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Reporting Summary

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Statistics

For	all statistical analyses, confirm that the following items are present in the figure legend, table legend, main text, or Methods section.
n/a	Confirmed
\ge	The exact sample size (<i>n</i>) for each experimental group/condition, given as a discrete number and unit of measurement
\boxtimes	A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
\boxtimes	The statistical test(s) used AND whether they are one- or two-sided Only common tests should be described solely by name; describe more complex techniques in the Methods section.
\ge	A description of all covariates tested
\boxtimes	A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
\boxtimes	A full description of the statistical parameters including central tendency (e.g. means) or other basic estimates (e.g. regression coefficient) AND variation (e.g. standard deviation) or associated estimates of uncertainty (e.g. confidence intervals)
\boxtimes	For null hypothesis testing, the test statistic (e.g. <i>F</i> , <i>t</i> , <i>r</i>) with confidence intervals, effect sizes, degrees of freedom and <i>P</i> value noted <i>Give P values as exact values whenever suitable.</i>
\ge	For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
\square	For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
\ge	Estimates of effect sizes (e.g. Cohen's <i>d</i> , Pearson's <i>r</i>), indicating how they were calculated
	, Our web collection on <u>statistics for biologists</u> contains articles on many of the points above.

Software and code

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Data collection	Cryo-electron tomography data were collected using commercial software FEI-tomo. NAMD v2.13 - parallel CPU/GPU code used for computer simulations
Data analysis	Cryo-electron tomographic volumes were initially reconstructed from tilt series using IMOD, and further processed using emClarity 1.0 (available and referenced in methods). VMD v1.9.4 - molecular visualization program used for basic analysis of computer simulations and rendering of molecular structures python v2.7 - version of python language and libraries used in this study matplotlib v2.2.3 - python module used for creating plot figures scipy v1.2.1 - python module used for performing hierarchical clustering analysis MDAnalysis v0.19.0 - python module used for basic analysis of computer simulations Chimera v1.13 - software used for map segmentation and and rigid-body docking

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors/reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Research guidelines for submitting code & software for further information.

Data

Policy information about availability of data

All manuscripts must include a <u>data availability statement</u>. This statement should provide the following information, where applicable: - Accession codes, unique identifiers, or web links for publicly available datasets

- A list of figures that have associated raw data
- A description of any restrictions on data availability

The cryoET density map of the E. coli 4Q core signaling unit is deposited in EMBD under accession code: EMD-10050. The raw tilt series will be deposited EMPIAR database. Coordinates for the corresponding atomic model are deposited in the RCSB Protein Data Bank under accession code 6S1K.

Field-specific reporting

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Life sciences Behavioural & social sciences Ecological, evolutionary & environmental sciences

For a reference copy of the document with all sections, see nature.com/documents/nr-reporting-summary-flat.pdf

Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

Sample sizeFor cryoEM structure determination, sample sizes were those required for the resolution. The details of datasets, including sample sizes, are
listed in table S1.Data exclusionsFor cryoET subtomogram averaging, standard 3D classification approaches in emClarity were used to exclude data.ReplicationFor cryoEM, two randomly divided half datasets were processed independently, and combined to give rise to the final structures. The
resolution of the structure is assessed by comparing the two independent maps.RandomizationcryoEM particles were randomly divided into ODD and EVEN datasets, as standard approach implemented in emClarity.BlindingN/A

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

Materials & experimental systems

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\boxtimes	ChIP-seq

- Flow cytometry
- MRI-based neuroimaging