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Molecular dissection of the actin-binding ability of the fission yeast α -actinin, Ain1, *in vitro* and *in vivo*

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Running title: in vitro and in vivo activities of S. pombe Ain1

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Summary

A contractile ring (CR) is involved in cytokinesis in animal and yeast cells. Although several types of actin-bundling proteins associate with F-actin in the CR, their individual roles in the CR have not yet been elucidated in detail. Ain1 is the sole α -actinin homologue in the fission yeast *Schizosaccharomyces pombe* and specifically localizes to the CR with a high turnover rate. *S. pombe* cells lacking the *ain1*⁺ gene show defects in cytokinesis under stress conditions. We herein investigated the biochemical activity and cellular localization mechanisms of Ain1. Ain1 showed weaker affinity to F-actin *in vitro* than other actin-bundling proteins in *S. pombe*. We identified a mutation that presumably loosened the interaction between two calponin-homology domains constituting the single actin-binding domain (ABD) of Ain1, which strengthened the actin-binding activity of Ain1. This mutant protein induced a deformation in the ring shape of the CR. Neither a truncated protein consisting only of an N-terminal ABD nor a truncated protein lacking a C-terminal region containing an EF-hand motif localized to the CR, whereas the latter was involved in the bundling of F-actin *in vitro*. We herein propose detailed mechanisms for how each part of the molecule is involved in the proper cellular localization and function of Ain1.

Keywords: actin, actin-bundling protein, contractile ring, cytokinesis, fission yeast

Introduction

The contractile ring (CR), which mainly consists of filamentous actin (F-actin) and myosin-II, induces the formation of a cleavage furrow during cytokinesis in animal and yeast cells. Several types of actin-modulating proteins control the properties and behavior of F-actin in the CR (1, 2). The bundling of F-actin may be indispensable for maintaining the microstructure and overall architecture of the CR. However, contraction of the CR, which is mediated by an interaction between F-actin and myosin II, may be suppressed if F-actin is too tightly bundled. This may apply not only to the CR, but also to the large number of actomyosin-sliding systems responsible for cell migration and movement. Moreover, the three-dimensional structure of the actin cytoskeleton and its mechanical properties may be significantly affected by the presence of actin-bundling proteins that directly determine the number of cross-linking sites, the distance and angle between filaments, and plasticity of the cytoskeletal structure. This may explain why various types of actin-bundling proteins are found in eukaryotic cells.

Several members of the calponin superfamily play important roles in the bundling of F-actin. Calponin is strongly expressed in smooth muscle cells and controls actomyosin activity (3). Approximately 100 a. a. in the N-terminal half of this protein is called the calponin-homology domain (CHD). Various actin-bundling proteins utilize their CHDs as an interface to interact with F-actin (4, 5). IQGAP is one of the actin-bundling proteins, the function of which has been characterized in detail both *in vivo* and *in vitro* (6–9). This protein forms a dimer and bundles F-actin by binding to different F-actins with the N-terminal CHD of each subunit (10). A previous study reported that IQGAP and its homologue localized to the CR of various types of cells including the sea urchin fertilized egg (11), cultured mammalian cells (12, 13), the budding yeast Saccharomyces cerevisiae (14), and the fission yeast Schizosaccharomyces pombe (15). Furthermore, another single CHD-containing protein Stg1, a homologue of the SM22/transgelin-like protein, was found in the division site of an S. pombe cell; however, unlike the IQGAP-homologue Rng2, its function was not essential for the progression of cytokinesis (16). On the other hand, many actin-bundling proteins utilize a double repeat of CHDs as a single actin-binding domain (ABD). Filamin forms a dimer and exerts F-actin-bundling activity through the N-terminal ABD of each subunit (17). This protein is found in a cleavage furrow in vertebrate cells (18, 19). The cellular slime mold *Dictyostelium discoideum* has homo- and hetero-dimeric actin-bundling proteins, called cortexillin-I, -II, and -III, which are involved in cytokinesis (20, 21). Cortexillin-I bundles F-actin with its C-terminal short tail instead of its N-terminal ABD (22). An ABD has also been found in the actin-bundling proteins fimbrin and α -actinin. Fimbrin consists of a tandem repeat of double ABDs in a single molecule, and each ABD binds to different F-actins. Previous studies demonstrated that fimbrin localizes to the CR in the ciliate *Tetrahymena* (23) and *S. pombe* cells (24, 25). α -Actinin forms an anti-parallel dimer via its spectrin-like repeats (SRs) located in the middle of the molecule and bundles F-actin. In mammalian cells, α -actinin is involved in controlling the ingression rate of the cleavage furrow (26). The α -actinin homologue Ain1 has also been identified in the CR of *S. pombe* cells (25).

The molecular mechanisms underlying the formation of the CR have been extensively investigated in the fission yeast *S. pombe*, and the cytokinetic functions of actin-bundling proteins have been studied in detail (*15*, *16*, *23*, *24*, *27*, *28*). Of these, only IQGAP Rng2 and α -actinin Ain1 show CR-limited localization (15, 25), while other actin-bundling proteins such as fimbrin Fim1 and Stg1 are also found in actin patches, a different actin cytoskeleton engaged in endocytosis, in this organism (*16*, *24*, *25*). Rng2 plays an essential role in the proper organization of F-actin in the CR (*15*, *27*). However, mutant cells, in which the chromosomal *rng2*⁺ gene is replaced with the truncated gene encoding N-terminal CHD-lacking Rng2, still remain and undergo cell division (*28*). Therefore, some actin-bundling proteins may redundantly function with Rng2 in cytokinesis in this organism. Functional relationships between Rng2, Ain1, and Fim1 have been detected in *S. pombe* (*24*, *25*). Similar to mammalian α -actinin, Ain1 contains three functional moieties: N-terminal ABD, SRs in the middle region, and the C-terminal EF-hand motif. A previous study demonstrated that cytokinesis failed in *ain1*-null cells exposed to stress conditions and synthetic lethality occurred with the simultaneous gene deletion of *fim1*⁺ (*25*). Moreover, Ain1 appears to cooperatively promote the formation of the CR with myosin-II Myo2 (29). However, the biochemical and structural characterization of Ain1 had not been performed until recently (30, 31). We herein evaluated the actin-binding activity of Ain1 using the recombinant protein. We also investigated the cellular localization of Ain1 and mutant proteins and elucidated the intra-molecular mechanisms underlying the CR-limited localization of Ain1. Our results indicate an elaborate mechanism involving Ain1 in the binding of F-actin in the CR.

