Molecular mechanism for the regulation of yeast separase by securin
Shukun Luo1 & Liang Tong1

Separase is a cysteine protease with a crucial role in the dissolution of cohesion among sister chromatids during chromosome segregation1–7. In human tumours separase is overexpressed, making it a potential target for drug discovery8. The protease activity of separase is strictly regulated by the inhibitor securin, which forms a tight complex with separase and may also stabilize this enzyme9–16. Separases are large, 140–250-kilodalton enzymes, with an amino-terminal α-helical region and a carboxy-terminal caspase-like catalytic domain. Although crystal structures of the C-terminal two domains of separase17 and low-resolution electron microscopy reconstructions of the separase–securn complex18,19 have been reported, the atomic structures of full-length separase and especially the complex with securin are unknown. Here we report crystal structures at up to 2.6 Å resolution of the yeast Saccharomyces cerevisiae separase–securn complex. The α-helical region of separase (also known as Esp1) contains four domains (I–IV), and a substrate-binding domain immediately precedes the catalytic domain and has tight associations with it. The separase–securn complex assumes a highly elongated structure. Residues 258–373 of securin (Pds1), named the separase interaction segment, are primarily in an extended conformation and traverse the entire length of separase, interacting with all of its domains. Most importantly, residues 258–269 of securin are located in the separase active site, illuminating the mechanism of inhibition. Biochemical studies confirm the structural observations and indicate that contacts outside the separase active site are crucial for stabilizing the complex, thereby defining an important function for the helical region of separase.

Saccharomyces cerevisiae separase contains 1,630 amino acid residues and the N-terminal α-helical region covers approximately 1,160 residues. We obtained good quality crystals for the complex of S. cerevisiae separase (residues 51–1630; Fig. 1a, Extended Data Figs 1–3) and securin (residues 258–373; Extended Data Fig. 4). The N-terminal segment of securin contains the KEN-box and D-box motifs, which are important for the destruction of securin but not for the interaction with separase20. The structure was determined and refined at 3.0 Å resolution (Extended Data Table 1). Under the same crystallization condition, we observed another crystal form and collected a dataset to 3.7 Å resolution. We readily solved the structure of this crystal form and did not observe any notable structural differences.

We produced crystals of another complex of S. cerevisiae separase (residues 71–1630) and securin (residues 258–373), and determined its structure at 2.6 Å resolution. The overall structure of this complex is essentially the same as that at 3.0 Å resolution (Extended Data Fig. 5). One difference is that residues 75–79 at the N-terminal end of separase form an anti-parallel β-sheet with the equivalent residues of another separase molecule in the crystal (Extended Data Fig. 5), which is an artefact owing to the longer truncation of separase in this complex.

The overall structure of the separase–securn complex has a highly elongated shape, with dimensions of $65 \times 70 \times 165$ Å (Fig. 1b, c). The catalytic domain (CD) is located at one end of the complex, far away from the N terminus of separase at the other end. This elongated shape is generally consistent with that observed for the human separase–securn complex by electron microscopy18, but distinct from a closed form reported for the Caenorhabditis elegans complex19.

The structure shows that the N-terminal helical region (residues 53–1163) of separase can be divided into four domains (I–IV) (Fig. 1a–c). Their structures are generally like HEAT repeats, with extensions that connect each domain to the next one. Domain I is folded back onto domain II such that the loops at one end of each domain are facing each other (Fig. 2a), with a buried surface area of 1,100 Å² for each domain, suggesting that this may be a stable association between them. Two consecutive helices (the fourth and fifth helices) in domain I are parallel to each other (Fig. 2a, Extended Data Fig. 6). The first 50 residues of separase are poorly conserved (Extended Data Fig. 1). Earlier studies showed that the first 155 residues of separase are important for function12. These residues include helices α1 to α5 of domain I. Their deletion would disrupt the structure of this domain and its interaction with domain II (Fig. 2a), and helix α3 is in the binding site for the C-terminal helix of securin (see below), thereby explaining their functional importance. Human separase is stimulated by DNA of greater than 100 base pairs21, and this stimulation is also likely to be mediated by the helical region.

