The three-dimensional positions of atoms in protein molecules define their structure and their roles in biological processes. The more precisely atomic coordinates are determined, the more chemical information can be derived and the more mechanistic insights into protein function may be inferred. Electron cryo-microscopy (cryo-EM) single-particle analysis has yielded protein structures with increasing levels of detail in recent years. However, it has proved difficult to obtain cryo-EM reconstructions with sufficient resolution to visualize individual atoms in proteins. Here we use a new electron source, energy filter and camera to obtain a 1.7 Å resolution cryo-EM reconstruction for a human membrane protein, the β3 GABA<sub>α</sub> receptor homopentamer. Such maps allow a detailed understanding of small-molecule coordination, visualization of solvent molecules and alternative conformations for multiple amino acids, and unambiguous building of ordered acidic side chains and glycans. Applied to mouse apoferritin, our strategy led to a 1.22 Å resolution reconstruction that offers a genuine atomic-resolution view of a protein molecule using single-particle cryo-EM. Moreover, the scattering potential from many hydrogen atoms can be visualized in difference maps, allowing a direct analysis of hydrogen-bonding networks. Our technological advances, combined with further approaches to accelerate data acquisition and improve sample quality, provide a route towards routine application of cryo-EM in high-throughput screening of small molecule modulators and structure-based drug discovery.

Multiple factors determine the attainable resolution of reconstructions from single-particle cryo-EM. However, for biological macromolecules, the radiation damage caused by electron interactions with the sample is a fundamental limitation. To preserve the molecular structure, damage is restricted by carefully limiting the number of electrons used for imaging. The resulting counting statistics lead to high levels of noise. The signal-to-noise ratio (SNR) of cryo-EM images drops rapidly with spatial frequency, and at higher spatial frequencies the noise is typically orders of magnitude higher than the signal.

High-resolution reconstructions can still be calculated by averaging over many images of individual particles, provided that their relative orientations can be determined. However, because noise reduction scales with the square root of the number of particles, and because higher SNRs lead to more accurate orientations, acquiring more particles is often less efficient than increasing the SNR in the images. Consequently, although microscope automation and faster image processing programs have allowed reconstructions from larger datasets in recent years, increasing the SNR of the raw data may lead to bigger improvements, as is illustrated by the sudden increase in cryo-EM resolutions with the introduction of more sensitive direct electron cameras in 2013.

Here we describe the effect of three technological developments that further increase the SNR of cryo-EM images: a new cold field emission electron gun (CFEG), a new energy filter and the latest generation Falcon direct electron camera (Fig. 1a). Combined, these developments lead to a marked increase in the achievable resolution that ultimately enables the visualization of individual protein atoms in optimized samples.

**Electron source optimized for energy spread**

The source inside the microscope emits electrons with a range of different wavelengths, or energies. Because not all of these electrons can be focused in the same plane owing to chromatic aberration in the objective lens, the energy spread of the electrons leads to a blur in the images. The corresponding loss in SNR increases with spatial frequency and is described by an envelope on the contrast transfer function (CTF). Many state-of-the-art electron microscopes are equipped with a field emission gun (FEG) that is operated at a temperature of 1,700–1,800 K.
Elastically scattered electrons maintain their energy and contribute to the signal in the images. Inelastically scattered electrons deposit part of their energy in the sample and contribute noise to the images. Therefore, SNRs can be improved by filtering away electrons that have experienced an energy loss. Gatan pioneered the use of energy filters with a 90° bending prism for biological cryo-EM, and their filters are in widespread use. JEOl microscopes use a different, omega-shaped filter design. We used a new energy filter developed by TFS using the experience that previously led to the energy filter described by Kahl et al. The filter is located below the projector column of the microscope and comprises a 90° bending prism with an adjustable energy-selecting slit and multiple electromagnetic lenses to correct for aberrations and to enlarge both the energy dispersion and the image. The mechanics of the new filter were designed to minimize the effect of temperature variations on the position of the optical elements, including the energy slit, with respect to the optical axis of the system. The large radius of the bending prism (135 mm compared to 75 mm in Gatan filters) reduces third- and fourth-order distortions. The new filter is stable over many days of operation: the position of the electrons that have not lost any energy, the so-called zero-loss peak, with respect to the centre of the energy slit changes by less than 3 eV in either direction (Fig. 1d).

A next-generation camera

The relatively large pixel size of Falcon direct-electron cameras, each measuring $14 \times 14 \mu m^2$, permits the use of a thick epilayer that increases the SNR. Noise in the resulting images arises mainly from the variation in height and shape of the charge distribution that is generated by individual incident electrons, or events. This noise can be reduced by a counting algorithm that positions a constant signal distribution at an estimated centre position for each event. However, in order to identify all individual events, the number of events that are in close proximity to each other on a single frame should be minimized. With a frame rate of 40 Hz, the Falcon-3 was the first Falcon camera that was fast enough for electron counting, but only when the electron dose was spread over typical exposure times of close to a minute.

The Falcon-4 camera operates at 248 Hz, while an improved counting algorithm estimates event positions with sub-pixel accuracy and its epilayer design optimizes the event size versus event signal strength and thereby reduces the number of events that are lost during counting (Fig. 1e). As a result, images with typical exposure times of a few seconds can be recorded, while the detective quantum efficiency remains similar to that of the Falcon-3 camera in counting mode (Fig. 1f). In addition, the resulting videos can be written out in a new data format, called electron-event representation (EER), which stores each detected electron event as an $x$, $y$ position on a four-times oversampled grid and its original time, or frame, of recording. We adapted our image processing software RELION (see Methods) to read the new data format, thus allowing motion correction by the Bayesian polishing algorithm and reconstructions of video frames with flexible dose fractionation schemes, up to the original frame rate of the camera.

