Hydrophobic signal sequences target secretory polypeptides to a protein-conducting channel formed by a heterotrimeric membrane protein complex, the prokaryotic SecY or eukaryotic Sec61 complex. How signal sequences are recognized is poorly understood, particularly because they are diverse in sequence and length. Structures of the inactive channel show that the largest subunit, SecY or Sec61α, consists of two halves that form an hourglass-shaped pore with a constriction in the middle of the membrane and a lateral gate that faces lipid1–10. The cytoplasmic funnel is empty, while the extracellular funnel is filled with a plug domain. In bacteria, the SecY channel associates with the translating ribosome in co-translational translocation, and with the SecA ATPase in post-translational translocation11. How a translocating polypeptide inserts into the channel is uncertain, as cryo-electron microscopy structures of the active channel have a relatively low resolution (~10 Å) or are of insufficient quality6–8. Here we report a crystal structure of the active channel, assembled from SecY complex, the SecA ATPase, and a segment of a secretory protein fused into SecA. The translocating protein segment inserts into the channel as a loop, displacing the plug domain. The hydrophobic core of the signal sequence forms a helix that sits in a groove outside the lateral gate, while the following polypeptide segment intercalates into the gate. The carboxy (C)-terminal section of the polypeptide loop is located in the channel, surrounded by residues of the pore ring. Thus, during translocation, the hydrophobic segments of signal sequences, and probably bilayer-spanning domains of nascent membrane proteins, exit the lateral gate and dock at a specific site that faces the lipid phase.

To determine the structure of an active SecY channel, we initially generated in Escherichia coli a translocation intermediate, consisting of SecA, SecY complex, and a short segment of a secretory protein fused to a fast-folding green fluorescent protein (GFP) (Extended Data Fig. 1a). Although this complex could be purified12, it failed to crystallize. We therefore reduced the complexity of the system by fusing a short segment of a secretory protein directly into SecA. The segment contains the signal sequence of OmpA and a short polypeptide segment. Similar results were obtained with Bacillus subtilis SecA-OAIns and Geobacillus thermodenitrificans SecYE (Extended Data Fig. 2c), a complex of increased thermostability that is functional in E. coli. After optimization (Extended Data Fig. 3a), we used for crystallization.

Figure 1 | Architecture of the active SecY channel. a, A secretory protein segment was inserted into the two-helix finger of the SecA ATPase (SecA-OAIns). The segment contains a linker (grey), the signal sequence of OmpA, consisting of the N-, H-, and C-regions (in red, black, and purple letters, respectively), and a region (in light green) that includes a unique cysteine (yellow star). Residues in the signal sequence are numbered backwards from the cleavage site. The fused segment inserts into the SecY channel in vivo and spontaneously forms a disulfide bridge with a cysteine in the plug. This complex was used for structure determination. b, Ribbon diagram of the complex, viewed from the side. The numbers refer to TMs of SecY. The lines indicate the membrane boundaries. A nanobody was used for crystallization.

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Figure 2 | Conformational changes of the SecY channel. a. Comparison of the lateral gate of the active *G. thermodenitrificans* channel (salmon, except for TM7 in cyan) with the closed SecY channel of *M. jannaschii* (in tan). The largest changes are indicated by arrows. The translocating peptide segment is omitted for clarity. b. As in a, but a comparison with that of the SecA-bound *T. maritima* channel lacking a translocating polypeptide (pink). c. Comparison of the plugs (orange for idle *M. jannaschii* channel and yellow for the active *G. thermodenitrificans* channel). Pore residues are shown as red spheres. d. As in c, but comparison with the inactive *T. maritima* channel (plug in magenta).

Fig. 2d–f, the construct chosen for crystallization contained 49 residues inserted into the two-helix finger of *B. subtilis* SecA, with a cysteine at position +7 in the region following the signal sequence of 20 residues. Channel insertion of the secretory protein segment was similar to that observed with the physiological system, containing wild-type SecA and a GFP fusion to a secretory protein fragment (Extended Data Fig. 1), except that the latter required an additional polypeptide segment to span the SecA molecule. Thus, our simplified system is a faithful mimic of normal initiation of protein translocation. Binding of SecA to the SecY complex seems to be sufficient to cause polypeptide chain insertion into the channel, similar to how ribosome binding allows nascent chain insertion in co-translational translocation. In our system, disulfide crosslinking at the periplasmic side made channel insertion irreversible.