The helical hairpins of domain III are arranged in a tight, right-handed super-helix (Fig. 2b). This domain is located in the middle of the structure and has contacts with domain II at one end (buried surface area of approximately 900 Å²) and domain IV at the other (1,100 Å²) (Fig. 1b). It also has a direct contact with a segment immediately following the catalytic Cys1531 residue in the CD (strands β4A, β4B and the connecting loop; Fig. 2c, Extended Data Figs 3, 6). The overall arrangement of the helices in this domain has similarity to that in several other structures based on a DaliLite search22, and the closest homologues include the TPR domain of the G-protein signalling modulator LGN23–25 and others (Extended Data Fig. 7).

Domain IV has extensive interactions with the domain immediately before the CD (substrate-binding domain (SD), Fig. 1a; see next paragraph), covering one face of its β-sheet (buried surface area of approximately 1,900 Å² for each domain) and providing one extra strand to this β-sheet (Fig. 2c). It also interacts with two of the surface helices of the CD. Therefore, domain IV could be considered as forming a module together with the SD–CD.

An $α + β$ domain is located just before the CD in separases18,26. The backbone fold of this domain has no similarity to proteases such as caspases17. The domain is tightly associated with the CD and contributes to substrate binding. We will refer to it as the substrate-binding domain (SD) (Fig. 1a). Other names for the SD and CD include PPD (pseudo-protease domain) and APD (active protease domain), respectively17,27. The SD contains a five-stranded $(β1–β5)$ mostly anti-parallel β-sheet, and there is a large insertion (residues 1200–1380) between the third and fourth strands, including a hairpin of two long helices ($αA$–$αB$) and a four-helical bundle ($αC$–$αF$) (Fig. 2c). The backbone fold of this β-sheet is RNase H-like,

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with close homologues including part of the catalytic PIWI domain of Argonaute27 and others (Extended Data Fig. 7).

The overall structure of the SD–CD of *S. cerevisiae* separase is similar to that of the *Chaetomium thermophilum* separase17 (Fig. 2c), with a root mean squared (r.m.s.) distance of 1.5 Å for their equivalent Ca atoms and amino acid sequence identity of 39%, suggesting that the binding of securin did not cause a large overall conformational change in these two domains. The β-sheets of the two domains are juxtaposed in the structure, although the two neighbouring strands at the interface (β5 of SD and β6 of CD) are at an angle of nearly 90°. The αA–αB hairpin from SD covers a portion of the active site in CD and helps to mediate substrate binding.

Securin is primarily in an extended conformation, with only two short helices, and runs along the entire length of separase (Figs. 1b, 3a). We will refer to this segment of securin, residues 258–373, as the separase interaction segment (SIS). The backbone direction of SIS is anti-parallel to that of separase, such that the N-terminal region of SIS interacts with the C-terminal region of separase. The securin SIS has contacts with every domain of separase, being positioned in prominent grooves in its surface (Fig. 3a). Approximately 4,600 Å² of the surface area of securin is buried in the interface with separase (3,900 Å² surface area burial), indicating the tight association and the stability of this complex.

Most importantly, residues 258–269 of the securin SIS, with good electron density (Fig. 3b), are located in the active site of separase, interacting with the SD and CD (Fig. 3a, c). Therefore, one mechanism for securin to inhibit the protease activity of separase is through blocking substrate access to the active site. Residues Ile259–Glu260–Ile261 of securin mimic the P2–P3 residues of the substrate17 (Fig. 3c). Notably, the side chain of Ile259 has hydrophobic interactions with Trp1249 and Trp1250 located near the tip of the αA–αB helical hairpin of SD, and that of Glu260 is positioned near the N terminus of helix E in CD. However, the P1 Arg residue of the substrate is replaced with Pro263 in securin. It assumes a different conformation and its main-chain carbonyl is 6 Å away from the thiol group of the catalytic Cys1531 residue (Fig. 3c, d). The position of Pro263 also clashes with that of His2083 in the *C. thermophilum* CD17, the second member of the catalytic machinery, and the conformation of the equivalent His1505 in the securin complex is probably an inactive one (Fig. 3d). These differences ensure that securin is not cleaved by separase even though it is located in the active site. Earlier studies showed that mutations of residues of securin equivalent to the P2, P1 and P1′ residues could turn it into a substrate14,17 (Extended Data Fig. 8).