Application to a human GABA<sub>R</sub> receptor sample

To evaluate the effect of the technological advances described above, we used a homopentameric β3 γ-aminobutyric acid type-A receptor (GABA<sub>R</sub>) variant, an approximately 200-kDa human membrane protein bound to the Mb25 megabody to overcome its severe preferential orientation in ice. GABA<sub>R</sub> are a large family of pentameric ligand-gated chloride channels that mediate inhibitory neurotransmission and are targeted by a wide range of psychoactive drugs. However, structure-based drug development has been hampered by the difficulty of solving GABA<sub>R</sub> structures, which are currently limited to 2.5–3 Å resolution in the best cases. Whereas most GABA<sub>R</sub> are heteromeric, β3-subunits form functional homomeric channels that are gated by histamine. Their fivefold symmetry allowed us to collect multiple datasets in a time-efficient manner, as one would need five times more data to reach the same resolution with an asymmetrical sample.
We collected seven datasets from three different GABA<sub>R</sub> grids on three different Titan Krios microscopes (Extended Data Table 1, Extended Data Fig. 2) and assessed their quality in terms of their estimated B-factors<sup>24</sup> (see Methods, Fig. 2a). To assess the current state of the art, we collected two datasets on a first grid on a microscope with an XFEG using either a Gatan K3 camera and a Gatan energy filter (with a slit width of 20 eV), or a bottom-mounted Falcon-3 camera. This resulted in B-factors of 84 and 72 Å<sup>2</sup>, respectively. We note that data acquisition on the K3 was three times faster than on the Falcon-3. Therefore, reconstructions at a given target resolution may be obtained in less time on the K3. When we imaged a second grid on a microscope with an XFEG, the new energy filter (with retractable slits) and a Falcon-4 camera, an improvement of 15 Å<sup>2</sup> in the B-factor over the Falcon-3 dataset was observed. Using the same grid and the same microscope, a further improvement of 7 Å<sup>2</sup> in the B-factor was achieved when we used the energy filter with a slit width of 5 eV. Varying the energy slit widths to 3 eV and 10 eV did not have a noticeable effect on the B-factors (Extended Data Fig. 2a). Imaging a third grid on yet another microscope, with a CFEG, the new energy filter (with a slit width of 5 eV) and a Falcon-4 camera, led to a small improvement for those reconstructions with resolutions better than 2 Å. The latter is expected from the relatively small gains in the CTF envelope at those resolutions when using a CFEG instead of an XFEG (Fig. 1c). Finally, we performed tomography on the same GABA<sub>R</sub> grid used in the CFEG experiment to determine its ice thickness, which we found to be in the range of 320–370 Å (Extended Data Fig. 2b).

By combining the datasets with the new filter on both the XFEG and the CFEG, we obtained a 1.7 Å resolution reconstruction from 371,693 particles (Fig. 2b–d, Extended Data Figs. 2c–g, 3, Extended Data Table 2). The resulting map provides unprecedented levels of detail for a protein structure, other than apoferritin, solved by single-particle cryo-EM. The N-linked glycan on Asn149 shows density of exceptional quality; alternative side-chain conformations are visible for many amino acids; lipid molecules are visible in the transmembrane region of the receptor; and in the most ordered regions, hydrogen-bonding networks can be visualized directly in difference maps calculated in REFMAC<sup>25</sup> with a refined model from which the hydrogen atoms were omitted. Each agonist pocket is occupied by a histamine molecule, which binds in a similar manner to that observed for benzamidine<sup>1</sup> and the neurotransmitter GABA<sup>21</sup>. However, it is now possible to visualize the coordinating carboxyl groups of Glu155 and Asp43, alongside a network of ordered water molecules that surround the ligands to fill the agonist pocket.

A video of the dose-dependent evolution of the cryo-EM density indicates that the structure of the agonist pocket may be affected by radiation damage (Supplementary Video 1). Similarly, the disulfide bond between Cys136 and Cys150, a signature of the Cys-loop receptor family to which GABA<sub>R</sub> belongs, appears to be partially reduced during the electron exposure (Supplementary Video 2).

**Atomic-resolution structure of apoferritin**

To explore the potential of the new technologies to yield even higher-resolution reconstructions, we tested a mouse apoferritin sample. Apoferritin is a popular cryo-EM benchmark because its molecular stability and its 24-fold symmetry allow high-resolution reconstructions to be calculated from relatively few particles. Using JEOL’s CFEG and omega energy filter with a K2 camera and the same mouse apoferritin sample as we use here, but a different grid, resulted in the best apoferritin map reported thus far, to our knowledge: 1.53 Å resolution from 120,295 particles, with a B-factor of 46 Å<sup>2</sup><sup>11</sup>. We previously also reported a reconstruction of human apoferritin with a resolution of 1.65 Å from 426,450 particles recorded on a Titan Krios microscope with an XFEG, a Gatan energy filter and a Gatan K2 camera<sup>19</sup>. The B-factor for that dataset was 66 Å<sup>2</sup>

We imaged mouse apoferritin on the CFEG microscope, using the Falcon-4 camera and a slit width of 10 eV in the energy filter. From 3,370 videos, we obtained a reconstruction with a resolution of 1.22 Å from 363,126 particles (Extended Data Fig. 4, Supplementary Video 3). Corrections during image processing, for third and fourth-order optical aberrations<sup>26</sup>, as well as for the curvature of the Ewald sphere<sup>27,28</sup>, were crucial in obtaining a B-factor of 32 Å<sup>2</sup> (Fig. 3a). After sharpening of the map, most atoms are resolved as individual blobs of density; C=O bonds and some of the C–N bonds in the peptide planes are less well separated (Fig. 3b–d). Many hydrogen atoms, including those from individual water molecules, are resolved in a REFMAC difference map, more clearly than in the case of the GABA<sub>R</sub> receptor and allowing a direct analysis of hydrogen-bonding networks between β-strands.

**Fig. 2 | GABA<sub>R</sub> reconstructions.** a, B-factor plots for five datasets using: the new CFEG, the new energy filter with a slit width of 5 eV and a Falcon-4 camera (orange); an XFEG, the new energy filter with a slit width of 5 eV and a Falcon-4 (blue); an XFEG, the new energy filter with the slits retracted and a Falcon-4 (grey); an XFEG, no energy filter and a (bottom-mounted) Falcon-3 detector (black); and an XFEG, a Gatan imaging filter (GIF) and a K3 detector (brown). B-factors estimated from the slopes of fitted straight lines are shown in the same colours. Numbers in parentheses and error bars represent estimated and sample s.d.s from sevenfold random resampling, respectively. b, Overview of map quality at the agonist binding pocket, illustrating histamine (HSM) coordination and multiple water molecules (red spheres). c, The N-acetyl glucosamine (NAG) moieties attached to Asn149. d, Difference map (green, positive; orange, negative) visualizes hydrogen atoms in the hydrogen-bonding network between β-strands.