The disulfide-bridged complex of *B. subtilis* SecA-OAIns and *G. thermodenitrificans* SecYE was purified and crystallized in the presence of ADP and BeFx (Extended Data Fig. 3), conditions that lock SecA into a conformation close to its ATP-bound state and maximize the affinity of SecA for the channel. The diffraction of the crystals was improved by the use of single-domain antibody fragments (nanobodies), raised against *G. thermodenitrificans* SecYE and selected for binding to periplasmic loops of the SecA-OAIns/SecYE complex, and by soaking crystals with a Ta₆Br₁₂ metal ion cluster. The structure was determined from multi-wavelength anomalous diffraction (MAD) data obtained with a crystal that diffracted to a resolution of 3.70 Å along one axis and 4.48 Å along the other two (Fig. 1b, Extended Data Fig. 4 and Extended Data Table 1). An initial experimental electron density map had a resolution of ~5.5 Å. This map was improved by density modification and molecular replacement using higher-resolution structures of SecA, SecYE, and the nanobody, followed by cycles of model building and refinement. Inclusion of the model-refined Ta₆Br₁₂ clusters as the resolved heavy atom substructure for recalculation of MAD phases did not further improve the map. Nevertheless, the final map allowed the unambiguous placement of all transmembrane segments (TMs) of SecY and SecE and of many other regions. The translocating polypeptide segment could be built into the map without model bias (Extended Data Fig. 4c). As expected, the nanobody bound to the periplasmic side of SecY, interacting with both the plug and the loop between TM3 and TM4 (Fig. 1b and Extended Data Fig. 5a).

The structure of the active channel shows that SecA undergoes relatively small conformational changes compared with a *Thermotoga maritima* complex lacking a translocating chain (Extended Data Fig. 6). SecA binds to the cytoplasmic loop between TM8 and TM9 and the C-terminal tail of SecY (Extended Data Fig. 5b, c). It probably binds only weakly to the loop between TM6 and TM7, as its tip is disordered. In contrast to the ribosome-6–8, SecA also binds to the amino (N)-terminal half of SecY; that is, the loop between TM2 and TM3 (Extended Data Fig. 5b). Thus, at least in its ATP-bound state, SecA prevents large relative movements of the two halves of SecY.

SecY also undergoes relatively small changes, except at the lateral gate (Fig. 2). Compared with the idle *Methanocaldococcus jannaschii* or *Thermus thermophilus* channels1–3, only TM7 (*M. jannaschii*) or TM7 and TM8 (*T. thermophilus*) significantly shift their positions (Fig. 2a and Extended Data Fig. 7a). Compared with the SecA-bound *T. maritima* channel9, the periplasmic ends of TM3 and TM7 move towards each other, and TM7 tilts by 10° relative to the plane of the membrane (Fig. 2b), changes that generate a pocket for the signal sequence (see below). In both structures, the lateral gate is partly open (compare Fig. 2a and Fig. 2b).

In the active *G. thermodenitrificans* SecY channel, the plug consists of two β-strands and therefore differs from the α-helical structures observed in other species1,4–9 (Extended Data Fig. 8a). Such variability is consistent with the fact that the amino-acid sequence of the plug region is least conserved1, and that plug deletions cause neighbouring polypeptide regions to form new plug domains15. Different plug structures can probably be tolerated, as long as they fill the extracellular cavity of the channel, so that the closed state of the channel is stabilized and small molecules cannot pass through it.
However, it is possible that the plug has different conformations in the closed and active channels.

Whereas the plug is close to the central constriction in the closed *M. jannaschii* channel, in the active *G. thermodenitrificans* channel it moves to the periplasmic side and towards the back of the channel, away from the lateral gate (Fig. 2c). The plug comes close to the TM of SecE, consistent with disulfide crosslinking experiments. In a SecA–SecY structure lacking a translocation substrate, the plug moves a smaller distance and towards the front (Fig. 2d), partly sealing the opened lateral gate (Extended Data Fig. 8b). In an intact membrane, this would prevent surrounding lipid molecules from moving through the lateral gate into the extracellular cavity. The plug is probably flexible in the active channel, but in our crystal structure it is confined both by the disulfide bond to the translocating chain and by the interaction with the nanobody (Extended Data Fig. 5a). Indeed, in a 6.5 Å-resolution structure determined without nanobody, the plug is shifted further towards the back, although there are otherwise only small differences (Extended Data Fig. 9a).

The signal sequence of the secretory protein segment forms a helix that is tilted ~45° relative to the plane of the membrane (Fig. 3a). The positively charged N terminus of the signal sequence (N-region) is in the same plane as hydrophilic residues of SecY. In an intact membrane, the N-region could interact with the negatively charged head groups of the phospholipid bilayer. This interaction may retain the N terminus on the cytoplasmic side of the membrane while the C-terminal end of the signal sequence moves through the channel, resulting in loop insertion of the translocating polypeptide.

The hydrophobic core of the signal sequence (H-region; residues –17 to –6 of the original OmpA sequence; Fig. 1a) sits in a groove outside the lateral gate of SecY and forms a helix that runs almost parallel to TM2 (Fig. 3a, b). Some residues make van der Waals contacts with hydrophobic amino acids in TM2 (Extended Data Fig. 7c), but most...
Figure 4 | Scheme of SecA-mediated protein translocation. Stage a corresponds to the closed channel\(^1\)–\(^3\), stage b to the structure of the inactive SecA–SecY complex\(^9\), stage c represents an intermediate to stage d, which corresponds to the structure of the active channel. Stage e is attained after signal sequence cleavage. The translocating polypeptide is shown in green and the signal sequence as a green cylinder. The lateral gate of the channel is shown as a broken line on the left. The clamp of SecA is indicated.