Residues 264–269 of securin have interactions with the CD, and the P′ side of the substrate may be recognized by this region of separase as well. In contrast to the sequence conservation of the P side of the substrate, the P′ side is much less conserved.
Figure 3 | Interactions between separase and securin. a, Molecular surface of separase, coloured by the domains. The active site of separase is indicated by the red asterisk. The view is as in Fig. 1c. b, Omit F$_o$ – F$_c$ electron density at 3.0 Å resolution for residues 258–269 of securin, contoured at 2σ. c, Interactions between residues 258–265 of securin SIS (magenta) with the active site of separase. The side chains of residues in contoured at 2σ, Interactions between residues 258–265 of securin SIS (magenta) with the active site of separase. The side chains of residues 258–269 of securin, contoured at 2σ. d, Close-up of the active site region showing the differences between the bound position of securin (magenta) and that of the inhibitor to C. thermophilum separase (grey). e, Interactions between residues 271–288 of securin SIS (magenta) with domain III (light blue) of separase.

(Extended Data Fig. 8). A P$_1$′ Arg residue, found in several fungal species but not in Schizosaccharomyces pombe or C. thermophilum, may mediate the degradation of this fragment by the N-end rule pathway. The structure shows that a groove in the active site region can accommodate the P′ residues, and the β4A–β4B hairpin following the catalytic Cys1531 may contribute to substrate binding here (Extended Data Fig. 8: see below). Residues 264–269 of securin are in contact with the rim of this groove rather than being located at its bottom, possibly owing to the shift in the position of the Pro263 residue relative to that of the P$_1$ Arg of the substrate (Fig. 3c).

Following the active site region, residues 271–288 of the securin SIS are in close contact with domain III of separase (Fig. 3e), and residues 290–296 are located in a deep groove at the interface between domains II and III (Fig. 3a, Extended Data Fig. 6). These interactions are mostly hydrophobic in nature, and also involve several aromatic (Phe and Tyr) residues. Residues 306–316 of the securin SIS are placed on the side of domain II of separase (Fig. 1c). Finally, residues 365–373 at the C terminus of securin form a short helix (Extended Data Fig. 4) and are located in a surface depression in domain I (Fig. 1b, Extended Data Fig. 6). Residues 299–305 and 317–362 of securin are disordered in the current structure.

The securin SIS contains three potential phosphorylation sites (29, 30) (Fig. 1c: Ser277 (Fig. 3e), Ser292 (Extended Data Fig. 6) and Thr304. The S277A/S292A/T304A triple mutant had weaker interaction with separase and reduced nuclear location of separase, whereas the single and double mutants had no growth phenotype. pSer277 should have favourable interactions with Arg1130 in domain IV of separase (Fig. 3e). The side chain of Ser292 is hydrogen-bonded to the main-chain amides of residues 294 and 295 in the SIS, and Thr304 is in a disordered region in the current structure, suggesting possible conformational rearrangements of Ser292 and Thr304 after phosphorylation.

Securin residues in contact with securin are generally well conserved (Extended Data Figs 1–3). The securin SIS is poorly conserved among species, but the residues in the major contact sites with securin are better conserved (Extended Data Fig. 4), suggesting that the securin homologues may share a common mechanism of interacting with and inhibiting separase. The observed binding mode of the SIS indicates that securin probably helps to stabilize the overall structure of separase, as suggested by previous biochemical studies. This might also be possible that once securin is removed by proteasome degradation, the structure of separase alone could undergo a conformational change to assume a different arrangement of its domains, especially for domains I–III. The β4A–β4B segment is a loop (L4) in the structure of the C. thermophilum SD–CD (Extended Data Figs 6, 8), and residues in this loop are important for catalytic activity. This segment is stabilized by contacts with domain III (Extended Data Figs 6, 8), and therefore a change in the position of domain III might be necessary for catalysis.
This would represent another, indirect mechanism for securin to inhibit the catalytic activity of separase.