Materials and Methods

Yeast strains and cell culture

We used *S. pombe* wild-type (h⁻ *leu1-32*) and $\Delta ain1$ (h⁻ *leu1-32 ain1::kan*^r) strains in this study. In order to observe the actin cytoskeleton, we used a strain that expressed a LifeAct peptide (*32*) fused with a four tandem repeat of the fluorescent protein mCherry (h⁻ *ura4-D18 ade6-M216*⁺ *leu1::cdc4-promoter-lifeact-4×mCherry*) (*33*). According to the method described by Forsburg and Rhind (*34*), rich medium YE and minimal medium EMM with supplements were used for cell cultures. In microscopic observations, cells were cultured in EMM liquid medium at 30°C to the log growth phase.

Protein Purification

The *ain1*⁺ gene was artificially synthesized and cloned in a bacterial plasmid (see Fig. S1) because our attempts to clone any cDNA encoding an entire region of the open reading frame (ORF) were unsuccessful. The codon usage of artificially synthesized cDNA was modified for the efficient translation of the gene product in *Escherichia coli* and *S. pombe*. After plasmid DNA was digested with *Eco*RI and *Sal*I, electrophoretically separated 1.2-kb DNA fragments covering the ORF of *ain1*⁺ were extracted from agarose gel and inserted into pGEX6P-1 (GE Healthcare (GE)). In order to prepare truncated or point-mutated recombinant proteins (Table S1), the *ain1*⁺ gene on the vector, pGEX6P-ain1, was modified using the PrimeSTAR*mutagenesis basal kit (Takara). *E. coli* strains, *XL1-Blue*, *BL21*, or Rosetta-gami (DE3) pLysSRARE2 (Takara) were transformed with the resultant vectors, and N-terminal glutathione *S*-transferase (GST)-tagged proteins were expressed according to the manufacturer's protocol (GE).

Bacterial cells expressing the recombinant proteins were lysed by sonication in 50 mM Tris-HCl (pH 8.0) containing 100 mM NaCl, 0.5% NP-40, 1 mM DTT, and protease inhibitors (2 mM PMSF and 1 μ g/mL Pepstatin A) on ice, and the cell lysate was centrifuged (10,000×g, 10 min) to remove cell debris. The supernatant was incubated with Glutathione Sepharose 4B beads (GE) for the affinity purification of GST-fusion proteins at 4°C for 18 h. The proteins that bound to the

beads were cleaved by PreScission Protease (GE) in 50 mM Tris-HCl (pH 6.8) containing 150 mM NaCl, 1 mM DTT, and 1 mM EDTA on ice. The eluted final product was collected and stored at -80°C. Before being examined, thawed proteins were centrifuged ($200,000 \times g$, 10 min) to remove aggregates.

G-actin from rabbit skeletal muscle was purified in our lab as described previously (27) or an equivalent product was kindly provided by Prof. Uyeda's lab (National Institute of Advanced Industrial Science and Technology, Japan (AIST)). Some experiments were performed using chicken skeletal muscle actin instead of rabbit skeletal muscle actin. Their primary sequences were identical. G-actin was dissolved in KMEI buffer (100 mM KCl, 1 mM Mg₂Cl, 0.2 mM EGTA, 0.2 mM DTT, 0.2 mM ATP, and 100 mM imidazole-HCl (pH 7.5)) to prepare F-actin.

Purified proteins used in this study are shown in Fig. 1A.

Co-sedimentation assay

F-actin with various concentrations of Ain1 or Ain1ABD was incubated in KMEI at 25°C for 30 min. In the bundling assay with Ca²⁺ ions, CaCl₂ was added at a final concentration of 0.2 mM to the reaction solution in consideration of the concentrations of Ca²⁺ chelators and salts. The mixtures were centrifuged at 200,000×g for 20 min (high-speed) or at 15,000×g for 15 min (low-speed). The supernatants and pellets were separated by SDS-PAGE, and gels were stained with Coomassie brilliant blue (CBB). CBB-stained gels were scanned as digital images in order to quantify the concentrations of actin and Ain1 using ImageJ 1.47 (National Institutes of Health, USA). Apparent dissociation constants (K_d) were calculated for F-actin from hyperbolic curves fit to the plot of [protein bound to F-actin] versus [protein free] using R (http://www.R-project.org/).

Actin-bundling assay

Dried rhodamine-phalloidin (Molecular Probes) in a microtube was dissolved in KMEI buffer, and F-actin was labeled overnight in the microtube. Labeled F-actin (4 μ M) and Ain1 or Ain1-mutant proteins (8 μ M) were mixed and incubated at 25°C for 10 min. The fluorescence microscope BX51

(Olympus) with a PlanApo 100x/1.40 NA objective lens (Olympus) was used for observations. Microscopic images were obtained by ORCAII-ER-1394, the cooled charge-coupled device (CCD) camera, (Hamamatsu Photonics) with the software Simple PCI (Compix Inc.).

Cellular localization of Ain1 and mutant proteins

The thiamine-repressible expression vector pREP1 (35) was used for the expression of the N-terminal yellow-fluorescence protein (YFP)-tagged Ain1 and mutant proteins in *S. pombe* (Table S1). Truncated or point mutated genes were prepared using the PrimeSTAR* mutagenesis basal kit (Takara). Cells were transformed with vectors using the lithium acetate method (36). SD medium (Becton, Dickinson and Company) was conveniently used as a minimal medium for the first auxotrophic selection of transformants. EMM medium with (repressed condition of the inserted gene) or without (induced condition) 5 μ M thiamine was used for microscopic observations of cells.

Cell images were acquired with the fluorescence microscopy system described above. Prior to measuring the florescence intensity of YFP-tagged proteins using ImageJ 1.47 software, mean background intensity was subtracted from the images acquired. Cytoplasmic areas, avoiding nuclei, were arbitrarily selected. Lines with a width of 3 pixels (0.19 μ m) were drawn on the CR and the average intensity of two peaks was measured for each cell.