The C-terminal region of the signal sequence (C-region; residues −5 to −1; Fig. 1a) replaces the periplasmic end of TM7 in the closed \(M.\ jannaschii\) channel (Extended Data Fig. 7b). This segment forms a distorted, amphipathic α-helix that is intercalated between TM7 and TM3 into the periplasmic side of the lateral gate (Fig. 3a, c). In our model, the side chains of Thr(−5) and Ser(−2) point into the periplasmic cavity previously occupied by the plug in the closed channel (Extended Data Fig. 7d). The hydrophobic residues Val(−4) and Ala(−3) face detergent/lipid. The lateral gate is thus sealed by the C-region from surrounding lipid molecules, which could otherwise pass through a large opening generated by the displacement of the plug from the front (Fig. 3c). After signal sequence cleavage, the periplasmic parts of TM7 and TM3 probably move towards each other and seal the lateral gate.

The hydrophilic polypeptide segment following the signal sequence adopts a partly extended conformation with a loop in the periplasmic cavity, centred on the cytosine used for crosslinking to the plug. The signal sequence cleavage site is located inside the channel (Fig. 3c), probably inaccessible to the periplasmically disposed active site of signal peptidase\(^18\). Thus, at some point during translocation, the C-region of the signal sequence probably must adopt a more extended conformation, a change also suggested by experiments with synthetic signal peptides\(^19\),\(^20\).

The polypeptide chain inside the channel is perpendicular to the plane of the membrane (Fig. 3a). The two strands of the hairpin formed by the translocating polypeptide do not interact with one another, so that during translocation the C-terminal part of the hairpin could move unimpeded through the centre of the channel. Our model places residue Gly(+19), or one of the neighbouring residues (Ala18 or Gly20), of the translocating polypeptide inside the pore ring (Fig. 3d). The density around Gly19 is particularly strong, indicating that this segment is confined by the surrounding four pore ring residues (Ile78, Ile183, Ile275, Ile404). The ring is wider than in the idle \(M.\ jannaschii\) or SecA-bound \(T.\ maritima\) channels (diameters 8.8 Å versus 5.6 Å or 6.6 Å respectively). Crystallization may have favoured the presence of small amino acids in the pore, minimizing its expansion by the presence of a translocating chain. However, even a small increase in pore diameter would allow the passage of amino acids with larger side chains. The pore ring residues fit snugly around the translocating polypeptide (Fig. 3e), confirming that they form a ‘gasket’ that maintains the permeability barrier for ions and other small molecules during translocation\(^21\). Consistent with disulfide crosslinking experiments\(^22\), only pore ring residues contact the translocating chain (Fig. 3d).

Thus, the hourglass-shape of the channel minimizes interactions with the translocation substrate, facilitating its movement through the channel.

The pore ring of the idle \(M.\ jannaschii\) channel contains two additional residues (Ile75 and Ile170; Fig. 3f). In the active channel, the corresponding residues (Val74, Leu179) are displaced (Fig. 3e). The pore ‘ring’ is thus open at the lateral gate between Ile78 in TM2 and Ile275 in TM7 (Fig. 3e). These features suggest that a translocating polypeptide segment continuously encounters the hydrocarbon chains of surrounding lipids; when sufficiently hydrophobic, the segment will partition into the lipid phase and become a TM domain of a membrane protein\(^23\),\(^24\).

Our crystal structure probably reflects the physiological situation of a translocating polypeptide. Five of the seven residues of the N-terminal linker are invisible and thus probably flexible, allowing unrestricted interaction of the signal sequence with the channel. In addition, most polypeptide segments following the signal sequence are in a relaxed conformation, unconstrained by fusion to SecA, or the disulfide bridge to the plug. The disulfide bridge helps to stabilize the signal sequence in the channel, but it probably does not lead to gross distortions, because the plug is mobile and the disulfide bridge is formed spontaneously in vivo.

The crystal structure of the active channel leads to a refined model for post-translational protein translocation in bacteria (Fig. 4). The SecY channel is initially in the idle state, with the plug in the centre and the lateral gate closed (Fig. 4a). Binding of SecA primes the channel for the arrival of a secretory protein precursor: the lateral gate is partly opened, the pore ring widened, and the plug domain moved towards the front (Fig. 4b). Next, the secretory protein inserts into the channel as a loop, with the C-terminal section of the polypeptide hairpin in the pore proper, surrounded by pore ring residues (Fig. 4c, d). During subsequent cycles of ATP hydrolysis, SecA uses a ‘push-and-slide’ mechanism to move the C-terminal part of the polypeptide loop through the pore\(^14\). Eventually, the signal sequence is cleaved by signal peptidase (Fig. 4e).

