Title
Mass-sensitive particle tracking (MSPT) to elucidate the membrane-associated MinDE reaction cycle

Authors
Tamara Heermann¹,², Frederik Steiert¹,², Beatrice Ramm¹,³, Nikolas Hundt⁴,⁵,∗ and Petra Schwille¹
# equal contribution

Affiliations
¹Department of Cellular and Molecular Biophysics, Max Planck Institute of Biochemistry, Am Klopferspitz 18, 82152 Planegg, Germany
²Physics Department, Technical University Munich, Garching, Germany.
³current affiliation: Department of Physics, Princeton University, 08544 Princeton, NJ, USA
⁴Department of Cellular Physiology, Biomedical Center (BMC), Ludwig-Maximilians-Universität München, Großhaderner Str. 9, 82152 Planegg, Germany

* corresponding authors:

Prof. Petra Schwille
ORCID: 0000-0002-6106-4847
schwille@biochem.mpg.de
Phone: +49 89 8578-2900
Fax: +49 89 8578-2903

Dr. Nikolas Hundt
ORCID: 0000-0001-8217-671X
nikolas.hundt@med.uni-muenchen.de
Phone: +49 (0) 89 2180 71563
Fax: +49 (0) 89 2180 71732

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Abbreviations
adenosine-5'-triphosphate (ATP), adenosine-5'-diphosphate (ADP), bovine serum albumin (BSA), 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-cap biotinyl (18:1 Biotinyl Cap PE),
dioleoyl-sn-glycero-3-phosphocholine (DOPC), dioleoyl-sn-glycero-3-phosphoglycerol (DOPG), horseradish peroxidase (HRP), interferometric scattering (iSCAT), phosphate binding loop (P-loop), point spread function (PSF), single-particle tracking (SPT), small unilamellar vesicle (SUV), supported lipid bilayer (SLB), mass photometry (MP), mass-sensitive particle tracking (MSPT), membrane targeting sequence (MTS)

Abstract
In spite of their great importance in biology, methods providing access to spontaneous molecular interactions with and on biological membranes have been sparse. So far, it has been consensus that their observation with sufficient sensitivity and time resolution requires the introduction of - predominantly fluorescent- labels to the system. However, the recent advent of mass photometry to quantify mass distributions of unlabelled biomolecules landing on surfaces raised hopes that this approach could be transferred to membranes. Here, we introduce mass-sensitive particle tracking (MSPT), enabling simultaneous label-free tracking and monitoring of molecular masses of single biomolecules diffusing on lipid membranes. We applied this approach to the highly non-linear reaction cycles underlying MinDE protein self-organisation. MSPT allowed us to determine the stoichiometry and turnover of individual membrane-bound MinD/MinDE protein complexes and to quantify their size-dependent diffusion. We found that MinD assembles into complexes larger than the commonly postulated dimer, through lateral interactions of membrane-bound complexes and subunit recruitment from solution. Furthermore, the ATPase-activating protein MinE interconnects MinD into high-molecular-weight heteromeric complexes and affects their subunit turnover and concerted membrane release. This study demonstrates the potential of MSPT to enhance our quantitative understanding of both prokaryotic and eukaryotic membrane-associated biological systems.

Introduction
The recruitment of proteins to lipid interfaces is crucial for various cell biological processes, such as the regulation of membrane trafficking⁴, mediation of signalling cascades⁵, and the establishment of cell polarity. These membrane-associated reactions often rely on short-lived complexes that coexist in a dynamic equilibrium with their respective cytosolic forms. Transient interactions on the membrane eventually serve as nucleation sites for the assembly of larger, more stable complexes. Additionally, the spatial distribution, stoichiometry and temporal dynamics of membrane-associated complexes are often heterogeneous. This combination of fast dynamics and compositional heterogeneity makes membrane-associated reactions difficult targets for conventional analytical techniques, which determine the composition and follow the dynamics of molecular systems based on ensemble-averaged measures. In recent decades, single-particle tracking (SPT) has revolutionised the analysis of membrane-associated
systems by following the dynamics of individual molecules at nanometre precision and millisecond time resolution\textsuperscript{6-8}, using fluorophores as labels in combination with highly sensitive microscopy. More recently, scattering-based detection using gold nanoparticles as labels has pushed the spatiotemporal resolution down to the sub-nanometre and microsecond range\textsuperscript{9-12}. The single-molecule nature of these approaches has provided detailed mechanistic insight into the dynamics of biomolecular systems\textsuperscript{13}. However, as label-based technique, SPT also suffers from label-induced artifacts: large particles, but also small fluorescent tags\textsuperscript{14} may perturb native protein function, and fluorescence-specific phenomena like photo-bleaching and -blinking hinder continuous particle tracking and limit the attainable spatiotemporal resolution\textsuperscript{15}. More importantly, although fluorescence-based SPT may provide access to interaction dynamics between tagged molecules through brightness changes, it has proven extremely complicated to extract the molecular composition of the tracked particles. In general, relating the signal of an external marker to the molecular stoichiometry of the labelled particle requires careful characterisation of labelling efficiencies and imaging conditions, which is very often hampered by quenching effects. Especially for multicomponent systems, access to the molecular composition of single particles has been prohibitive for standard biological applications.

Recent advances in interferometric scattering (iSCAT) microscopy\textsuperscript{11,16} made it possible to detect individual biomolecules label-free based on light scattering. The linear relationship between iSCAT contrast and molecular mass of a biomolecule led to the development of mass photometry (MP)\textsuperscript{17,18}. This technique allows the determination of size distributions of biomolecules in solution by measuring the individual masses of molecules landing on a glass surface and counting their relative abundances, to determine both, molecular identity and composition. For purely shot noise-limited detection of a single molecule landing on a glass surface, on top of the static scattering pattern, MP applies a sophisticated background removal strategy optimised for this type of experiment, which is not compatible with the detection of mobile molecules diffusing on lipid membranes. Here, by introducing a new iSCAT image processing and analysis strategy, we enable mass-sensitive particle tracking (MSPT) of single unlabelled biomolecules on a supported lipid bilayer (SLB). We show that the iSCAT signal-to-mass linearity holds true for membrane-associated proteins and that we can relate molecular composition, accessible via mass, to diffusive behaviour. Moreover, we can follow the mass time course along individual trajectories, making it possible to observe the (dis-)assembly of biomolecular complexes in real-time. The approach is fast and provides high particle statistics within minutes, all without need for protein labelling and its caveats.