A video of the dose-dependent evolution of the cryo-EM density indicates that the structure of the agonist pocket may be affected by radiation damage (Supplementary Video 1). Similarly, the disulfide bond between Cys136 and Cys150, a signature of the Cys-loop receptor family to which GABA<sub>R</sub> belongs, appears to be partially reduced during the electron exposure (Supplementary Video 2).
Finally, we calculated reconstructions for each of the 434 camera frames that were acquired per exposure and were able to follow the dose-dependent evolution of the cryo-EM density with an unprecedented resolution of 0.1 electrons (e⁻) per Å² (Supplementary Video 4). Following the initial dose of 1.5 e⁻ Å⁻², the resolution of the single-frame reconstructions extended to 1.7 Å and remained better than 2 Å up to an accumulated dose of 11.5 e⁻ Å⁻². However, the first few reconstructions were of relatively poor quality, probably because initial beam-induced motions were not modelled adequately. As radiation damage is minimal in the earliest frames, further improvements in beam-induced motion correction, possibly by considering particle rotations and height changes²⁸, may therefore extend the attainable resolution even further.

Outlook

The three developments in electron microscopy hardware described in this paper provide a step-change in the resolution that can be achieved using single-particle cryo-EM. Using apoferritin, we have shown that these developments allow the determination of the structure of a protein to true atomic resolution, as per the Sheldrick criterion²⁹,³⁰. The improved energy spread of the CFEG is crucial for extending resolutions to better than 1.5 Å, consistent with similar observations made with the JEOL CFEG³⁰. There are two alternatives to the CFEG strategy to improve the CTF envelope function at atomic resolution. First, one could use objective lenses with a lower chromatic aberration, which are common in material science, although their narrow gap polepieces would limit the amount by which the sample could be tilted. Second, one could reduce the energy spread by using a monochromator³¹. The disadvantage of this strategy would be a considerably more complicated experimental setup.

Our results for the GABA⁺R illustrate how the new technology can improve cryo-EM structures at lower resolutions, beyond the highly stable test samples that are represented by apoferritin. Although variations between microscopes and grids can complicate direct comparisons of B-factors, our results for the GABA⁺R suggest that the Falcon-4 camera in combination with the new filter provides the best image quality (Fig. 2a). At this stage, it is unclear whether the removal of inelastically scattered electrons from our thin samples alone would be enough to account for this improvement, or whether other effects—for example, increased amplitude contrast—³²—are also involved. In the future, it will also be of interest to explore whether, instead of removing them, inelastically scattered electrons could still be used to add to the signal in a chromatic-aberration-corrected microscope³³,³⁴.

The increased SNR of cryo-EM images enabled by the technology described here will expand the applicability of both single-particle analysis and electron tomography to more difficult samples, including membrane proteins in lipid bilayers, small proteins and structurally heterogeneous macromolecular complexes. Moreover, higher-resolution reconstructions will allow improved visualization of hydrogen-bonding networks, ordered waters and alternative side-chain conformations, which will further our understanding of protein function and facilitate cryo-EM fragment-based drug discovery.

Online content

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**Methods**

**Characterization of the new microscopy hardware**

The energy spread of the field emission gun was measured using the new energy filter. Post-slit multi-poles of the filter were set to spectroscopy mode in order to form an enlarged image of the spectrum on the camera. The intensity of the incoming beam was carefully reduced so as not to saturate or damage the camera. The energy dispersion at the camera was 8 meV/pixel. The exposure time was 31 ms. The energy spread as FWHM of $\Delta E = 0.71$ for XFEG, and $\Delta E = 0.28$ for CFEG (Fig. 1b) were used to calculate the CTF envelope function, defocus spread $f_C$, using the following equation:\(^4\)

$$
\Delta f = C_e \sqrt{(\Delta E/eV)^2 + (\Delta V/V)^2 + (2\Delta I/I)^2},
$$

using chromatic aberration constant $C_e = 2.7$ mm; gun energy spread $\Delta V/V = 0.02$ ppm and objective lens current instability $\Delta I/I = 0.1$ ppm. Note that the latter two contributions account for less than a 1% drop in the CTF envelope.

The stability of the energy filter was assessed by repeatedly measuring the position of the zero-loss peak with respect to the energy-selecting slit. To this end, we mechanically moved one slit edge to the optical axis and fine-tuned an offset on the high tension supply such that the slit edge blocked half of the beam intensity. The resulting offset of the high tension was read out as the shift in the zero loss position. Measurements, each taking approximately 40 s, were repeated continuously for 14 days.

Dose–response curves for the Falcon-3 and Falcon-4 cameras were measured using the flu-screen and fixed magnification steps. Before each experiment, the flu-screen was carefully calibrated with a Faraday cup in its linear range (0.4–5.0 nA at 300 kV). The read-out from the flu-screen was interpreted as the expected electron dose and plotted against the counted number of electrons on the camera.

DQE measurements were performed as described\(^3\) at 300 keV on a Titan Krios G2 microscope at the LMB for the Falcon-3 and on a Titan Krios G3i microscope at the Materials Science Department of the University of Cambridge, UK for the Falcon-4. For calibration of the flu-screen on the microscope with the Falcon-4, a Faraday cage and a Keithley 6485 picoammeter (provided by G. van Hoften, Thermo Fisher Scientific) were used. For the microscope with the Falcon-3, the drift tube of a Gatan energy filter (BioQuantum 686e) was used as a Faraday cup and the current was measured using a Deben 087-001 SEM probe current meter. Because the beam currents used in counting mode imaging are too low to be accurately measured, a defined beam with a measurable current was set at a low magnification and the desired electron flux on the camera was obtained by increasing the magnification. The relative magnification change was measured by using a higher current to fully illuminate the flu-screen at the low magnification and noting the change in current upon going to the higher magnification.

To account for the nonlinearity that is introduced by coincidence loss, DQEs were first measured at a low dose rate (0.19 e⁻/pixel/s for the Falcon-3; 0.32 e⁻/pixel/s for the Falcon-4) and then extrapolated to a typical dose rate (0.5 e⁻/pixel/s for the Falcon-3; 3.6 e⁻/pixel/s for the Falcon-4) using the measured response versus dose-rate curves (Fig. 1e). Modulation transfer functions (MTFs) were measured using the knife-edge method from a straight part of a shadow of the pointer on the camera. The actual area of the images used was chosen carefully by looking at the difference between the observed image and that predicted from the inferred MTF, to ensure that the edge in the selected area was straight and free of defects. In the case of the Falcon-4, the only part of the pointer edge that was suitable was at a 15° angle to the pixel array of the camera, the effect of which was accounted for when calculating the MTF. Even at the lowest dose rates used, there was still a small decrease in the power spectra at low spatial frequencies due to coincidence loss. To avoid overestimating the resulting DQE at low spatial frequency, the noise power spectra were extrapolated from a Gaussian fit of the power between 0.2 and 0.8 of Nyquist. The amplitudes and widths of fitted Gaussians to the noise power spectra were in excellent agreement with those expected from the number of counted electrons per frame and estimates from the point spread functions of images of individual incident electrons.