Results

Ain1 binds weakly to F-actin

We measured the actin-binding activity of the full-length molecule of Ain1 (Ain1FL) using a co-sedimentation assay. F-actin predominantly accumulated in pellets with ultracentrifugation (Fig. 1B, bottom panel). On the other hand, Ain1FL alone was detected in the supernatant; however, this molecule was also detected in the pellet fraction under our experimental conditions (Fig. 1B, upper panel). Although the reason for this remains unclear, Ain1FL has been suggested to adhere to the internal walls of centrifuge tubes. By mixing Ain1FL with F-actin, the proportion of Ain1FL in the pellet fraction increased (Fig. 1B, bottom panel). This result indicates that Ain1FL possesses F-actin-binding activity. We then estimated the K_d of Ain1FL (Ain1FL, 6.53 ± 0.55; mean ± standard deviation (SD)) (Fig. 1C); however, the saturated binding ratio of Ain1 to F-actin was not accurately measured. This apparent K_d was similar to the value of 4.3 μ M reported by Li and colleagues (31). In fission yeast, the K_d of three other actin-binding proteins were as follows; Stg1, 4.4 μ M (16); Rng2CHD, 0.21 ± 0.08 μ M (27); Fim1, 0.645 ± 0.15 μ M (37). Regarding the K_d of Rng2CHD, Tebbs and Pollard recently obtained a 100-fold higher K_d , 20 μ M (28). They indicated that the lower K_d in a previous study (27) may have been an artifact of the effect of a His tag that was not removed from recombinant Rng2CHD. On the other hand, Uyeda and his colleagues independently purified Rng2CHD and measured its K_d against F-actin after removing a His tag from the recombinant protein. Their measured value was 0.42 μ M (our personal communication). Thus, Ain1 appears to be an actin-binding protein with relatively low affinity to F-actin in S. pombe. It should be noted that all of the measurements cited above and this study have been performed with actin purified from vertebrate skeletal muscle but not S. pombe cells.

We next attempted to assess the biochemical activity of the ABD in the N terminus of Ain1. The ratios of the truncated protein Ain1ABD in a pellet did not significantly change with or without F-actin (Fig. 1D), suggesting that Ain1ABD hardly bound to F-actin. We discussed the reason for this later.

Ain1 bundles F-actin independently of Ca²⁺ ions

The actin-bundling activity of Ain1 was examined. F-actin formed thick and long bundles in the presence of Ain1FL, but not Ain1ABD (Fig. 2A). In addition, F-actin with Ain1FL, but not F-actin alone dominantly appeared in the pellet fraction after low-speed centrifugation (Fig. 2B). These results demonstrated that Ain1 exhibits actin-bundling activity *in vitro*. Moreover, a significant amount of F-actin was found in the pellets also at relatively lower concentrations of Ain1FL (Fig. 2C) even though the actin-binding affinity of Ain1FL was not so high as mentioned before (*see* Fig. 1C), suggesting that F-actin could be effectively precipitated at low degree of crosslinking in this assay. On the other hands, Ain1ABD did not exhibit actin-bundling activity in the same condition (Fig. 2A, C).

The EF-hand is the representative Ca²⁺-binding motif of eukaryotic proteins (38). A structural analysis revealed that the C-terminal EF-hand of α -actinin was adjacent to the N-terminal ABD of the paired molecule in a dimer of α -actinin (39). In addition, several types of α -actinin including mammalian isoforms 1 and 4 expressed in non-muscle tissues have been shown to exhibit Ca²⁺-sensitive actin-binding activity (40). The interaction between the EF-hand motif and ABD regulates the actin-binding activity of α -actinin. A low-conservative EF-hand was detected in Ain1, but may be insensitive to Ca²⁺ ions (25). In order to assess the effects of Ca²⁺ ions on the actin-bundling activity of Ain1, we performed a low-speed co-sedimentation assay (Fig. 2C). The results obtained showed that Ca²⁺ ions did not significantly affect the actin-bundling activity of Ain1.

Three actin-binding sites of Ain1 may have different roles in a cell

The ABD of α -actinin consists of two-tandem CHDs. The N-terminal CHD (CH1) contains two actin-binding sites, ABS1 and ABS2, whereas the other CHD (CH2) has a single site, ABS3 (41–43). In dividing *S. pombe* cells, Ain1 specifically localizes to the CR in a manner that depends on F-actin (25). We investigated the cellular localization of YFP-tagged Ain1-truncated proteins in wild-type cells in order to characterize the roles of two CHDs in Ain1 in a cell (Fig. 3A). Consistent

with biochemical activity, Ain1ABD did not localize to any structures known as the actin cytoskeleton (Fig. 3A). On the other hand, CH1 had the ability to localize to the CR (Fig. 3A, S3). However, the cellular localization of CH1 appeared to be abnormal because several cortical dots and cables were detected besides the medial ring. Most of these excrescent structures were not labeled with lifeact-4×mCherry, which was used as a marker for F-actin structures (Fig. S3). CH1 and the Lifeact peptide may bind to F-actin in a competitive manner (44). Moreover, the overproduction of CH1 induced the formation of thick actin fibers that were significantly resistant to the actin-polymerization inhibitor, Latrunculin-A (Fig. S4). Therefore, CH1 may bind to F-actin in the actin cytoskeleton of a cell; however, this appeared to affect its organization and dynamics. On the other hand, CH2 was not observed in any actin structures (Fig. 3A). We speculated that CH2 retained actin-binding activity because Ain1 Δ CH1, the N-terminal truncated Ain1 that lacks CH1, remained localized to the CR, although its activity appeared to be significantly weaker than that of Ain1FL (Fig. 3A). Moreover, a point mutation in ABS3 abolished the CR localization of Ain1 ACH1 (see below), and neither Ain1 ABD nor Ain1 EF localized to the CR (Fig. 3A). In addition, the overexpression of Ain1 Δ CH1 produced elongated cells with multiple nuclei, possibly as a result of interfering with the actin cytoskeletal organization during cytokinesis (Fig. 3B), which is reminiscent of the defect in cytokinesis caused by the overexpression of Ain1FL (25). Taken together, both CHDs in Ain1 appear to have the ability to bind to F-actin in a living cell, but may exhibit different activities. We then attempted to assess their biochemical activities, but were unsuccessful because their corresponding functional recombinant proteins were not available as they were insoluble (data not shown).