We showcase the abilities of this method for the detailed analysis of complex biological systems by analysing the membrane-associated reaction cycle of the \textit{Escherichia coli} Min system.
This system consists of three proteins - MinC, MinD and MinE - and is essential for the spatiotemporal regulation of the division site in *E. coli*\(^{19}\). To perform this task, the ATPase MinD and the ATPase-activating protein MinE oscillate between the cell poles, forming a concentration gradient of the passenger protein MinC. This gradient supposedly enables MinC to inhibit FtsZ polymerisation at the poles and directs Z-ring formation to the mid-cell\(^{20,21}\). To this end, the phospholipid bilayer acts as a catalytic interface and membrane interaction of MinD and MinE is mediated by amphipathic helices, i.e. membrane targeting sequences (MTS)\(^{22-25}\). The interaction of MinD and MinE with the membrane decreases their diffusion rates and facilitates lateral molecular interactions enabling the system to self-organise\(^{26}\). Despite its compositional simplicity the system exhibits surprisingly complex, non-linear dynamics. Its self-organisation can be reconstituted *in vitro*\(^{26}\) and the underlying mechanism has been probed by various techniques such as SPT\(^{13}\), high-speed atomic force microscopy\(^{27}\), nuclear-magnetic resonance\(^{28}\), electron microscopy\(^{29}\), plasmonic nano sensors\(^{30}\) and mutational analysis\(^{31,32}\). Despite this intense characterisation, molecular details about MinDE self-assembly, such as the presumed cooperativity in bilayer-attachment of MinD, have remained poorly understood. Thus, the system was set to benefit from the unique ability of MSPT to characterise dynamics as well as molecular composition of membrane-attached protein complexes. Using MSPT, we dissected the membrane-associated MinDE reaction cycle by determining the stoichiometry, turnover, and diffusion of individual membrane-bound MinD/MinDE protein complexes. Our results indicate that MinD, in contrast to the classical model, assembles not only into dimers, but also forms larger oligomers, confirming recent findings\(^{27}\). We furthermore show that MinE promotes MinD self-assembly on the lipid bilayer due to its ability to interconnect MinD into so far unresolved higher-order heteromeric complexes. We believe our experiments on the Min system demonstrate that MSPT is a powerful, widely applicable tool for the mechanistic analysis of both pro- and eukaryotic membrane-associated systems.

Results

**Mass-sensitive dynamic imaging of membrane-associated protein complexes**

The major obstacle for detecting single macromolecules with iSCAT microscopy (Fig. 1a) is separating their comparatively small signal from the dominant scattering background. In a standard mass photometry experiment, molecules landing on a glass surface are detected through continuous comparison of an averaged image with the average of its preceding frames as background (Supplementary Fig. 1a)\(^{17,18}\). In this image processing approach, a molecule appears as a dark spot at the moment of landing and disappears when it becomes part of the static background\(^{17}\). The mass of the particle can then be determined by fitting the detected peak signal with a model point spread function (PSF). However, for a moving molecule, this
strategy produces distorted images of the molecule’s PSF. The missing particle density at its previous location produces a bright spot, while the added density at the new position generates a dark spot. The spatial overlap of these patterns causes moving objects to appear as dark fronts carrying bright tails, thus hampering the determination of their mass and location (Supplementary Fig. 1b). To address this issue, we employed the temporal median of an image sequence as background estimate\(^{10,33,34}\), which includes only features that do not move during the median period. To retain shot-noise limited detection, which is strongly affected by sample drift, we calculated the median of an image window as background estimate for its central frame and moved this window from frame to frame throughout the video (Supplementary Fig. 1c). Due to the clear separation of static background from moving objects, background corrected movies showed clear, undistorted images of moving PSFs (Fig. 1a, Supplementary Fig. 1c, Supplementary Table 1 and Supplementary Movie 1), a prerequisite that allows mass-sensitive particle tracking (MSPT) of single unlabelled biomolecules diffusing on lipid interfaces.

First, we set out to assess the quality of mass determination of mobile molecules compared to landing particles in conventional mass photometry. To this end, we used a bilayer supplemented with biotinylated lipids and attached a set of biotinylated standard proteins with known mass via divalent streptavidin\(^{35}\). This system has several advantages, such as the ability to cover different protein size regimes, standardised membrane binding, and the added benefit of simplified complex stoichiometries by using divalent streptavidin. For each molecule, we determined its median contrast throughout its trajectory. Analogous to conventional mass photometry (Fig. 1b, left column), histograms of the contrasts of all molecules determined with MSPT revealed the particle size distribution (Fig. 1b, right column, Supplementary Movie 2). For our standard proteins, ISCAT contrast as a function of mass exhibited the expected linear relationship\(^{17}\). The calibration line obtained from standards diffusing on SLBs was in fact indistinguishable from the one determined with molecules landing on glass (Fig. 1c). This result suggests that mass calibrations performed with landing assays can be transferred to particles diffusing on membranes. However, it is recommended to verify this for any new lipid/buffer combination.

Besides mass determination, MSPT also enables the analysis of the diffusive behaviour of membrane-bound molecules. To determine the diffusion coefficients of individual molecules, we performed a jump-distance analysis\(^{36}\) (Supplementary Fig. 2). As a verification of the diffusion coefficients obtained in this manner, we again made use of streptavidin attached to a membrane via biotinylated lipids. In line with literature values ranging from 0.8 to 2.0 µm\(^2\) s\(^{-1}\)\(^{37–39}\), the lateral diffusion coefficient of streptavidin was found to be 1.2 ± 0.4 µm\(^2\) s\(^{-1}\) (Fig. 1d).
Similar diffusion coefficients were obtained for all standard proteins attached via divalent streptavidin (Supplementary Fig. 3). To highlight a unique advantage of MSPT, we plotted the diffusion coefficient versus the respective molecular mass obtained from individual trajectories, enabling an unprecedented direct connection of these two parameters. As displayed in Figure 1d, the unimodal distribution of membrane-bound streptavidin indicates a distinct population of tetramers undergoing Brownian motion. Having validated the method with membrane-attached streptavidin, we thought the method offers the potential of detailed insight into the dynamics of molecular interactions within more complex membrane-bound systems. To further explore the method’s capabilities, we turned to the membrane-associated E. coli Min system.

Figure 1 - Principle of mass-sensitive particle tracking (MSPT). (a) Schematic displaying the interferometric scattering (iSCAT)-based measurement principle of MSPT. In this setup, proteins freely diffuse on a supported lipid bilayer and can be tracked and identified according to the linear relationship between their interferometric scattering contrast and molecular weight. Exemplary structures of three aldolase oligomer states (PDB: 4S1F\textsuperscript{16}) are shown in the top panel, and their respective iSCAT images at the bottom. Scale bar: 1 µm. (b) Probability density distributions of standard proteins determined using the conventional MP landing assay (left panel) or using MSPT (right panel). All data represent pooled distributions of three independent experiments per condition: alcohol dehydrogenase (ADH; particle number n = 9828), bovine serum albumin (BSA; n = 11,408), TEV protease (TEV; n = 1,705), β-amylase (bAm; n = 10,043), protein A (prA; n = 12,720), divalent streptavidin (Strep; n = 28,126), divalent streptavidin with biotinylated aldolase (Strep-ALD; n = 26,419), divalent streptavidin with biotinylated bovine serum albumin (Strep-BSA; n = 15,127) and divalent streptavidin with biotinylated protein A (Strep-prA; n = 33,774). Dashed lines mark peaks not considered for mass calibration (left panel). Continuous lines represent oligomer states included in the mass calibration. 2D plots of mass vs. diffusion coefficient for the four proteins measured with MSPT (right panel).
are shown in Supplementary Fig. 3. (c) Comparison of the contrast-to-mass calibration for MP and MSPT, derived from peak contrasts in panel (b) and their assigned sequence masses (see Supplementary Table 2, 3). Error bars represent the standard error of the peak locations estimated by bootstrapping. (d) 2D kernel density estimation of 1.25 nM tetravalent streptavidin bound to biotinylated lipids on a supported lipid bilayer (n = 130 325 trajectories of three independent replicates). Marginal probability distributions of the molecular mass and the diffusion coefficient are presented at the top and right, respectively.

