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Novel strategy for anchorage position control of GPI-attached proteins in the yeast cell wall using different GPI-anchoring domains

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Abstract

The yeast cell surface provides space to display functional proteins. Heterologous proteins can be covalently anchored to the yeast cell wall by fusing them with the anchoring domain of glycosylphosphatidylinositol (GPI)-anchored cell wall proteins (GPI-CWPs). In the yeast cell-surface display system, the anchorage position of the target protein in the cell wall is an important factor that maximizes the capabilities of engineered yeast cells because the yeast cell wall consists of a 100- to 200-nm-thick microfibrillar array of glucan chains. However, knowledge is limited regarding the anchorage position of GPI-attached proteins in the yeast cell wall. Here, we report a comparative study on the effect of GPI-anchoring domain-heterologous protein fusions on yeast cell wall localization. GPI-anchoring domains derived from well-characterized GPI-CWPs, namely Sed1p and Sag1p, were used for the cell-surface display of heterologous proteins in the yeast Saccharomyces cerevisiae. Immunoelectron-microscopic analysis of enhanced green fluorescent protein (eGFP)-displaying cells revealed that the anchorage position of the GPI-attached protein in the cell wall could be controlled by changing the fused anchoring domain. eGFP fused with the Sed1-anchoring domain predominantly localized to the external surface of the cell wall, whereas the anchorage position of eGFP fused with the Sag1anchoring domain was mainly inside the cell wall. We also demonstrate the application of the anchorage position control technique to improve the cellulolytic ability of cellulase-displaying yeast. The ethanol titer during the simultaneous saccharification and fermentation of hydrothermally-processed rice straw was improved by 30% after repositioning the exo- and endo-cellulases using Sed1- and Sag1-anchor domains. This novel anchorage position control strategy will enable the efficient utilization of the cell wall space in various fields of yeast cell-surface display technology.

Keywords	Saccharomyces cerevisiae; yeast surface display; glycosylphosphatidylinositol anchored cell wall protein; anchorage position; Sed1p; Sag1p	
Taxonomy	Cellulase, Saccharomyces cerevisiae, Applied Microbiology, Fungal Cell Wall, Cell Surface Protein	
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October 15, 2019

Dear Professor Jay Keasling,

MBE_2019_253: Kentaro Inokuma, Hiroki Kurono, Riaan den Haan, Willem Heber van Zyl, Tomohisa Hasunuma, and Akihiko Kondo; Novel strategy for anchorage position control of GPI-attached proteins in the yeast cell wall using different GPI-anchoring domains.

Thank you very much for your mail dated on September 25, 2019 together with the comments of reviewers. We have studied their comments carefully, conducted an additional experiment, and have made necessary corrections.

The following changes are our response to the comments of reviewers. The text has been revised to accommodate the comments. These changes were highlighted using red fonts in the revised manuscript.

Sincerely, Akihiko Kondo Graduate School of Science, Technology and Innovation, Kobe University 1-1 Rokkodai-cho, Nada-ku, Kobe 657-8501, Japan Telephone: +81-78-803-6196, Fax: +81-78-803-6196 E-mail: akondo@kobe-u.ac.jp In response to the comments of reviewer #1 Comments

(1) Although the authors claimed significant increase in ethanol titer, I have different opinion from the authors. The increase should be 30%, not 1.3-fold. The ethanol titer using the engineering strain IS 1.3-fold of that of the control, not increased 1.3-fold.

Author's response: Thank you for pointing that out. "1.3-fold" was corrected to "30%" (line 44 in the revised manuscript).

(2) My major concern is that the results in this study only focused on two enzymes and two anchoring domains, it will be nice if the authors provide more solid data on the mechanisms or on more applications using their anchoring domain control technology. More in-depth discussion should be added on why such results were obtained, can the authors expand their study to other cell wall proteins as the source of anchoring domains? How can other researchers benefit from this study when they used other enzymes?

Author's response: We appreciate your valuable suggestion and accordingly the following sentences were added to the Discussion section (lines 362-375 and 429-435 in the revised manuscript).

"The results presented in this study suggest that yeast cells recognize GPIanchoring domains attached to target proteins and control their anchorage positions in the cell wall. Although the anchorage mechanism of yeast GPI-CWPs liberated from the plasma membrane to the cell wall remains unclear, recent studies have suggested that plasma membrane-anchored GPI proteins Dfg5p and Dcw1p are potential candidates for cross-linking the GPI-anchor remnant and cell wall β -(1 to 6) glucan (Gonzalez et al., 2010; Orlean, 2012). These proteins are putative glycosidase/transglycosidases homologous to bacterial family 75 (Cantarel et al., 2009) and depletion of these enzymes by repressing their expression in the double-null background led to secretion of a GPI-CWP into the medium (Kitagaki et al., 2002). These enzymes might recognize differences in GPI anchoring domains and be involved in controlling the anchorage position of GPI-attached proteins. Further analysis using GPI-anchoring domains derived from other GPI-CWPs are urgently required to identify the determinants of the anchorage position of GPI-attached proteins. On the other hand, in order to expand this research to a wide range of GPI-CWPs, it will be necessary to develop a novel method for high-throughput anchorage position analysis."

"The anchorage position control technique demonstrated in this study will also benefit applications of yeast cell-surface display other than the construction of cellulasedisplaying yeast. The hydrolysis efficiency of other plant-derived polysaccharides such as hemicellulase and starch may be improved by this technique because the complete hydrolysis of these polysaccharides also requires the cooperation of endo- and exo-type enzymes. In addition, the Sed1-anchoring domain that can expose the target protein to the external surface of the cell wall will also be a potential anchor candidate for protein screening requiring contact with large ligands."

In accordance with these additions, new references were added to the reference list (lines 473-475, 493-495, and 525-528 in the revised manuscript).

(3) How many cells have the authors observed for cell wall localization studies?

Author's response: In the immunoelectron-microscopic analysis, we observed 12 cells for each of strains BY-eGFP-SSS and BY-eGFP-SSA, and confirmed that the number and localization tendency of eGFP in the cell wall were clearly different between these strains. To make it clearer, the following sentence was added to section 3.2 (lines 249-250 in the revised manuscript).

"We observed 12 cells for each of strains BY-eGFP-SSS and BY-eGFP-SSA."

(4) How did the author evaluate the ethanol fermentation results with the previous studies? Can the authors integrate their results with their previous ones displaying more cellulosic enzymes?

Author's response: As the reviewer mentioned, we previously reported simultaneous saccharification and fermentation from pretreated rice straw using a recombinant yeast strain, in which BGL1, EGII, CBH1, and CBH2 were co-displayed using the Sed1-anchoring domain (Liu et al., 2016). Although it is not possible to fully integrate our

result (Fig. 4) with the result in the previous study due to differences in fermentation scale and agitation procedure, the BGL, EG, and CBHs co-displaying strain achieved approximately 9.5 g/L of ethanol production after 96 h fermentation with 0.2 FPU/g biomass of commercial cellulase cocktail (Liu et al., 2016). This ethanol titer is higher than that of BY-ESBA strain with 0.4 FPU/g biomass of commercial cellulase cocktail shown in Fig. 4 (7.3 g/L at 96 h). These results suggest the importance of co-display of BGL, EG, and CBHs for efficient hydrolysis of lignocellulosic biomass.

Some descriptions to contextualize this result with those we reported previously were added to the Discussion section (lines 416-428 in the revised manuscript).

(5) Line 239-240, more fluorescence was observed in the intracellular vacuoles of strain BY-eGFP-SSA than in those of strain BY-eGFP-SSS, please confirm, is the word "vacuoles" correct?

Author's response: Thank you for pointing that out. We checked several published papers and found that the word "vacuoles" was commonly used as the plural form of vacuole. As shown in Fig. 2A, we observed multiple cells containing a vacuole. Therefore, we would like to keep the word "vacuoles" in this sentence.

(6) Fig. S1, the C-terminal GPI attachment site (the ω site) marked in bold...here the sentence should be ...was marked in bold.

Author's response: "marked in bold" was corrected to "was marked in bold" according to the suggestion (line 649 in the revised manuscript and page 6 line 2 in the revised Supplementary materials).

In response to the comments of reviewer #2 Comments

(1) As EG II is endoglucanase, it will have limited action on cellulose. Both CBH and EG are required for efficient hydrolysis of insoluble cellulose. Since only EG II has been used in this work, it will produce cellodextrins which may be poorly hydrolysed by beta-G. This may be the reason for low yield of ethanol by BY-ESBA strain in SSF. The authors may like to mention this point in

Discussion.

Author's response: We appreciate this valuable suggestion. The following sentences were subsequently added to the Discussion section (lines 409-428 in the revised manuscript).

"In this study, we used EG and BGL co-displaying strains for the simultaneous saccharification and fermentation of pretreated rice straw. It has been demonstrated that synergistic cooperation of EG and cellobiohydrolases (CBHs) is essential for efficient degradation of insoluble cellulose (Jalak et al. 2012). CBHs are chain end-specific processive exo-glucanases. EG randomly hydrolyzes amorphous regions of insoluble cellulose and generates reducing and non-reducing ends that can be attacked by CBHs, while CBHs recognize the cellulose chain ends and continuously hydrolyze crystalline regions between the amorphous parts into cellobiose units (Jalak et al. 2012). Previously, we reported a simultaneous saccharification and fermentation from pretreated rice straw using a recombinant yeast strain, in which BGL1, EGII, and CBHs (CBH1 and CBH2) were displayed using the Sed1-anchoring domain (Liu et al., 2016). Although direct comparison with the result shown in Fig. 4 is not possible due to differences in fermentation scale and agitation procedure, the BGL, EG, and CBHs codisplaying strain achieved approximately 9.5 g/L of ethanol production after 96 h fermentation with 0.2 FPU/g biomass of commercial cellulase cocktail (Liu et al., 2016). This ethanol titer is higher than that of BY-ESBS strain with 0.4 FPU/g biomass of commercial cellulase cocktail (7.3 g/L at 96 h, Fig. 4). These results also suggest the importance of co-display of BGL, EG, and CBHs for efficient hydrolysis of insoluble cellulosic materials. Additional display of CBHs on the cell surface of BY-ESBA strain will be required for further improvement of its ethanol yield from pretreated biomass. Furthermore, it will be necessary to verify the optimal anchorage position for CBHs in the cell wall to maximize synergies between cellulases."

In accordance with this change, a new reference was added to the reference list (lines 523-524 in the revised manuscript).

(2) Line 217: Did the yeast grow at 38C or 30C?

Author's response: First of all, we apologize for the incorrect description of

fermentation temperature. We mistakenly stated that the fermentation temperature was 38 °C, but it was carried out at 37 °C. We have already corrected this error (lines 217 and 296 in the revised manuscript).

In this study, cultivation to obtain yeast cells was performed at 30 °C, while simultaneous saccharification and fermentation using the obtained cells was performed at 37 °C to promote the activity of cellulases displayed on the cell surface (optimal temperature of *Aspergillus aculeatus* BGL1 and *Trichoderma reesei* EGII is 65 °C). To make it clearer, the following sentence was added to the section 3.5" (lines 296-298 in the revised manuscript).

"The fermentation was performed at 37 °C to promote the activity of cellulases displayed on the cell surface because the optimal temperature of *A. aculeatus* BGL1 and *T. reesei* EGII is 65 °C (Decker et al., 2000; Trudeau et al., 2014)."

In accordance with this change, new references were added to the reference list (lines 478-480 and 586-588 in the revised manuscript).

(3) Fig 4: Although BY-ESBA strain has been shown to be 1.3 fold better than BY-ESBS strain for ethanol production in SSF process, it would also be useful to compare the effectiveness of both BY-ESBA and BY-ESBS strains with that of control yeast strain not having any surface-displayed enzyme (but containing 0.4 FPU/g biomass). This may be suitably incorporated in M&M, results and discussion sections.

Author's response: We appreciate your valuable suggestion. According to the suggestion, we performed the SSF process from the pretreated rice straw with a control yeast strain BY4741, which is the parental strain of BY-ESBA and BY-ESBS strains, and 0.4 FPU/g biomass of commercial cellulase cocktail. As expected, the control strain showed lower ethanol production from the pretreated biomass than those of both BY-ESBA and BY-ESBS strains.

The fermentation result with BY4741 was included in Fig. 4 in the revised manuscript and some descriptions were added to section 3.5 (lines 294 and 298-301) and Figure legends (line 634), respectively.