The structure of the separase–securin complex provides a molecular basis for the large amount of experimental data on this complex. This structure is also supported by our biochemical observations. The complex from the fungus *Zygosaccharomyces rouxii* (Extended Data Figs 1–4). *Zygosaccharomyces rouxii* securin contains only 282 residues owing to a shorter N-terminal segment (Fig. 4a). We found that separase was produced in the insoluble fraction in insect cells unless an appropriate securin was co-expressed with it (Fig. 4b, Extended Data Fig. 9), which was probably due to the instability/misfolding of securin. On the basis of this observation, we assessed the importance of various segments of securin for producing soluble separase, which would reflect both its binding and chaperone functions. The N-terminal segment of securin, up to residue 166 (equivalent to residue 258 of *S. cerevisiae* securin, Extended Data Fig. 4), could not help produce soluble separase, and deleting these 165 N-terminal residues had little effect on the production of soluble separase (Fig. 4b, c). Additional experiments showed that deleting residue 231 to the C terminus of securin led to a relatively small reduction in the solubility of separase (Fig. 4b, c). These data indicate that residues 166–230 of *Z. rouxii* securin, equivalent to residues 258–316 of *S. cerevisiae* securin, are required for interactions with separase, in agreement with our structural information.

We also observed that the segment covering residues 189 to the C terminus of *Z. rouxii* securin was able to produce a small amount of soluble separase (Fig. 4b, c). This segment would be missing the residues located in the active site of separase, suggesting that binding to the active site is not essential for complex formation with separase. Conversely, gel-filtration studies of this complex showed that it migrated in the void volume, indicating that it was mostly aggregated. Therefore, even though this segment of securin could bind separase, it might not be sufficient for its chaperone function to promote the correct separase folding and/or conformation.

The segment covering residues 166–188 of *Z. rouxii* securin was not able to help produce soluble separase, indicating that binding to the active site region of securin alone is not sufficient to produce a stable complex and that the N-terminal helical region of separase has a crucial role in the regulation by securin. In addition, residues 189–250 of securin produced a much lower amount of soluble separase compared to that for residues 189–282 of securin, indicating an appreciable role for the C-terminal residues of securin.

Overall, our studies have produced the first crystal structure information on the separase–securin complex and defined the molecular basis for the tight regulation of separase by securin. The structure provides a foundation for understanding the biochemical and biological functions of this complex from *S. cerevisiae*. While the N-terminal helical region is poorly conserved among separases at the amino acid sequence level, they are better conserved at the secondary structure level (Extended Data Figs 1, 2). Therefore, it might be possible that the separase–securin complexes from other organisms could assume a similar overall architecture as observed here, which is supported by the similar shape for the human separase–securin complex. The information obtained from this structure may also be useful for deciphering the molecular mechanisms of other separase–securin complexes.

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 Information 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.T. (ltong@columbia.edu).

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**METHODS**

No statistical methods were used to predetermine sample size. The experiments were not randomized, and investigators were not blinded to allocation during experiments and outcome assessment.

**Protein expression and purification.** Separase and securin were co-expressed in insect cell system using Multibac technology (Geneva Biotech)\(^{31}\). Separase and securin gene fragments were cloned into the pFL dual plasmid through its PH and p10 promoters, respectively. The construct used for crystallization contained residues 51–1630 or 71–1630 of *S. cerevisiae* separase and residues 258–373 of securin. A hexa-histidine tag was added to the N terminus of separase, whereas securin did not carry a tag. Bacmids were generated by transforming constructs to DH10EMBacY competent cells (Geneva Biotech) and isolated by Bac-to-Bac protocol (ThermoFisher). Bacmids were further transfected into Sf9 cells using Cellfectin II (ThermoFisher) following the manufacturer’s instructions to produce P1 baculoviruses. P2 viruses were obtained by amplifying P1 viruses in Sf9 once and were used for large-scale protein expression.