**Cooperative Membrane-Catalysed Association Dynamics of MinD**

The MinDE system is known for its ability to self-organise into mesoscopic protein patterns on lipid membranes. To generate these patterns, MinD and MinE are generally assumed to undergo a canonical membrane binding-unbinding cycle displayed in Figure 2a. In brief, upon ATP-complexation cytosolic MinD dimerises and localises to the membrane interface. After homodimeric MinE binds to MinD, nucleotide hydrolysis is stimulated and MinD dissociates from the lipid bilayer to return to its monomeric state. Despite this established model, it has remained rather enigmatic how ATP-dependent dimerisation and the resulting increase in MinD membrane affinity alone can confer the non-linear attachment required for pattern formation. In recent years, it has been proposed that the presence of higher MTS valences and thus higher-order oligomer structures might contribute to the local self-enhancement of MinD at the membrane. Due to the fast diffusion and attachment/detachment dynamics, however, it has remained challenging to provide convincing evidence of their existence.

Taking advantage of the ability to monitor the mass of membrane-attached and diffusing protein complexes, we thus applied our MSPT approach to investigate potential oligomeric species formed by the MinDE system. We started by reconstituting the membrane recruitment of MinD and compared the mass distribution of MinD in solution, measured in an MP landing assay, with the mass distribution of MinD on a lipid bilayer (Fig. 2b, Supplementary Fig. 4). In the presence of ATP, MinD monomers were detected as the predominant species in solution. In contrast, the major species on the membrane was a MinD dimer already at sparse densities corresponding to one particle per frame in the field of view (38 µm²). However, it should be noted that at the chosen imaging conditions for MSPT, the signal-to-noise ratio of monomers (33 kDa) was too low for their quantitative detection. Hence, an adequate estimate of their membrane abundance cannot be stated.

One of the major advantages of iSCAT-based imaging is that it provides a direct estimate of the molecular density on a bilayer from the number of detected particles, as compared to single-molecule fluorescence where factors like labelling efficiency and photo bleaching have to
be taken into account. Accordingly, we could classify video sections of membrane-bound MinD into conditions of different particle densities and observed the resulting change in the oligomeric distribution (Fig. 2c, Supplementary Fig. 5, 6 and Supplementary Movie 3). While MinD was mainly present in the dimer state at low particle densities (0.1 µm\(^2\)), the MinD population shifted towards a broad distribution with higher-order complexes on crowded bilayers (0.9 µm\(^2\)). This data directly demonstrates that MinD indeed assembles into complexes larger than a dimer as previously suggested\(^{27,31}\). To further investigate the structural determinants for higher-order MinD assemblies, we performed the same MSPT experiment with a MinD mutant (D40A) that is reported to predominantly reside in the dimeric state due to its impaired ability to hydrolyse ATP\(^{43}\). According to the prevailing mechanism, it should not be possible for a locked dimer mutant to participate in MinD self-assembly, due to its inability to switch between the monomeric and dimeric state\(^{23,41,44}\). Nevertheless, we found a distinct population of MinD D40A tetramers in our contour plot (Fig. 2d, Supplementary Fig. 7), suggesting that MinD is capable of forming higher-order oligomers using an interface distinct from the canonical dimerisation site.

Another striking aspect of the D40A mutant was its clear separation into two distinct species (dimer and tetramer), whereas the distribution for the WT protein appeared unresolved. This result suggested that the WT was able to also recruit MinD monomers forming trimeric species as intermediates, which were not as abundant for the D40A mutant. Based on the assumption that all MinD oligomer states can be populated, we fitted the mass distributions of MinD WT and MinD D40A with Gaussian mixture models and quantified the relative abundance of each oligomer as a function of bilayer molecule density (Fig. 2e, g, Supplementary Fig. 8), an information hardly obtainable with any other method so far. These graphs show a sequential appearance of increasingly larger oligomers of MinD for higher molecule densities (Fig. 2f). Notably, compared to the WT, the D40A mutant had a higher tendency to populate stoichiometries with even numbers of subunits and to transform its dimer state into the tetramer state, likely due to the increased stability of the dimer (Fig. 2h).

Furthermore, MSPT confers the unique possibility to determine oligomer-specific lateral diffusion coefficients. This knowledge can be used to deduce structural information about the observed molecules when considering the theory of Evans-Sackmann\(^{45}\), which postulates the relation of an object’s membrane inclusion size with its respective diffusion coefficient. Assuming a similar membrane viscosity as for pure DOPC membranes\(^{46}\), we can thus estimate that a MinD D40A dimer with a diffusion coefficient of 0.8 µm\(^2\)/s has an inclusion size of 2 nm, and a tetramer with 0.4 µm\(^2\)/s an inclusion size of 5 nm (Supplementary Fig. 9). These estimates
imply that all monomers of the tetramer are able to insert their MTS into the bilayer, suggesting certain geometrical constraints for subunit orientation.

**Figure 2 - Lateral MinD–MinD interactions lead to self-assembly into large homo-oligomers.** (a) Schematic of the canonical membrane binding-unbinding cycle of MinDE. Upon ATP-complexation, MinD dimerises (1) and attaches to the membrane interface (2). In the event of MinE binding (3), MinE stimulates the intrinsic ability of MinD to hydrolyse ATP, which leads to the release of MinD from the membrane in its monomeric form (4). (b) MinD mass distribution in solution (light blue line; n = 16,101 particles) and upon attachment to the supported lipid bilayer (blue line; n = 7,012 particles). For solution experiments, 175 nM MinD with 0.5 µM ATP were measured in the conventional MP landing assay. The membrane mass distribution of MinD was determined using MSPT at a particle density of 0.03 µm$^2$. (c) 2D kernel density estimation of membrane-attached MinD and (d) MinD D40A at particle densities of 0.1 µm$^2$ (light blue, n = 158,102 particles) and 0.9 µm$^2$ (dark blue, n = 8,507 particles) and 0.1 µm$^2$ (light green, n = 15,374 particles) and 0.5 µm$^2$ (dark green, n = 31,978 particles), respectively. Marginal probability distributions of both molecular mass and diffusion coefficient are presented at the top and right, respectively. (e) Representative mass distributions (grey) of MinD and MinD D40A (g) and Gaussian mixture model estimations (black line, coloured lines highlight underlying Gaussian components) with five components for three different membrane particle densities. MinD: 0.1 µm$^2$ – light blue, 0.5 µm$^2$ – blue, 0.9 µm$^2$ – dark blue; MinD D40A: 0.1 µm$^2$ – light green, 0.3 µm$^2$ – green, 0.5 µm$^2$ – dark green. (f,h) Relative oligomer abundance and its corresponding mass (inset) as a function of bilayer density. Error bars are the standard deviation of results from 1000 random fit initialisations. The oligomer analysis is based on a total of n = 1,455,585 particles for MinD and n = 253,717 for MinD D40A.
Time-resolved mass analysis of single MinD trajectories

Aside from measuring a particle’s location frame by frame, MSPT also allows to determine its respective mass in a time-resolved fashion, thus enabling the detection of attachment and detachment events along the trajectory of a single particle (Fig. 3a, Supplementary Fig. 10). For MinD, the minimal expected mass increment (33 kDa) for monomer-wise turnover was close to the measurement uncertainty (28 kDa standard deviation) of the mass for a single frame. To minimise user bias, we employed a step-finding algorithm that locates mass change points along trajectories based on statistical criteria47.