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26 Abstract

27 The yeast cell surface provides space to display functional proteins. Heterologous proteins can 28 be covalently anchored to the yeast cell wall by fusing them with the anchoring domain of 29 glycosylphosphatidylinositol (GPI)-anchored cell wall proteins (GPI-CWPs). In the yeast cell-30 surface display system, the anchorage position of the target protein in the cell wall is an important 31 factor that maximizes the capabilities of engineered yeast cells because the yeast cell wall consists 32 of a 100- to 200-nm-thick microfibrillar array of glucan chains. However, knowledge is limited 33 regarding the anchorage position of GPI-attached proteins in the yeast cell wall. Here, we report a 34 comparative study on the effect of GPI-anchoring domain-heterologous protein fusions on yeast 35 cell wall localization. GPI-anchoring domains derived from well-characterized GPI-CWPs, namely 36 Sed1p and Sag1p, were used for the cell-surface display of heterologous proteins in the yeast 37 Saccharomyces cerevisiae. Immunoelectron-microscopic analysis of enhanced green fluorescent 38 protein (eGFP)-displaying cells revealed that the anchorage position of the GPI-attached protein in 39 the cell wall could be controlled by changing the fused anchoring domain. eGFP fused with the 40 Sed1-anchoring domain predominantly localized to the external surface of the cell wall, whereas 41 the anchorage position of eGFP fused with the Sag1-anchoring domain was mainly inside the cell 42 wall. We also demonstrate the application of the anchorage position control technique to improve 43 the cellulolytic ability of cellulase-displaying yeast. The ethanol titer during the simultaneous 44 saccharification and fermentation of hydrothermally-processed rice straw was improved by 30% 45 after repositioning the exo- and endo-cellulases using Sed1- and Sag1-anchor domains. This novel 46 anchorage position control strategy will enable the efficient utilization of the cell wall space in 47 various fields of yeast cell-surface display technology.

48

49 Keywords: Saccharomyces cerevisiae, yeast surface display, glycosylphosphatidylinositol-

50 anchored cell wall protein, anchorage position, Sed1p, Sag1p

52	Abbreviations: BGL, β -glucosidase; BSA, bovine serum albumin; EG, endoglucanase; eGFP,
53	enhanced green fluorescent protein; ER, endoplasmic reticulum; FPU, filter paper unit; GPCR, G
54	protein-coupled receptor; GPI, glycosylphosphatidylinositol; GPI-CWP, GPI-anchored cell wall
55	protein; GRAS, generally regarded as safe; nano-UPLC-MS ^E , nanoscale ultra-pressure liquid
56	chromatography electrospray ionization quadrupole time-of-flight tandem mass spectrometry; pAb,
57	polyclonal antibody; pNPG, p-nitrophenyl-β-D-glucopyranoside; YP, yeast extract peptone

59 1. Introduction

The expression of functional proteins on the cell surface is a promising approach to construct cell-surface-engineered microorganisms with special functions. Cell-surface display technology can be used to address a wide range of applications such as the engineering and screening of enzymes, antibodies, or peptides (Angelini et al., 2015; Grzeschik et al., 2017; Li et al., 2007), the bioadsorption of specific molecules (Shibasaki and Ueda, 2014), and the production whole cell catalysts for bioconversion (Inokuma et al., 2018; Liu et al., 2016), biodegradation (Richins et al., 1997; Shibasaki et al., 2009), and biosensing (Tang et al., 2014; Wang et al., 2013).

67 Among host microorganisms employed for cell-surface display, baker's yeast (Saccharomyces 68 *cerevisiae*) is the most frequently used to develop such systems because of the vast knowledge of 69 its genetics, physiology, and fermentation characteristics, as well as its generally regarded as safe 70 (GRAS) status. In yeast cell-surface display, the glycosylphosphatidylinositol (GPI)-anchoring 71 system is the typical and most widely-used technique to immobilize heterologous proteins. In this 72 method, a yeast cell is transformed by introducing fusion genes coding proteins of interest and the 73 anchoring domain of the GPI-anchored cell wall protein (GPI-CWP). In the recombinant yeast cells, 74 the fused proteins are synthesized on endoplasmic reticulum (ER)-bound ribosomes, cleaved at the 75 C-terminal GPI attachment site (the ω site), and modified by the addition of a pre-assembled GPI 76 anchor in the ER. Subsequently, the GPI-attached proteins leave the ER in COPII-coated vesicles 77 and travel via the Golgi to the plasma membrane (Doering and Schekman, 1996). Finally, these 78 proteins are liberated from the plasma membrane and become immobilized in the cell wall through 79 covalent linkage to a β -(1 to 6) glucan via a remnant of the anchor structure (Klis et al., 1997; Lu et 80 al., 1994).

81 It has been reported that the anchoring domains from different GPI-CWPs exhibit different 82 efficiencies for the cell-surface display of target enzymes (Andreu and Del Olmo, 2018; Hamada et 83 al., 1999). Therefore, selection of the appropriate anchoring domain for fusion with target proteins

84 is important for efficient cell-surface display. In previous studies, indeed, the activities of some 85 cellulolytic enzymes fused with the Sed1-anchoring domain were found to be higher than those 86 fused with the α-agglutinin (Sag1)-anchoring domain (Inokuma et al., 2014). However, the degree 87 to which activity was improved by changing the anchoring domain varied greatly depending on the 88 enzymes displayed. For the cell-surface display of *Aspergillus aculeatus* β-glucosidase 1 (BGL1), 89 the activity of the enzyme fused with the Sed1-anchoring domain was approximately 2-fold higher 90 than that upon fusion with the Sag1-anchoring domain. In contrast, the hydrolytic activity of 91 Trichoderma reesei endoglucanase II (EGII) for water-insoluble cellulose was improved 60-fold 92 when using the Sed1-anchoring domain compared to that with the Sag1-anchoring domain 93 (Inokuma et al., 2014). Based on these results, we hypothesized that selection of the anchoring 94 domain would affect not only the display efficiency of the target protein but also its localization in 95 the cell wall.

96 The yeast cell wall is composed of a microfibrillar array of β -(1 to 3) glucan and β -(1 to 6) 97 glucan chains with a thickness of 100 to 200 nm (Dupres et al., 2010). Therefore, cell wall proteins 98 exposed to the external surface represent only a portion of the whole and the remainder are buried in the glucan layer (Van der Vaart et al., 1997). Small substrates such as cellobiose and p-99 100 nitrophenyl-B-D-glucopyranoside (pNPG) are accessible to all integrated enzymes because these 101 substrates penetrate the cell wall. In contrast, large substrates such as water-insoluble cellulose can 102 only access enzymes exposed on the external surface. However, to our knowledge, no comparative 103 analysis has been reported concerning the effect of the anchoring domain on target protein 104 localization in the yeast cell wall.

In the present study, we performed a comparative analysis of the effect of different anchoring domains on the cell wall localization of fused heterologous proteins in *S. cerevisiae*. First, the intracellular localization of enhanced green fluorescent protein (eGFP) fused with Sed1 or Sag1anchoring domains was analyzed using a confocal fluorescence microscope. Subsequently, immunoelectron-microscopic analysis of ultra-thin sections of the eGFP-displaying yeast cells was carried out to investigate the effect of the anchoring domains on the anchorage position of GPIattached proteins in the cell wall. Finally, by applying the information obtained from this novel system, we successfully demonstrate improved ethanol production from pretreated lignocellulosic biomass by cellulase-displaying yeast after controlling the anchorage position of exo- and endocellulases using different anchoring domains.

115

116 **2. Materials and Methods**

117 **2.1. Strains and media**

Escherichia coli strain DH5α (Toyobo, Osaka, Japan) was used as the host for recombinant
DNA manipulation. *E. coli* medium was prepared as described (Inokuma et al., 2016). The genetic
properties of all yeast strains used in this study are shown in Table 1. The gene cassettes for the
cell-surface display of heterologous proteins were expressed in the haploid yeast strain *S. cerevisiae*BY4741 (Life Technologies, Carlsbad, CA, USA).

The *S. cerevisiae* transformants were screened and cultivated as previously described (Inokuma et al., 2016). After 48 h of cultivation, yeast cells were harvested by centrifugation at $1000 \times g$ for 5 min, washed twice with distilled water, and again centrifuged at $1000 \times g$ for 5 min. The wet cell weight of the washed yeast cells was determined by weighing the cell pellet. The estimated dry cell weight of a yeast cell is approximately $0.15 \times$ its wet cell weight (Inokuma et al., 2014). Cell pellets were used for microscopic observation, immunoelectron-microscopy, enzyme assays, and ethanol fermentation.

130

131 **2.2. Plasmid construction and yeast transformation**

The plasmids and primers used in this study are listed in Supplementary Tables S1 and S2,
respectively. The integrative plasmids for the expression of eGFP, *T. reesei* EGII, and *A. aculeatus*

BGL1 were transformed into *S. cerevisiae* by the lithium acetate method (Chen et al., 1992) and integrated into the *HIS3* locus or the 3' noncoding region of YFL021W and YFL020C genes (I2 region) of the chromosomal DNA by homologous recombination. Details on the construction of plasmids and yeast transformation have been provided as Supplementary Text S1.

138

139 **2.3. Fluorescence microscopy**

140 Cell pellets of eGFP-expressing yeast strains were resuspended in 15 mM FM4-64 (Invitrogen 141 Carlsbad, CA, USA) diluted in culture medium and incubated for 15 min at 150 rpm and 30 °C in 142 the dark to stain vacuolar membranes. The cells were washed and resuspended in culture medium 143 followed by a further 2-h incubation at 150 rpm and 30 °C in the dark. After washing twice with 144 distilled water, the cells were observed using a confocal fluorescence microscope BZ-X810 145 (Keyence, Osaka, Japan) with a Nikon Plan Apo λ 100x/1.45 oil-immersion objective lens (Nikon, 146 Tokyo, Japan) and appropriate filters for eGFP and FM4-64.

147

148 **2.4. Sample preparation for immunoelectron-microscopy**

149 Washed cell pellets were sandwiched between two copper disks and frozen in liquid propane at -175 °C. The frozen samples were freeze-substituted with acetone containing 0.2% glutaraldehyde 150 151 and 2% distilled water at -80 °C for 2 days. The substituted samples were then transferred to -20 °C for 3 h and then warmed to 4 °C over 90 min. Next, they were dehydrated in anhydrous 152 153 acetone and anhydrous ethanol at 4 °C. Infiltration was performed with LR white resin (London 154 Resin Co. Ltd., Berkshire, UK) at 4 °C [ethanol:resin 50:50 for 2 h; 100% resin for 30 min; 100% resin for 30 min]. The samples were then transferred to a fresh 100% resin for embedding and the 155 resins were polymerized at 50 °C overnight. The polymerized resins were cut into ultrathin sections 156 157 of 90 nm thickness using an ultramicrotome (Ultracut CUT; Leica, Vienna, Austria) and the 158 sections were placed on nickel grids.

159

160 **2.5. Immunostaining**

161 Ultrathin sections were incubated with the primary antibody [rabbit anti-GFP polyclonal 162 antibody (pAb)] in blocking solution [PBS containing 1% bovine serum albumin (BSA)] at 4 °C 163 overnight and washed three times with the blocking solution. Subsequently, they were incubated 164 with secondary antibody conjugated to 10-nm gold particles (goat anti-rabbit IgG pAb; BBI 165 Solutions, Cardiff, UK) at room temperature for 90 min and washed with PBS. The sections in the nickel grids were placed in 2% glutaraldehyde in 0.1 M phosphate buffer (pH 7.4). After the grids 166 167 were dried, the sections were stained with 2% uranyl acetate for 15 min and Lead stain solution 168 (Sigma-Aldrich, St. Louis, MO, USA) for 3 min at room temperature.

169

170 **2.6. Immunoelectron-microscopy**

Ultrathin sections were observed using a transmission electron microscope (JEM-1400Plus;
JOEL Ltd., Tokyo Japan) at an acceleration voltage of 80 kV. Digital images (2048 × 2048 pixels)
were taken with a CCD camera (VELETA; Olympus Soft Imaging Solutions, Münster, Germany).

174

175 **2.7. Enzyme assays**

BGL and EG activities of washed yeast cell pellets were evaluated as described previously (Inokuma et al., 2016). Briefly, BGL activity was assayed at pH 5.0 and 30 °C with 2 mM *p*NPG as the substrate. One unit of BGL activity was defined as the amount of enzyme required to liberate 1 μ mol of *p*-nitrophenol per min. EG activity for water-insoluble cellulose was assayed at pH 5.0 and 38 °C using AZCL-HE-Cellulose (Cellazyme C tablets; Megazyme, Bray, Ireland) as the substrate.

182

183 **2.8.** Quantification of the transcript levels of cellulase-encoding genes by real-time PCR

The transcript levels of the genes encoding BGL1 and EGII were quantified by real-time PCR as described previously (Liu et al., 2017). The primers used are listed in Supplementary Table S2. Gene expression levels of target genes were normalized to those of the housekeeping actin gene, *ACT1*.