During loop insertion, the H-region of the signal sequence moves through the cytoplasmic part of the lateral gate and ends up in a hydrophobic groove on the outside, while the following hydrophilic segment crosses the lateral opening of the pore ring. A signal sequence might move through a partly open gate in an extended conformation, or it could move through a widened gate as a preformed helix. The latter possibility is suggested by a \(≈10\) Å-resolution electron microscopy (EM) structure of a ribosome/nascent chain/channel complex, in which a signal sequence helix was seen inside the lateral gate\(^6\), probably prevented from exit by a disulfide bridge between a cysteine at the end of the signal sequence and a cysteine in the plug. The groove on
the outside of the lateral gate appears to be a general binding site for hydrophobic sequences, as indicated by a 8.5 Å resolution structure of a complex in which the OmpA signal sequence was replaced by that of DsbA (Extended Data Fig. 9b, c), and EM structures of ribosome–channel complexes, in which TMs of nascent membrane proteins are located at about the same position2,25 (Extended Data Fig. 7c). Like TMs, signal sequences appear to be recognized mostly by lipid partitioning, consistent with their ability to be crosslinked to lipids26,27, and the correlation between the partitioning of synthetic peptides into hydrophobic solvents and their function as signal sequences in vivo28. Nevertheless, amino-acid interactions with TM2 of the channel may contribute to the recognition of signal sequences. This may explain why some amino acids occur more frequently than others in signal sequences, even when they have about the same hydrophobicity: leucine is preferred over isoleucine and valine29, perhaps because its extended side chain can make tighter van der Waals contacts with residues in TM2. Whereas signal sequences of translocating secretory proteins would tend to stay in the binding pocket until they are cleaved off, the more hydrophobic TMs of membrane proteins could move away once the connecting loop to the polypeptide segment inside the channel pore attains adequate length.

While this paper was under review, a cryo-EM structure was published describing an active ribosome-bound mammalian Sec61 channel containing a short secretory polypeptide segment30. The authors conclude that the signal sequence helix replaces TM2 of Sec61α in the idle channel, implying that the signal sequence is intercalated into the lateral gate and raising the possibility that the conformational changes differ greatly from those in our system. However, a superposition of the two active channels based on secondary structure matching shows that they are actually very similar, with only moderate differences at the periplasmic/luminal ends of some TMs (Extended Data Fig. 7f). Importantly, in both cases the signal sequence helix docks to the same site outside the lateral gate and runs parallel to TM2 (Extended Data Fig. 7f). Thus, regardless of the organism and mode of translocation, lipid partitioning appears to be the major mechanism by which signal sequences are recognized.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

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Author Contributions E.P. designed the SecA–substrate fusion constructs, performed biochemical tests, and obtained the initial crystals; L.L. optimized and determined the crystal structures of the SecY complex; J.I. and H.P. generated and cloned nanobodies; T.A.R., L.L., and E.P. interpreted the structure and wrote the manuscript. T.A.R. supervised the project.

Author Information The crystal structure determined in this study has been deposited in the Protein Data Bank (PDB) under accession number SEUL. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to L.L. (long.i@hms.harvard.edu) or T.A.R. (tomp.rapoport@hms.harvard.edu).
Letter

Protein translocation and disulfide crosslinking assays. To test in vivo OmpA–GFP translocation by E. coli SecA and E. coli SecY complex (three-component system), E. coli strain EP52 or EP62 (∆rnm ΔompT secY-CBP) was transformed with pACYC-secY expressing E. coli SecY complex with SecY containing a unique Cys at position 68 (ref. 21). The cells also expressed OmpA–GFP from pBAD-OmpA–GFP under an arabinose-inducible promoter21. In all constructs, including those used in the two-component system, position 1 of the signal sequence was mutated to Tyr to prevent signal peptide cleavage21. The cells were grown at 37 °C to log phase in lysogeny broth (LB) medium supplemented with 100 μg ml−1 ampicillin and 40 μg ml−1 chloramphenicol. OmpA–GFP expression was induced by addition of 0.1% t-arabinose for 1 h. Where indicated, 0.3 mM copper(II) (1.10-phenanthroline (CuPPh) was added to the bacterial culture for 15 min at room temperature (22 °C). The culture was treated with 10 mM N-ethyl maleimide (NEM) for 30 min on ice to block free cysteines. The cells were lysed in SDS sample buffer. Samples (equivalent amounts based on an absorbance at 600 nm, A600 of E. coli cultures) were subjected to non-reducing SDS–PAGE and analysed by western blotting with anti-SecY- or GFP-antibodies. Where indicated, the samples were treated with 2% β-mercaptoethanol (3-ME) at 50 °C for 20 min before SDS–PAGE. To test the translocation of E. coli SecA-OAIns fusion with E. coli SecY complex (two-component system), EP52 cells were transformed with pACYC-SecY and pBAD-EsEc-SecA-OAIns. The cells were grown to log phase in LB medium supplemented with 100 μg ml−1 ampicillin and 40 μg ml−1 chloramphenicol, and SecA-OAIns expression was induced by addition of 0.1% t-arabinose for 4 h at room temperature or 30 min at 37 °C. Disulfide crosslinking and non-reducing SDS–PAGE analysis were performed as described above.