As described above, we were unable to clearly detect the actin-binding activity of Ain1ABD in vitro and in vivo, which suggests that CH1 and CH2 mutually affect each other in a suppressive manner. In *H. sapiens* α -actinin ACTN4, the single point mutation that substitutes the lysine residue at 225 with glutamate causes a change in the configuration of two CHDs and increases affinity to F-actin (42). We examined the K_d of Ain1 R216E that possessed a mutation presumably equivalent to the K225E mutation in ACTN4, against F-actin, and estimated it to be 1.80 ± 0.47 (mean ± SD),

suggesting that the substitution of the single amino acid residue greatly increased the actin-binding activity of Ain1 (Fig. 1C). In addition, it was found that Ain1 R216E bundled F-actin (Fig. 2A) although the activity was somewhat weak as compared with the wild-type Ain1FL at the lower concentrations of those proteins to actin (Fig. 2C). We also found that although Ain1 R216E proteins encircled the middle region of mitotic cells, the ring shape was deformed (Fig. 3A, S3). This phenotype resembled the abnormal organization of the CR in Ain1-overproducing cells (29). Independent of this study, Li et al. recently reported that the formation and constriction of the CR were affected by the R216E mutation in $ain1^+$ (31). In order to compare the accumulation of Ain1R216E with the wild-type Ain1, Ain1FL, in the middle of the cell, the intensities of these YFP-fusion proteins were quantified (Fig. 3C). The accumulation of Ain1R216E in the cell middle $(7.39 \pm 3.01, \text{ mean} \pm \text{SD})$ was approximately 2-fold greater than that of Ain1FL (3.78 ± 0.92) . These results suggest that the structural interaction between CH1 and CH2 moderates the actin-binding activity of Ain1 in a similar manner to ACTN4 (42). However, the introduction of the R216E mutation did not restore the actin cytoskeleton-binding activity of the ABD in vivo (see Ain1ABD R216E in Fig. 3A). Therefore, the dimerization of protein or the intra- or inner-molecular interaction of the ABD with another part of Ain1 may be important for the actin-binding activity of the full-length molecule.

We also investigated the functional relationship of each ABS using the point mutations, ABS1-2A, ABS2-1A, and ABS3-2A (*see* Fig. 6A, Table S1). These mutations substitute the charged amino acid(s) to alanine in the corresponding ABS. The ABSs of Ain1 were deduced from the alignment of the primary sequence of Ain1 with the human skeletal muscle α -actinin ACTN3 and *S. pombe* fimbrin (41, 43). ABS1-2A did not significantly affect the cellular localization of the full-length protein or truncated CH1 domain (Fig. 3A), whereas the corresponding mutation significantly reduced the actin-binding activity of ACTN4 (42). In contrast, ABS2-1A abolished the CR localization of the full-length protein (Fig. 3A). Moreover, CH1 with the ABS2-1A mutation did not display marked cellular localization (Fig. 3A). Therefore, ABS2 appears to be the major actin-binding site of Ain1 in *S. pombe* cells. However, the importance of ABS1 and ABS2 in the

cellular localization of Ain1 was affected by the R216E mutation. Ain1 containing the ABS2-1A mutation simultaneously with R216E localized to the CR, whereas Ain1 containing ABS1-2A and R216E did not (Fig. 3A). Therefore, the R216E mutation may induce a steric conformational change in the CH1 domain. On the other hand, Ain1ΔCH1 with the ABS3-2A mutation lost its localization to the CR, suggesting that ABS3 is the sole actin-binding site in the CH2 domain (Fig. 3A). Furthermore, Ain1 ABS3-2A localized not only to the CR, but also cortical dots (Fig. 3A); however, excrescent dot-like localization was not observed in cells co-expressing Lifeact-4×mCherry presumably because of their competitive binding to F-actin (Fig. S3). Thus, ABS3 appears to play an important role in the CR-specific localization of Ain1.

We then evaluated the cell growth abilities of *ain1*-null cells expressing each Ain1 mutant protein under stress conditions. At 18°C in the presence of 1 M potassium chloride, a condition described previously (25), *ain1*-null cells expressing Ain1ABS3-2A grew as well as wild-type cells and *ain1*-null cells expressing Ain1FL (Fig. 4). This result suggested that Ain1ABS3-2A was able to function as well as the wild-type Ain1FL *in vivo* even though it localizes to other actin structures besides CR. Thus, CR-limited localization of Ain1 is not essential for cytokinesis. On the other hand, neither the expression of Ain1 ABS1-2A nor Ain1 ABS2-1A restored growth defects in *ain1*-null cells under the same conditions. Therefore, not only ABS2, but also ABS1 may be required for the binding of Ain1 to F-actin in the CR. In support of this, Ain1 ABS1-2A was excluded from the CR when Lifeact-4×mCherry was simultaneously expressed (Fig. S3).

The C-terminal region is required for the cellular localization of Ain1

In the course of our study on the cellular localization of truncated Ain1 proteins, we found that Ain1 lacking the C-terminal region, Ain1 Δ EF, did not localize to the CR (Figs. 3A, S5B). The expression level of Ain1 Δ EF was similar to that of the wild-type protein in control cells (Fig. S5A, C). Coincidentally, the expression of Ain1 Δ EF from the plasmid failed to rescue the growth defect in

ain1-null cells under stress conditions (Fig. 4). A previous study suggested that the actin-binding activity of mammalian α -actinin is controlled by the C-terminal region, containing an EF-hand motif, of the paired molecule in the dimer (*45*). Similarly, the intersubunit interaction between the N-terminal ABD and C-terminal region may be required for the actin-binding activity of Ain1 in *S. pombe* cells. Ain1 \Delta EF containing the R216E mutation localized to the medial ring and cortical dots (Figs. 3A, S3). Therefore, the C-terminal region of Ain1 may positively function in the actin binding of the paired molecule in the α -actinin dimer by controlling conformational changes in the ABD. The results of biochemical experiments showed that Ain1 Δ EF bound F-actin with a slightly higher affinity than Ain1FL; the apparent K_d of Ain1 Δ EF with F-actin was 4.92 ± 1.08 (mean ± SD) (Fig. 1C). Moreover, Ain1 Δ EF induced the formation of actin bundles *in vitro*, but is important *in vivo*.

Investigation of the role of PIP₂ in the cellular localization of Ain1

Phosphatidylinositol 4,5-bisphosphate (PIP₂) has been identified as a regulatory molecule of α -actinin. α -actinin has a single PIP₂-binding site (46), and PIP₂ enhances the formation of F-actin gels induced by α -actinin (47). A previous study reported that the putative PIP₂ binding-site in Ain1 had 47 % identity with chicken skeletal-muscle α -actinin (25). In addition, PI (4) P-5-kinase Its3 and the product PIP₂ were both present in the division site in *S. pombe* (48). We then investigated whether Its3 is required for the localization of Ain1 in the middle of mitotic cells. However, Ain1 was still localized there in *its3* mutant cells, even at the restrictive temperature 36°C (Fig. 5). Therefore, PIP₂ is unlikely to regulate the mitotic-specific interaction of Ain1 with F-actin in the CR.