High5 cells were grown in Fernbach flasks (PYREX) in ESF 921 medium (Expression Systems) by shaking at 120 r.p.m. at 27°C until the density reaches 2 × 10⁶ cells ml⁻¹. Cells were infected by P2 virus and collected 48 h after infection by centrifugation at 500g. The harvested cells were resuspended in lysis buffer containing 50 mM phosphate, pH 7.6, 500 mM NaCl, 5% (v/v) glycerol, 20 mM imidazole and protease inhibitor cocktail (Sigma) and lysed by sonication. Cell lysates were centrifuged at 24,000g for 40 min at 4°C before incubating with nickel beads (Qiagen). After 1 h, beads were transferred to a gravity flow column (Biorad) and washed extensively with lysis buffer. Protein was eluted with a buffer containing 50 mM phosphate (pH 7.6), 500 mM NaCl, 5% (v/v) glycerol and 500 mM imidazole. Protein eluate was further purified by gel filtration using Sephacryl S-300 column (GE Healthcare) equilibrated in a buffer containing 50 mM phosphate, pH 7.6, 500 mM NaCl and 2 mM DTT. The protein samples were concentrated to 10 mg ml⁻¹ and stored at −80°C.

**Protein crystallization.** Crystals were grown using the sitting-drop method at 20°C by mixing 1.0 μl protein solution at 3 mg ml⁻¹ with 1.0 μl well solution containing 0.25 M Na/K-phosphate (pH 6.0) and 14% (w/v) polyethylene glycol 3,350. Crystals appeared in 3 days and were transferred to the well solution plus 30% (v/v) glycerol as cryoprotectant before being flash-frozen in liquid nitrogen for data collection at 100 K. For phasing, a gold derivative was obtained by soaking crystals in mother liquor with 2 mM KAu(CN)₂ for 5 h and transferring to cryoprotectant for 2 min. The crystals belong to space group P3₂₁ with one separase–securin complex in the asymmetric unit.

**Data collection and structure determination.** X-ray diffraction datasets up to 3.0 Å resolution were collected at beamline 24-ID-E of the Advanced Photon Source (APS). The data were processed with the HKL package\(^{32}\). The structure was solved using the single isomorphous replacement with anomalous scattering method with the gold derivative with AutoSharp\(^{33}\), which located 10 gold sites. The atomic model was built manually with Coot\(^{34}\) and refined with PHENIX\(^{35}\). The crystallographic information is summarized in Extended Data Table 1.

From the same crystallization condition, crystals belonging to space group C₂ were also obtained, with unit cell parameters of \(a = 238.4 \AA, b = 89.5 \AA, c = 119.0 \AA\) and \(β = 105.2°\). There is one separase–securin complex in the asymmetric unit. The best diffraction dataset that we could collect on this crystal form extended to 3.7 Å resolution. The structure could be readily solved with the molecular replacement method using the structure in P3₂₁ as the model, and no significant structural differences were observed.

X-ray diffraction datasets for the crystal containing residues 71–1630 of separase were collected at beamline 24-ID-C of the Advanced Photon Source (APS). The structure was refined at 2.6 Å resolution, and the crystallographic information is summarized in Extended Data Table 1.

For the 2.6 Å structure, 94.8% of the residues are in the favoured region of the Ramachandran plot. For the 3.0 Å structure, 96.9% of the residues are in the favoured region of the Ramachandran plot. No residues are in the disallowed region in either structure.

**Mutagenesis studies.** Site-specific and deletion mutations were introduced with the QuikChange kit (Agilent) and sequenced for confirmation. The mutants were expressed in insect cells following the same protocol as that for the wild-type protein.

**Data availability.** The atomic coordinates and the diffraction datasets have been deposited at the Protein Data Bank with accession codes SU1S and SU1T. All other data are available from the corresponding author upon reasonable request.