Protein expression and purification. E. coli strain EP51 (∆rnm ΔompT) was transformed with pTet-G. thermodenitrificans SecEHis8/Y and pBAD-OmpA–GFP under an arabinose-inducible promoter21. The cells also were transformed with pBAD-OmpA–GFP/B. subtilis SecA, a modified version of pBAD-OmpA, which contains an additional ribosome binding site and the B. subtilis secA gene (for bicistronic expression) following OmpA–GFP. In the case of the two-component system, EP51 cells were transformed with pTet-G. thermodenitrificans SecEHis8/Y and pBAD-B. subtilis SecA-OAIns. The cells were grown to log phase at 37 °C, and the expression of G. thermodenitrificans SecYE was induced by addition of 200 ng ml−1 α-hydroxytetracycline for 1.5–2 h at 37 °C. Then, the expression of OmpA–GFP, B. subtilis SecA or B. subtilis SecA-OAIns was induced by addition of 0.2% t-arabinose for 1–3 h. The cells were collected and subjected to SDS–PAGE followed by western blotting as described above. G. thermodenitrificans SecY was detected using anti-His antibodies.

Nanobody library generation. Purified G. thermodenitrificans SecYE in n-dodecyl β-d-maltoside (DDM)-containing buffer was injected into an alpaca to elicit an immune response. A male alpaca (Vicugna pacos) was purchased locally, maintained in pasture, and immunized following a protocol authorized by the Tufts University Cummings Veterinary School Institutional Animal Care and Use Committee. After five rounds of immunization, total RNA was isolated from 1 L of culture. After suspension in 15 ml of TES buffer (200 mM Tris-HCl pH 7.5, 150 mM NaCl, 10% glycerol, 0.02% DDM) for SecYE, and with 20 mM HEPES-KOH pH 7.0, 150 mM NaCl, 10% glycerol, 0.02% DDM, 5 mM MgCl2, 0.1 mM ADP/BeFx for SecA-OAIns/SecYE. For each protein, 95 clones were sequenced, and sequences that appeared more than 5 times were selected for subsequent validation.

Nanobody screening. Thirteen distinct families of nanobodies directed against G. thermodenitrificans SecYE were identified by DNA sequencing. Twenty-two nanobody clones were sub-cloned into the pHEΝ6 vector23, which adds an N-terminal pelB sequence and C-terminal sortase and His tags (LPETGGS-His). From these cultures, small nanobodies (100 μg/ml Tris-HCl buffer) were expressed in microfluidic M13 culture. After Ni-resin purification, the labelled nanobodies were labelled with Alexa 555 (Invitrogen) by sortase reaction24, and were captured on B. subtilis SecA-OAIns/SecY complex or with SecYE alone at a molar ratio of 2:1. Nanobody binding was monitored by a shift of the peak in size-exclusion chromatography coupled with a fluorescence detector (Shimadzu). All 22 nanobodies bound to SecYE, but only nanobody AYC08 had a high affinity for SecA-OAIns/SecYE. The binding of AYC08 to free SecYE was weaker than to SecA-OAIns/SecYE, indicating that AYC08 interacts with the periplasmic side of SecY and that the binding epitope is only fully exposed in the active channel.
of monomolecular antibodies generated against SecY. However, crystals with improved diffraction were only obtained when the complex was co-crystallized with nanobody AYC08.

The complex of *G. thermodenitrificans* SecYE and *B. subtilis* SecA-OAlls was mixed with nanobody AYC08 at a molar ratio of 1:1.2 with addition of 1 mg ml\(^{-1}\) lipids (42 mg ml\(^{-1}\) 1,2-dipalmitoyl-sn-glycero-3-phosphoglycerol (DPPG) plus 1.2-dipalmitoyl-sn-glycero-3-phosphoethanolamine (DPPE) (3:1) suspension in 0.5% DDM\(^{47}\)). The mixture was incubated at 4°C overnight and clarified by ultra-centrifugation before setting up crystallization trays. The initial crystal screening yielded several crystal forms. Three of them were readily reproducible, but all diffractioned to a minimum of ~6 Å resolution. Heavy atom complexes were screened for crystal soaking and the Ta\(_6\)Br\(_{12}\) cluster improved the resolution limit of one crystal form. The best crystals were obtained with the hanging drop method, mixing 0.5 μl of the protein solution and 0.5 μl of well solution (21–24% polyethylene glycol (PEG) 1500, 100 mM Tris-HCl pH 8.5, 50–100 mM MgAc\(_2\), 2% 2-methyl-2,4-pentandiol (MPD) and 24-well VDX plates with 500 μl well solution. The crystals were grown at 22°C over a week. The Ta\(_6\)Br\(_{12}\) powder (Jena Bioscience) was suspended in buffer at a concentration of 20 mM and added to the crystallization drops at a final concentration of ~2 mM. After overnight incubation, the crystals were flash-frozen in liquid nitrogen.