188

189 2.9. Relative quantitative analysis of cell wall-associated heterologous proteins

190 The identification and relative quantification of heterologous proteins in the veast cell wall were 191 performed based on precise mass measurements of tryptic peptides from each protein using 192 nanoscale ultra-pressure liquid chromatography electrospray ionization quadrupole time-of-flight 193 tandem mass spectrometry (nano-UPLC-MS^E). The extraction of cell wall-associated proteins, 194 sample preparation, and protein identification using nano-UPLC-MS^E were conducted as described 195 previously (Bamba et al., 2018) with a minor modification in which an ACOUITY UPLC Peptide 196 BEH C18 nanoACQUITY Column (75 μm × 100 mm; particle size, 1.7 μm; Waters Corporation, 197 Milford, MA, USA) was used as the analytical column.

198 LC-MS^E data processing and the relative quantitative analysis of cell wall-associated 199 heterologous proteins were performed using ProteinLynx Global SERVER v3.0 (Waters 200 Corporation) as described previously (Bamba et al., 2018).

201

202 2.10. Simultaneous saccharification and fermentation of pretreated rice straw

Rice straw was pretreated with the liquid hot water method and its insoluble fraction was then subjected to four cycles of ball milling as described previously (Sasaki et al., 2015). The composition of the pretreated rice straw was 43% (w/w) glucan, 2% (w/w) xylan, 42.3% (w/w) ash and lignin, and 12.7% (w/w) other materials (Matano et al., 2012). The pretreated rice straw was used as the substrate for simultaneous saccharification and fermentation in this study.

208 S. cerevisiae strains used for fermentation were cultivated at 30 °C for 48 h in 500 mL YPD medium. The veast cells were collected by centrifugation at $1000 \times g$ for 10 min at 20 °C, and then 209 210 washed twice with distilled water. The cells were then resuspended in 10 mL yeast extract peptone 211 (YP) medium (10 g/L of yeast extract and 20 g/L of Bacto-peptone) containing 50 mM sodium 212 citrate buffer (pH 5.0), 100 g/L of pretreated rice straw, and 0.4 filter paper units (FPU)/g-biomass of commercial cellulase (Cellic CTec2; Novozymes Inc., Bagsvaerd, Denmark) in a 50-mL 213 214 polypropylene tube (Corning Inc., Corning, NY, USA) at an initial cell concentration of 100 g wet 215 cells/L. Fermentation was initiated by the addition of yeast cells into the tube followed by axial 216 rotation using a heat block (Thermo Block Rotator SN-06BN; Nissin, Tokyo, Japan) at 35 rpm and 217 37 °C. The ethanol concentration in the fermentation medium was determined by HPLC 218 (Shimadzu, Kyoto, Japan), as described previously (Hasunuma et al., 2011).

219

3. Results

221 3.1. Construction of eGFP-displaying or secreting S. cerevisiae strains

To verify the localization of heterologous proteins fused with GPI-anchoring domains, gene 222 223 cassettes for the cell-surface display of eGFP were constructed using the S. cerevisiae SED1 224 promoter and two different GPI-anchoring regions derived from S. cerevisiae SED1 or SAG1 (Fig. 225 1A). We also constructed a gene cassette without the GPI-anchoring region for the secretory 226 production of eGFP. The plasmids containing these cassettes were integrated into the HIS3 locus of 227 the chromosomal DNA of S. cerevisiae BY4741 by homologous recombination. The constructed 228 eGFP-displaying strains were used for microscopic observation and immunoelectron-microscopy. 229 For all gene cassettes used in this study, we used the secretion signal sequence derived from S. 230 *cerevisiae SED1* because it showed high performance with respect to the cell-surface display and 231 secretory production of heterologous proteins in our previous study (Inokuma et al., 2016).

233 **3.2.** Fluorescence and immunoelectron microscopic observations of eGFP-displaying cells

To evaluate the localization of heterologous proteins fused with GPI-anchoring domains, the 234 235 fluorescence of eGFP-displaying strains (BY-eGFP-SSS and BY-eGFP-SSA) was observed using 236 a confocal fluorescence microscope (Fig. 2A). In the strain carrying the Sed1-anchoring domain (BY-eGFP-SSS), most green fluorescence was observed on the cell surface. In contrast, in the 237 238 strain carrying the Sag1-anchoring domain (BY-eGFP-SSA), less green fluorescence was observed 239 on the cell surface compared to that in BY-eGFP-SSS. However, more fluorescence was observed 240 in the intracellular vacuoles of strain BY-eGFP-SSA than in those of strain BY-eGFP-SSS. For 241 comparison, we also conducted the same experiment on an eGFP-secreting strain (BY-eGFP-SSn). 242 No significant fluorescence was observed either on the cell surface or in the intracellular vacuoles 243 of cells of this strain.

244 To further evaluate the localization of heterologous proteins fused with GPI-anchoring domains in the cell wall, we performed an immunoelectron microscopy analysis of eGFP-displaying strains. 245 246 Fixed BY-eGFP-SSS and BY-eGFP-SSA cell samples were cut into ultrathin sections, which were 247 then immunostained with the primary antibody (rabbit-anti GFP) and the secondary antibody (goat 248 anti-rabbit IgG) conjugated with 10-nm gold particles, as described in the Materials and Methods 249 section. Immunoelectron micrographs of these strains are shown in Fig. 2B. We observed 12 cells 250 for each of strains BY-eGFP-SSS and BY-eGFP-SSA. In both strains, most gold particles, 251 indicating the eGFP fusion proteins, were detected on the cell surface. The number of detected gold 252 particles associated with the cell wall was higher in BY-eGFP-SSS cells than in BY-eGFP-SSA 253 cells. This result was in good agreement with the fluorescence observations shown in Fig. 2A. 254 Furthermore, these strains showed different localization tendencies with respect to the eGFP fusion 255 proteins. In strain BY-eGFP-SSS (expressing eGFP-Sed1), most gold particles were detected on 256 the external side of the cell wall, whereas in strain BY-eGFP-SSA (expressing eGFP-Sag1), a large proportion of gold particles was detected on the internal side of the cell wall. 257

258

259 3.3. Construction of S. cerevisiae strains co-displaying exo- and endo-cellulases

260 To demonstrate the effect of anchorage position control using different anchoring domains, we 261 applied this technology to the co-display of exo- and endo-cellulases. Gene cassettes for the cellsurface display of A. aculeatus BGL1 with the S. cerevisiae SED1 promoter and GPI-anchoring 262 263 regions derived from S. cerevisiae SED1 or SAG1 were constructed (Fig. 1B). The plasmids 264 containing these cassettes were integrated into the 3' noncoding region of YFL021W and YFL020C 265 genes in the chromosomal DNA of the BY-EG-SSS strain (Inokuma et al., 2016), which is a 266 recombinant S. cerevisiae strain displaying T. reesei EGII fused with the Sed1-anchoring domain 267 (Fig. 1C), by homologous recombination. The constructed EG and BGL co-displaying strains, 268 designated BY-ESBS and BY-ESBA (i.e., containing combinations of EGII-Sed1 + BGL1-Sed1 269 and EGII-Sed1 + BGL1-Sag1, respectively), were used for enzyme assays and direct ethanol 270 production from pretreated rice straw.

271

272 **3.4.** Enzyme activity and relative quantity of cell wall-associated EGII and BGL1

273 The EG and BGL co-displaying strains (BY-ESBS and BY-ESBA) and their parental strain 274 (BY-EG-SSS) were cultivated at 30 °C for 48 h and cell-surface EG and BGL activities were 275 evaluated as described in the Methods section (Fig. 3A). In the BY-ESBS strain, in which both EG 276 and BGL were displayed using the Sed1-anchoring domain, cell-surface EG activity was 277 approximately 40% lower compared to that in its parental strain (BY-EG-SSS). In contrast, no 278 significant difference in cell-surface EG activity was observed between the parental strain and the 279 BY-ESBA strain displaying EG and BGL using Sed1- and Sag1-anchoring domains, respectively. 280 The cell-surface BGL activity of BY-ESBS was approximately 1.5-fold higher than that of BY-281 ESBA. We also investigated the transcriptional expression levels of T. reesei EGII and A. aculeatus BGL1 genes in these strains by quantitative real-time PCR analysis. In these strains, no significant
difference was observed in the expression levels of these genes after 48 h of cultivation (Fig. 3B).

We also performed the relative quantification of cell wall-associated cellulases in BY-ESBS and BY-ESBA strains by nano-UPLC-MS^E. The amount of cell wall-associated BGL1 per unit dry cell-weight of BY-ESBS was 1.67 ± 0.14 -fold higher than that in BY-ESBA (Fig. 3C), which was similar to the fold-change in cell-surface BGL activity between these strains. In contrast, cell wallassociated EGII was not detected by nano-UPLC-MS^E analysis in either strain.

289

290 3.5. Simultaneous saccharification and fermentation of pretreated rice straw

291 To further verify the effect of the anchorage position control of enzymes on cellulase-displaying 292 veast, we performed the simultaneous saccharification and fermentation of pretreated rice straw, 293 which was subjected to hydrothermal and ball milling treatments, using strains BY-ESBS, BY-294 ESBA, and their parental strain BY4741. A small amount of a commercial cellulase cocktail (0.4 295 FPU/g-biomass) was added to the fermentation mixture to supply auxiliary cellulolytic enzymes. 296 The fermentation was performed at 37 °C to promote the activity of cellulases displayed on the cell 297 surface because the optimal temperature of A. aculeatus BGL1 and T. reesei EGII was 65 °C 298 (Decker et al., 2000; Trudeau et al., 2014). As shown in Fig. 4, the use of the EG and BGL co-299 displaying strains resulted in increased ethanol production from the pretreated biomass compared to 300 that with their parental strain. Furthermore, BY-ESBA improved the ethanol production more 301 significantly than BY-ESBS.

302

303 4. Discussion

As mentioned in the Introduction, cell-surface display systems can be utilized for a wide range of applications in *S. cerevisiae*. However, as the yeast cell wall has a thickness of 100 to 200 nm (Dupres et al., 2010), the optimal position of functional proteins in the cell surface might vary 307 depending on each application. For example, in protein screening, the exposure of target proteins to 308 the external surface of the cell wall is necessary to put them in contact with large ligands. 309 Conversely, localization close to the plasma membrane might be advantageous for screening 310 procedures utilizing signaling pathways through transmembrane proteins such as G protein-coupled 311 receptors (GPCRs) (Hara et al., 2012). Furthermore, in plant biomass degradation requiring multiple enzymes, proper segregation of each enzyme in the cell wall enables the efficient 312 313 utilization of its limited protein loading capacity. Therefore, a technology to control the localization of functional proteins in the cell wall is essential to further develop yeast cell-surface display 314 315 systems.

316 Although several studies on the localization control of GPI-attached proteins in the cell surface 317 have been reported over the past few decades, most have focused on whether GPI proteins are 318 retained on the plasma membrane or translocated to the cell wall (Hamada et al., 1998; Nuoffer et 319 al., 1991; Orlean, 2012). It has been suggested that the distribution of GPI proteins between the 320 plasma membrane and cell wall depends on the amino acid residues within the upstream region of 321 the GPI-attachment site (the ω -minus region). If the ω -minus region includes two basic amino acids, 322 the protein will be mostly retained in the plasma membrane in a lipid-anchored form, but if the 323 dibasic motif is absent or replaced by hydrophobic residues, the primary localization of the protein 324 is the glucan layer in the cell wall (Frieman and Cormack, 2003; Hamada et al., 1999). Another 325 determinant of the distribution of GPI proteins between the plasma membrane and cell wall is the 326 presence of longer regions rich in serine and threonine residues. Amino acid stretches that are rich 327 in serine and threenine can override the dibasic motif in the ω -minus region and promote 328 localization to the cell wall (Frieman and Cormack, 2004). Terashima et al. (2003) reported a 329 change in the localization of the GPI protein Ecm33p, from the plasma membrane to the cell wall, 330 after replacing its authentic ω-minus region with that of cell wall-localized GPI proteins, Fit1p and 331 Egt2p. In contrast, Hara et al. (2012) efficiently localized a GPCR-specific peptide ligand to the

plasma membrane by fusing it with the minimum length (six amino acids including the ω site) of the membrane-associated GPI protein Yps1p and activated the yeast pheromone response pathway. To our knowledge, however, no comparative study on the final anchorage position of GPI-attached proteins liberated from the yeast plasma membrane has been reported.