Discussion

Ain1 may ensure actin dynamics of F-actin bundles in the CR

We herein biochemically demonstrated that Ain1 exhibited actin-bundling activity, whereas it bound to F-actin with lower affinity than other actin-bundling proteins previously studied in S. *pombe*. The mechanisms underlying cytokinesis in fission yeast have been examined in detail. The SCPR model represents one of the convincing theories for the formation of the CR from nodes (49). S. pombe cells have approximately 50-60 nodes consisting of CR-precursor proteins in the cell middle before mitosis (50). After G2/M transition, each node recruits myosin II and formin, an actin-polymerizing protein essential for CR formation, and F-actin newly polymerized from each node connects them into a single network. The interaction of F-actin and myosin II between each node induces their redistribution together with their associated proteins, and the equatorial actomyosin network finally forms a ring. Ain1 has been suggested to subserve a function in CR formation from nodes by cross-linking the F-actin network (29). Moreover, Ain1 may bundle F-actin in the CR in order to ensure the maintenance of the ring shape after its formation. A previous study demonstrated that mammalian α -actinin, which is involved in cytokinesis, shows a high affinity for F-actin ($K_d = 0.20 \pm 0.01 \mu M$) (51). On the other hand, the extremely low affinity of Ain1 to F-actin indicates that Ain1 is unlikely to bundle F-actin stably. Differences in the a-actinin-mediated F-actin-bundling activity in the CR between mammalian cells and fission yeast cells may be attributed to the sizes of their CRs. In addition, mammalian α -actinin has been shown to decelerate the constriction rate of the CR (26). On the other hand, the constriction rate of the CR is not changed with or without Ain1 (29), suggesting that Ain1 is an actin cross-linker that does not affect actin dynamics in the CR. The point mutation R216E resulted in the prominent accumulation of Ain1 in the cell middle (Fig. 3C). The excessive F-actin-binding activity of Ain1 may deform the ring shape of the CR, which is consistent with the findings of computational simulations performed previously (29). In proper cytokinesis in fission yeast, Ain1 may ensure the dynamic constriction of the CR by weakly bundling F-actin in the CR.

Mechanism of the CR-specific localization of Ain1

Fission yeast has three types of actin structures (*i.e.* actin patches, actin cables, and the CR). Ain1 specifically localizes to the CR in a manner that is dependent on F-actin, as demonstrated previously (25). The spatial and temporal regulatory mechanisms underlying the localization of Ain1 have not yet been elucidated. In mammalian cells, the actin-binding activity of α -actinin decreases with Ca^{2+} , and Ca^{2+} sensitivity is required for the accumulation of α -actinin in the cleavage furrow (52). In contrast, the results of our *in vitro* experiment indicated that Ca²⁺ does not inhibit the actin-bundling activity of Ain1 (Fig. 2C). On the other hand, Ain1 Δ EF did not localize to any actin structures in vivo. A recent structural study on mammalian α -actinin identified an interaction between the EF-hand with the ABD (45) and the neck region connecting the ABD to SRs (39). Regardless of Ca²⁺ insensitivity, the C-terminal region including the EF-hand in Ain1 may be necessary for maintaining the functional dimer form of Ain1 with actin-binding activity. However, Ain1 AEF bundled F-actin in vitro (Fig. 2A, C). Thus, the C-terminal region-dependent regulation of Ain1 may exert a strong influence on F-actin undergoing a dynamic turnover and interacting with various types of actin-modulating proteins in vivo. In future studies, it may be important to investigate whether the structure and function of the C-terminal region are controlled by cell cycle machinery.

We successfully demonstrated the functional diversity of CH1 and CH2 in Ain1 using truncated and point mutated genes (Figs. 3A, 4). In a cell, CH1 localizes to the CR and other types of actin structures in a manner that may be mediated by the interaction of ABS2 with F-actin (Fig. 6). The non-selective association of CH1 with the actin cytoskeleton resembles the case of Rng2. The corresponding truncated protein, Rng2CHD, localizes not only to the CR, but also to other actin cytoskeletons including actin cables and actin patches (27). In Rng2, the C-terminal region adjacent to the CHD appears to play an important role in the recognition of F-actin in the CR because Rng2Ns consists of the CHD and the subsequent short region specifically localizes to CR, similar to the full-length protein. On the other hand, our results suggest that CH2 plays a critical role in specific targeting of Ain1 to the CR (Figs. 3A, 6). Therefore, the actin-binding property of CH1

may be modulated by the adjacent CH2. Coincidently, structural studies have indicated a regulatory interaction between CH1 and CH2 in mammalian α -actinin (41). The ABD of *S. pombe* Ain1 was also shown to form a similar structure (30). The R216E mutation, corresponding to the single point mutation *H. sapiens* α -actinin ACTN4 that causes a configurational change in two CHDs (42), caused the prominent accumulation of Ain1 in the middle region of mitotic cells, possibly by abrogating the interaction between CH1 and CH2 (31; Fig. 3C). However, the steric behaviors of CH1 and CH2 in the ABD of Ain1 have not yet been elucidated. Their interaction with the C-terminal region of the paired molecule may also be important. A role-sharing relationship between CHDs in the ABD may generally be conserved in α -actinin proteins because fungal α -actinin is considered to be a prototype of animal α -actinin proteins (53).

As discussed above, we successfully demonstrated that CH2 plays a critical role in the CR-specific localization of Ain1. CH2 may favor actin structures decorated with CR-specific actin-binding proteins. Previous studies demonstrated that the N-terminal-acetylated Cdc8, the sole tropomyosin homologue in *S. pombe*, specifically localized to the CR (*54*, *55*). The preference of CH2 for F-actin with acetylated Cdc8 warrants further study. In addition, actin-binding proteins may not only collaboratively but also competitively modulate their localization on the actin cytoskeleton. In *S. pombe*, Fim1 has been shown to dominantly localize to actin patches (*24*, *25*). Fim1 and Ain1 both bind to F-actin via their ABDs, and Fim1 has markedly higher affinity to F-actin than Ain1, as described above (*31*, *56*, this study). Therefore, Fim1 may competitively exclude Ain1 from actin patches and confine Ain1 to interact with F-actin in the CR.