Figure 3b displays examples of mass time courses for individual particles and the mass steps detected by the algorithm. Note that in a majority of cases, MinD complexes retained their size throughout their entire trajectory (Fig. 3b, upper panel). However, in ~10% of trajectories, the mass of a tracked particle changed during its trajectory (Figure 3b, bottom panel), suggesting the attachment or detachment of MinD subunits to and from the membrane-bound complex. By analysing the sizes of mass steps along all trajectories, it was possible to obtain a distribution of subunit sizes attaching and detaching from membrane-bound MinD complexes (Fig. 3c), which appeared to mostly (dis-)assemble in one- or two-subunit increments at low particle densities. However, for higher MinD bilayer densities, when larger oligomers had accumulated, these complexes often turned-over greater subunits, as indicated by the shift of the distribution towards higher mass steps27 (Fig. 3c, dark blue profile). Moreover, the combined information of a mass plateau level and its dwell time as annotated in Fig. 3b could be used to extract the subunit turnover rates of each oligomer species (Fig. 3d). Here, the dwell time preceding a mass increase could be used to deduce subunit attachment rates (Fig. 3d – upper panel), whereas the dwell time followed by a mass decrease provided an estimate of detachment rates (Fig. 3d – lower panel). The resulting lifetime plots suggested that membrane-attached dimers had a faster subunit turnover than trimers and tetramers, implying higher stability of these larger complexes.

Plateaus at the end of trajectories (Fig. 3b, bottom panel, last plateau), along with trajectories without any mass change (Fig. 3b, top panel), could be used to identify the molecular weight of particles completely released (rl) from the membrane (Fig. 3e). Notably, this time-resolved mass analysis of the individual trajectories significantly improved the mass resolution, as compared to the median-based particle mass estimates used in Figure 2. Hence, MinD dimers, trimers and tetramers were now fully resolved as separate peaks (Fig. 3e). Accordingly, one could now recognize that the major species released from the membrane was a dimer at low particle densities and a tetramer at high particle densities. The corresponding dwell time plot showed that tetramers stayed associated to the bilayer significantly longer than dimers, in line
with a higher avidity in membrane binding conferred by additional MTS (Fig. 3f). For comparison, the dimer-arrested mutant MinD D40A almost exclusively dissociate as dimers or tetramers (Supplementary Fig. 11, 12). Taken together, our detailed trajectory analysis confirms our previous observations suggesting that MinD WT assembles into species of higher order with an intermediate trimer state. This indicates that subunits bind at a location different from the canonical dimerisation site.

**Figure 3 - Subunit (dis-)assembly of MinD particles diffusing on membranes.** (a) Schematic representation of the time-resolved mass analysis of single MinD trajectories, which reveals attachment (at) and detachment (dt) events along the trajectory as well as a particle’s full membrane release (rl) at the end of its trajectory. (b) Representative mass time traces of MinD trajectories (grey line) and corresponding step fits (black line) determined by a step-finding algorithm that locates mass change points within a trajectory. (c) Mass step size distribution derived from step fits as depicted in (b) revealing MinD subunit sizes for attachment (at) and detachment (dt) events at particle densities: 0.1 µm⁻² – pale blue (n = 19,974 plateaus), 0.3 µm⁻² – light blue (n = 24,102 plateaus), 0.5 µm⁻² – blue (n = 21,236 plateaus), 0.9 µm⁻² – dark blue (n = 3,207 plateaus). (d) Dwell time plots for MinD particles before attachment (at) events (top plot), representing the lengths of mass plateaus preceding a mass increase, and before detachment (dt) events (bottom plot), representing the lengths of mass plateaus preceding a mass decrease. Dwell times are shown for the MinD dimer state (66 kDa, light blue line) and the MinD tetramer (132 kDa, dark blue line). Plateau numbers for (at): dimer - n = 25,193; tetramer - n = 4,510; for (dt): dimer - n = 4,020; tetramer - n = 10,766. Insets: Mean dwell times of the two oligomer states. Error bars represent the standard deviation of the dwell times estimated by bootstrapping. (e) MinD mass distribution for membrane release (rl) events. MinD particle densities: 0.1 µm⁻² – pale blue (n = 157,944 plateaus), 0.3 µm⁻² – light blue (n =
227,506 plateaus), 0.5 μm² – blue (n = 371,911 plateaus), 0.9 μm² – dark blue (n = 85,713 plateaus). (f) Plot of the dwell times before membrane release (rl) for the MinD dimer and tetramer state. Plateau numbers: dimer - n = 827,833; tetramer - n = 57,099. Inset: Mean dwell times of the two oligomer states. Error bars represent the standard error of dwell times estimated by bootstrapping.

**MinE-induced formation of large heteromeric complexes**

In the past decade, several different mechanistic models have been proposed to explain the role of the ATPase-activating protein MinE for MinDE detachment dynamics. Some of these models are based on the cooperation of both MinE and MinD dimer to prompt membrane-release through MinD ATPase activity stimulation. However, this effect alone cannot explain the recently observed cooperative membrane-detachment of MinDE filaments. To address this issue, we used MSPT to determine the stoichiometry of the membrane-bound MinDE complex and followed its membrane dynamics on a molecular level.

In accordance with previous structural studies that suggest a conformational switch of MinE allowing MinD binding only upon the encounter of membrane-bound MinD, we found no indication for MinDE interaction in solution (Supplementary Fig. 13). In the presence of a supported lipid bilayer, however, the MinDE complex existed predominantly in a stable double-dimeric state (Fig. 4a – light pink, Supplementary Fig. 14). Furthermore, if the MinDE complex encountered more proteins on a crowded bilayer, MinE promoted the interconnection into very large heteromeric MinDE complexes, a behaviour unexpected considering the common models (Fig. 4a – magenta). One possible explanation for this behaviour is the ability of a MinE dimer to symmetrically bind to both sides of a MinD dimer, thus effectively acting as a bridge between MinD assemblies. Accordingly, our time-resolved mass step analysis revealed that during subunit turnover on membrane-bound particles, predominantly dimeric and tetrameric subunits attached (at) and detached (dt) at high particle densities of 0.5 and 0.9 μm² (Fig. 4b, Supplementary Fig. 15). This effect required a critical minimum density on the bilayer though, since MinDE complexes in sparsely populated environments (0.1 and 0.3 μm²) mainly exhibited conversion in their minimum subunit increments. Hence, their final oligomeric state during membrane release (rl) resembled a MinDE dimer (Fig. 4c, light pink). Strikingly, on bilayers with high protein density, MinDE complex sizes released from the membrane increased beyond the sizes observed for MinD alone and reached masses of >350 kDa (Fig. 4c, purple).