336 In this study, we investigated the effect of the fusion of GPI-anchoring domains to heterologous 337 proteins on their localization in yeast cells using two GPI-anchoring domains derived from well-338 characterized GPI-CWPs, namely Sed1p and Sag1p (Supplementary Fig. S1). As the Sed1- and 339 Sag1-anchoring domains used in this study have hydrophobic amino acids in their ω -minus region 340 and the serine and threonine contents are high (41.8 and 40.3%, respectively), the proteins fused 341 with these domains were expected to be predominantly localized to the cell wall. Confocal 342 microscopy observations using a reporter protein (eGFP) indicated that fusing the GPI-anchoring 343 domain to eGFP promotes intracellular transportation efficiency of the fusion protein. This result is 344 in good agreement with the results of cell-surface BGL activity measurements in a previous study 345 (Inokuma et al., 2014). Similar anchoring domain-dependent changes in the intracellular 346 accumulation of GPI-attached proteins were also reported in the methylotrophic yeast Pichia 347 pastoris (Zhang et al., 2013). Furthermore, immunoelectron-microscopic analysis of ultra-thin 348 sections of the eGFP-displaying yeast cells clearly indicated that the fusion of GPI-anchoring 349 domains with eGFP also determined its final immobilized location, and in particular, the depth in 350 the cell wall. To our knowledge, this is the first report comparing the final destination of a 351 heterologous protein fused with different GPI-anchoring domains in the yeast cell wall. 352 Immunoelectron-microscopic analyses of yeast cells displaying enzymes (glucoamylase and 353 carboxymethylcellulase) fused with the Sag1-anchoring domain have been reported previously 354 (Murai et al., 1997a; Murai et al., 1997b). In these reports, the fusion proteins were detected only on 355 the external surface of the cell wall. These results are not consistent with our observation shown in Fig. 2B, which is likely due to a difference in the analytical methods adopted. In the current study, 356

immunostaining was carried out after the embedding and ultrathin sectioning of the cells (see Materials and Methods section), whereas in previous reports, immunostaining was performed prior to embedding and sectioning (Murai et al., 1997a; Murai et al., 1997b). Therefore, enzymes fused with the Sag1-anchoring domain buried in the glucan layer might not have been detected in these previous reports.

362 The results presented in this study suggest that yeast cells recognize GPI-anchoring domains attached to target proteins and control their anchorage positions in the cell wall. Although the 363 364 anchorage mechanism of yeast GPI-CWPs liberated from the plasma membrane to the cell wall 365 remains unclear, recent studies have suggested that plasma membrane-anchored GPI proteins 366 Dfg5p and Dcw1p are potential candidates for cross-linking the GPI-anchor remnant and cell wall β -(1 to 6) glucan (Gonzalez et al., 2010; Orlean, 2012). These proteins are putative 367 368 glycosidase/transglycosidases homologous to bacterial family 75 (Cantarel et al., 2009) and 369 depletion of these enzymes by repressing their expression in the double-null background led to 370 secretion of a GPI-CWP into the medium (Kitagaki et al., 2002). These enzymes might recognize 371 differences in GPI anchoring domains and be involved in controlling the anchorage position of 372 GPI-attached proteins. Further analysis using GPI-anchoring domains derived from other GPI-373 CWPs are urgently required to identify the determinants of the anchorage position of GPI-attached 374 proteins. On the other hand, in order to expand this research to a wide range of GPI-CWPs, it will 375 be necessary to develop a novel method for high-throughput anchorage position analysis.

In this study, we also demonstrated the application of the localization control technique for the construction of cellulase-displaying yeast. EGII, which requires contact with bulky insoluble cellulose, was preferentially localized to the external surface of the cell wall by fusing it with the Sed1-anchoring domain. Concomitantly, BGL1 was immobilized on the inside of the cell wall using the Sag1-anchoring domain, which avoided competition with EGII for space on the outer surface. As a result of the reallocation of cell wall space, cell-surface EG activity in BY-ESBA (containing combinations of EGII-Sed1 + BGL1-Sag1) was almost twice that of BY-ESBS (containing combinations of EGII-Sed1 + BGL1-Sed1) (Fig. 3A). Despite lower BGL1 activity (Fig. 3A), BY-ESBA achieved a higher ethanol titer after the simultaneous saccharification and fermentation of pretreated lignocellulosic biomass, as compared to that with BY-ESBS (Fig. 4); this is likely due to the enhanced access of EGII to its polymeric substrate. These results indicate the importance of the anchorage position control of target proteins in yeast cell-surface display systems.

389 To investigate the status of cellulases immobilized in the yeast cell wall in more detail, we 390 performed relative quantitative analysis of cell wall-associated cellulases in BY-ESBS and BY-391 ESBA strains by nano-UPLC-MS^E. The amount of cell wall-associated BGL1 per unit dry cell-392 weight of BY-ESBS was 1.67-fold higher compared to that with BY-ESBA. This result indicates 393 that the difference in cell-surface BGL activity between these strains is due to differences in the 394 abundance of cell wall-associated BGL1. Although we attempted the relative quantification of cell 395 wall-associated EGII, this protein was not detected in the cell wall fractions of both strains. One 396 possible reason for this result could be the hyperglycosylation of EGII in S. cerevisiae. It was 397 previously reported that recombinant T. reesei EGII expressed in S. cerevisiae has a larger 398 molecular mass compared to the native enzyme produced by T. reesei (48 kDa) due to different 399 levels of glycosylation; moreover, a portion of recombinant EGII presents as hyperglycosylated 400 isoforms with a broad molecular mass up to 200 kDa (Qin et al., 2008). In contrast, it was reported 401 that the glycosylation level of recombinant Aspergillus kawachii BGLA (Genbank annotation No. 402 BAA19913), which has significant similarity (81.8%) to A. aculeatus BGL1 (Genbank annotation 403 No. BAA10968) produced by S. cerevisiae, is fairly homogenous and that this protein has an 404 apparent molecular mass of 120 kDa (Iwashita et al., 1999). In the nano-UPLC-MS^E analysis, 405 protein identification is conducted based on precise mass measurements of tryptic peptides from 406 each protein. The masses of tryptic peptides derived from EGII displayed in this study might have 407 been altered by variable glycosylation, and therefore, it might not have been possible to identify this
408 enzyme by the nano-UPLC-MS^E analysis.

409 In this study, we used EG and BGL co-displaying strains for the simultaneous saccharification 410 and fermentation of pretreated rice straw. It has been demonstrated that synergistic cooperation of 411 EG and cellobiohydrolases (CBHs) is essential for efficient degradation of insoluble cellulose 412 (Jalak et al. 2012). CBHs are chain end-specific processive exo-glucanases. EG randomly 413 hydrolyzes amorphous regions of insoluble cellulose and generates reducing and non-reducing ends that can be attacked by CBHs, while CBHs recognize the cellulose chain ends and 414 415 continuously hydrolyze crystalline regions between the amorphous parts into cellobiose units (Jalak 416 et al. 2012). Previously, we reported a simultaneous saccharification and fermentation from 417 pretreated rice straw using a recombinant yeast strain, in which BGL1, EGII, and CBHs (CBH1 418 and CBH2) were displayed using the Sed1-anchoring domain (Liu et al., 2016). Although direct 419 comparison with the result shown in Fig. 4 is not possible due to differences in fermentation scale 420 and agitation procedure, the BGL, EG, and CBHs co-displaying strain achieved approximately 9.5 421 g/L of ethanol production after 96 h fermentation with 0.2 FPU/g biomass of commercial cellulase 422 cocktail (Liu et al., 2016). This ethanol titer is higher than that of BY-ESBS strain with 0.4 FPU/g 423 biomass of commercial cellulase cocktail (7.3 g/L at 96 h, Fig. 4). These results also suggest the 424 importance of co-display of BGL, EG, and CBHs for efficient hydrolysis of insoluble cellulosic 425 materials. Additional display of CBHs on the cell surface of BY-ESBA strain will be required for 426 further improvement of its ethanol yield from pretreated biomass. Furthermore, it will be necessary 427 to verify the optimal anchorage position for CBHs in the cell wall to maximize synergies between 428 cellulases.

The anchorage position control technique demonstrated in this study will also benefit
applications of yeast cell-surface display other than the construction of cellulase-displaying yeast.
The hydrolysis efficiency of other plant-derived polysaccharides such as hemicellulase and starch

432 may be improved by this technique because the complete hydrolysis of these polysaccharides also 433 requires the cooperation of endo- and exo-type enzymes. In addition, the Sed1-anchoring domain 434 that can expose the target protein to the external surface of the cell wall will also be a potential 435 anchor candidate for protein screening requiring contact with large ligands.

436

437 **5.** Conclusions

438 In the present study, we provide the first experimental evidence that the anchorage position of 439 GPI-attached heterologous proteins in the yeast cell wall can be controlled by the specific 440 anchoring domain fused to them. A reporter protein (eGFP) was predominantly localized to the 441 external surface of the cell wall when fused with the Sed1-anchoring domain, whereas the 442 anchorage position of eGFP fused with the Sag1-anchoring domain was mainly inside of the cell 443 wall. By applying this anchorage position control technique, the cellulolytic ability of the 444 recombinant yeast strain co-displaying EG and BGL was successfully improved. Although further 445 analyses using GPI-anchoring domains derived from a wide-range of GPI-CWPs are required to identify the determinants of GPI-attached protein anchorage positions, our novel strategy for 446 447 anchorage position control will enable the efficient utilization of the cell wall space for various 448 fields of yeast cell-surface display.

449

450 **Declaration of interest**

451 The authors declare that they have no competing interests.

452

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- 460

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Strains	Relevant genotype	Source
S. cerevisiae BY4741	$MATa$ his3 $\Delta 1$ leu2 $\Delta 0$ met15 $\Delta 0$ ura3 $\Delta 0$	Invitrogen
BY-eGFP-SSS	BY4741/pIeGFP-SSS	This study
BY-eGFP-SSA	BY4741/pIeGFP-SSA	This study
BY-eGFP-SSn	BY4741/pIeGFP-SS2	This study
BY-BG-SSS	BY4741/pIBG-SSS	Inokuma et al. (2016)
BY-EG-SSS	BY4741/pIEG-SSS	Inokuma et al. (2016)
BY-ESBS	BY-EG-SSS/pIL2BG-SSS	This study
BY-ESBA	BY-EG-SSS/pIL2BG-SSA	This study

Table 1 Characteristics of yeast strains used in this study

608 Figure legends

609

Fig. 1 Schematic summary of the construction of gene cassettes used in this study. (A) Gene cassettes for cell-surface display and the secretory production of eGFP. (B) Gene cassettes for the cell-surface display of BGL1. (C) Gene cassettes for cell-surface display of EGII.

613

Fig. 2 Localization analyses of eGFP fused with Sed1- or Sag1-anchoring domains. **(A)** Fluorescence images of strains BY-eGFP-SSS, BY-eGFP-SSA, and BY-eGFP-SSn. The cells were incubated in YPD medium for 48 h, stained with FM4-64 (red) to visualize vacuolar membranes, and then observed using a confocal microscope. **(B)** Immunoelectron micrographs of strains BY-eGFP-SSS and BY-eGFP-SSA. The cells were immunogold-labeled with an antibody against GFP. The arrowheads indicate gold particles.

620

621 Fig. 3 Effects of anchorage position control on enzyme activities of cellulase-displaying yeasts. (A) 622 Comparison of cell-surface EG and BGL activities in strains BY-BG-SSS, BY-ESBS, and BY-623 ESBA after cultivation in YPD medium for 48 h. The relative EG activity of each strain is shown 624 as a fold-change in EG activity relative to the average level observed with the parental strain BY-625 EG-SSS. (B) Comparison of transcript levels of EGII- and BGL1-encoding genes in strains BY-626 ESBS and BY-ESBA after cultivation in YPD medium for 48 h. The relative transcript level of 627 each gene is shown as a fold-change in mRNA levels relative to the average level detected in strain 628 BY-ESBS. (C) Relative quantification of BGL1 in the cell walls of strains BY-ESBS and BY-629 ESBA by nanoscale ultra-pressure liquid chromatography electrospray ionization quadrupole time-630 of-flight tandem mass spectrometry (nano-UPLC-MS^E). The amount of BGL1 was normalized to 631 the dry cell weight of each strain. Data are presented as the means \pm standard deviation (n = 3).
- **Fig. 4** Time course of the simultaneous saccharification and fermentation of 100 g dry weight/L of
- 634 pretreated rice straw by strains BY-ESBS, BY-ESBA, and their parental strain (BY4741). A small
- 635 amount of a commercial cellulase cocktail (0.4 FPU/g-biomass) was added to the fermentation
- 636 mixture. Data are presented as the means \pm standard deviation (n = 3).

641	Supplementary materials
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Text S1 Plasmid construction and yeast transformation.

- **Table S1** Characteristics of integrative plasmids used in this study.

Table S2 PCR primers used in this study.

- 648 Fig. S1 Amino acid sequence of Sed1- and Sag1-anchoring domains used in this study. The C-
- 649 terminal GPI attachment site (the ω site) was marked in bold. The hydrophobic amino acid residues
- 650 in the ω -minus region are underlined.