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Legends to figures

Fig. 1 Ain1 binds weakly to F-actin

(A) Purified recombinant proteins used in this study. After being subjected to SDS-PAGE, purifies were measured by the densitometry of the CBB-stained gel. (B) High-speed co-sedimentation of full-length Ain1 (Ain1FL) (0.25–8 μ M) with (bottom) or without F-actin (2 μ M) (top). The supernatant (S) and pellet (P) were separated by SDS-PAGE and stained with CBB. (C) Plots of free vs. F-actin-bound proteins (Ain1FL, circle; Ain1FL R216E, triangle; Ain1\DeltaEF, square). Three independent experiments using the high-speed co-sedimentation assay were performed. The average of three experiments is shown with SD (error bar). Each plot was fit to hyperbolic curves. Apparent dissociation constants (K_d) were calculated from each curve. (D) High-speed co-sedimentation of the actin-binding domain of Ain1 (Ain1ABD) (2–24 μ M) with (bottom) or without (top) F-actin (4 μ M). Experiments were performed as in B. Half amounts of the supernatant were applied on lanes with an asterisk.

Fig. 2 Ain1 bundles F-actin independently of Ca²⁺ ions

(A) Fluorescent micrographs of Rhodamine-phalloidin-labeled F-actin incubated with Ain1FL, Ain1ABD, Ain1FL R216E, or Ain1 Δ EF. Note that thick actin bundles were formed in the presence of Ain1FL, Ain1FL R216E, and Ain1 Δ EF. Images were converted into gray-scale and inverted. (B) Low-speed co-sedimentation of F-actin (2 μ M) with Ain1FL or Ain1ABD (0.25–2 μ M). The supernatant (S) and pellet (P) were separated by SDS-PAGE and stained with CBB. Three independent experiments were performed. Control experiment with Ain1 alone was shown in the supplemental data (Fig. S2). (C) Ratios of actin in the pellet fractions estimated from the CBB-stained SDS-PAGE gels in B (mean ± SD, n = 3).

Fig. 3 Cellular localization of Ain1-mutant proteins

(A) Localization of Ain1FL and mutant proteins. Wild-type cells transformed with pREP1-derived vectors containing constructs indicated on the left side were logarithmically grown in EMM with 5

 μ M thiamine at 30°C. The cellular localization of N-terminal YFP-tagged proteins was observed. Mitotic cells displaying the representative protein localization were shown. The same result was obtained using *ain1*-null cells (data not shown). Images were converted into gray-scale and inverted. Asterisks indicate point mutation site(s) including R216E, ABS1-3A, ABS2-1A, and ABS3-2A. Arrows and arrowheads indicate the CR and cortical dots, respectively. The bar is 10 μ m. (**B**) Overexpression of N-terminal-truncated Ain1 lacking CH1. Wild-type cells containing pREP1YFP-ain1 Δ CH1 were incubated as in B without thiamine. Multinucleated cells were frequently observed. (**C**) Ain1FL R216E excessively accumulates in CR. The average intensity of the cytosol and maximum intensity of the CR were measured from a fluorescent micrograph. Data show the ratio of the ring and cytosol (mean \pm SD, n > 30). A two-tailed Student's *t*-test was performed assuming unequal variance.

Fig. 4 Functional analysis of Ain1-mutant proteins in cell growth under stress conditions

(A) Wild-type and *ain1*-null cells were transformed with the pREP1-derived vectors indicated above. Cells were incubated on a YE plate at 30°C for 3 days (right) or YE-containing 1M KCl plate at 18°C for one week (left). The viability of cells was indicated; ++, normal sized colonies; +, small colonies; -, no colony.

Fig. 5 Localization of Ain1 in the its3-mutant

its3-mutant cells containing pREP1-YFP-ain1FL were incubated at 25°C and shifted to the restrictive temperature of 36°C for 3 h. The bar shows 10 μ m.

Fig. 6 Functional relationship of each domain in Ain1

(A) Structural model of the Ain1 ABD (PDBID: 5BVR) (*30*). Two CHDs (right: CH1, left: CH2), three ABSs (black), and four mutation sites are shown. The rotated model (bottom) shows the presumed F-actin binding surface consisting of ABS2 and ABS3. (**B**) Summary of the domain analysis in this study. See the main text for details.



Fig. 1



Fig. 2



Fig. 3



Fig. 4









Supplemental Methods

Construction of strains

C-terminal GFP-tagged ain1 strains (ain1⁺-gfp::kan^r and ain1 ΔEF -gfp::kan^r) were constructed by PCR-based homologous recombination (1). The following PCR primer pairs were used to amplify 5'-GAAGCATTGCAAGGATGCAGGAGTACTGGC-3' (F1) the F-fragments; and 5'-TTAATTAACCCGGGGGATCCGAACTATTTCTTTGTCTTCGG-3' (R2) for ain1+-gfp::kan', and F1 and 5'-TTAATTAACCCGGGGGATCCGTTTAGTAATTCCGTCTAGTTC-3' (R3) for $ain1\Delta EF$ -gfp::kan^r. The R-fragment amplified was by 5'-GTTTAAACGAGCTCGAATTCAACGGAGCATTTTGAATCAC-3' (F2) and 5'-GTTATGCTATGCGAGCCATGTATATTCTTG-3' (R1). F- and R-fragments were both fused by PCR with SalI-digested pFA6a-GFP(S65T)-KanMX6. S. pombe cells were transformed with amplified constructs using the lithium acetate method. YE medium containing 100 μ g/mL G418 (Sigma-Aldrich) was used for the selection of drug-resistant transformants.

Western blot

Log-phase incubated cells were harvested and extracted by the NaOH treatment method (2, 3). Protein extracts were subjected to 10% SDS-PAGE. Gels were transferred to PVDF membranes and probed with a mouse anti-GFP antibody (Roche). Horseradish peroxidase (HRP)-conjugated goat anti-mouse IgG (Santa Cruz Biotechnology) was used as a secondary antibody. Membranes were developed using ECL reagents (GE).

Supplemental data

Fig. S1 Artificially synthesized ain1⁺ gene

The nucleotide sequence of the open reading frame of an artificially synthesized $ain l^+$ gene is shown. Several triplets are replaced with another synonymous codon for stronger expression Ε. according to the codon usage table of S. pombe (4)and coli (http://www.kazusa.or.jp/codon/cgi-bin/showcodon.cgi?species=83333).

Fig. S2 Control experiment of the low-speed co-sedimentation assay

Ain1FL (0.5–8 μ M) alone was centrifuged at low-speed as in Fig. 2B. The supernatant (S) and pellet (P) were separated by SDS-PAGE and stained with CBB (A). Amounts of Ain1 FL in the supernatant versus pellet were plotted (B). About 16% of Ain1 FL was detected in the pellet fractions probably because of their adhesion to the internal walls of centrifuge tubes.