Based on our mass dwell time analysis, we found that the presence of MinE generally reduced the diffusion coefficients (Fig. 4d) of MinDE oligomers and slowed down turnover rates (Fig. 4e, pink), when compared to their respective MinD versions (Fig. 4e, blue). In addition, MinDE
complexes were found to reside significantly longer on the membrane before full release (Fig. 4f). This suggests that MinE can effectively stabilise membrane-bound MinD when present in equimolar amounts as previously suggested\textsuperscript{13,49}.

Figure 4 - MinE interconnects MinD oligomers into large complexes with a prolonged membrane dwell time. (a) 2D kernel density estimation of membrane-attached MinDE complexes at particle densities of 0.1 µm\(^{-2}\) (light pink, n = 263,594 particles) and 0.9 µm\(^{-2}\) (purple, n = 8,681 particles). Marginal probability distributions of both molecular mass and diffusion coefficient are presented at the top and right, respectively. (b) Mass step size distribution revealing MinDE subunit turnover (at and dt events) on membrane-bound particles at particle densities of 0.1 µm\(^{-2}\) – pale pink (n = 120,643 plateaus), 0.3 µm\(^{-2}\) – light pink (n = 81,197 plateaus), 0.5 µm\(^{-2}\) – pink (n = 9,650 plateaus), 0.9 µm\(^{-2}\) – dark purple (n = 2,942 plateaus). (c) MinDE mass distribution for membrane release (rl) events at the same particle densities as in (b). (d) Analysis of oligomer specific diffusion coefficients for MinD (blue lines) and MinDE complexes (pink lines). light blue/pink – dimer; dark blue/purple – tetramer. Inset: Mean diffusion coefficients of each oligomer state. Error bars represent the standard error of the dwell times estimated by bootstrapping. (e) Dwell time plots for MinD and MinDE attachment (at) events (top plot) as well as for MinD and MinDE detachment (dt) events (bottom plot). Dwell times are shown for the MinD dimer (light blue line) and MinD tetramer (dark blue line) as well as for their respective MinDE versions (hetero-dimer state – light pink, hetero-tetramer state – purple). Plateau numbers for (at): MinD dimer – n = 25,193, MinDE dimer – n = 43,558, MinD tetramer – n = 4,510, MinDE tetramer – n = 12,069; (dt): MinD dimer – n = 4,020, MinDE dimer – n = 5,189, MinD tetramer – n = 10,766, MinDE tetramer – n = 24,394. Insets: Mean dwell times of the two oligomer states. Error bars represent the standard error of the dwell times.
estimated by bootstrapping. (f) Plot of the dwell times before membrane release (rl) for the MinDE dimer and tetramer state and the respective MinD versions for comparison. Plateau numbers: MinD dimer – n = 827,833, MinDE dimer – n = 395,737, MinD tetramer – n = 57,099, MinDE tetramer – n = 65,941. Inset: Mean dwell times of the two oligomer states. Error bars represent the standard error of the dwell times estimated by bootstrapping.

Conclusions
In this work, we presented a simple and versatile single-molecule-based method for the determination of membrane-associated oligomer distributions, subunit turnover and mass-resolved residence times in the context of the E. coli Min system. Based on our results, we propose the extension of the established membrane binding-unbinding models for MinDE self-organisation\textsuperscript{44,48} (Fig. 5). Our data support the original model regarding nucleotide exchange from ADP to ATP triggering membrane-dependent dimerisation of MinD, thus coupling the process to energy dissipation. On the membrane, we find that lateral MinD interactions as well as recruitment of MinD subunits from solution lead to the formation of a dynamic mixture of MinD oligomeric states that assemble through attachment and detachment of subunits at a location different from the canonical dimerisation site\textsuperscript{31}, a behaviour unexpected by common models. We assume that the ability of MinD to assemble into these complexes is generally required for its local self-accumulation and in combination with the initial nucleotide-dependent membrane recruitment explains the observed attachment cooperativity during MinDE self-organisation. At elevated membrane densities, a small percentage of MinD self-assembles into tetramers, which either fall apart or interact with MinE. In contrast to the previously assumed cooperation of the MinE and MinD dimer to prompt membrane-release\textsuperscript{44,48}, we found that MinE promotes the interconnection\textsuperscript{49} of heteromeric MinDE complexes. We assume that this behaviour is based on the ability of two MinE dimers to symmetrically bind to both sides of a MinD dimer, thus effectively acting as a bridge between MinD assemblies\textsuperscript{49}. The existence of multivalent membrane-bound structures would likewise explain the prolonged residence times of MinDE complexes compared to their respective MinD oligomer variants. Notably, opposed to the previously assumed monomer detachment\textsuperscript{51}, we found MinDE to detach from the membrane interface in complexes with sizes beyond 350 kDa, which could correspond to eight MinDE subunits. This corroborates that MinE could induce nucleotide-conversion of MinD subunits and thereby weakens the overall membrane avidity of the MinDE complex prior to its full dissociation from the membrane interface\textsuperscript{27}. Upon membrane release, larger complexes might remain temporarily stable, stay in the vicinity of the bilayer, and can potentially rebind close to their dissociation spot, once MinD subunits have exchanged their nucleotide. To conclude, we believe that MinDE self-organisation arises from an interplay of cooperative MinD membrane attachment into higher-order oligomers, anisotropy of the local MinD concentration through
quick re-binding of oligomers to the membrane, and ATP-dependent membrane release of MinD assemblies coordinated by the ATPase-stimulating activity of MinE.

The mechanistic dissection of the MinDE membrane-cycle described in this study constitutes a detailed technical demonstration of the capabilities of mass-sensitive particle tracking (MSPT). By combining the advantages of mass photometry\textsuperscript{17,18}, where biomolecular complex stoichiometries are determined from the mass of individual particles, with single-particle tracking, MSPT provides invaluable insights into the complex dynamics of biomolecules on lipid membranes. Advantages of the label-free detection of membrane complexes over conventional fluorescence-based single-particle tracking approaches are: 1) the possibility to determine particle densities, i.e. local concentration on the bilayer; 2) the correlation of mass and diffusion coefficient which, in combination with a derived membrane inclusion size, provides information about the stoichiometry and arrangement of a membrane-bound complex; 3) the analysis of time-resolved subunit turnover and its kinetics by analysing mass changes along a particle’s trajectory; 4) extended observation times due to the absence of photo-bleaching, thus enabling the collection of high particle statistics in a short period of time. Ultimately, we believe that mass-sensitive particle tracking will make a strong contribution to the quantitative understanding of both prokaryotic and eukaryotic membrane-associated biological systems.

