature $H(\theta)$, surface area $A(\theta)$, rim length $\mathcal{E}(\theta)$ and projected area $A_p(\theta)$. A: Surface area fitted with CAM;
 955 R: Radius fitted with CCM; γ : Constant rate of curvature increase fitted with CoopCM; H_0 : Preferred curvature of
 956 the clathrin coat fitted with CoopCM; R_0 : preferred radius of the clathrin coat fitted with CoopCM; A_0 : Fraction of
 957 surface area growing as a flat lattice before curvature initiation, defined as $A_0 = A(\theta = 0.01)/A(\theta = \pi)$; k_{on} :
 958 local growth rate obtained from $\theta(t)$ fitted with CoopCM and measured in nm per pseudotime units \tilde{s}^{-1} . $N = 3$
 959 cells; $n = 241$ sites.

		Curvature $H(\theta)$	Surface area $A(\theta)$	Rim length $\mathcal{E}(\theta)$	Projected area $A_p(\theta)$
CAM	A	$80800 \pm 1600 \text{ nm}^2$	$76900 \pm 1700 \text{ nm}^2$	$65900 \pm 1600 \text{ nm}^2$	$65400 \pm 1600 \text{ nm}^2$
CCM	R	$119.1 \pm 2.5 \text{ nm}$	$101.6 \pm 1.5 \text{ nm}$	$122.8 \pm 2.7 \text{ nm}$	$115 \pm 2.4 \text{ nm}$
CoopCM	γ	$(7.9 \pm 0.2) \cdot 10^{-3} \text{ nm}^{-1}$	$(7.7 \pm 0.2) \cdot 10^{-3} \text{ nm}^{-1}$	$(8.1 \pm 0.2) \cdot 10^{-3} \text{ nm}^{-1}$	$(7.9 \pm 0.2) \cdot 10^{-3} \text{ nm}^{-1}$
	H_0	$(11.8 \pm 0.2) \cdot 10^{-3} \text{ nm}^{-1}$	$(11.5 \pm 0.2) \cdot 10^{-3} \text{ nm}^{-1}$	$(11.0 \pm 0.3) \cdot 10^{-3} \text{ nm}^{-1}$	$(11.1 \pm 0.3) \cdot 10^{-3} \text{ nm}^{-1}$
	R_0	$85.0 \pm 1.8 \text{ nm}$	$87.1 \pm 1.7 \text{ nm}$	$90.5 \pm 2.7 \text{ nm}$	$90.2 \pm 2.6 \text{ nm}$
	A_0	0.52	0.52	0.45	0.47
	k_{on}	$89.3 \text{ nm}\tilde{s}^{-1}$	$91.5 \text{ nm}\tilde{s}^{-1}$	$110.3 \text{ nm}\tilde{s}^{-1}$	$105.5 \text{ nm}\tilde{s}^{-1}$

960 **Table 3 – Summary of clathrin coat growth model fits in 3T3 mouse fibroblasts.** Fitted parameter values for the
 961 constant area model (CAM), the constant curvature model (CCM) and the cooperative curvature model (CoopCM)
 962 when fitting curvature $H(\theta)$, surface area $A(\theta)$, rim length $\mathcal{E}(\theta)$ and projected area $A_p(\theta)$. A: Surface area
 963 fitted with CAM; R: Radius fitted with CCM; γ : Constant rate of curvature increase fitted with CoopCM; H_0 :
 964 Preferred curvature of the clathrin coat fitted with CoopCM; R_0 : preferred radius of the clathrin coat fitted with
 965 CoopCM; A_0 : Fraction of surface area growing as a flat lattice before curvature initiation, defined as $A_0 = A(\theta =$
 966 $0.01)/A(\theta = \pi)$; k_{on} : local growth rate obtained from $\theta(t)$ fitted with CoopCM and measured in nm per
 967 pseudotime units \tilde{s}^{-1} . $N = 7$ cells; $n = 688$ sites.