Fig. S3 Co-localization of an Ain1-mutated protein with actin

Cells expressing Lifeact-4×mCherry were transformed with vectors for the expression of the proteins indicated above and observed as in Figure 3A. Images of YFP-Ain1 (green) and Lifeact-4×mCherry (red) were merged. Asterisks indicate a mutation site. Arrows indicate the CR, and arrowheads indicate cortical dots.

Fig. S4 Overexpression of the CH1 of Ain1

Lifeact-4×mCherry-expressing cells that were transformed with pREP1YFP-ain1FL or -ain1CH1 were incubated at 30°C for 18 h after removing thiamine from the cell culture to induce the overproduction of the protein. Cells treated with 1% DMSO with or without 8 μ M Latrunculin-A (Lat-A) were observed. Note that abnormal cables decorated with Ain1 CH1 and Lifeact-4×mCherry remained, even in the presence of Lat-A. The bar shows 10 μ m.

Fig. S5 Expression of fluorescent protein-tagged Ain1AEF

(A, C) Western blot of N-terminal YFP- or C-terminal GFP-tagged-Ain1. Arrowheads indicate tagged Ain1FL (98 kDa) and tagged Ain1 Δ EF (83 kDa). (B) Localization of Ain1FL-GFP or Ain1 Δ EF-GFP in mitotic cells. The bar shows 10 μ m.

Table S1. Plasmids used in this study

plasmid	gene product (a. a.)	comment								
pREP1		empty vector								
pREP1YFP-ain1FL	1–621	Full-length protein								
pREP1YFP-ain1ABD	1–235	actin-binding domain only								
pREP1YFP-ain1CH1	1–122	CH1 domain only								
pREP1YFP-ain1CH2	123–235	CH2 domain with an extra 5 residues GGRSR in the N terminus								
pREP1YFP-ain1 <u>A</u> CH1	123–621	Ain1 Δ CH1 with GGRSR in the N terminus								
pREP1YFP-ain1FL R216E	1–621, R216E	Arg216 was replaced with Glu								
pREP1YFP-ain1ABD R216E	1–235, R216E	actin-binding domain only								
		Arg216 was replaced with Glu								
pREP1YFP-ain1 \DEF	1–490	C-terminal region containing the EF-hand was removed								
pREP1YFP-ain1EF	491–621	C-terminal region containing the EF-hand with GGRSR in the								
		terminus								
pREP1YFP-ain1 \DABD	236–621	Ain1 Δ ABD with GGRSR in the N terminus								
pREP1YFP-ain1 ΔEF R216E	1–490, R216E	The EF-hand was removed								
		Arg216 was replaced with Glu								
pREP1YFP-ain1FL ABS1-2A	1–621,	Q10 and T13 were replaced with Ala								
	Q10A, T13A									
pREP1YFP-ain1CH1 ABS1-2A	1–122,	CH1 domain only								
	Q10A, T13A	Q10 and T13 were replaced with Ala								
pREP1YFP-ain1FL R216E ABS1-2A	1–621, R216E,	Q10 and T13 were replaced with Ala								
	Q10A, T13A	Arg216 was replaced with Glu								
pREP1YFP-ain1FL ABS2-1A	1–621, K100A	K100 was replaced with Ala								
pREP1YFP-ain1CH1 ABS2-1A	1–122, K100A	CH1 domain only								
		K100 was replaced with Ala								
pREP1YFP-ain1FL R216E ABS2-1A	1–621, R216E, K100A	K100 was replaced with Ala								
		Arg216 was replaced with Glu								
pREP1YFP-ain1FL ABS3-2A	1–621, R135A, K136A	R135, and K136 were replaced with Ala								
pREP1YFP-ain1 ACH1 ABS3-2A	123–621, R135A, K136A	Ain1 Δ CH1 with GGRSR in the N terminus								
		R135 and K136 were replaced with Ala								