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Extended Data Figure 1 | Sequence alignment of domains I and II of separase. The secondary structure elements in the *S. cerevisiae* separase structure are shown and labelled. The boundaries of the domains are indicated. Residues in contact with securin are indicated by the purple dots. *Ag*, *Ashbya gossypii*; *Kl*, *Kluyveromyces lactis*; *Sc*, *Saccharomyces cerevisiae*; *Zr*, *Zygosaccharomyces rouxii*. Modified from an output from ESPript.© 2017 Macmillan Publishers Limited, part of Springer Nature. All rights reserved.
Extended Data Figure 2 | Sequence alignment of domains III and IV of separase. The secondary structure elements in the *S. cerevisiae* separase structure are shown.
Extended Data Figure 3  |  Sequence alignment of the SD and CD of separase. The catalytic Cys1531 and His1505 residues are indicated by the red dots. Ct, Chaetomium thermophilum; Hs, Homo sapiens; Sp, Schizosaccharomyces pombe.
Extended Data Figure 4 | Sequence alignment of securin. The SIS is indicated. Residues in contact with separase are indicated by the blue dots. Residue 263 is equivalent to the P1 residue of separase substrates, and is indicated by the red asterisk. Residues 317–360 of \textit{S. cerevisiae} securin are disordered in the current structures and are poorly conserved in sequence.
Extended Data Figure 5 | Overlay of the structures of the separase–securin complexes. a, The complex formed by residues 71–1630 of separase and 258–373 of securin is shown in colour, and that by residues 51–1630 of separase and 258–373 of securin in grey. Residues 73–80 from another molecule of separase in the crystal is shown in orange, forming a β-sheet with the N-terminal segment of separase. b, Panel a viewed after a 50° rotation around the vertical axis.
Extended Data Figure 6 | Additional structural information on the separase–securin complex. a, 2Fo − Fc electron density for helices 4 and 5 of domain I at 2.6 Å resolution, contoured at 1σ. Helices 6 and 7 are also shown for reference. The directions of the helices are indicated by the red arrows. b, The β4A–β4B segment of CD (cyan) is immediately after the catalytic Cys1531 residue, and has interactions with domain III (light blue). The equivalent segment in the *C. thermophilum* SD–CD-free enzyme structure is a loop (L4), shown in dark grey. c, Interactions between residues 290–296 of securin SIS (magenta) with domains III (light blue) and II (light brown) of separase. d, Interactions between the C-terminal segment of securin SIS (magenta) and domain I of separase (green). Deletion of the first 155 residues of separase would remove helix α3 in this binding site.
Extended Data Figure 7 | Structural homologues of domains III and SD of separase. a, Overlay of the structures of domain III of separase (colour from N (blue) to C (red) terminus) and the TPR domain of LGN (grey, PDB accession 4WNG; 11% sequence identity, Z-score of 14.4)\textsuperscript{23–25}. The Frmpd4 ligand (black) of LGN is bound to a different region of the structure compared to securin. b, Overlay of the structures of domain III of separase and the subunit 7 of the APC/C (PDB accession 5G04; 5% identity, 14.3 Z-score\textsuperscript{37}). c, Overlay of the structures of the SD of separase (green) and a part of the PIWI domain of Argonaute (grey, PDB accession 4N76; 10% sequence identity, 5.5 Z-score). As a comparison, matching this \(\beta\)-sheet to that in \textit{C. thermophilum} separase produced a Z-score of 5.7 (refs 27, 38). Residues in the helical insert between \(\beta\)3 and \(\beta\)4 of separase are removed for clarity. d, Overlay of the structures of the SD of separase and the Yqgf domain of Tex (grey, PDB accession 3BZK; 4% sequence identity, 5.5 Z-score\textsuperscript{39}).
Extended Data Figure 8 | Possible binding groove for the P' residues of the substrate. a, Alignment of the separase cleavage sites in Scc1 substrates. The two cleavage sites in each protein are named 'a' and 'b'. The equivalent residues in securin are also shown. Asterisks indicate securin mutants (mutations in green) that become substrates of separase. The P and P' residues are labelled at the top, and the cleavage site is indicated by the vertical line. b, The overall binding mode of residues 258–271 of securin in separase. The β4A–β4B segment of CD (cyan) is a loop (L4, dark grey) in the *C. thermophilum* SD–CD structure. c, A groove in the active site of separase (red arrows) can accommodate the P' residues. The blue arrow indicates another groove in this region, but the binding mode of securin suggests that the groove indicated by the red arrows is more likely.
Extended Data Figure 9 | Biochemical characterizations of the interactions between separase and securin. *Zygosaccharomyces rouxii* separase was co-expressed with various segments of *Z. rouxii* securin, with truncations at the N and/or C terminus. The insoluble fraction was run on SDS–PAGE. The position of separase (with an N-terminal His tag) is indicated with the black arrowhead. WT, full-length *Z. rouxii* separase; WT-CS, full-length *Z. rouxii* separase with C1497S mutation; WT', *Z. rouxii* separase with an internal deletion of residues 952–1010, corresponding to a segment in domain IV of the *S. cerevisiae* separase structure. For gel source data, see Supplementary Fig. 1.
### Extended Data Table 1 | Data collection and refinement statistics