**X-ray data collection and structure determination.** Hundreds of crystals were screened at NE-CAT and GM/CA-CAT of the Advanced Photon Source (Argonne National Laboratory). The diffraction of the crystals decayed rapidly and was weak, caused by strong heavy atom absorption. However, a complete set of three-wavelength MAD data from a single crystal could be collected at GM/CA-CAT. The data were processed with the XDS package\(^{48}\) and analysed by the program Aimless\(^{49}\). The crystal belongs to the P6\(_1\)2\(_2\)2 space group. The diffraction was anisotropic. Along axis \(c\), the diffraction went beyond 3.70 Å (\(f/\sigma = 2.0\)), whereas along the axes \(a\) and \(b\), the diffraction went only to limited 4.48 Å (\(f/\sigma = 1.5\)). The data were initially processed to 3.9 Å. Anisotropic correction was applied in the different programs used in the following calculations. Molecular replacement was used to locate SecA, SecYE, and the nanobody, employing as search models *B. subtilis* SecA (PDB accession number 1TF5), *T. maritima* SecYE (PDB accession number 3DIN), and an anti-DHFR nanobody (PDB accession number 4EIG), respectively. The crystal contained one complex per asymmetric unit with a solvent content of 69%. Heavy atom sites were identified with the molecular replacement results by MR-SAD in Phaser\(^{46}\) and refined without molecular replacement models in Sharp (Global Phasing). The experimental map based on the positions of the heavy metal ions clustered contained useful phase information to 5.5 Å (judged upon deletion of any of the TMs of SecY or of the signal sequence helix. A similar resolution\(^{48}\) was used are PDB 1RH5, 3MP7, 2ZJS, 3DIN, SAWW, JFC2, 4CG6. The new structure described in this paper has accession number 5EUL.

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Extended Data Figure 1 | Generation of translocation intermediates with a three-component system. a, Strategy to generate SecA-dependent translocation intermediates in E. coli cells. The intermediates are assembled from E. coli SecA, E. coli SecY complex, and substrate containing an N-terminal OmpA signal sequence and C-terminal superfolder GFP51 (sfGFP). After loop insertion into the SecY channel, translocation of the C terminus is stalled by the folded sfGFP. Insertion is monitored by disulfide bond formation between a pair of cysteines introduced into the substrate and the plug of SecY (yellow stars).

b, Scheme showing the simplified system, in which a secretory protein segment is fused into the two-helix finger of SecA.

c, Sequence of the substrate used in a. The −1 position of the original signal sequence was changed to Tyr to prevent signal sequence cleavage. The position of the cysteine and the length of the translocated segment were varied (here shown for Cys at position +3 and 58 amino acids in length).

d, Variation of the Cys position with a translocating segment of 58 residues. Where indicated, disulfide crosslinks to SecY with a Cys at position 68 (OmpA–GFP×SecY) were induced by the oxidant copper phenanthroline (CuPh) before harvesting the cells. The samples were analysed by non-reducing SDS–PAGE, followed by western blotting (WB) with anti-SecY and anti-GFP antibodies. e, As in d, but in the absence of oxidant, with Cys at different positions and variation of the length of the translocated segment. Asterisks indicate non-specific bands.

f, As in e, but with E. coli cells expressing B. subtilis (B.s.) SecA and G. thermodenitrificans (G.t.) SecYE. The substrate contained a Cys at position +7, and SecY a Cys at position 60. The red arrows indicate spontaneously generated disulfide crosslinks (GFP sometimes does not unfold in SDS, resulting in two bands). The OmpA–GFP constructs contained a C-terminal Strep-tag that was detected by StrepTactin conjugated with horseradish peroxidase (HRP).
Extended Data Figure 2 | See next page for caption.
Extended Data Figure 2 | Generation of translocation complexes with SecA–substrate fusion constructs. 

a, Translocation complexes were generated as indicated in the scheme in Extended Data Fig. 1b. An E. coli SecA–substrate fusion (SecA-OAIns74 (L12)) was overexpressed together with E. coli SecY complex in E. coli cells. SecA-OAIns74 (L12) contains 74 amino acids inserted into the two-helix finger of SecA, including a linker of 12 residues, and a GFP tag following SecA. Translocation of the substrate segment was monitored by spontaneous disulfide crosslinking between a cysteine at position +7 (with respect to the original signal peptidase cleavage site) and a cysteine at position 68 in the plug of SecY. Where indicated, β-mercaptoethanol (β-ME) was added to reduce the disulfide bond. The samples were analysed by non-reducing SDS–PAGE and western blotting with anti-SecY antibodies. The overexpression of SecA-OAIns was monitored by the fluorescence of GFP (data not shown) and staining with Coomassie blue (CBB, lower panel).