Sp ain1 Sp ain1 artificial	a a a		cat	ATG													TTT	ACA			51 60
Sp ain1artificial	TTC	AAC	ACA	AAA	СТТ	TCA	TCG	AGA	GAC	TTA	CCA	тст	GTG	TTT	GAC	TTG	AGA	AAG	GAT	TTA	111
Sp ain1_artificial	TTT	AAT	ACT	AAG	СТТ	AGT	AGC	AGA	GAC	СТТ	ССТ	AGT	GTT	TTT	GAC	СТС	CGT	AAG	GAT	СТТ	120
Sp ain1 Sp ain1_artificial		GAT	GGT	ATT	CTT	CTA	ATC	CAA	СТС	CTG	GAA GAA	ATA	ATC	GGA GGA	GAT	GAA		CTT	GGG		171 180
Sp ain1 Sp ain1_artificial	T A C T A T	AA <mark>T</mark> AA <mark>C</mark>	C G A C G G	AA <mark>C</mark> AA <mark>T</mark>	C C G C C T	CGT CGT	ATG ATG	C G T C G A	GTT GTT		AGG CGT	TTG CTT	G <mark>A</mark> G G <mark>A</mark> A	A A T A A T	GTC GTA	AAC AAC		GCT GCT	T T A C T G	GAA GAG	231 240
Sp ain1 Sp ain1 artificial		ATA					ATG	CCG	TT G CT T	ACC	AAC	ATT	GG T GG C		GCT		ATT	GT G GT C	GAT GAT		291 300
Sp ain1 Sp ain1 artificial		CTG	AAG AAA	CTA	ATT	CTG	GG A			TGG		СТС			CGT		ACT		GCT GCT		351 360
Sp ain1 Sp ain1 artificial	ATC			GAA GAA	GGT	CTT	ACC	GCA	AAG	GAA	GGT		TTG	CTT	TGG	TGT				ACA	411
Sp ain1_artificial	GCA				CCA	GAG	GTT	GAT	GTG		GAT	TTT	ACT	CGG	AGT	TGG			GGC	TTG	471
Sp ain1_artificial	GCT	TTC	TGT	GCA	TTA					CGT	CCT	GAT	TTA	CTG	GAT			AAG		GAC	531 540
Sp ain1_artificial	AAG	AAG		CAT	AGA	GCT		ATG	CAA	TTG	GCT	TTT	GAC	ATT	GCG	CAA	AAA	TCT	ATT	GGA	591
Sp ain1_artificial		ССТ	CGT	TTG	ATA	GAG	GTC	GAA	GAT	GTT	TGT	GAC	GTT	GAC	GGA	CCC	GAT	GAG	CGT	AGT	651 660
Sp ain1_artificial		ATG	ACT		GTC	GCC	GAA	TAC	TTC		GCG	TTT	TCT	ACC	TTG	GAT		GTG	GAA	ACT	711
Sp ain1_artificial	GCG	GCT		CGT	GTA	GAG		TTT		GAA	GTT	TTG		AGC	ACT		GAT			ATT	771
Sp ain1 Sp ain1 artificial	GAC			TCT	CGA	ATG		CGA	CTG	TTG	GGA		ATT	GCA				GAG			831 840
Sp ain1 Sp ain1_artificial			G T C G T T		TTT					ACC	GAC GAT	GT G GT T		TCT		TCT			TTT	GCC	891 900
Sp ain1 Sp ain1_artificial				GCT GCT		GAG GAA		AGA AGA		TGG	GTA GTT	AAG AAG	GAA GAA			GAC GAT	TTG	GAA GAG	T C T A G T		951 960
Sp ain1 Sp ain1_artificial	TTG CTG	GGA GGT				ACC	AA T AA T	T T G C T T					СТТ		AAG AAG		GAA GAA	CCG CCT		GCG GCT	1011 1020
Sp ain1 Sp ain1_artificial	GG A GG T	TTA	AAG AAA	A T C A T T	GTC GTC	GAT GAT	TTG CTT	GAA GAA	A G A C G T	CAG CAA	TGG TGG	AAA AAG	G <mark>A T</mark> G <mark>A C</mark>	TTT TTT	C T T C T T	T CA Ag T	GAA GAA	GAA GAG	G C C G C A	AAT AAT	1071 1080
Sp ain1 Sp ain1_artificial	CAA CAA	T C G A G C	AAA AAA	T T G C T C	A T T A T T	AAT AAT	ACC ACT		ATG ATG	AGA CGT	GAA GAA	ATA Atc	AAA AAA	GAA GAG	AGT AGC	ATG ATG	A G A C G C	A T T A T A	GCT GCA	TTT TT C	1131 1140
Sp ain1 Sp ain1_artificial	GCT GCT	GAT GAT	CGT CGT	G C C C	AAC AAC	AGT AGT	TTC TTT	T C A A G T	AAA AAA	ATG ATG	T T A	TCT AGC	ACG ACT	ATA ATC	T C A A G T	AA T AA C	G A G G A A	A T T A T T	ACC ACT	AAT AAC	1191 1200
Sp ain1 Sp ain1_artificial	T T A C T T	C A G C A A	GGT GGT	GAT GAT	T G G T G G	C G T A G A	G <mark>A</mark> C G <mark>A</mark> C	CAA CAA	C T C C T T	GAC GAT	C A T C A C	GTT GTT	GAA GAA	TTT TTT	T T G C T C	CAA CAA	GAA GAA	CAC CAT	СТ <mark>С</mark> СТТ	GGG GGT	1251 1260
Sp ain1 Sp ain1_artificial	ССТ ССТ	CTT	GAA GAA	GTT GTT	GAA GAA	СТ <mark>С</mark> СТТ	GCT GCT	AGT AGT	GT G GT C	AAG AAA	GT G GT T	TTG CTA	T A T T A T	GAT GAT	AAC AAC	TGT TGT	TTT TTT	CAA CAA	GCT GCT	GG <mark>A</mark> GGT	1311 1320
Sp ain1 Sp ain1_artificial	ATT	GAA GAG	GAA GAA		GAT GAC	TAC TAC	ACC ACT	ATG ATG	TTT TTC		T A C T A T	GAA GAA	GAC GAT	CTG CTT	G A G G A G	CAT CAT	GAA GAA	TTT TTT	GG T GG A	ATT	1371 1380
Sp ain1 Sp ain1_artificial	ACT	GCC GCT	AAT AAT	ATC	A T T A T A	GCC	AA T AA T	AAG AAG	ATC	AAG AAA	T A C T A T	CTC	GAG GAA	AAT AAC	GAA GAA	C T C	TT A CTT	GAA GAA	AGG CGT	GAA GAG	1431 1440
Sp ain1 Sp ain1_artificial	AAG AAA	A G A C G A	ACA ACT	C T C C T T	T C C A G T	AAA AAG	CAA CAA	GAA GAA	CTC	GAC GAT	GGA GGT	ATT ATT	ACT ACT	AAA AAA	GT C GT T	TTT TTT	C G C C G T	CAT CAT	TTT TT <mark>C</mark>	GAA GAA	1491 1500
Sp ain1 Sp ain1_artificial	AAG AAA	AAA AAG		T CA AG C	AAT AAT	ATG ATG	T T A C T T	AAT AAC	GAG GAA	GTC GTC	GAG GAA	TTT TTT	T A T T A T	GCC GCT	G C A G C C	T T G C T C	GCC GCA	TCA AGT	TTG CTT	GG <mark>C</mark> GGT	1551 1560
Sp ain1 Sp ain1_artificial	TTG CTC	GTT		GAT GAT	ACC	GAA GAA	GAA GAG	GGA GGT	ACT	GCT GCT	TTG CTT	TTT TTT		AGG CGT	GCT GCC	GCT GCT		TCG AGT	GAA GAG	GAG GAA	1611 1620
Sp ain1 Sp ain1_artificial	GG T GG T	GTT GTT	ACT		GAG GAA		TTC	ACA ACT	GAA GAA	ATC	GT G GT T	ATG ATG	GAA GAA	GAA GAG	CT C	GAG GAA	GAT GAT	CGT CGT	GAT GAC	AGT AGT	1671 1680
Sp ain1 Sp ain1_artificial	GCT GCT	CGT		GTT GTT	СТТ		GCG GCT	TTT	TGT TGT	GA T GA T	GTT GTA	GCA GCT	GAC GAT	GGA GG T		TCT AGT	TAT	GT C GT T	ACG ACT		1731 1740
Sp ain1 Sp ain1_artificial	GAT		СТА	CTC		AGT		GTT GTT	CGT				GTA		TTC	CTC	GAA GAA	TGC TGT		ATG ATG	1791 1800
Sp ain1 Sp ain1_artificial		AAA AAG		T CA AG T	GAA GAA	GGA GG <mark>T</mark>	CTG	GAT GAT		CTT	ACA ACT	TGG TGG	ATT	AAA AAA		T T A C T T	C T C	GCC GCT	GAA GAA	GAC GAC	1851 1860
Sp ain1 Sp ain1_artificial	AAA AAA	G A A G A G	A T A A T T	GTT GTT	TAA tag	t <mark>c g</mark>	1 ac 1	866 880													

Fig. S1



Fig. S2



Fig. S3



Fig. S4



Fig. S5