**GABA<sub>\text{A} \text{R} \text{ receptor production, purification and nanodisc reconstitution****

Human GABA<sub>\text{A} \text{R} \text{ receptor} β<sub>2</sub>-subunit with a truncated M3–M4 loop was transiently expressed in HEK293S GnTI- cells (which lack N-acetylglucosaminyltransferase I activity) as previously described\(^1\). The sample used for XFEG datasets collected on a Falcon-4 contained a point mutation (K279T) that is thought to stabilize the receptor and a C-terminal ID4 tag for purification. The sample used for the CFEG dataset, as well as the Falcon-3 and K3 datasets, contained the wild-type sequence and an N-terminal streptavidin-binding peptide (SBP) tag for purification. Subsequent comparison of the resulting cryo-EM maps revealed that the K279T mutation had no structural impact, so the Falcon-4 datasets could be merged during data processing. Frozen cell pellets were resuspended in 30 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) pH 7.6, 300 mM NaCl, ≥1 mM histamine supplemented with 1% (v/v) mammalian protease inhibitor cocktail (Sigma-Aldrich) and solubilized with 1% (w/v) lauryl maltose neopentyl glycol (LMNG) and 0.1% (w/v) cholesteryl hemisuccinate Tris salt (CHS). No CHS was added to the sample that was used for the CFEG dataset. Receptors were purified by ID4 and SBP affinity chromatography. Nanodisc reconstitution was performed as previously described\(^2\).

In brief, after the binding step, while the receptor was still bound to the affinity beads, an excess of bovine brain extract (BBE; type I, Folch fraction I, Sigma-Aldrich, 20 mg/ml stock) and phosphatidylcholine (POPC, Avanti, 10 mg/ml) mixture (15:85, v/v) was added to the samples and incubated for 30 min. Receptors were then reconstituted into membrane scaffold protein 2N2 (MSP2N2) lipid nanodiscs by incubating with an excess of MSP2N2 (0.6 mg/ml final concentration) and bio-beds for 2 h. The resin was then washed extensively and the GABA<sub>\text{A} \text{R} \text{ samples were eluted with a buffer containing 12.5 mM HEPES pH 7.6, 75 mM NaCl, 1 mM histamine and 2 mM ID4 peptide or 2.5 mM biotin.**

**Cryo-EM grid preparation**

Purified GABA<sub>\text{A} \text{R} \text{ was incubated with ~1 μM Megabody-251} and a 3.5-µl sample was applied to glow-discharged 300 mesh 1.2/1.3 and 2/2 UltraAuFoil gold grids (Quantifoil) for 30 s then blotted for 5.5 s before plunge-freezing with liquid ethane cooled by liquid nitrogen. Grid vitrification was performed using a Leica plunger (Leica Microsystems; XFEG datasets) or Vitrobot Mark IV (Thermo Fisher Scientific; CFEG datasets) at 100% humidity and 4 °C. A frozen aliquot of 7 mg/ml mouse apoferritin in 20 mM HEPES pH 7.5, 150 mM NaCl, 1 mM dithiothreitol (DTT) and 5% trehalose, which we received from the Kikkawa laboratory at Tokyo University, was thawed at room temperature and cleared by centrifugation at 10,000g for 10 min. The supernatant was diluted to 5 mg/ml with 20 mM HEPES pH 7.5, 150 mM NaCl, and 3 µl of the diluted sample was applied onto glow-discharged R1.2/1.3 300 mesh UltrAuFoil gold grids (Quantifoil) for 30 s and then blotted for 5 s before plunge-freezing the grids into liquid ethane cooled by liquid nitrogen. Plunge-freezing was performed using a Vitrobot Mark IV (Thermo Fisher Scientific) at 100% humidity and 4 °C.

**Cryo-EM data acquisition**

All cryo-EM data were collected on Falcon or K3 cameras in electron counting mode using Titan Krios microscopes (Thermo Fisher Scientific) operating at 300 kV. Before data acquisition, twofold astigmatism was corrected and beam tilt was adjusted to the coma-free axis using the autoCTF program (Thermo Fisher Scientific). All datasets
were acquired automatically using EPU software (Thermo Fisher Scientific). Detailed data acquisition parameters for all datasets are given in Extended Data Table 1.

For GABA\textsubscript{R}, two datasets were acquired on a Titan Krios at the Department of Biochemistry, University of Cambridge, UK. This microscope is equipped with an XFEG and a bottom-mounted (BM) Falcon-3 camera and a K3 camera behind the Gatan energy filter. All other datasets were acquired at the Thermo Fisher Scientific R&D division in Eindhoven, The Netherlands. Four GABA\textsubscript{R} datasets were collected on a Titan Krios equipped with an XFEG and a prototype Falcon-4 camera that was mounted behind the new energy filter. These datasets were collected consecutively over a period of five days from the same grid to study the effect of varying energy slit widths. To keep the ice thickness similar between these experiments, all the exposure areas were selected at the start using a constant ice filter within EPU software. One GABA\textsubscript{R} dataset was acquired on a microscope with a CFEG and a prototype Falcon-4 camera that was mounted behind the new energy filter. The slit width of the energy filter was set to 5 eV. For this dataset, 5,573 videos were acquired using a 100-μm objective aperture and 3,160 videos were collected without an objective aperture. The CFEG was automatically flashed every 5 h using the EPU software.

For the apoferritin dataset, we used the same microscope with the CFEG, the new filter and the prototype Falcon-4 camera. No objective aperture was used and the slit width of the energy filter was set to 10 eV. Flashing of the CFEG had no noticeable effect on data quality (Extended Data Fig. 4f).