		Curvature $H(\theta)$	Surface area $A(\theta)$	Rim length $\mathcal{E}(\theta)$	Projected area $A_p(\theta)$
CAM	A	$90100 \pm 1000 \text{ nm}^2$	$83700 \pm 1000 \text{ nm}^2$	$65800 \pm 1000 \text{ nm}^2$	$66200 \pm 1000 \text{ nm}^2$
CCM	R	$115.8 \pm 1.5 \text{ nm}$	$98.3 \pm 0.7 \text{ nm}$	$121.1 \pm 2 \text{ nm}$	$108.5 \pm 1.5 \text{ nm}$
CoopCM	γ	$(8.2 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(7.8 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(7.9 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(7.8 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$
	H_0	$(11.3 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(11.2 \pm 0.1) \cdot 10^{-3} \text{ nm}^{-1}$	$(11.2 \pm 0.2) \cdot 10^{-3} \text{ nm}^{-1}$	$(11.2 \pm 0.2) \cdot 10^{-3} \text{ nm}^{-1}$
	R_0	$88.8 \pm 0.7 \text{ nm}$	$89.3 \pm 0.7 \text{ nm}$	$89.1 \pm 1.4 \text{ nm}$	$89.5 \pm 1.3 \text{ nm}$
	A_0	0.45	0.49	0.47	0.49
	k_{on}	$128.4 \text{ nm}\tilde{s}^{-1}$	$120.8 \text{ nm}\tilde{s}^{-1}$	$123.9 \text{ nm}\tilde{s}^{-1}$	$121 \text{ nm}\tilde{s}^{-1}$

968 **Supplementary data**

969 Supplementary data can be downloaded from
970 <https://oc.embl.de/index.php/s/sszVT1ZAUB2LOFW> and will be uploaded to the BioStudies
971 image data repository (<https://www.ebi.ac.uk/biostudies/>) upon publication.

972

973 ***Supplementary movie (mp4 format, 1 file)***

974 Pseudo-temporal movie of clathrin coat remodeling during endocytosis in SK-MEL-2 cells.
975 Each frame corresponds to a sliding window average of 30 sites, with a frame-to-frame
976 increment of 20 sites. The median pseudotime of those 30 sites is indicated.

977

978 ***Localization data (csv format, 23 files)***

979 Localization coordinates for all superresolution images, grouped by cell line (SK-MEL-2, 13
980 files; U2OS, 3 files; 3T3, 7 files).

981 The localizations were grouped (localizations that persisted over several adjacent frames and
982 thus resulted from the same single fluorophore were combined into one localization) and
983 filtered with a maximum localization precision of 20 nm in xy, and 30 nm in z. The first 2000
984 frames of each acquisition are excluded, since the blinking density is too high for reliable
985 localization. Each file contains the columns *xnm* (x coordinate in nm); *ynm* (y coordinate in
986 nm); *znm* (z coordinate in nm); *locprecnm* (localization precision in xy in nm); *locprecznm*
987 (localization precision in z in nm); *frame*; *site_ID* (marks all localizations belonging to the
988 respective segmented clathrin structure; localizations outside of analyzed sites are put to 0);
989 *filename* (identifier for acquisition file, unique for each cell line).

990 Identifiers for acquisition files and clathrin-coated structures are kept consistent between all
991 supplementary data.

992

993 ***Model fit results (csv format, 23 files)***

994 Results from LocMoFit analysis of individual clathrin coated structures, grouped by cell line.
995 Columns are *ID* (identifier for individual structures); *file_number* (identifier for each
996 acquisition file); *cell_line*; *disconnected_sites* (true for those sites that are disconnected from
997 the main data cloud, see Supplementary Figures 2, 3). Model fit parameters calculated for
998 each structure using LocMoFit include theta θ ; radius R ; curvature H ; surface_area A ;
999 rim_length ϵ ; and projected_area A_p . Further included in the table are the corrected values
1000 theta_corrected and curvature corrected, where each structure fitted with a negative radius
1001 was manually put to $\theta = 0.0001^\circ$ and curvature = 0 nm⁻¹.

1002

1003 ***Site gallery (PDF format, 3 files)***

1004 Snapshots of all analyzed clathrin coated structure and their respective model fit results.

1005 For each structure, shown are the top view (left), as well as 50 nm side-view cross-sections
1006 through the structure at angles 0° , 45° , 90° and 135° (left to right). Shown are site number,
1007 theta θ , surface area A , and radius R . Scale bar is 100 nm.

1008

1009 ***Movie gallery (mp4 format, 49 files)***

1010 Rotating movies of the 3D SMLM data (left) and renderings of the corresponding models
1011 (right) of all clathrin-coated structures in SK-MEL-2 cells. Shown are site identifiers and model
1012 fit results (theta θ , surface area A , and radius R). Scale bar is 100 nm.