|                           | Separase (51-1630)-securin (258-373) complex crystal form I | Separase (71-1630)-securin (258-373) complex crystal form I | Separase (51-1630)-securin (258-373) complex crystal form II |
|---------------------------|-------------------------------------------------------------|-------------------------------------------------------------|-------------------------------------------------------------|
| **Data collection**        |                                                             |                                                             |                                                             |
| Space group               | $P3_21$                                                     | $P3_21$                                                     | $C2$                                                        |
| Cell dimensions           |                                                             |                                                             |                                                             |
| $a, b, c$ (Å)              | 126.3, 126.3, 273.9                                         | 125.9, 125.9, 271.9                                         | 238.4, 89.5, 119.0                                          |
| $\alpha, \beta, \gamma$ (°) | 90, 90, 120                                                | 90, 90, 120                                                | 90, 105.2, 90                                               |
| Resolution (Å)            | 50-3.0 (3.11-3.0)*                                          | 50-2.6 (2.69-2.6)*                                         | 50-3.7 (3.93-3.7)                                           |
| $R_{merge}$ (%)           | 14.4 (>100)                                                | 14.5 (>100)                                                | 16.6 (232)                                                  |
| $CC_{1/2}$                | (0.480)                                                     | (0.476)                                                     | (0.271)                                                     |
| $I/|\sigma|$                | 15.0 (1.5)                                                  | 14.4 (1.2)                                                  | 4.43 (0.49)                                                 |
| Completeness (%)          | 99.8 (99.6)                                                 | 100 (99.9)                                                  | 96.2 (90.9)                                                 |
| Redundancy                | 10.4 (7.6)                                                  | 10.0 (8.2)                                                  | 2.7 (2.5)                                                   |
| **Refinement**            |                                                             |                                                             |                                                             |
| Resolution (Å)            | 50-3.0 (3.11-3.0)                                           | 50-2.6 (2.69-2.6)                                          |                                                             |
| No. reflections           | 51,130 (4,818)                                              | 77,428 (7,588)                                             |                                                             |
| $R_{work}$ / $R_{free}$ (%) | 19.5 (31.4) / 25.2 (36.3)                                   | 22.1 (34.6) / 26.4 (37.0)                                 |                                                             |
| No. atoms                 |                                                             |                                                             |                                                             |
| Protein                   | 12,609                                                      | 12,633                                                      |                                                             |
| Ligand/ion                | 0                                                           | 0                                                           |                                                             |
| Water                     | 0                                                           | 16                                                          |                                                             |
| B-factors                 |                                                             |                                                             |                                                             |
| Protein                   | 97.1                                                        | 72.0                                                        |                                                             |
| Ligand/ion                | -                                                           | -                                                           |                                                             |
| Water                     | -                                                           | 63.2                                                        |                                                             |
| R.m.s deviations          |                                                             |                                                             |                                                             |
| Bond lengths (Å)          | 0.003                                                       | 0.011                                                       |                                                             |
| Bond angles (°)           | 0.65                                                        | 1.2                                                         |                                                             |

*Two crystals were used for data collection for crystal form I, and one crystal for crystal form II.*

*Highest resolution shell is shown in parenthesis.*
Corrigendum: Molecular mechanism for the regulation of yeast separase by securin
Shukun Luo & Liang Tong

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In Fig. 3c of this Letter, the bottom left helix should have been labelled ‘αE’ rather than ‘αA’. This helix is also mentioned in the main text, where “positioned near the N terminus of helix αA in CD” should have read “positioned near the N terminus of helix αE in CD”. The original Letter has been corrected online.