Data acquisition was not optimized in terms of speed and the prototype Falcon-4 version did not write videos at the highest possible rate. Typically, two videos were collected per hole, with physical stage movements in between each hole. The resulting data acquisition rate was -96 videos per hour (including Dewar filling times and CFEG flashing). The 3,370 apoferritin videos were collected within 36 h; GABA\textsubscript{R} datasets allowed -2 Å reconstructions in -9 h. Fringe-free imaging (FFI) and aberration-free image shift (AFIS), which allow the acquisition of multiple images per hole and multiple holes per stage position, would allow considerably higher throughputs.

The EER movie format

Electron event representation (EER) is a video format that is introduced with the Falcon-4 camera\textsuperscript{28}. Whereas in the existing MRC image format groups of subsequent camera frames, or dose fractions, are summed and represented as images, EER stores the location and time format groups of subsequent camera frames, or dose fractions, are produced with the Falcon-4 camera\textsuperscript{18}. Whereas in the existing MRC image representation (EER) is a video format that is introduced as binary pixels in a 8k × 8k grid, which is then Fourier cropped into a 4k × 4k grid. Using the modified version, EER movies can be used for frame alignment using RELION’s implementation of the MotionCor2 algorithm\textsuperscript{36} and for per-particle beam-induced motion correction using its Bayesian polishing program\textsuperscript{25}. For both programs, the user needs to specify a dose fractionation rate.

Ice-thickness measurement

The ice thickness was measured on a GABA\textsubscript{R} grid using tomography. Tilt series were collected on three different grid-squares with a 10-eV slit on a Falcon-4 camera. Data were acquired covering an angular range from -45° to +45° with 3° angular increments recorded automatically using the dose-symmetric tilting scheme under low-dose conditions using TEM tomography software (TFS). Each tilt series was collected with a nominal defocus value of 5 μm in electron counting mode using a dose of 4 e\textsuperscript{-/Å\textsuperscript{2}} per tilt and a calibrated pixel size of 1.898 Å. Tomograms were reconstructed with patch tracking and ice thickness measured with IMOD software\textsuperscript{27}.

Apoferritin image processing

A total of 3,370 videos in EER format were motion corrected with RELION’s implementation of the MotionCor2 algorithm\textsuperscript{38}. For this purpose, original hardware video frames were dose-fractionated into groups of 14 frames, corresponding to an accumulated dose of 1.3 e\textsuperscript{-/Å\textsuperscript{2}} per fraction. CTF estimation was performed with CTFFIND-4.1.13\textsuperscript{38} using the sums of power spectra from combined fractions corresponding to an accumulated dose of 4 e\textsuperscript{-/Å\textsuperscript{2}}. Micrographs whose estimated resolution from CTFFIND was worse than 5 Å were removed, leaving 3,080 micrographs for further processing.

Using RELION’s template-based algorithm, 428,590 particles were picked and subjected to 2D classification. After selection of good classes, 380,236 particles were subjected to standard 3D auto-refinement to give an initial reconstruction with an estimated resolution of 2.3 Å. Subsequently, three runs of CTF refinement\textsuperscript{38} were performed: first refining magnification anisotropy; then refining optical aberrations (up to the fourth order); and finally refining per-particle defocus and per-micrograph astigmatism. A 3D auto-refinement with the refined values yielded an estimated resolution of 1.7 Å. Bayesian polishing to optimise per-particle beam-induced motion tracks, followed by another round of auto-refinement, resulted in a 1.43 Å map. CTF refinement was then repeated for optical aberration correction, magnification anisotropy, per-particle defocus and per-micrograph astigmatism. To compensate for changes in the tilt of the electron beam during the data acquisition session, separate optics groups were created for particles belonging to consecutive groups of 125 micrographs, or whenever the CFEG was flashed. At this point, 201 micrographs with refined scale factors (rlnGroupScaleCorrection) with values below 0.5 were also removed, resulting in 363,126 particles in the final set (Extended Data Fig. 4f). Following a last round of auto-refinement, these particles were subjected to a second round of Bayesian polishing with a large particle extraction box (spanning 320 Å), such that high-resolution signals that were delocalized by the CTF were still contained in the images. At this point, seven hardware frames were grouped into each dose fraction to be able to capture more rapid motions. As Ewald sphere curvature was predicted to limit the resolution of a particle with a diameter of 120 Å to approximately 1.3 Å\textsuperscript{39}, the particles were then used for reconstruction with Ewald sphere correction\textsuperscript{27}. The estimated resolution of the final map, following standard post-processing procedures in RELION, was 1.22 Å.

GABA\textsubscript{R} receptor image processing

GABA\textsubscript{R} datasets were processed following the same strategy as for apoferritin, with the following exceptions. All videos, except for those acquired on the microscope with a CFEG, were recorded in MRC format. For the CFEG data, hardware video frames were initially dose-fractionated into groups of 24 frames, corresponding to an accumulated dose of 0.86 e\textsuperscript{-/Å\textsuperscript{2}} per fraction. For the second round of Bayesian polishing, a dose fractionation of 12 hardware frames was used. Particle picking was performed using an in-house generated Linux port of the of BoxNet2D neural network in Warp\textsuperscript{40}. Prior to the last auto-refinement, 3D classification into three classes without alignment
was performed to select the particles contributing to the best reconstruction. The first part of the CFEG data was acquired with a 100-μm objective aperture. Charging of the aperture probably led to third- and fourth-order optical aberrations, which were estimated during CTF refinement in optics groups of 160 micrographs, or whenever the CFEG was flashed or liquid nitrogen was refilled. The second part of the CFEG data was acquired without an objective aperture. For these images, no significant higher-order optical aberrations were observed and creating multiple optics groups was deemed unnecessary. When merging the energy-filtered CFEG and XFEG datasets, refinement of per-micrograph scale factors and per-particle normalization corrections were unstable, and these were switched off using the –no_norm and –no_scale arguments. The mask used for resolution estimation during post-processing contained density for GABAAR only; all density corresponding to the megabodies was removed.

B-factor estimation

For each GABAAR dataset, a random subset of 32,000 particles was selected from the final set of particles after the second round of Bayesian polishing. This subset was subjected to 3D auto-refinement and the resulting orientations were used to calculate reconstructions for each of the two random halves used in the auto-refinement. Similar half-set reconstructions were also calculated for random subsets of 16,000, 8,000, 4,000, 2,000, 1,000 and 500 particles. This procedure was repeated seven times. For each subset and repeat, resolution estimation was again performed using standard post-processing functionality in RELION. The inverse square of the resulting estimated resolutions were then plotted against the natural logarithm of the number of particles in the subset, and B-factors were calculated from the slope of a straight line fitted through all points in the plot. For apoferritin, the same procedure was used, starting from a subset of 180,000 particles and halving the subset size eight times.