b, As in a, but with E. coli SecA-OAIns constructs containing from 6 to 12 residues in the linker (L6–L12) or mutations (3Q) in the H-region of the signal sequence. Expression of SecA-OAIns was verified by the strong green fluorescence of cell lysates, caused by GFP fused to the C terminus of SecA (not shown). The lower panel shows the sequences of the SecA-inserted segments. c, As in a, but with B. subtilis SecA-OAIns74 (L12) and G. thermodenitrificans SecYE. SecY and SecA were detected by western blotting with anti-His antibodies and Ponceau staining, respectively. d, As in c, but with B. subtilis SecA-OAIns containing different inserted segments. SecA-OAIns was expressed under different conditions, as indicated. Expression of SecA-OAIns was verified by green fluorescence of GFP fused to the C terminus of SecA (not shown) and Ponceau staining (second panel). The sequences of the constructs are shown in the lowest panel. e, As in d, but with different constructs, the sequences of which are shown in the lower panel. Where indicated, copper phenanthroline (CuPh) was added to the cells to induce disulfide bridge formation. SecA-OAIns49(L7) was used for crystallization. f, As in e, but with a Cys at position −1 (the last residue of the OmpA signal sequence) instead of position +7. Note that in this case disulfide formation does not occur spontaneously.
Extended Data Figure 3 | Purification of an active translocation complex. 

**a.** Scheme of the purification protocol. 

- Co-expression of *G. thermodenitrificans* SecY/E_{E392} and *B. subtilis* SecA-OAIns-GFP in *E. coli* cells
- Lysis of *E. coli* cells
- Enrichment of membranes (by ultracentrifugation) and detergent solubilization
- Immobilized metal (Co²⁺) affinity chromatography (IMAC)
- Treatment with HRV 3C protease to cleave off the GFP tag from SecA-OAIns
- Anion exchange chromatography to remove free SecY/E_{E392} and GFP (MonoQ column)
- Size-exclusion chromatography (Superdex200 column)

**b.** Elution of the *G. thermodenitrificans* SecYE/B. subtilis SecA-OAIns complex from a Superdex200 column during the last chromatography step.

**c.** Non-reducing SDS–PAGE analysis of samples taken during the purification procedure and of fractions indicated with red numbers in **b.** Lane 1, molecular mass markers. Lane 2, sample analysed after immobilized metal ion affinity chromatography. Lane 3, sample after cleavage of the GFP tag. Lane 4, sample after anion exchange chromatography (MonoQ).
Extended Data Figure 4 | Electron density map and refined model.

a, Stereo view of the unsharpened density map (2Fo – Fc; 1σ) of the entire complex. Heavy metal ion clusters are shown in yellow. b, As in a, but with the density map derived from MAD phasing after density modification.

c, SigmaA-weighted phase-combined 2Fo – Fc density maps of the translocating peptide region. Left: omit map calculated without a model for the translocating peptide. Right: map calculated with the model. Phe (–7) is one of the residues used to determine the registry of the helix.

d, A side view of the density for TM3 and TM4. e, Density showing the disulfide crosslink between the plug and translocating chain. f, Top view of Gly19 of the translocating chain surrounded by pore residues.
Extended Data Figure 5 | Interactions of the nanobody and SecA with SecY.  

**a.** The nanobody binds to the plug and to the loop between TM3 and TM4 (L3/4).  

**b.** The polypeptide crosslinking domain (PPXD; in yellow) of SecA interacts with the loop between TM8 and TM9 of SecY (L8/9; in red), and the long helix of the helical scaffold domain (HSD; in blue) with the loop between TM2 and TM3 (L2/3; in red). The loop between TM6 and TM7 of SecY (L6/7; in red) does not seem to make strong contact with SecA.  