1 Highlights

- The GPI-anchoring domain fused with a heterologous protein determines its anchorage
 position in yeast cell wall.
- Proteins fused with the Sed1-anchoring domain predominantly localize to the external surface
 of the cell wall.
- The anchorage position of proteins fused with the Sag1-anchoring domain is mainly inside of
 the cell wall.
- 8 By repositioning exo- and endo-cellulases in cellulase-displaying yeast, the ethanol titer from
- 9 pretreated rice straw was improved by 30%.

1	Novel strategy for anchorage position control of GPI-attached proteins in the yeast cell wall
2	using different GPI-anchoring domains
3	
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26 Abstract

27 The yeast cell surface provides space to display functional proteins. Heterologous proteins can 28 be covalently anchored to the yeast cell wall by fusing them with the anchoring domain of 29 glycosylphosphatidylinositol (GPI)-anchored cell wall proteins (GPI-CWPs). In the yeast cell-30 surface display system, the anchorage position of the target protein in the cell wall is an important 31 factor that maximizes the capabilities of engineered yeast cells because the yeast cell wall consists 32 of a 100- to 200-nm-thick microfibrillar array of glucan chains. However, knowledge is limited 33 regarding the anchorage position of GPI-attached proteins in the yeast cell wall. Here, we report a 34 comparative study on the effect of GPI-anchoring domain-heterologous protein fusions on yeast 35 cell wall localization. GPI-anchoring domains derived from well-characterized GPI-CWPs, namely 36 Sed1p and Sag1p, were used for the cell-surface display of heterologous proteins in the yeast 37 Saccharomyces cerevisiae. Immunoelectron-microscopic analysis of enhanced green fluorescent 38 protein (eGFP)-displaying cells revealed that the anchorage position of the GPI-attached protein in 39 the cell wall could be controlled by changing the fused anchoring domain. eGFP fused with the 40 Sed1-anchoring domain predominantly localized to the external surface of the cell wall, whereas 41 the anchorage position of eGFP fused with the Sag1-anchoring domain was mainly inside the cell 42 wall. We also demonstrate the application of the anchorage position control technique to improve 43 the cellulolytic ability of cellulase-displaying yeast. The ethanol titer during the simultaneous 44 saccharification and fermentation of hydrothermally-processed rice straw was improved by 30% 45 after repositioning the exo- and endo-cellulases using Sed1- and Sag1-anchor domains. This novel 46 anchorage position control strategy will enable the efficient utilization of the cell wall space in 47 various fields of yeast cell-surface display technology.

48

49 Keywords: Saccharomyces cerevisiae, yeast surface display, glycosylphosphatidylinositol-

50 anchored cell wall protein, anchorage position, Sed1p, Sag1p

52	Abbreviations: BGL, β -glucosidase; BSA, bovine serum albumin; EG, endoglucanase; eGFP,
53	enhanced green fluorescent protein; ER, endoplasmic reticulum; FPU, filter paper unit; GPCR, G
54	protein-coupled receptor; GPI, glycosylphosphatidylinositol; GPI-CWP, GPI-anchored cell wall
55	protein; GRAS, generally regarded as safe; nano-UPLC-MS ^E , nanoscale ultra-pressure liquid
56	chromatography electrospray ionization quadrupole time-of-flight tandem mass spectrometry; pAb,
57	polyclonal antibody; pNPG, p-nitrophenyl-β-D-glucopyranoside; YP, yeast extract peptone

59 1. Introduction

The expression of functional proteins on the cell surface is a promising approach to construct cell-surface-engineered microorganisms with special functions. Cell-surface display technology can be used to address a wide range of applications such as the engineering and screening of enzymes, antibodies, or peptides (Angelini et al., 2015; Grzeschik et al., 2017; Li et al., 2007), the bioadsorption of specific molecules (Shibasaki and Ueda, 2014), and the production whole cell catalysts for bioconversion (Inokuma et al., 2018; Liu et al., 2016), biodegradation (Richins et al., 1997; Shibasaki et al., 2009), and biosensing (Tang et al., 2014; Wang et al., 2013).

67 Among host microorganisms employed for cell-surface display, baker's yeast (Saccharomyces 68 *cerevisiae*) is the most frequently used to develop such systems because of the vast knowledge of 69 its genetics, physiology, and fermentation characteristics, as well as its generally regarded as safe 70 (GRAS) status. In yeast cell-surface display, the glycosylphosphatidylinositol (GPI)-anchoring 71 system is the typical and most widely-used technique to immobilize heterologous proteins. In this 72 method, a yeast cell is transformed by introducing fusion genes coding proteins of interest and the 73 anchoring domain of the GPI-anchored cell wall protein (GPI-CWP). In the recombinant yeast cells, 74 the fused proteins are synthesized on endoplasmic reticulum (ER)-bound ribosomes, cleaved at the 75 C-terminal GPI attachment site (the ω site), and modified by the addition of a pre-assembled GPI 76 anchor in the ER. Subsequently, the GPI-attached proteins leave the ER in COPII-coated vesicles 77 and travel via the Golgi to the plasma membrane (Doering and Schekman, 1996). Finally, these 78 proteins are liberated from the plasma membrane and become immobilized in the cell wall through 79 covalent linkage to a β -(1 to 6) glucan via a remnant of the anchor structure (Klis et al., 1997; Lu et 80 al., 1994).

81 It has been reported that the anchoring domains from different GPI-CWPs exhibit different 82 efficiencies for the cell-surface display of target enzymes (Andreu and Del Olmo, 2018; Hamada et 83 al., 1999). Therefore, selection of the appropriate anchoring domain for fusion with target proteins

84 is important for efficient cell-surface display. In previous studies, indeed, the activities of some 85 cellulolytic enzymes fused with the Sed1-anchoring domain were found to be higher than those 86 fused with the α-agglutinin (Sag1)-anchoring domain (Inokuma et al., 2014). However, the degree 87 to which activity was improved by changing the anchoring domain varied greatly depending on the 88 enzymes displayed. For the cell-surface display of *Aspergillus aculeatus* β-glucosidase 1 (BGL1), 89 the activity of the enzyme fused with the Sed1-anchoring domain was approximately 2-fold higher 90 than that upon fusion with the Sag1-anchoring domain. In contrast, the hydrolytic activity of 91 Trichoderma reesei endoglucanase II (EGII) for water-insoluble cellulose was improved 60-fold 92 when using the Sed1-anchoring domain compared to that with the Sag1-anchoring domain 93 (Inokuma et al., 2014). Based on these results, we hypothesized that selection of the anchoring 94 domain would affect not only the display efficiency of the target protein but also its localization in 95 the cell wall.

96 The yeast cell wall is composed of a microfibrillar array of β -(1 to 3) glucan and β -(1 to 6) 97 glucan chains with a thickness of 100 to 200 nm (Dupres et al., 2010). Therefore, cell wall proteins 98 exposed to the external surface represent only a portion of the whole and the remainder are buried in the glucan layer (Van der Vaart et al., 1997). Small substrates such as cellobiose and p-99 100 nitrophenyl-B-D-glucopyranoside (pNPG) are accessible to all integrated enzymes because these 101 substrates penetrate the cell wall. In contrast, large substrates such as water-insoluble cellulose can 102 only access enzymes exposed on the external surface. However, to our knowledge, no comparative 103 analysis has been reported concerning the effect of the anchoring domain on target protein 104 localization in the yeast cell wall.

In the present study, we performed a comparative analysis of the effect of different anchoring domains on the cell wall localization of fused heterologous proteins in *S. cerevisiae*. First, the intracellular localization of enhanced green fluorescent protein (eGFP) fused with Sed1 or Sag1anchoring domains was analyzed using a confocal fluorescence microscope. Subsequently, immunoelectron-microscopic analysis of ultra-thin sections of the eGFP-displaying yeast cells was carried out to investigate the effect of the anchoring domains on the anchorage position of GPIattached proteins in the cell wall. Finally, by applying the information obtained from this novel system, we successfully demonstrate improved ethanol production from pretreated lignocellulosic biomass by cellulase-displaying yeast after controlling the anchorage position of exo- and endocellulases using different anchoring domains.

115

116 **2. Materials and Methods**

117 **2.1. Strains and media**

Escherichia coli strain DH5α (Toyobo, Osaka, Japan) was used as the host for recombinant
DNA manipulation. *E. coli* medium was prepared as described (Inokuma et al., 2016). The genetic
properties of all yeast strains used in this study are shown in Table 1. The gene cassettes for the
cell-surface display of heterologous proteins were expressed in the haploid yeast strain *S. cerevisiae*BY4741 (Life Technologies, Carlsbad, CA, USA).

The *S. cerevisiae* transformants were screened and cultivated as previously described (Inokuma et al., 2016). After 48 h of cultivation, yeast cells were harvested by centrifugation at $1000 \times g$ for 5 min, washed twice with distilled water, and again centrifuged at $1000 \times g$ for 5 min. The wet cell weight of the washed yeast cells was determined by weighing the cell pellet. The estimated dry cell weight of a yeast cell is approximately $0.15 \times$ its wet cell weight (Inokuma et al., 2014). Cell pellets were used for microscopic observation, immunoelectron-microscopy, enzyme assays, and ethanol fermentation.

130

131 **2.2. Plasmid construction and yeast transformation**

The plasmids and primers used in this study are listed in Supplementary Tables S1 and S2,
respectively. The integrative plasmids for the expression of eGFP, *T. reesei* EGII, and *A. aculeatus*

BGL1 were transformed into *S. cerevisiae* by the lithium acetate method (Chen et al., 1992) and integrated into the *HIS3* locus or the 3' noncoding region of YFL021W and YFL020C genes (I2 region) of the chromosomal DNA by homologous recombination. Details on the construction of plasmids and yeast transformation have been provided as Supplementary Text S1.

138

139 **2.3. Fluorescence microscopy**

140 Cell pellets of eGFP-expressing yeast strains were resuspended in 15 mM FM4-64 (Invitrogen 141 Carlsbad, CA, USA) diluted in culture medium and incubated for 15 min at 150 rpm and 30 °C in 142 the dark to stain vacuolar membranes. The cells were washed and resuspended in culture medium 143 followed by a further 2-h incubation at 150 rpm and 30 °C in the dark. After washing twice with 144 distilled water, the cells were observed using a confocal fluorescence microscope BZ-X810 145 (Keyence, Osaka, Japan) with a Nikon Plan Apo λ 100x/1.45 oil-immersion objective lens (Nikon, 146 Tokyo, Japan) and appropriate filters for eGFP and FM4-64.

147

148 **2.4. Sample preparation for immunoelectron-microscopy**

149 Washed cell pellets were sandwiched between two copper disks and frozen in liquid propane at -175 °C. The frozen samples were freeze-substituted with acetone containing 0.2% glutaraldehyde 150 151 and 2% distilled water at -80 °C for 2 days. The substituted samples were then transferred to -20 °C for 3 h and then warmed to 4 °C over 90 min. Next, they were dehydrated in anhydrous 152 153 acetone and anhydrous ethanol at 4 °C. Infiltration was performed with LR white resin (London 154 Resin Co. Ltd., Berkshire, UK) at 4 °C [ethanol:resin 50:50 for 2 h; 100% resin for 30 min; 100% resin for 30 min]. The samples were then transferred to a fresh 100% resin for embedding and the 155 resins were polymerized at 50 °C overnight. The polymerized resins were cut into ultrathin sections 156 157 of 90 nm thickness using an ultramicrotome (Ultracut CUT; Leica, Vienna, Austria) and the 158 sections were placed on nickel grids.

160 **2.5. Immunostaining**

161 Ultrathin sections were incubated with the primary antibody [rabbit anti-GFP polyclonal 162 antibody (pAb)] in blocking solution [PBS containing 1% bovine serum albumin (BSA)] at 4 °C 163 overnight and washed three times with the blocking solution. Subsequently, they were incubated 164 with secondary antibody conjugated to 10-nm gold particles (goat anti-rabbit IgG pAb; BBI 165 Solutions, Cardiff, UK) at room temperature for 90 min and washed with PBS. The sections in the nickel grids were placed in 2% glutaraldehyde in 0.1 M phosphate buffer (pH 7.4). After the grids 166 167 were dried, the sections were stained with 2% uranyl acetate for 15 min and Lead stain solution 168 (Sigma-Aldrich, St. Louis, MO, USA) for 3 min at room temperature.

169

170 **2.6. Immunoelectron-microscopy**

Ultrathin sections were observed using a transmission electron microscope (JEM-1400Plus;
JOEL Ltd., Tokyo Japan) at an acceleration voltage of 80 kV. Digital images (2048 × 2048 pixels)
were taken with a CCD camera (VELETA; Olympus Soft Imaging Solutions, Münster, Germany).