![Figure 5](image-url)

**Figure 5 - Schematic of the proposed membrane-associated MinDE reaction cycle.** (1) Upon nucleotide exchange from ADP to ATP, membrane-dependent dimerisation of MinD is triggered. (2) On the membrane, lateral MinD interactions and recruitment of MinD subunits from solution lead to MinD higher-order structures that assemble through attachment of subunits at a location different from the canonical dimerisation site. MinD assemblies then either dissociate from the membrane (3), or encounter MinE (4). MinE promotes the interconnection of very large heteromeric MinDE complexes that, due to their multivalent MTS structure, reside significantly longer on the membrane interface (5). (6) However, MinE also induces nucleotide-
conversion of MinD subunits, thereby weakening the overall membrane avidity of the MinDE complex, prior to its full release from the membrane interface in complexes >350 kDa.

Methods

ADP/ATP stock solution
Both ADP and ATP stocks were prepared from their respective salt hydrates (#A2754 and #A2383, Sigma Aldrich, St. Louis, USA), supplemented with an equal molar amount of MgCl₂ and adjusted to pH 7.5 with 1 M Tris-HCl. Final nucleotide concentration was spectroscopically determined (λ = 259 nm; V-650, Jasco, Pfungstadt, Germany) using an extinction coefficient of 15,400 M⁻¹ cm⁻¹.

Protein Purification and Modification
Purifications of MinD, MinD D40A and MinE were performed as previously described⁵² and all used plasmid constructs are indicated in Supplementary Table 4. Divalent streptavidin was assembled as outlined in Howarth et al.⁵³, both pET21a-Streptavidin-Alive (Addgene plasmid #20860) and pET21a-Streptavidin-Dead (Addgene plasmid #20859) were a gift from Alice Ting⁵³. Biotin-modification of aldolase (#28403842, Cytiva, Marlborough, USA) with EZ-Link™ Maleimide-PEG2-Biotin (#A39261, Thermo Fisher Scientific, Waltham, USA) was achieved through incubation at RT for 1 h and subsequent size-exclusion chromatography on a 16/600 Superdex 200 pg column (GE Healthcare, Pittsburgh, USA), equilibrated in storage buffer (50 mM HEPES pH 7.25, 150 mM KCl, 0.1 mM EDTA, 10% glycerol, 0.4 mM TCEP), using an Äkta Pure chromatography system (GE Healthcare, Pittsburgh, USA). LC-MS and SDS-PAGE was performed to assess purity and integrity of all purified or modified proteins. A customized Bradford assay (#5000006, Bio-Rad Protein Assay; Bio-Rad Laboratories Inc., Hercules, USA) was used to determine protein concentrations and single-use aliquots were flash frozen in liquid nitrogen and stored at -80 °C.

Small Unilamellar Vesicles (SUVs) and Supported Lipid Bilayer (SLB) Formation
For the formation of small unilamellar vesicles (SUV), dioleoyl-sn-glycero-3-phosphocholine (DOPC; #850375, Avanti Polar Lipids, Alabaster, USA), dioleoyl-sn-glycero-3-phosphoglycerol (DOPG; #840475, Avanti Polar Lipids, Alabaster, USA) and 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-cap biotinyl (18:1 Biotinyl Cap PE; #870273, Avanti Polar Lipids, Alabaster, USA) were dissolved in chloroform (Sigma Aldrich, St. Louis, USA) and mixed in a ratio of 70 mol % DOPC to 30 mol % DOPG or 70 mol % DOPC with 29.99 mol % DOPG and 0.01 mol % Biotinyl Cap PE. After solvent evaporation through nitrogen, residual chloroform was removed for 1 h in a vacuum desiccator. Lipid film hydration was achieved with Min buffer (25
mM Tris/HCl pH 7.5, 150 mM KCl, 5 mM MgCl₂) and SUVs were formed through consecutive freeze-thaw cycles (8-10) using liquid nitrogen and a 90 °C water bath. For monodisperse vesicle distribution, lipid mixtures were extruded across a Whatman nuleopore membrane (#110603, GE Healthcare, Chicago, USA) with a pore size of 50 nm for 37 passes.

SLBs were formed by fusion of SUVs on cleaned glass cover slides (Paul Marienfeld GmbH & Co. KG, Lauda-Königshofen, Germany) that were assembled into a flow chamber through double-sided sticky tape (Scotch, Conrad Electronic SE, Germany). Prior to their assembly, cover slides (#0102242, #1.5, 24 x 60 mm; #0102062, #1.5, 24 x 24 mm) were cleaned by sequential sonication in Milli-Q water, isopropanol, and Milli-Q water (15 min each), and subsequently dried under a nitrogen stream. Slides were then activated for 30 s (30% power, 0.3 mbar) in a Zepto plasma cleaner (Diener electronic GmbH, Ebhausen, Germany) using oxygen as the process gas. After flow chamber assembly, SUVs were added to each reaction chamber at a final concentration of 0.4 mg/mL in Min buffer with additional 2 mM CaCl₂ to promote vesicle rupture. Unfused SUVs were removed through subsequent washing with Min buffer.

**Mass Calibration Curves and Diffusion Control**

To convert interferometric scattering contrast into the respective protein mass, the contrast of a set of mass standards was measured for each experimental setup – MP landing assays and MSPT on the lipid interface. In line with Young et al.¹⁷, mass calibration for landing assays was performed in flow chambers filled with filtered (sterile syringe filters 0.45 µm cellulose acetate membrane, VWR International, Radnor, USA) SLB buffer (25 mM Tris/HCl pH 7.5, 150 mM KCl). As mass standards, either 50 nM TEV protease (MPIB Core Facility), 50 nM Pierce™ recombinant protein A (#21184, Thermo Fisher Scientific, Waltham, USA) 50 nM bovine serum albumin (#A4612, Sigma Aldrich, St. Louis, USA), 20 nM alcohol dehydrogenase (#A8656, Sigma Aldrich, St. Louis, USA), or 20 nM β-amylase (#A8781, Sigma Aldrich, St. Louis, USA) were injected and landing events recorded. To also enable mass calibration on supported lipid bilayers, membranes containing 0.01 mol % Biotinyl Cap PE were used. As linker for the attachment of biotinylated standard proteins, 2.5 nM divalent streptavidin were incubated for 10 min (RT) prior to the addition of 100 nM biotin labeled bovine albumin (#A8549, Sigma Aldrich, St. Louis, USA), biotinylated Pierce™ protein A (#29989, Thermo Fisher Scientific, Waltham, USA) or custom biotinylated aldolase (#28403842, Cytiva, Marlborough, USA).

For diffusion coefficient verification, 1.25 nM unconjugated tetravalent Streptavidin (#SNN1001, Thermo Fisher Scientific, Waltham, USA) were incubated for 10 min (RT) on a supported lipid membrane containing biotinylated lipids.
Mass Photometry Landing Assay

To determine the nucleotide-dependent solution-state of MinD (175 nM) and MinDE (175 nM), all proteins were diluted in filtered (sterile syringe filters 0.45 µm cellulose acetate membrane, VWR International, Radnor, USA) Min buffer in the presence of either 0.5 mM ADP or ATP. Measurements were performed in flow chambers and 50 µL protein solution were flushed in five times consecutively, to collect sufficient landing events.