**c.** Two helices of the HSD interact with the C-terminal tail of SecY (C-tail; in red).
Extended Data Figure 6 | Comparison of the conformations of SecA in the active *G. thermodenitrificans* and inactive *T. maritima* complexes. The domains of SecA in the *G. thermodenitrificans* complex are labelled with different colours (nucleotide binding domain 1 (NBD1), blue; nucleotide binding domain 2 (NBD2), cyan; helical scaffold domain (HSD), brown; helical wing domain (HWD), grey; polypeptide crosslinking domain (PPXD), yellow). SecA in the *T. maritima* complex is shown in pink. Left: a top view (the channel would be underneath); right: a side view with the two-helix finger (part of helical scaffold domain) indicated.
Extended Data Figure 7 | Localization of signal sequences in the *G. thermodenitrificans* SecY and mammalian Sec61 channels and of a TM domain in the mammalian Sec61 channel. **a**, In the active channel (salmon), the signal sequence displaces TM7 and TM8 in the idle *T. thermophilus* channel (cyan). **b**, As in **a**, but comparison with the idle *M. jannaschii* channel (tan). The C-region of the signal sequence takes the position of TM7. **c**, Side view of the interactions of the H-region of the signal sequence with TM2 of *G. thermodenitrificans* SecY. Interacting amino acids are indicated. **d**, Stereo view showing the intercalation of the C-region into the periplasmic side of the lateral gate. Residues of the amphipathic helix are indicated. **e**, The active *G. thermodenitrificans* channel (in salmon) was aligned with a mammalian channel (in grey) containing a nascent membrane protein (PDB accession number 4CG6) using secondary structure matching. The signal sequence in the bacterial channel is shown in green, and the TM segment of the nascent membrane protein in yellow. **f**, As in **e**, but comparison of the active *G. thermodenitrificans* channel with a mammalian Sec61 channel (light blue) containing a secretory protein fragment (PDB accession number 3JC2). The signal sequences are shown in green and yellow, respectively.
Extended Data Figure 8 | The plug domains in SecY channels. a, The plugs in channels of different organisms have different structures. Shown are side views with the plugs in yellow and pore residues as red spheres. PDB accession numbers are given below the names of the organisms. For the G. thermodenitrificans channel, the translocating peptide segment was omitted. b, In the inactive T. maritima SecY channel, the plug (in yellow) is at the front of the channel, partly sealing the periplasmic side of the lateral gate. Shown is a side view in a surface representation, with hydrophilic and hydrophobic residues in blue and orange, respectively.
Extended Data Figure 9 | Structures of the active *G. thermodenitrificans* complex determined without nanobody or with a different signal sequence. a, Stereo view of density maps at 6.5 Å resolution for the active complex in the absence of nanobody. Shown is a 2Fo − Fc density map at 1σ (blue mesh) and a difference map (Fo − Fc) at 3σ (green mesh), both calculated by molecular replacement with a model lacking the plug. Strong positive density is seen close to SecE, probably corresponding to parts of the plug. The arrow indicates the movement of the plug from the position in the structure with nanobody to the density seen in the structure without nanobody. b, Stereo views of density maps at ~8.5 Å resolution for the active *G. thermodenitrificans* complex in which the OmpA signal sequence was replaced by that of DsbA. Shown is a side view of the 2Fo − Fc density map at 1σ (blue mesh) and a difference map (Fo − Fc) at 3σ (green mesh), both calculated by molecular replacement with a model lacking the signal sequence. Note that the model for the OmpA signal sequence fits well into the density corresponding to the DsbA signal sequence. c, As in b, but top view and not in stereo.
Extended Data Table 1 | Data collection and refinement statistics for MAD structures

| Data collection | SecA-OAIns/SecYE + Ta₂Br₄₂⁻ |
|-----------------|--------------------------------|
| Space group     | GM/CAT-CAT @APS               |
| Cell dimensions |                                |
| α, β, γ (°)     | 127.798, 127.798, 554.772     |
|                 | 90, 90, 120                   |

|                      | Remote  | Inflection | Peak    |
|----------------------|---------|------------|---------|
| Wavelength (Å)       | 1.2782  | 1.2552     | 1.2548  |
| Resolution (Å)       | 3.70    | 4.00       | 4.50    |
|                      | (3.80-3.70) | (4.10-4.00) | (4.62-4.50) |
| R_sym or R_merge     | 0.084 (>-1) | 0.095 (>1) | 0.128 (>1) |
| I/σI                | 9.30 (0.4) | 9.69 (0.9) | 11.90 (1.0) |
| Completeness (%)     | 99 (100) | 99 (100)   | 99 (100) |
| CC1/2                | 0.999 (0.57) | 0.998 (0.80) | 0.999 (0.92) |
| Redundancy           | 10.6 (10.2) | 10.6 (10.4) | 21.3 (21.6) |

**Refinement**

|                      |         |
|----------------------|---------|
| Resolution (Å)       | 3.7     |
| No. reflections      | 53812   |
| Rwork/Rfree          | 0.306/0.325 |
| No. atoms            | 9888    |
| Protein              | 9532    |
| Ligand/ion           | 356     |
| Water                | N/A     |
| B-factors            | 194.00  |
| Protein              | 190.24  |
| Ligand/ion           | 300.34  |
| Water                | N/A     |
| R.m.s. deviations    |         |
| Bond lengths (Å)     | 0.007   |
| Bond angles (°)      | 1.15    |
| Ramachandran         |         |
| Favored / allowed /  | 91.3 / 6.8 / |
| outliers (%)         | 1.9     |

*Highest resolution shell is shown in parenthesis.*