174

175 **2.7. Enzyme assays**

BGL and EG activities of washed yeast cell pellets were evaluated as described previously (Inokuma et al., 2016). Briefly, BGL activity was assayed at pH 5.0 and 30 °C with 2 mM *p*NPG as the substrate. One unit of BGL activity was defined as the amount of enzyme required to liberate 1 μ mol of *p*-nitrophenol per min. EG activity for water-insoluble cellulose was assayed at pH 5.0 and 38 °C using AZCL-HE-Cellulose (Cellazyme C tablets; Megazyme, Bray, Ireland) as the substrate.

182

183 **2.8.** Quantification of the transcript levels of cellulase-encoding genes by real-time PCR

The transcript levels of the genes encoding BGL1 and EGII were quantified by real-time PCR as described previously (Liu et al., 2017). The primers used are listed in Supplementary Table S2. Gene expression levels of target genes were normalized to those of the housekeeping actin gene, *ACT1*.

188

189 2.9. Relative quantitative analysis of cell wall-associated heterologous proteins

190 The identification and relative quantification of heterologous proteins in the veast cell wall were 191 performed based on precise mass measurements of tryptic peptides from each protein using 192 nanoscale ultra-pressure liquid chromatography electrospray ionization quadrupole time-of-flight 193 tandem mass spectrometry (nano-UPLC-MS^E). The extraction of cell wall-associated proteins, 194 sample preparation, and protein identification using nano-UPLC-MS^E were conducted as described 195 previously (Bamba et al., 2018) with a minor modification in which an ACOUITY UPLC Peptide 196 BEH C18 nanoACQUITY Column (75 μm × 100 mm; particle size, 1.7 μm; Waters Corporation, 197 Milford, MA, USA) was used as the analytical column.

198 LC-MS^E data processing and the relative quantitative analysis of cell wall-associated 199 heterologous proteins were performed using ProteinLynx Global SERVER v3.0 (Waters 200 Corporation) as described previously (Bamba et al., 2018).

201

202 2.10. Simultaneous saccharification and fermentation of pretreated rice straw

Rice straw was pretreated with the liquid hot water method and its insoluble fraction was then subjected to four cycles of ball milling as described previously (Sasaki et al., 2015). The composition of the pretreated rice straw was 43% (w/w) glucan, 2% (w/w) xylan, 42.3% (w/w) ash and lignin, and 12.7% (w/w) other materials (Matano et al., 2012). The pretreated rice straw was used as the substrate for simultaneous saccharification and fermentation in this study.

208 S. cerevisiae strains used for fermentation were cultivated at 30 °C for 48 h in 500 mL YPD medium. The veast cells were collected by centrifugation at $1000 \times g$ for 10 min at 20 °C, and then 209 210 washed twice with distilled water. The cells were then resuspended in 10 mL yeast extract peptone 211 (YP) medium (10 g/L of yeast extract and 20 g/L of Bacto-peptone) containing 50 mM sodium 212 citrate buffer (pH 5.0), 100 g/L of pretreated rice straw, and 0.4 filter paper units (FPU)/g-biomass of commercial cellulase (Cellic CTec2; Novozymes Inc., Bagsvaerd, Denmark) in a 50-mL 213 214 polypropylene tube (Corning Inc., Corning, NY, USA) at an initial cell concentration of 100 g wet 215 cells/L. Fermentation was initiated by the addition of yeast cells into the tube followed by axial 216 rotation using a heat block (Thermo Block Rotator SN-06BN; Nissin, Tokyo, Japan) at 35 rpm and 217 37 °C. The ethanol concentration in the fermentation medium was determined by HPLC 218 (Shimadzu, Kyoto, Japan), as described previously (Hasunuma et al., 2011).

219

3. Results

221 3.1. Construction of eGFP-displaying or secreting S. cerevisiae strains

To verify the localization of heterologous proteins fused with GPI-anchoring domains, gene 222 223 cassettes for the cell-surface display of eGFP were constructed using the S. cerevisiae SED1 224 promoter and two different GPI-anchoring regions derived from S. cerevisiae SED1 or SAG1 (Fig. 225 1A). We also constructed a gene cassette without the GPI-anchoring region for the secretory 226 production of eGFP. The plasmids containing these cassettes were integrated into the HIS3 locus of 227 the chromosomal DNA of S. cerevisiae BY4741 by homologous recombination. The constructed 228 eGFP-displaying strains were used for microscopic observation and immunoelectron-microscopy. 229 For all gene cassettes used in this study, we used the secretion signal sequence derived from S. 230 *cerevisiae SED1* because it showed high performance with respect to the cell-surface display and 231 secretory production of heterologous proteins in our previous study (Inokuma et al., 2016).

233 **3.2.** Fluorescence and immunoelectron microscopic observations of eGFP-displaying cells

To evaluate the localization of heterologous proteins fused with GPI-anchoring domains, the 234 235 fluorescence of eGFP-displaying strains (BY-eGFP-SSS and BY-eGFP-SSA) was observed using 236 a confocal fluorescence microscope (Fig. 2A). In the strain carrying the Sed1-anchoring domain (BY-eGFP-SSS), most green fluorescence was observed on the cell surface. In contrast, in the 237 238 strain carrying the Sag1-anchoring domain (BY-eGFP-SSA), less green fluorescence was observed 239 on the cell surface compared to that in BY-eGFP-SSS. However, more fluorescence was observed 240 in the intracellular vacuoles of strain BY-eGFP-SSA than in those of strain BY-eGFP-SSS. For 241 comparison, we also conducted the same experiment on an eGFP-secreting strain (BY-eGFP-SSn). 242 No significant fluorescence was observed either on the cell surface or in the intracellular vacuoles 243 of cells of this strain.

244 To further evaluate the localization of heterologous proteins fused with GPI-anchoring domains in the cell wall, we performed an immunoelectron microscopy analysis of eGFP-displaying strains. 245 246 Fixed BY-eGFP-SSS and BY-eGFP-SSA cell samples were cut into ultrathin sections, which were 247 then immunostained with the primary antibody (rabbit-anti GFP) and the secondary antibody (goat 248 anti-rabbit IgG) conjugated with 10-nm gold particles, as described in the Materials and Methods 249 section. Immunoelectron micrographs of these strains are shown in Fig. 2B. We observed 12 cells 250 for each of strains BY-eGFP-SSS and BY-eGFP-SSA. In both strains, most gold particles, 251 indicating the eGFP fusion proteins, were detected on the cell surface. The number of detected gold 252 particles associated with the cell wall was higher in BY-eGFP-SSS cells than in BY-eGFP-SSA 253 cells. This result was in good agreement with the fluorescence observations shown in Fig. 2A. 254 Furthermore, these strains showed different localization tendencies with respect to the eGFP fusion 255 proteins. In strain BY-eGFP-SSS (expressing eGFP-Sed1), most gold particles were detected on 256 the external side of the cell wall, whereas in strain BY-eGFP-SSA (expressing eGFP-Sag1), a large proportion of gold particles was detected on the internal side of the cell wall. 257

259 3.3. Construction of S. cerevisiae strains co-displaying exo- and endo-cellulases

260 To demonstrate the effect of anchorage position control using different anchoring domains, we 261 applied this technology to the co-display of exo- and endo-cellulases. Gene cassettes for the cellsurface display of A. aculeatus BGL1 with the S. cerevisiae SED1 promoter and GPI-anchoring 262 263 regions derived from S. cerevisiae SED1 or SAG1 were constructed (Fig. 1B). The plasmids 264 containing these cassettes were integrated into the 3' noncoding region of YFL021W and YFL020C 265 genes in the chromosomal DNA of the BY-EG-SSS strain (Inokuma et al., 2016), which is a 266 recombinant S. cerevisiae strain displaying T. reesei EGII fused with the Sed1-anchoring domain 267 (Fig. 1C), by homologous recombination. The constructed EG and BGL co-displaying strains, 268 designated BY-ESBS and BY-ESBA (i.e., containing combinations of EGII-Sed1 + BGL1-Sed1 269 and EGII-Sed1 + BGL1-Sag1, respectively), were used for enzyme assays and direct ethanol 270 production from pretreated rice straw.

271

272 **3.4.** Enzyme activity and relative quantity of cell wall-associated EGII and BGL1

273 The EG and BGL co-displaying strains (BY-ESBS and BY-ESBA) and their parental strain 274 (BY-EG-SSS) were cultivated at 30 °C for 48 h and cell-surface EG and BGL activities were 275 evaluated as described in the Methods section (Fig. 3A). In the BY-ESBS strain, in which both EG 276 and BGL were displayed using the Sed1-anchoring domain, cell-surface EG activity was 277 approximately 40% lower compared to that in its parental strain (BY-EG-SSS). In contrast, no 278 significant difference in cell-surface EG activity was observed between the parental strain and the 279 BY-ESBA strain displaying EG and BGL using Sed1- and Sag1-anchoring domains, respectively. 280 The cell-surface BGL activity of BY-ESBS was approximately 1.5-fold higher than that of BY-281 ESBA. We also investigated the transcriptional expression levels of T. reesei EGII and A. aculeatus BGL1 genes in these strains by quantitative real-time PCR analysis. In these strains, no significant
difference was observed in the expression levels of these genes after 48 h of cultivation (Fig. 3B).

We also performed the relative quantification of cell wall-associated cellulases in BY-ESBS and BY-ESBA strains by nano-UPLC-MS^E. The amount of cell wall-associated BGL1 per unit dry cell-weight of BY-ESBS was 1.67 ± 0.14 -fold higher than that in BY-ESBA (Fig. 3C), which was similar to the fold-change in cell-surface BGL activity between these strains. In contrast, cell wallassociated EGII was not detected by nano-UPLC-MS^E analysis in either strain.

289

290 3.5. Simultaneous saccharification and fermentation of pretreated rice straw

291 To further verify the effect of the anchorage position control of enzymes on cellulase-displaying 292 veast, we performed the simultaneous saccharification and fermentation of pretreated rice straw, 293 which was subjected to hydrothermal and ball milling treatments, using strains BY-ESBS, BY-294 ESBA, and their parental strain BY4741. A small amount of a commercial cellulase cocktail (0.4 295 FPU/g-biomass) was added to the fermentation mixture to supply auxiliary cellulolytic enzymes. 296 The fermentation was performed at 37 °C to promote the activity of cellulases displayed on the cell 297 surface because the optimal temperature of A. aculeatus BGL1 and T. reesei EGII was 65 °C 298 (Decker et al., 2000; Trudeau et al., 2014). As shown in Fig. 4, the use of the EG and BGL co-299 displaying strains resulted in increased ethanol production from the pretreated biomass compared to 300 that with their parental strain. Furthermore, BY-ESBA improved the ethanol production more 301 significantly than BY-ESBS.

302

303 4. Discussion

As mentioned in the Introduction, cell-surface display systems can be utilized for a wide range of applications in *S. cerevisiae*. However, as the yeast cell wall has a thickness of 100 to 200 nm (Dupres et al., 2010), the optimal position of functional proteins in the cell surface might vary 307 depending on each application. For example, in protein screening, the exposure of target proteins to 308 the external surface of the cell wall is necessary to put them in contact with large ligands. 309 Conversely, localization close to the plasma membrane might be advantageous for screening 310 procedures utilizing signaling pathways through transmembrane proteins such as G protein-coupled 311 receptors (GPCRs) (Hara et al., 2012). Furthermore, in plant biomass degradation requiring multiple enzymes, proper segregation of each enzyme in the cell wall enables the efficient 312 313 utilization of its limited protein loading capacity. Therefore, a technology to control the localization of functional proteins in the cell wall is essential to further develop yeast cell-surface display 314 315 systems.

316 Although several studies on the localization control of GPI-attached proteins in the cell surface 317 have been reported over the past few decades, most have focused on whether GPI proteins are 318 retained on the plasma membrane or translocated to the cell wall (Hamada et al., 1998; Nuoffer et 319 al., 1991; Orlean, 2012). It has been suggested that the distribution of GPI proteins between the 320 plasma membrane and cell wall depends on the amino acid residues within the upstream region of 321 the GPI-attachment site (the ω -minus region). If the ω -minus region includes two basic amino acids, 322 the protein will be mostly retained in the plasma membrane in a lipid-anchored form, but if the 323 dibasic motif is absent or replaced by hydrophobic residues, the primary localization of the protein 324 is the glucan layer in the cell wall (Frieman and Cormack, 2003; Hamada et al., 1999). Another 325 determinant of the distribution of GPI proteins between the plasma membrane and cell wall is the 326 presence of longer regions rich in serine and threonine residues. Amino acid stretches that are rich 327 in serine and threenine can override the dibasic motif in the ω -minus region and promote 328 localization to the cell wall (Frieman and Cormack, 2004). Terashima et al. (2003) reported a 329 change in the localization of the GPI protein Ecm33p, from the plasma membrane to the cell wall, 330 after replacing its authentic ω-minus region with that of cell wall-localized GPI proteins, Fit1p and 331 Egt2p. In contrast, Hara et al. (2012) efficiently localized a GPCR-specific peptide ligand to the

plasma membrane by fusing it with the minimum length (six amino acids including the ω site) of the membrane-associated GPI protein Yps1p and activated the yeast pheromone response pathway. To our knowledge, however, no comparative study on the final anchorage position of GPI-attached proteins liberated from the yeast plasma membrane has been reported.