In Vitro Reconstitution of Min Complexes on a Lipid Bilayer

MinD

For MSPT of single MinD complexes on the lipid interface, we added increasing protein concentrations (50, 75, 100, 125, 150, 175 and 200 nM, n = 3 flow chambers each) to a flow chamber with a supported lipid bilayer in the presence of 0.5 mM ATP in filtered Min buffer. The same experiments were performed likewise for MinD D40A. Videos of 0.5 mM ATP in Min buffer without protein were recorded as image background control (0.8% of particles detected as compared to videos containing MinD).

MinDE

Co-reconstitution of MinDE was performed in a similar manner as for MinD, except for the addition of equimolar amounts of both reactants (50, 75, 100, 125, 150, 175 and 200 nM, n = 3 flow chambers each) to the sample chamber. For MinD D40AE, experiments were performed with 200 nM of each protein.

Microscopy

Imaging of MinDE landing assays was performed on a custom-built interferometric scattering microscope described in17.54 with a 445 nm laser diode for illumination and 635 nm for focus-stabilization. Image acquisition was controlled using custom-written software in Labview described in17. Landing events were recorded at a frame rate of 1 kHz for 60 s per video. Videos were saved 5-fold frame-averaged (200 Hz effective frame rate) and 3-fold pixel-binned (70.2 nm effective pixel size).

SLB experiments were carried out on a commercial Refeyn OneMP mass photometer (Refeyn Ltd., Oxford, UK). Movies were acquired for either 45 s (landing assay of mass standards, Fig. 1) or 350 s (SLB assays) with the AquireMP (Refeyn Ltd., v2.3) software at a frame rate of 1 kHz. Movies were saved 5-fold frame-averaged (200 Hz effective frame rate) and 4-fold pixel-binned (84.4 nm effective pixel size).
Data analysis

Image Processing of Conventional MP Landing Assays

Videos of the proteins MinD, MinE and their equimolar mixture MinDE landing on glass cover slides were analysed with the software Discover\textsuperscript{MP} (version 2.1.0, Refeyn Ltd.), using a rolling background removal strategy\textsuperscript{17}.

MinD monomers have a molecular mass of 33 kDa, which approaches the lower detection limit of the mass photometer. We have therefore systematically determined the optimal frame averaging factor $n_{avg}$ and filter thresholds $T1$ and $T2$ for the detection of MinD monomers and dimers. To this end, we generated a semi-synthetic movie that used frames from a video recorded in a chamber with Min buffer alone to reconstruct the experimental background and added simulated PSFs (same as fitting model PSF) as landing events that had the expected scattering contrast of MinD monomers (contrast = $1.9 \times 10^{-3}$) or MinD dimers (contrast = $3.5 \times 10^{-3}$). We then varied $n_{avg}$ as well as $T1$ and $T2$, ran the analysis procedure, and evaluated the number of true positive and false positive detections. To determine the maximum number of true positive detections possible at the respective signal-to-noise ratio, we simulated 1000 frames with 100 landing events that were not allowed to spatially overlap closer than 12 pixels and 26 frames temporally. Based on the simulation, we chose $n_{avg} = 12$, $T1 = 1.1$ and $T2 = 0.2$ to process the experimental videos. Using these parameters, the number of true positive detections of monomers was $68.8 \pm 6.6\%$ (mean $\pm$ STD, $n = 5$ simulated videos), while the total of false positive detections was $9.4 \pm 2.1\%$. For dimers, a simulation with the same parameters gave $95.8 \pm 0.8\%$ true positive detections and $4.2 \pm 1.1\%$ false positive ones.

Mass Calibration for Landing Assays

Landing events imaged in three independent experiments per standard protein were pooled and kernel density estimations were calculated. For each standard protein, the contrast associated with a probability density maximum was determined and plotted against its nominal mass (see Supplementary Table 1). In case of alcohol dehydrogenase and $\beta$-amylase, the peak with higher contrast was used.

Error bars in Fig 1c represent the standard error of the peak position calculated from 10,000 bootstrap resamples.

Image Processing and Analysis Procedure for Single-Particle Tracking and Mass Determination of Biomolecules on SLBs

To detect and analyse diffusing biomolecules on a lipid bilayer, we applied a new image processing strategy that removed the dominant static scattering background, while conserving the shape of mobile features and displaying them on top of a shot-noise limited background.
Around each frame, a pixel-wise temporal median image was calculated that only contains the features which did not move in the median period. This static background, calculated for each frame, was then removed from the respective frame with index \( i \) according to equation (1) and with \( n \) denoting the median half-size.

\[
(1) \quad \text{mobile features frame}_i = \left( \frac{\text{frame}_i}{\text{median(\text{frame}_{i-n:n}\text{frame}_{i+n})}} \right) - 1
\]

Through this process, moving objects in the mobile features frame appeared as undistorted PSFs and displayed ISCAT contrasts similar to those obtained for molecules in the conventional landing assay. We kindly received a custom-written Python script from Philipp Kukura and Gavin Young which automatised this image analysis procedure and we modified it to load and process videos in the Refeyn format (.mp files). To track individual proteins diffusing on supported lipid bilayers in the background-corrected movies, the Python script also included a single-particle tracking routine. For particle detection, a Laplace filter was applied to each frame (scipy.ndimage.filters.gaussian_laplace function), to suppress shot noise and highlight potential particles. The Laplace-filtered images were thresholded, such that only objects with at least the size of 53 kDa remained, a threshold that would preserve streptavidin as our smallest standard protein and MinD dimers. Local maxima were then found (scipy.ndimage.filters.maximum_filter function) as candidate pixels that contained a particle. Around each of these candidate pixels in the non-Laplace-filtered image, a region of interest (ROI) of 13 by 13 pixels (84.4 nm/pixel) was excised and fit by the model PSF used in the DiscoverMP software to extract particle contrast and location at sub-pixel resolution. Particle locations were linked into trajectories using the Trackpy package (version 0.4.2, link_df function)\(^5^\). Only trajectories of particles with a lifetime of at least 5 frames (25 ms) were considered for subsequent analyses. Supplementary Table 1 summarises all parameters used for particle detection, fitting and trajectory linking.

### Determination of Diffusion Coefficients

To determine translational diffusion coefficients from individual trajectories, a jump-distance analysis was performed\(^3^6\). Cumulative frequency distributions of jump distances were fitted (scipy.optimize.least_squares, Trust Region Reflective algorithm) with either one or two species depending on which model resulted in a reduced chi-squared statistic. For comparison, an effective diffusion coefficient was calculated in the case of a two-component model according to

\[
(2) \quad D_t = a_1 D_1 + a_2 D_2,
\]
where \( a_i \) and \( D_i \) are the fraction and diffusion coefficient of component \( i \).

**Kernel Density Estimation in 1D**

For univariate distributions, kernel density estimations (KDEs) were computed with the Python package *KDEpy* (version 1.0.10, FFTKDE function) [https://github.com/tommyod/KDEpy], using a Gaussian kernel and the Improved Sheather-Jones plug-in bandwidth selector\(^ {\text{A6}} \).