Atomic modelling

The atomic models for GABAAR and apoferritin were derived from PDB-4cof and PDB-6s61, respectively. Both models were manually adjusted in COOT. Difference maps were used to rebuild the model and to add alternative conformations and water molecules. Real-space refinement was performed using PHENIX, version 1.16.3549. Both models were refined with anisotropic atomic B-factors in reciprocal space using REFMAC, version 5.8.0258. For cross-validation, the atomic models refined against the full map were perturbed by random shifts of the particles of up to 0.3 Å and the perturbed model was refined in REFMAC against one of the two half-maps. FSC curves for the resulting model against that half-map (FSCwork) and against the other half-map (FSCref) were calculated for each of the two random halves used in the auto-refinement. FSC was calculated from half maps. Structural figures were prepared using PyMOL (Schrödinger) and UCSF ChimeraX.

Hydrogen position calculations

X-ray scattering and electron scattering factors are related through the Mott–Bethe formula, which is the solution of Poisson equations in Fourier space under certain conditions:

\[ f_c(s) = c \frac{Z - f_x(s)}{s^2} \]

Where \( c = \frac{2\pi m e^2}{h^2} \), \( m \) is the mass of an electron; \( e \) is the charge of an electron; \( \hbar \) is the Planck constant; \( \epsilon_0 \) is the dielectric constant of the vacuum; \( Z \) is the nuclear charge; \( f_x(s) \) is the X-ray scattering factor; \( f_c(s) \) is the electron scattering factor; and \( s = 1/d \) is the resolution. Adding the position-independent, isotropic atomic B-factor gives:

\[ f_{c,B}(s) = c \frac{Z - f_x(s)}{s^2} e^{-\frac{B_s^2}{4}} \]

This formula is valid if the centre of electron density coincides with the position of the nucleus. If this is not the case, the formula should be modified according to the vector \( \Delta x \) between the centre of electron density and the nuclear position (assuming that the centre of electron density is at the origin):

\[ f_{c,B}(s) = c \frac{Z e^{-2m\Delta x} - f_x(s)}{s^2} e^{-\frac{B_s^2}{4}} \]

Note that, in general, the form factor for electron scattering becomes a complex quantity. For all non-hydrogen atoms, it is safe to assume that the centre of electron density and the nuclear position coincide. For hydrogen atoms, the centre of electron density and the nuclear position do not coincide and we are interested in the peak height for the density of electrostatic potential along the bond between a hydrogen and its parent atom. The formula below ignores the fact that electron densities will be non-spherical, as they are elongated along the bond, thus adding additional complexity to the Mott–Bethe formula.

The densities for the proton and the electron of a hydrogen atom at a resolution \( s_{max} \) and with a given B value is given by:

\[ \psi_p(x) = c \int_{|s|=s_{max}} \frac{Z e^{-\frac{B_s^2}{4}} e^{-2m\Delta x}}{2\pi |s||x|} ds = 4\pi c \int_{0}^{s_{max}} f_x(s) e^{-\frac{B_s^2}{4}} \frac{\sin(2\pi s|s||x|)}{2\pi |s||x|} ds, \]

\[ \psi_e(x) = 4\pi c \int_{0}^{s_{max}} f_x(s)e^{-\frac{B_s^2}{4}} \frac{\sin(2\pi s|s||x|)}{2\pi |s||x|} ds \]

To calculate the density at a position \( x \) we need to calculate:

\[ \psi_{p+e}(x) = \psi_p(x - \Delta x) - \psi_e(x) \]

Even if both densities corresponding to the proton and the electron are spherically symmetric, the combined density will not be spherical. Extended Data Fig. 5 shows density profiles and their maxima for a hydrogen bonded to a carbon atom for different resolutions and B values. These profiles were calculated with the assumption that the electron–carbon distance is 0.98 Å and the proton–carbon distance is 1.09 Å.

Reporting summary

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

Data availability

Atomic coordinates for the GABAAR construct and apoferritin have been deposited in the Protein Data Bank with accession codes 7A5V and 7A4M, and the cryo-EM density maps have been deposited in the Electron Microscopy Data Bank with accession codes EMD-11657 and EMD-11638, respectively. Raw movies of all datasets have been deposited in the Electron Microscopy Public Image Archive (https://www.ebi.
ac.uk/pdbe/emdb/epmir/) with accession codes EMPIAR-10500 for GABA\(_R\) and EMPIAR-10424 for apoferritin.

**Code availability**

Both the implementation of reading EER movies into RELION and the EERformat itself have been distributed as free, open-source software, with downloads available from https://github.com/3dem/relion and https://github.com/fei-company/EerReaderLib. The in-house-generated Linux port of the BoxNet2D neural network in Warp is available from https://gist.github.com/biochem-fan/76ff5934495585da56ab2c0af4fe2563.

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**Author contributions** E.d.J., J.K., M.B., J. McCormack and P.T. contributed to microscope hardware and software developments; G. McMullan performed DQE analysis; D.K., A.S. and S.M. prepared cryo-EM grids; T.U. produced megalodons; A.K., L.Y., E.V.P., S.W.H. and D.Y.C. acquired cryo-EM data; T.N., A.K., A.S., L.Y., S.M., P.M.G.E.B., I.T.G., L.M., J. Miehling and S.H.W.S. analysed cryo-EM data; T.N., A.R.A., T.M. and G. Murshudov prepared atomic models; T.N., A.K., A.R.A. and S.H.W.S. conceived the project and designed the experiments; G.M., A.R.A. and S.H.W.S. wrote the manuscript, with input from all authors.

**Competing interests** A.K., S.M., L.Y., D.K., E.V.P., E.d.J., J.K., M.B., J. McCormack and P.T. are employees of Thermo Fisher Scientific. The other authors declare no competing interests.