Supplementary note to: Superresolution microscopy reveals partial preassembly and subsequent bending of the clathrin coat during endocytosis

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Marko Kaksonen, Ori Avinoam, Ulrich S. Schwarz, and Jonas Ries

I. COOPERATIVE CURVATURE MODEL

We derive a kinetic model for clathrin-mediated endocytosis that is based on minimal assumptions based on our experimental observations. We first assume that the area A of clathrin coats grows mainly by addition of new triskelia at the edges [1]. With a local growth rate k_{on} and an edge length \mathcal{E} , we have the simple growth law

$$\frac{d}{dt}A = \dot{A} = k_{\text{on}}\mathcal{E}. \quad (1)$$

We also know from our experimental observations that all clathrin-coated pits have the geometry of a spherical cap with area $A_{\text{cap}} = 2\pi R^2(1 - \cos\theta)$, where R is the cap radius and θ the closing angle. The edge length then is $\mathcal{E}_{\text{cap}} = 2\pi R \sin\theta$. Using these formulas on Eq. (1) we find

$$2\dot{R} \tan \frac{\theta}{2} + R\dot{\theta} = k_{\text{on}}. \quad (2)$$

Because we parametrize the shape of the growing clathrin coat by the two dynamical variables $R(t)$ and $\theta(t)$, but have only one growth equation up to now, we have to make additional assumptions to completely define our kinetic model.

As the main function of clathrin is to generate curvature, we now focus on coat curvature $H = 1/R$ and assume that it increases with a basal rate γ . Experimentally we observe that clathrin patches first grow flat and then start to curve. Curvature generation in clathrin lattices is the combination of a preferred curvature of the single triskelion and cooperative effects in the lattice [2, 3]. Several mechanisms exist that might generate curvature during coat growth. First, clathrin triskelia are likely to be geometrically frustrated within the flat clathrin coat. To overcome this frustration, they would have to adjust their positions relative to their neighbors, such that their free energy is getting more favorable. Second, lattice vacancies could be filled up with new triskelia, as predicted theoretically [4]. Note that these vacancies would not necessarily relate to patch area, because the clathrin lattice consists of many overlapping arms. Filling of vacancies would however increase clathrin density and coat stiffness, and thus could also drive invagination of the coat [5]. We note that it was indeed confirmed experimentally that the clathrin density increases within the coat during invagination [2, 6]. Third, lattice pentagons, associated with lattice curvature, could form at the edge of the clathrin lattice and diffuse into the coat, similar to lattice defects on curved surfaces [7]. All three mechanisms could generate curvature with a certain rate which we assume to be constant during the initial stages of growth.

At late stages, increase of curvature has to stop and therefore we assume curvature saturation at a characteristic value H_0 , which is similar to, but different from the radius measured for clathrin cages. A simple estimate would be 40 nm for a typical radius of the membranes in the pits [8] plus 15 nm thickness of the clathrin coat [9]. These 55 nm would be larger than the 40 nm for clathrin cages, which are expected to be frustrated. If all triskelia in the lattice had achieved their optimal positions, a preferred or spontaneous curvature H_0 should emerge. Thus a growth equation for H should have a stable fixed point generated by a higher order term. Since the corresponding mechanism is known to be of cooperative nature, larger groups of triskelia should be involved and the mechanism should start to dominate relatively late in the process, but then dominate quickly. Here we assume that the mechanism is proportional to H^2 (rather than to H , with a linear law corresponding to a non-cooperative effect that starts to show early). Assuming that curvature generation is a geometrical effect, its time development should depend on the closing angle and we get

$$\frac{dH}{d\theta} = \gamma \left(1 - \frac{H^2}{H_0^2} \right). \quad (3)$$

The stable fixed point at $H = H_0$ can then be interpreted as corresponding to the preferred curvature of the mature coat.

Eq. (3) can be solved with the initial condition $H(\theta = 0) = 0$ by

$$H(\theta) = H_0 \tanh \left(\frac{\gamma}{H_0} \theta \right), \quad (4)$$

TABLE I. Coat curvature, coat area, and edge length summarized for the CCM, the CAM, and the CoopCM.