**Kernel Density Estimation in 2D**

2D maps of the diffusion coefficient and mass were computed with the Python package *fastkde* (version 1.0.14, fastkde.pdf function)\(^ {\text{A7}} \). The probability density for the diffusion coefficient was calculated in base 10 logarithmic space. Marginal distributions displayed on top and on the right of the graph were calculated by summing up the bivariate probability densities along the y- and x-axis, respectively.

The resulting 2D kernel densities were plotted as filled contours with either six (if an overlay of two conditions is shown) or eight linearly spaced levels. The opacity of the levels was reduced linearly from 100% for the highest to 0% for the lowest level, rendering the lowest level fully transparent.

**Mass Calibration for MSPT**

Analogous to the mass calibration for landing assays, peak contrasts were related to the nominal mass of streptavidin (Strep) or to the mass of streptavidin and the standard protein (Strep-ALD, Strep-BSA, Strep-prA). For the streptavidin-protein complexes, the peak with lowest contrast was excluded from analysis as it resembles streptavidin without an attached standard protein (blue crosses, Supplementary Fig. 3). The two other peaks with higher contrast were assigned to dimeric and tetrameric aldolase, and monomeric and dimeric BSA and protein A (Supplementary Table 3).

Error bars in Fig 1c represent the standard error of the peak position calculated from 10,000 bootstrap resamples.

**Analysis of MinD and MinD40A Oligomerisation**

Subpopulations in the mass distributions of MinD and MinD D40A were assessed with Gaussian mixture modelling using the Python package *scikit-learn* (version 0.23.2, sklearn.mixture.GaussianMixture class). Based on the Bayesian information criterion (BIC), the oligomeric mixture was best characterised by five components assuming that all components have equal variance (Supplementary Fig. 8a, b).

Each trajectory was assigned to a membrane density determined as the median of particle numbers detected during the trajectory divided by the area of the field of view (FOV, 38.5 \( \mu m^2 \)).
For densities ranging from 0.03 to 0.96 µm⁻² (corresponding to 1 to 37 particles in the FOV), the best-fit Gaussian mixture was estimated. The densities were binned in an overlapping manner such that all trajectories were selected that had particle numbers within a range of plus or minus two (e.g. at a density of 0.1 µm⁻², corresponding to 4 particles/FOV, trajectories with membrane densities between 2 and 6 particles/FOV were pooled). This procedure was repeated with 1000 random initialisations for mass and weights as a means to estimate the robustness of the algorithm’s solution. The results in Fig 2e-h correspond to the average and standard error of 1000 runs. The progression of the mean mass of the components as well as the width of the distributions versus density is shown in Supplementary Fig. 8c, d, e, f.

**Step Detection**

To detect mass change points during trajectories, we employed a MATLAB (R2020a, The MathWorks, Natick, MA) implementation of the Kalafut-Visscher algorithm⁵⁸ described in⁴⁷. While the algorithm has no parameters except the time series itself, we noticed that the results depended on the length of the time series and the position of potential change points if they were located close to the beginning or the end of a trajectory. Thus, change points were more likely to be inserted, the shorter a time series was and the closer a point was to the boundaries of the time series. To address this ambiguity, we concatenated the trajectories of a specific imaging condition (e.g. of all videos containing MinD, Supplementary Fig. 10a) and divided this concatenated series into \( n \) subsets of equal length \( l \), which were then analysed with the step detection algorithm (Supplementary Fig. 10b). This procedure enabled an equal treatment of trajectories of different lengths and avoided bias in change point detection at the beginning and the end of trajectories. Additionally, step identification was repeated \( l - 1 \) times shifting the start point of the linked time series circularly by one increment in each iteration. This second procedure ensured that the detected steps would not depend on the start points of a subset. From the \( l \) outputs generated in total, the relative significance of a step was reflected by the fraction \( f \) of iterations, in which a change point was found at a particular location, ranging from 0 (never) to 1 (in every iteration, Supplementary Fig. 10c).

The choice of fraction \( f \) as well as the length \( l \) is somewhat arbitrary yet interconnected. Increasing \( l \) leads to fewer detected steps which can be compensated for by reducing \( f \). Since the longest trajectory in all datasets has a length of 558 frames and an increasing length leads to a finer grating of the tunable parameter \( f \), we set \( l \) to 1000 frames. By visual inspection, we chose a fraction \( f \) of 0.25 as an appropriate tradeoff between under- and over-identification of steps considering the noise level in our trajectories (Supplementary Fig. 10d).
Subunit Attachment/Detachment During Membrane Diffusion and Release of Complexes

The concatenated mass traces and step-fitting results were split into their original trajectories (Supplementary Fig. 10e). For each trajectory, step heights (equivalent to mass changes) and dwell lengths (equivalent to time intervals without a change of mass) were extracted and categorised into three classes (Fig. 3b). Depending on whether the sign of the succeeding mass change is positive or negative, the preceding level is classified as an attachment (at) or as a detachment plateau (dt), respectively. The last plateau in a trajectory prior to the dissociation of the whole particle from the membrane is regarded separately as release (rl). If no step is detected in the mass time series as it was the case in ~90% of all trajectories, the single plateau is also classified as release (rl). Particles leaving the FOV have been excluded from the analysis. Therefore, it can be assumed that the end of a trajectory represents the particle’s release from the membrane. Generally, the mass which is initially binding to the membrane is not included in the list of steps for attachment or detachment events.

The distribution of mass changes during the trajectory as well as the distribution of complex mass released from the membrane for different densities are displayed as univariate kernel density estimates. To characterise the residence times of different oligomers, histograms of the observed dwell lengths were generated for plateaus with masses that correspond to dimeric (58 ± 17 kDa) or tetrameric (113 ± 17 kDa) MinD complexes. For each variant the average dwell times were calculated according to equation (3)

\[
\bar{t} = \frac{\sum_{i=m}^{n} N_i t_i}{\sum_{i=m}^{n} N_i} - t_{mf},
\]

where \(N_i\) is the total number of plateaus with dwell time \(t_i\), and \(t_{mf}\) represents the most frequent dwell time. Due to the noise level of mass detections in our trajectories, short dwell times could not be quantitatively detected. Hence, dwell times shorter than 5 frames (25 ms) were excluded from the analysis.

The standard error of the mean dwell time was estimated from 10,000 bootstrap resamples.

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Author Contributions
T.H., F.S., N.H. contributed equally to this work. T.H., N.H., F.S. and P.S. conceived the study. T.H. designed and performed all experiments. B.R. designed and purified proteins. F.S. and N.H. analysed the data. T.H., F.S. and N.H. wrote the manuscript draft. All authors discussed and interpreted results. All authors revised, reviewed and approved the manuscript.

Competing Interests Statement
The authors declare no conflict of interest.

Data Availability Statement
The datasets generated and analysed during the current study are available from the corresponding authors on reasonable request.

Code Availability Statement
The custom-written Python script for particle detection (as binary executable) and particle analysis (source code accessible) is available from the corresponding authors on reasonable request.

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