**Additional information**

**Supplementary information** is available for this paper at https://doi.org/10.1038/s41586-020-2829-0.

**Correspondence and requests for materials** should be addressed to A.R.A. or S.H.W.S.

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Extended Data Fig. 1 | Characteristics of the new cryo-EM technology. **a**, Four consecutive measurements of the CFEG beam current over a period of nine hours. The FEG tip was flashed just before the start of each measurement. **b**, The dose at the sample as measured for the same four experiments in a.
Extended Data Fig. 2 | Cryo-EM for GABA<sub>R</sub>. a, B-factor plots for three datasets using an X-FEG, the new energy filter with a slit width of 3 eV (orange), 5 eV (blue) and 10 eV (grey) and a Falcon-4 camera. b, Two orthogonal views of an electron tomogram for ice thickness measurement. Scale bar is 50 nm. c, Representative electron micrograph from the CFEG dataset. Scale bar is 30 nm. d, A selection of 2D class average images. e, Local resolution map. f, Fourier Shell Correlation (FSC) between the two independently refined half-maps. g, FSC between the model and the map calculated for the model refined against the full reconstruction (black); the model refined in the first half-map against that half-map (FSC<sub>0.5</sub>, blue); and the model refined in the first half-map against the second half-map (FSC<sub>0.5</sub>, dashed orange). Atomic models were refined including spatial frequencies up to 1.63 Å.
Extended Data Fig. 3 | GABA$_\text{A,R}$ reconstruction details. a, The GABA$_\text{A,R}$ cryo-EM map viewed from the extracellular space (top view). The density of one subunit of the homopentamer is highlighted in yellow; glycans are orange; the nanobody domain of Mb25 green; and the lipid nanodisc light blue. b, As in a, but viewed parallel to the plasma membrane space (side view). c, Stacking of histamine with aromatic residues in the ligand-binding pocket. Water molecules removed for clarity. d, Radiation damage in the ligand-binding pocket illustrated by difference maps of Asp43 and Glu155 carboxyl groups. e, Radiation damage causes partial reduction of the disulfide bond between Cys136 and Cys150. Left panel: oxidized state, right panel: reduced state. Only Cα, Cβ and sulfur atoms depicted for clarity. f, Difference map revealing the hydrogen bonding network between β-strands. g, A close-up view of the lipids (blue) surrounding the transmembrane region of the receptor. Compared to a, b, the contour level is decreased. Red arrow indicates section level as depicted in h. The general anaesthetics pocket and the neurosteroid modulation site are indicated by the red circle and rectangle, respectively. h, Top-view of a transverse section through the GABA$_\text{A,R}$ transmembrane region shows semi-ordered lipids surrounding individual subunits. i, Close-up view of the general anaesthetics pocket. j, Close-up view of the neurosteroid modulation site. Lipids in i and j could not be identified unambiguously and therefore are modelled as aliphatic chains.
Extended Data Fig. 4 | See next page for caption.
Extended Data Fig. 4 | Cryo-EM for apoferritin. 

a, Representative electron micrograph. Scale bar is 20 nm. 
b, 2D class average images. 
c, Local resolution map. 
d, Fourier Shell Correlation (FSC) between the two independently refined half-maps. 
e, FSC between the model and the map as calculated for the model refined against the full reconstruction against that map (black); the model refined in the first half-map against that half-map (FSCwork; blue); and the model refined in the first half-map against the second half-map (FSCtest; dashed orange). FSCwork and FSCtest curves are shown for three different models: with hydrogens and anisotropic B-factors (solid lines); with hydrogens and isotropic B-factors (dashed lines); and without hydrogens and with isotropic B-factors (dotted lines). Including hydrogens and using anisotropic B-factors gives a better fit to the data at medium resolution, but lead to a small amount of overfitting at high resolution. Atomic models were refined including spatial frequencies up to 1.15 Å. 
f, The beam tilt in X (blue) and Y (orange); the resolution (res.) for subsets of 5,000 consecutive particles (black); and RELION’s rlnGroupScaleCorrection values for individual micrographs (blue dots) during the data acquisition experiment. Events like CFEG flashing (red), liquid nitrogen filling (black) and moving the grid to another grid square (dark blue) are indicated with vertical lines. The cutoff in scale correction values used to discard part of the micrographs is indicated with a dashed grey line.
Extended Data Fig. 5 | Electrostatic potential of hydrogen atoms.

a, Calculated profile (see Methods) of the electrostatic scattering potential along the bond between a carbon and a hydrogen atom, for a $B$-value of 10 Å$^2$ and resolutions ($d$) of 1.0 Å (orange), 1.2 Å (blue), 1.5 Å (grey) and 1.8 Å (black). The two vertical lines indicate the electron-carbon distance of 0.98 Å and the proton-carbon distance of 1.09 Å. b, As in a, but for a resolution of 1.2 Å and $B$-values of 5 Å$^2$, 10 Å$^2$, 15 Å$^2$ and 20 Å$^2$. c, Distances from the carbon atom (in Å) at which the calculated profile of the electrostatic scattering potential along the bond between a carbon and a hydrogen atom is at its maximum, for different resolutions ($d$) and $B$-values.
## Extended Data Table 1 | Cryo-EM data collection parameters

| Data collection and processing | Apoferritin | GABAα-β3 | GABAα-β3 | GABAα-β3 | GABAα-β3 | GABAα-β3 | GABAα-β3 | GABAα-β3 |
|-------------------------------|-------------|-----------|-----------|-----------|-----------|-----------|-----------|-----------|
| Electron Gun Detector        | CFEG        | CFEG      | XFEG      | XFEG      | XFEG      | XFEG      | XFEG      | XFEG      |
| Magnification                | 270k        | 165k      | 165k      | 165k      | 165k      | 165k      | 96k       | 130k      |
| Energy filter slit width (eV)| 10          | 5         | open      | 10        | 5         | 3         | NA        | 20        |
| Voltage (kV)                 | 300         | 300       | 300       | 300       | 300       | 300       | 300       | 300       |
| Flux on detector (e-/pix/sec)| 4.5         | 4.7       | 2.7       | 2.8       | 2.8       | 2.8       | 0.7       | 14.9      |
| Electron exposure on sample (e-/Å²)| 40 | 40 | 35 | 35 | 35 | 35 | 37 | 40 |
| Target defocus range (µm)    | 0.3-0.9     | 0.3-1.1   | 0.9-1.8   | 0.3-1.1   | 0.3-1.1   | 0.3-1.1   | 0.5-1.5   | 0.3-1.5   |
| Observed defocus range (95th percentile in µm) | 0.27-0.97 | 0.30-1.11 | 0.80-1.83 | 0.31-0.95 | 0.32-0.87 | 0.34-1.00 | 0.58-1.51 | 0.48-1.46 |
| Calibrated pixel size (Å)    | 0.457       | 0.727     | 0.735     | 0.735     | 0.735     | 0.735     | 0.827     | 0.65482   |
| Symmetry imposed             | O           | C5        | C5        | C5        | C5        | C5        | C5        | C5        |
| Number of collected movies   | 3370        | 8733      | 1244      | 1268      | 1556      | 1952      | 1501      | 1507      |
| Initial particle images (no.)| 428590      | 1105069   | 192053    | 199867    | 242059    | 264675    | 412913    | 215957    |
| Final particle images (no.)  | 363126      | 233576    | 33166     | 42780     | 45009     | 50328     | 129810    | 46441     |
| Map resolution at FSC=0.143 (Å) | 1.22       | 1.73      | 1.97      | 1.89      | 1.87      | 1.88      | 1.97      | 2.32      |
### Extended Data Table 2 | Refinement and validation statistics