	CCM	CAM	CoopCM
H	$1/R_{\text{CCM}}$	$\sqrt{\frac{2\pi(1-\cos\theta)}{A_{\text{CAM}}}}$	$1/R_{\text{coat}}$
A	$2\pi R_{\text{CCM}}^2(1-\cos\theta)$	A_{CAM}	$2\pi R_{\text{coat}}^2(1-\cos\theta)$
\mathcal{E}	$2\pi R_{\text{CCM}} \sin\theta$	$\sqrt{\frac{2\pi A_{\text{CAM}}}{(1-\cos\theta)}} \sin\theta$	$2\pi R_{\text{coat}} \sin\theta$

and defines the coat radius as a function of the closing angle

$$R_{\text{coat}} = R(\theta) = \frac{1}{H(\theta)} = R_0 \frac{1}{\tanh(\gamma R_0 \theta)}, \quad (5)$$

with $R_0 = 1/H_0$. We note that the expansion of Eq. (5) gives $R \propto 1/\theta$ in leading order around the flat state.

We can combine Eq. (2) and Eq. (3) to find the full dynamics of coat invagination. Therefore, we compute the derivative of R_{coat}

$$\frac{dR_{\text{coat}}}{d\theta} = -\frac{\gamma R_0^2}{\sinh^2(\gamma R_0 \theta)}. \quad (6)$$

We use Eq. (5) and Eq. (6) and the chain rule $dR/dt = dR/d\theta d\theta/dt$ to rewrite Eq. (2). Solving for $\dot{\theta}$ yields

$$\dot{\theta} = k_{\text{on}} \frac{1}{2 \frac{dR(\theta)}{d\theta} \tan \frac{\theta}{2} + R(\theta)}. \quad (7)$$

After expanding Eq. (7) up to leading order in θ we find

$$\dot{\theta} = \frac{12\gamma k_{\text{on}}}{8\gamma^2 R_0^2 - 1} \frac{1}{\theta}, \quad (8)$$

which is solved with the initial condition $\theta(t=0) = 0$ by

$$\theta(t) = \sqrt{\frac{24\gamma k_{\text{on}}}{8\gamma^2 R_0^2 - 1}} t. \quad (9)$$

Thus the coat initially is flat ($\theta = 0$) and then starts to generate curvature, increasing the closing angle with a square root dependence in time, which reflects the slowing down of curvature generation as preferred curvature is approached. We refer to this model as the cooperative curvature model (CoopCM) due to the assumption of a non-linear growth law.

II. FIT RESULTS

To contrast the CoopCM with existing models, we fit it to experimental data for $H(\theta)$, $A(\theta)$ and $\mathcal{E}(\theta)$. We compare the fitted curves to the constant curvature model (CCM), defined through the constant coat curvature $1/R_{\text{CCM}}$, and to the constant area model (CAM), defined through the constant coat area A_{CAM} . The equations for the curvature $H(\theta) = 1/R(\theta)$, the area $A(\theta)$, and the edge length $\mathcal{E}(\theta)$ are summarized in Table I for all three models. In order to fit, we first filter the data according to curvature H , as described in the methods section of the main text. We then fit the different models to each of the data sets separately. For the CCM and CAM we get values for one parameter each, namely R_{CCM} and A_{CAM} , respectively. For the CoopCM we obtain two parameter values γ and H_0 . For the CoopCM we also determine the area when invagination occurs $A_0 = A(\theta = 0.01)/A(\theta = \pi)$, which is the relative transition size where the flat-to-curved-transition occurs. By definition, invagination in the CCM starts simultaneously with coat assembly ($A_0 = 0$), while invagination in the CAM starts at the maximum area ($A_0 = 1$). Moreover, for the CoopCM we also compute R_0 . For all parameters we also determine the relative fit errors, given by σ_x/x , where σ_x is the standard deviation of the parameter x and x is the value of the parameter. This procedure allows us to compare the fit results to each other. The fitted parameter values for the different models and cell lines are summarized in the Supplementary Tables 1–3.

We then fit Eq. (9) to the data, describing $\theta(t)$. Using the values of γ and H_0 , we determine k_{on} from the fit. The resulting values, measured in units of the pseudo time \bar{s} , are summarized in the Supplementary Tables 1–3. We do not show the fit errors of k_{on} , because they are small and result from the same fit of $\theta(t)$ and the errors of γ and H_0 , which does not allow us to discriminate between the different fits.

TABLE II. BIC for the curvature fits $H(\theta)$ of the CCM, CAM and CoopCM for the analyzed cell lines.