336 In this study, we investigated the effect of the fusion of GPI-anchoring domains to heterologous 337 proteins on their localization in yeast cells using two GPI-anchoring domains derived from well-338 characterized GPI-CWPs, namely Sed1p and Sag1p (Supplementary Fig. S1). As the Sed1- and 339 Sag1-anchoring domains used in this study have hydrophobic amino acids in their ω -minus region 340 and the serine and threonine contents are high (41.8 and 40.3%, respectively), the proteins fused 341 with these domains were expected to be predominantly localized to the cell wall. Confocal 342 microscopy observations using a reporter protein (eGFP) indicated that fusing the GPI-anchoring 343 domain to eGFP promotes intracellular transportation efficiency of the fusion protein. This result is 344 in good agreement with the results of cell-surface BGL activity measurements in a previous study 345 (Inokuma et al., 2014). Similar anchoring domain-dependent changes in the intracellular 346 accumulation of GPI-attached proteins were also reported in the methylotrophic yeast Pichia 347 pastoris (Zhang et al., 2013). Furthermore, immunoelectron-microscopic analysis of ultra-thin 348 sections of the eGFP-displaying yeast cells clearly indicated that the fusion of GPI-anchoring 349 domains with eGFP also determined its final immobilized location, and in particular, the depth in 350 the cell wall. To our knowledge, this is the first report comparing the final destination of a 351 heterologous protein fused with different GPI-anchoring domains in the yeast cell wall. 352 Immunoelectron-microscopic analyses of yeast cells displaying enzymes (glucoamylase and 353 carboxymethylcellulase) fused with the Sag1-anchoring domain have been reported previously 354 (Murai et al., 1997a; Murai et al., 1997b). In these reports, the fusion proteins were detected only on 355 the external surface of the cell wall. These results are not consistent with our observation shown in Fig. 2B, which is likely due to a difference in the analytical methods adopted. In the current study, 356

immunostaining was carried out after the embedding and ultrathin sectioning of the cells (see Materials and Methods section), whereas in previous reports, immunostaining was performed prior to embedding and sectioning (Murai et al., 1997a; Murai et al., 1997b). Therefore, enzymes fused with the Sag1-anchoring domain buried in the glucan layer might not have been detected in these previous reports.

362 The results presented in this study suggest that yeast cells recognize GPI-anchoring domains attached to target proteins and control their anchorage positions in the cell wall. Although the 363 364 anchorage mechanism of yeast GPI-CWPs liberated from the plasma membrane to the cell wall 365 remains unclear, recent studies have suggested that plasma membrane-anchored GPI proteins 366 Dfg5p and Dcw1p are potential candidates for cross-linking the GPI-anchor remnant and cell wall β-(1 to 6) glucan (Gonzalez et al., 2010; Orlean, 2012). These proteins are putative 367 368 glycosidase/transglycosidases homologous to bacterial family 75 (Cantarel et al., 2009) and 369 depletion of these enzymes by repressing their expression in the double-null background led to 370 secretion of a GPI-CWP into the medium (Kitagaki et al., 2002). These enzymes might recognize 371 differences in GPI anchoring domains and be involved in controlling the anchorage position of 372 GPI-attached proteins. Further analysis using GPI-anchoring domains derived from other GPI-373 CWPs are urgently required to identify the determinants of the anchorage position of GPI-attached 374 proteins. On the other hand, in order to expand this research to a wide range of GPI-CWPs, it will 375 be necessary to develop a novel method for high-throughput anchorage position analysis.

In this study, we also demonstrated the application of the localization control technique for the construction of cellulase-displaying yeast. EGII, which requires contact with bulky insoluble cellulose, was preferentially localized to the external surface of the cell wall by fusing it with the Sed1-anchoring domain. Concomitantly, BGL1 was immobilized on the inside of the cell wall using the Sag1-anchoring domain, which avoided competition with EGII for space on the outer surface. As a result of the reallocation of cell wall space, cell-surface EG activity in BY-ESBA

(containing combinations of EGII-Sed1 + BGL1-Sag1) was almost twice that of BY-ESBS (containing combinations of EGII-Sed1 + BGL1-Sed1) (Fig. 3A). Despite lower BGL1 activity (Fig. 3A), BY-ESBA achieved a higher ethanol titer after the simultaneous saccharification and fermentation of pretreated lignocellulosic biomass, as compared to that with BY-ESBS (Fig. 4); this is likely due to the enhanced access of EGII to its polymeric substrate. These results indicate the importance of the anchorage position control of target proteins in yeast cell-surface display systems.

389 To investigate the status of cellulases immobilized in the yeast cell wall in more detail, we 390 performed relative quantitative analysis of cell wall-associated cellulases in BY-ESBS and BY-391 ESBA strains by nano-UPLC-MS^E. The amount of cell wall-associated BGL1 per unit dry cell-392 weight of BY-ESBS was 1.67-fold higher compared to that with BY-ESBA. This result indicates 393 that the difference in cell-surface BGL activity between these strains is due to differences in the 394 abundance of cell wall-associated BGL1. Although we attempted the relative quantification of cell 395 wall-associated EGII, this protein was not detected in the cell wall fractions of both strains. One 396 possible reason for this result could be the hyperglycosylation of EGII in S. cerevisiae. It was 397 previously reported that recombinant T. reesei EGII expressed in S. cerevisiae has a larger 398 molecular mass compared to the native enzyme produced by T. reesei (48 kDa) due to different 399 levels of glycosylation; moreover, a portion of recombinant EGII presents as hyperglycosylated 400 isoforms with a broad molecular mass up to 200 kDa (Qin et al., 2008). In contrast, it was reported 401 that the glycosylation level of recombinant Aspergillus kawachii BGLA (Genbank annotation No. 402 BAA19913), which has significant similarity (81.8%) to A. aculeatus BGL1 (Genbank annotation 403 No. BAA10968) produced by S. cerevisiae, is fairly homogenous and that this protein has an 404 apparent molecular mass of 120 kDa (Iwashita et al., 1999). In the nano-UPLC-MS^E analysis, 405 protein identification is conducted based on precise mass measurements of tryptic peptides from 406 each protein. The masses of tryptic peptides derived from EGII displayed in this study might have 407 been altered by variable glycosylation, and therefore, it might not have been possible to identify this
 408 enzyme by the nano-UPLC-MS^E analysis.

409 In this study, we used EG and BGL co-displaying strains for the simultaneous saccharification 410 and fermentation of pretreated rice straw. It has been demonstrated that synergistic cooperation of 411 EG and cellobiohydrolases (CBHs) is essential for efficient degradation of insoluble cellulose 412 (Jalak et al. 2012). CBHs are chain end-specific processive exo-glucanases. EG randomly 413 hydrolyzes amorphous regions of insoluble cellulose and generates reducing and non-reducing 414 ends that can be attacked by CBHs, while CBHs recognize the cellulose chain ends and 415 continuously hydrolyze crystalline regions between the amorphous parts into cellobiose units (Jalak 416 et al. 2012). Previously, we reported a simultaneous saccharification and fermentation from 417 pretreated rice straw using a recombinant veast strain, in which BGL1, EGII, and CBHs (CBH1 418 and CBH2) were displayed using the Sed1-anchoring domain (Liu et al., 2016). Although direct 419 comparison with the result shown in Fig. 4 is not possible due to differences in fermentation scale 420 and agitation procedure, the BGL, EG, and CBHs co-displaying strain achieved approximately 9.5 421 g/L of ethanol production after 96 h fermentation with 0.2 FPU/g biomass of commercial cellulase 422 cocktail (Liu et al., 2016). This ethanol titer is higher than that of BY-ESBS strain with 0.4 FPU/g 423 biomass of commercial cellulase cocktail (7.3 g/L at 96 h, Fig. 4). These results also suggest the 424 importance of co-display of BGL, EG, and CBHs for efficient hydrolysis of insoluble cellulosic 425 materials. Additional display of CBHs on the cell surface of BY-ESBA strain will be required for further improvement of its ethanol yield from pretreated biomass. Furthermore, it will be necessary 426 427 to verify the optimal anchorage position for CBHs in the cell wall to maximize synergies between 428 cellulases.

429 The anchorage position control technique demonstrated in this study will also benefit 430 applications of yeast cell-surface display other than the construction of cellulase-displaying yeast. 431 The hydrolysis efficiency of other plant-derived polysaccharides such as hemicellulase and starch 432 may be improved by this technique because the complete hydrolysis of these polysaccharides also 433 requires the cooperation of endo- and exo-type enzymes. In addition, the Sed1-anchoring domain 434 that can expose the target protein to the external surface of the cell wall will also be a potential 435 anchor candidate for protein screening requiring contact with large ligands.

436

437 **5.** Conclusions

In the present study, we provide the first experimental evidence that the anchorage position of 438 GPI-attached heterologous proteins in the yeast cell wall can be controlled by the specific 439 440 anchoring domain fused to them. A reporter protein (eGFP) was predominantly localized to the external surface of the cell wall when fused with the Sed1-anchoring domain, whereas the 441 442 anchorage position of eGFP fused with the Sag1-anchoring domain was mainly inside of the cell 443 wall. By applying this anchorage position control technique, the cellulolytic ability of the 444 recombinant yeast strain co-displaying EG and BGL was successfully improved. Although further 445 analyses using GPI-anchoring domains derived from a wide-range of GPI-CWPs are required to identify the determinants of GPI-attached protein anchorage positions, our novel strategy for 446 447 anchorage position control will enable the efficient utilization of the cell wall space for various 448 fields of yeast cell-surface display.

449

450 **Declaration of interest**

451 The authors declare that they have no competing interests.

452

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- 460

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Strains	Relevant genotype	Source
S. cerevisiae BY4741	$MATa$ his3 $\Delta 1$ leu2 $\Delta 0$ met15 $\Delta 0$ ura3 $\Delta 0$	Invitrogen
BY-eGFP-SSS	BY4741/pIeGFP-SSS	This study
BY-eGFP-SSA	BY4741/pIeGFP-SSA	This study
BY-eGFP-SSn	BY4741/pIeGFP-SS2	This study
BY-BG-SSS	BY4741/pIBG-SSS	Inokuma et al. (2016)
BY-EG-SSS	BY4741/pIEG-SSS	Inokuma et al. (2016)
BY-ESBS	BY-EG-SSS/pIL2BG-SSS	This study
BY-ESBA	BY-EG-SSS/pIL2BG-SSA	This study

Table 1 Characteristics of yeast strains used in this study

608 Figure legends

609

Fig. 1 Schematic summary of the construction of gene cassettes used in this study. (A) Gene cassettes for cell-surface display and the secretory production of eGFP. (B) Gene cassettes for the cell-surface display of BGL1. (C) Gene cassettes for cell-surface display of EGII.

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Fig. 2 Localization analyses of eGFP fused with Sed1- or Sag1-anchoring domains. **(A)** Fluorescence images of strains BY-eGFP-SSS, BY-eGFP-SSA, and BY-eGFP-SSn. The cells were incubated in YPD medium for 48 h, stained with FM4-64 (red) to visualize vacuolar membranes, and then observed using a confocal microscope. **(B)** Immunoelectron micrographs of strains BY-eGFP-SSS and BY-eGFP-SSA. The cells were immunogold-labeled with an antibody against GFP. The arrowheads indicate gold particles.

620

621 Fig. 3 Effects of anchorage position control on enzyme activities of cellulase-displaying yeasts. (A) 622 Comparison of cell-surface EG and BGL activities in strains BY-BG-SSS, BY-ESBS, and BY-623 ESBA after cultivation in YPD medium for 48 h. The relative EG activity of each strain is shown 624 as a fold-change in EG activity relative to the average level observed with the parental strain BY-625 EG-SSS. (B) Comparison of transcript levels of EGII- and BGL1-encoding genes in strains BY-626 ESBS and BY-ESBA after cultivation in YPD medium for 48 h. The relative transcript level of 627 each gene is shown as a fold-change in mRNA levels relative to the average level detected in strain 628 BY-ESBS. (C) Relative quantification of BGL1 in the cell walls of strains BY-ESBS and BY-629 ESBA by nanoscale ultra-pressure liquid chromatography electrospray ionization quadrupole time-630 of-flight tandem mass spectrometry (nano-UPLC-MS^E). The amount of BGL1 was normalized to 631 the dry cell weight of each strain. Data are presented as the means \pm standard deviation (n = 3).