| Refinement | Apoferritin (EMD-11638) (PDB 7A4M) (EMPIAR-10424) | GABA_A-β3 (EMD-11657) (PDB 7A5V) (EMPIAR-10500) |
|------------|--------------------------------------------------|--------------------------------------------------|
| Map resolution range refined against (Å) | 1.22 - 136.5 | 1.6 - 291.1 |
| Map sharpening B factor (Å²) | 2.3 | 23 |
| Model composition in the asymmetric unit | | |
| Protein residues | 189 | 480 |
| Non-hydrogen atoms | 1668 | 4231 |
| Protein atoms | 1556 | 3850 |
| Waters | 110 | 204 |
| Ions | 2 | 2 |
| Glycans | - | 103 |
| HSM | - | 8 |
| Lipids | - | 64 |
| Isotropic B factors (Å²) | | |
| Protein | 23.3 | 51.2 |
| Waters | 36.2 | 45.7 |
| Ions | 26.6 | 67.2 |
| Glycans | - | 79.1 |
| HSM | - | 28.1 |
| Lipids | - | 110.0 |
| R.m.s. deviations | | |
| Bond lengths (Å) | 0.009 | 0.009 |
| Bond angles (°) | 1.549 | 1.300 |
| Validation | | |
| MolProbity score | 1.10 | 1.10 |
| Clashscore | 2.61 | 2.32 |
| Poor rotamers (%) | 1.18 | 0.48 |
| Ramachandran plot | | |
| Favored (%) | 98.24 | 97.56 |
| Allowed (%) | 1.76 | 2.44 |
| Outliers (%) | 0.00 | 0.00 |
Reporting Summary

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- Confirmed
- The exact sample size (n) for each experimental group/condition, given as a discrete number and unit of measurement
- A statement on whether measurements were taken from distinct samples or whether the same sample was measured repeatedly
- 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.
- A description of all covariates tested
- A description of any assumptions or corrections, such as tests of normality and adjustment for multiple comparisons
- 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)
- For null hypothesis testing, the test statistic (e.g. F, t, r) with confidence intervals, effect sizes, degrees of freedom and P value noted
  - Give P values as exact values whenever suitable.
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g. Cohen's d, Pearson's r), indicating how they were calculated

Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about availability of computer code

- **Data collection**: EPU v2.5 (Thermo Fisher Scientific)
- **Data analysis**: RELION v3.1 (https://github.com/3dem/reliion ), EER reader (https://github.com/fei-company/EerReaderLib), CTFIND-4.1.13, COOT v0.9-pre, REFMAC v5.8.0258, PHENIX version 1.16.3549 and v1.17.1, Pymol v1.8.4, UCSF ChimeraX v1.0, MolProbity, IMOD 4.10, Warp v1.06, in-house generated Linux port of the BoxNet2D neural network in Warp (available from https://gist.github.com/biochem-fan/76ff5934495585da56ab2c0aaf4fe2563)

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 and 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.

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Policy information about availability of data

All manuscripts must include a data availability statement. 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

Atomic coordinates for the GABAAR construct and apo-ferritin have been deposited in the Protein Data Bank with accession codes 7A5V and 7A4M and the cryo-EM density maps have been deposited in the Electron Microscopy Data Bank with accession codes EMD-11657 and EMD-11638, respectively. Raw movies of all datasets have been deposited in the Electron Microscopy Public Image Archive with accession codes EMPIAR-10500 for GABAAR and EMPIAR-10424 for apo-ferritin.
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- Life sciences
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Life sciences study design

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

| Sample size | No statistical method was used to determine sample size. Sample sizes were determined by available electron microscopy time and the number of particles on electron microscopy grids. The sample size is sufficient to obtain a structure at the reported resolution, as assessed by Fourier shell correlation. |
| Data exclusions | Micrographs whose estimated resolution from CTFFIND was worse than 5 Å were removed (rlnCtfMaxResolution). Pre-established common image classification procedures (S.H.W. Scheres, J. Struc. Biol. 180: S19-530, (2012)) were employed to select particle images with the highest resolution content in the cryo-EM reconstruction process. Micrographs with refined scale factors (rlnGroupScaleCorrection) with values below 0.5 were also removed. Details of the number of selected images are given in Supplemental Data Table 1. |
| Replication | Two independent maps were refined for both apoferritin and GABAA in order to estimate resolution according to the recommended procedures in the field (the ‘gold standard’). |
| Randomization | Division of datasets into two random halves was done based on standard approach in RELION 3.1. |
| Blinding | Blinding was not applicable to this study because this type of study does not use group allocation. |

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

| Antibodies | Involved in the study |
| Palaeontology and archaeology | |
| Animals and other organisms | |
| Clinical data | |
| Dual use research of concern | |

### Methods

| Involved in the study |
| ChIP-seq |
| Flow cytometry |
| MRI-based neuroimaging |

### Antibodies

| Antibodies used | High Capacity Streptavidin Agarose Resin was purchased from Thermo Scientific. Rho-1D4 antibody was purchased from the University of British Columbia. The megabody Mb25 is available upon request. |
| Validation | The nanobody Nb25, used to design the megabody Mb25 as described in methods, was characterised and published elsewhere (PMID: 28991263). Mb25 has been characterised and published elsewhere (https://www.biorxiv.org/content/10.1101/812230v1.full.pdf). |

### Eukaryotic cell lines

Policy information about cell lines

| Cell line source(s) | HEK 293S GnTI- cells were obtained from ATCC. |
| Authentication | Further authentication was not performed for this study. |
| Mycoplasma contamination | Mycoplasma testing was not performed for this study. |
No commonly misidentified cell lines were used in this study.