	CCM	CAM	CoopCM
SK-MEL-2 cells	-19079	-21218	-21670
c3T3 cells	-8119	-9179	-9498
U2OS cells	-2784	-3097	-3190

III. BAYESIAN INFORMATION CRITERION

Apparently, the CoopCM agrees better with the data compared to the CCM and the CAM. However, while the CCM and the CAM both only include one parameter, the CoopCM has two parameters. In order to base model selection on a statistical criterion and to account for the additional parameter of the CoopCM, we compute the Bayesian information criterion (BIC). The BIC takes into account both how well a model fits the data but also how many free parameters it includes to avoid overfitting. The BIC can be expressed by [10]

$$\text{BIC} = N \ln \left(\frac{\text{RSS}}{N - k} \right) + k \ln N \approx N \ln \left(\frac{\text{RSS}}{N} \right) + k \ln N, \quad (10)$$

where we used that $N \gg k$ in the last step. Here RSS is the residual sum of squares $\text{RSS} = \sum_i^N (f(x_i) - y_i)^2$ with x_i and y_i the measured data and $f(x_i)$ the fitted data, N is the number of data points and k is the number of free parameters. In general, when comparing different models, the model with the lowest BIC is preferred. In Table II the BIC is calculated from Eq. (10) for the curvature fits $H(\theta)$ for the different cell lines. For all cell lines, the BIC for the CoopCM is lowest. Therefore, we conclude that the additional parameter of the CoopCM is justified and that the CoopCM agrees best with the data.

IV. LINEAR CURVATURE MODEL

As an alternative to the CoopCM model we now consider a linear curvature model. As in the CoopCM model, we assume that the coat curvature $H = 1/R$ increases at a basal rate γ . In contrast to the CoopCM model however, we now assume that the increase in curvature at late stages is slowed down proportional to H and must stop at a value H_0

$$\frac{dH}{d\theta} = \gamma \left(1 - \frac{H}{H_0} \right). \quad (11)$$

Eq. (11) can be solved by

$$H(\theta) = H_0 \left(1 - e^{-\frac{\gamma}{H_0} \theta} \right). \quad (12)$$

Eq. (12) also defines the coat radius $R = 1/H$ as a function of the invagination angle θ

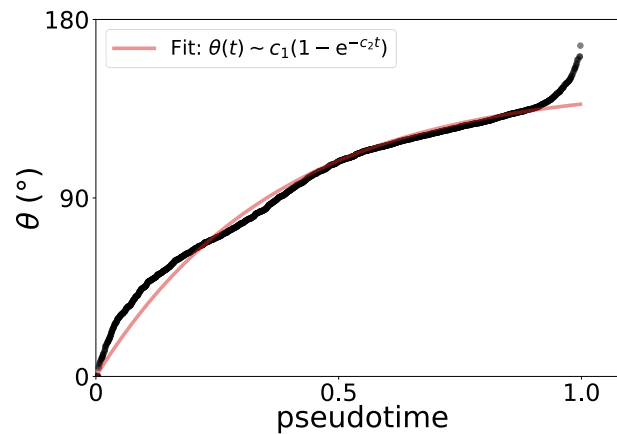
$$R(\theta) = R_0 \frac{1}{(1 - e^{-\gamma R_0 \theta})}, \quad (13)$$

with $R_0 = 1/H_0$. The derivative of $R(\theta)$ reads

$$\frac{dR}{d\theta} = -\frac{\gamma R_0^2}{4 \left(\sinh \frac{\gamma R_0 \theta}{2} \right)^2}. \quad (14)$$

Now we use Eq. (13) and Eq. (14) on Eq. (7). After expanding up to second order in θ we find

$$\dot{\theta} = \frac{2k_{\text{on}}}{R_0} \left(1 - \frac{1}{6} \left(2\gamma R_0 - \frac{1}{\gamma R_0} \right) \theta \right), \quad (15)$$



Supplementary Figure 9. Invagination angle θ as a function of the pseudotime t . Red line: the fit to the same data as in Fig. 4B according to Eq. (16) for the linear curvature model.

which is solved by

$$\theta(t) = \frac{1}{R_0} \left(\frac{3\gamma}{2\gamma^2 - \frac{1}{R_0^2}} \right) \left(1 - e^{-\left(\frac{2\gamma^2 - \frac{1}{R_0^2}}{3\gamma} \right) k_{\text{on}} t} \right). \quad (16)$$

For small t , Eq. (16) can be expanded to $\theta(t) \sim (k_{\text{on}}/R_0)t$. The linear increase of θ with t does not fit the data for small t (cf. Supplementary Figure 9).

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