- 633 Fig. 4 Time course of the simultaneous saccharification and fermentation of 100 g dry weight/L of
- 634 pretreated rice straw by strains BY-ESBS, BY-ESBA, and their parental strain (BY4741). A small
- 635 amount of a commercial cellulase cocktail (0.4 FPU/g-biomass) was added to the fermentation
- 636 mixture. Data are presented as the means \pm standard deviation (n = 3).

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640

641	Supplementary materials	
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Text S1 Plasmid construction and yeast transformation.

- **Table S1** Characteristics of integrative plasmids used in this study.

Table S2 PCR primers used in this study.

- 648 Fig. S1 Amino acid sequence of Sed1- and Sag1-anchoring domains used in this study. The C-
- 649 terminal GPI attachment site (the ω site) was marked in bold. The hydrophobic amino acid residues
- 650 in the ω -minus region are underlined.

Graphical abstract





Fig. 1



Fig. 2

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В



Fig. 3








Supplementary materials

Novel strategy for anchorage position control of GPI-attached proteins in the yeast cell wall using different GPI-anchoring domains

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Text S1 Plasmid construction and yeast transformation

The plasmids and primers used in this study are listed in supplemental data Tables S1 and S2, respectively. The plasmids for cell-surface display of the enhanced green fluorescent protein (eGFP) using Sed1-anchoring domain were constructed based on the plasmid pIEG-SS (Inokuma et al., 2014) containing the sequences for the S. cerevisiae SED1 promoter, the signal peptide (SP) sequence derived from Rhizopus oryzae glucoamylase (GLUASP), Trichoderma reesei EGII, S. cerevisiae SED1anchoring region, and SAG1 terminator as follows: Inverse PCR with the SED1a-F and GLUASP-R primers was performed to replace the EGII of pIEG-SS to eGFP. The DNA fragment encoding the eGFP was amplified from pGK426EGFP (Ishii et al., 2009) by PCR using the eGFP-F1 and eGFP-R1 primers. These fragments were ligated by the isothermal assembly method (Gibson et al., 2009), and the resulting plasmid was named pleGFP-SGS. Then, Inverse PCR with the SAG1t-F1 and eGFP-R2 primers was performed to replace the SED1-anchoring region of pIeGFP-SGS to the SAG1-anchoring region. The DNA fragment encoding the SAG1-anchoring region amplified from pIEG-TA (Inokuma et al., 2014) by PCR using the SAG1a-F1 and SAG1a-R primers. These fragments were ligated by the isothermal assembly method (Gibson et al., 2009), and the resulting plasmid was named pleGFP-SGA. Similarly, inverse PCR with the SAG1t-F2 and eGFP-R3 primers was performed to remove the SED1anchoring region of pIeGFP-SGS. This fragment was self-ligated by the isothermal assembly method (Gibson et al., 2009), and the resulting plasmid was named pleGFP-SGn. Finally, inverse PCR with the eGFP-F2 and Vector-R1 primers was performed to replace GLUASP of the plasmids pleGFP-SGS, pleGFP-SGA, and pleGFP-SGn to the SP sequence derived from Saccharomyces cerevisiae SED1 (SED1SP). The DNA fragment encoding the SED1 promotor and SED1SP was amplified from pIBG-SSS (Inokuma et al., 2016) by PCR using the SED1p-F and SED1SP-R primers. Then, the SED1 promotor-SED1SP fragment was ligated the vector fragments derived from pIeGFP-SGS, pIeGFP-SGA, and pIeGFP-SGn by the isothermal assembly method (Gibson et al., 2009), respectively. The resulting plasmids were named pleGFP-SSS, pleGFP-SSA, and pleGFP-SSn, respectively.

The plasmid for cell-surface display of *Aspergillus aculeatus* β-glucosidase 1 (BGL1) using Sed1anchoring domain was constructed as follows: Inverse PCR with the Vector-F and Vector-R2 primers was performed to replace the *HIS3* of pIBG-SSS (Inokuma et al., 2016) to the I2 region (a part of the 3' noncoding region of *YFL021W* and *YFL020C* genes) and *LEU2*. The DNA fragment encoding the I2 region and *LEU2* was amplified from pIL2GA-SS (Inokuma et al., 2015) by PCR using the I2-F and Leu2-R primers. These fragments were ligated by the isothermal assembly method (Gibson et al., 2009), and the resulting plasmid was named pIL2BG-SSS. Then, Inverse PCR with the SAG1t-F1 and BGL1-R primers was performed to replace the *SED1*-anchoring region of pIL2BG-SSS to the *SAG1*anchoring region. The DNA fragment encoding the *SAG1*-anchoring region amplified from pIBG-SA (Inokuma et al., 2014) by PCR using the SAG1a-F2 and SAG1a-R primers. These fragments were ligated by the isothermal assembly method (Gibson et al., 2009), and the resulting plasmid was named pIL2BG-SSA.

The plasmids pleGFP-SSS, pleGFP-SSA, and pleGFP-SSn were digested with *NdeI* within *HIS3*. Then, the linearized plasmids were transformed into *S. cerevisiae* BY4741 by the lithium acetate method (Chen et al., 1992) and integrated into the *HIS3* locus of the chromosomal DNA by homologous recombination, respectively. The resulting strains were designated BY-eGFP-SSS, BY-eGFP-SSA, and BY-eGFP-SSn, respectively. Similarly, the plasmids pIL2BG-SSS and pIL2BG-SSA were digested with *NdeI* within I2 region. Then, the linearized plasmids were transformed into the BY-EG-SSS strain (Inokuma et al., 2016) by the lithium acetate method (Chen et al., 1992) and integrated into the I2 region of the chromosomal DNA by homologous recombination, respectively. The resulting strains were designated BY-ESBS and BY-ESBA, respectively. A single integration of each plasmid into the chromosomal DNA was verified by diagnostic PCR (colony PCR) using primers with upstream and downstream sequences of *HIS3* locus (the His3-534-F and His3-1866-R primers) and I2 region (the I2-420-F and I2-1160-R primers), respectively (data not shown).

Plasmids	Relevant genotype	Source/references
pGK426-EGFP	URA3 PGK1 _P -eGFP-PGK1 _T	Ishii et al. (2009)
pleGFP-SSS	HIS3 SED1 _P -SED1 _{SP} -eGFP-SED1 _A -SAG1 _T	This study
pleGFP-SSA	HIS3 SED1 _P -SED1 _{SP} -eGFP-SAG1 _A -SAG1 _T	This study
pleGFP-SSn	HIS3 SED1 _P -SED1 _{SP} -eGFP-SAG1 _T	This study
pIEG-SS	HIS3 SED1 _P -GLUA _{SP} -T. reesei EGII-SED1 _A -SAG1 _T	Inokuma et al. (2014)
pIEG-TA	HIS3 TDH3 _P - GLUA _{SP} -T. reesei EGII-SAG1 _A -SAG1 _T	Inokuma et al. (2014)
pIBG-SS	HIS3 SED1 _P - GLUA _{SP} -A. aculeatus BGL1-SED1 _A -SAG1 _T	Inokuma et al. (2014)
pIBG-SA	HIS3 SED1 _P - GLUA _{SP} -A. aculeatus BGL1-SAG1 _A -SAG1 _T	Inokuma et al. (2014)
pIBG-SSS	HIS3 SED1 _P -SED1 _{SP} -A. aculeatus BGL1-SED1 _A -SAG1 _T	Inokuma et al. (2016)
pIEG-SSS	HIS3 SED1 _P SED1 _{SP} T. reesei EGIISED1 _A SAG1 _T	Inokuma et al. (2016)
pIL2GA-SS	LEU2 SED1 _P -GLUA _{SP} -SED1 _A -SAG1 _T	Inokuma et al. (2015)
pIBG-SA	HIS3 SED1 _P -GLUA _{SP} -A. aculeatus BGL1-SAG1 _A -SAG1 _T	Inokuma et al. (2014)
pIL2BG-SSS	LEU2 SED1 _P -SED1 _{SP} -A. aculeatus BGL1-SED1 _A -SAG1 _T	This study
pIL2BG-SSA	LEU2 SED1 _P -SED1 _{SP} -A. aculeatus BGL1-SAG1 _A -SAG1 _T	This study
1 aniloatus	Agnousillus agulantus T upperi Tuichedoung upperi	CLUA Dhi-orang orange

 Table S1 Characteristics of integrative plasmids used in this study.

A. aculeatus, Aspergillus aculeatus; T. reesei, Trichoderma reesei; GLUA, Rhizopus oryzae

glucoamylase; P, promoter; SP, signal peptide sequence; A, anchoring region; T, terminator

Table S2 PCR primers used in this study.

Primers	Sequence
SED1a-F	gcatggacgagctgtacaagggctcgagtaaattatcaactgtcc
GLUASP-R	ageteetegeeettgeteacacecatggagateteege
eGFP-F1	ccgcggagatctccatgggtgtgagcaagggcgagga
eGFP-R1	gttgataatttactcgagcccttgtacagctcgtccatgc
SAG1t-F1	tgctattctaaaacgggtactgtacagttagtacattgagtctaa
eGFP-R2	gagettttggegetegageeettgtacagetegtecatge
SAG1a-F1	gcatggacgagctgtacaagggctcgagcgccaaaa
SAG1a-R	ctcaatgtactaactgtacagtacccgttttagaatagcagg
SAG1t-F2	tggacgagctgtacaagtaaaacgggtactgtacagttagtacattgag
eGFP-R3	actgtacagtacccgttttacttgtacagctcgtccatgc
eGFP-F2	cctcgactactttggcccaagtgagcaagggcgagga
Vector-R1	gttaattttctatatccaatctggcgtaatagcgaagagg
SED1p-F	gaaatcggcaaaatccctta
SED1SP-R	ageteetegeeettgeteacttgggeeaaagtagtegagg
Vector-F	gaaacggccttacgacgtagcggatctatgcggtgtgaaatac
Vector-R2	tgttttgacgaggtattccctatggtgcactctcagtacaatctg
I2-F	tgtactgagagtgcaccatagggaatacctcgtcaaaacaagac
Leu2-R	tttcacaccgcatagatccgctacgtcgtaaggccgtttct
BGL1-R	gttgcaccttcgggagcg
SAG1a-F2	cgtcagetgecectteae
His3-534-F	gctttgtcttcattcaacgtttcc
His3-1866-R	cttgecacctatcaccacaactaac
I2-420-F	gaagcegegagtaegaacaatgatg
I2-1160-R	tggtattttcgtgagcaaacccaac
rt-ACT1-F	tggattccggtgatggtgtt
rt-ACT1-R	tcaaaatggcgtgaggtagaga
rt-BGL-F	cttccagggctttgtgatgtc
rt-BGL-R	aggtgatatcgccaggcatt
rt-EG-F	ggttgtttgtctttgggtgcttac
rt-EG-R	aattgagcatttgttggaccacctt

Fig. S1 Amino acid sequence of Sed1- and Sag1-anchoring domains used in this study. The C-terminal GPI attachment site (the ω site) was marked in bold. The hydrophobic amino acid residues in the ω -minus region are underlined.

Sed1-anchoring domain (337 a. a.)

KLSTVLLSAGLASTTLAQFSNSTSASSTDVTSSSSISTSSGSVTITSSEAPESDNGTSTAAPTETS TEAPTTAIPTNGTSTEAPTTAIPTNGTSTEAPTDTTTEAPTTALPTNGTSTEAPTDTTTEAPTTGL PTNGTTSAFPPTTSLPPSNTTTTPPYNPSTDYTTDYTVVTEYTTYCPEPTTFTTNGKTYTVTEPTT LTITDCPCTIEKPTTTSTTEYTVVTEYTTYCPEPTTFTTNGKTYTVTEPTTLTITDCPCTIEKSEA PESSVPVTESKGTTTKETGVTTKQTTANPSLTVSTVVPVSSSASSHS<u>VVI</u>NS**N**GANVVVPGALGLA GVAMLFL*

Sag1-anchoring domain (320 a. a.)

SAKSSFISTTTDLTSINTSAYSTGSISTVETGNRTTSEVISHVVTTSTKLSPTATTSLTIAQTSI YSTDSYITVGTDIHTTSEVISDVETISRETASTVVAAPTSTTGWTGAMNTYISQFTSSSFATINST PIISSSAVFETSDASIVNVHTENITNTAAVPSEEPTFVNATRNSLNSFCSSKQPSSPSSYTSSPLV SSLSVSKTLLSTSFTPSVPTSNTYIKTKNTGYFEHTALTTSSVGLNSFSETAVSSQGTKIDTFLVS SLIAYPSSASGSQLSGIQQNFTSTSLM<u>I</u>ST<u>Y</u>E**G**KASIFFSAELGSIIFLLLSYLLF*

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