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Title	The relationship between the fine structure of amylopectin and the 1 type of crystalline allomorph of starch granules in rice endosperm
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Description	



1	The relationship between the fine structure of amylopectin and the
2	type of crystalline allomorph of starch granules in rice endosperm
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- 37 Abstract
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Background and Objectives: It is known that any wild-type varieties of cereals so far examined have A-type starch crystals in their endosperm whereas their *starch branching enzyme 2b* (*be2b*) mutants that are usually referred to as *amylose-extender* (*ae*) mutants produce B-type starch crystals. The present study aimed to examine the structural features of amylopectin which are responsible for the A-type or B-type crystalline allomorphs of starch granules by using starches in wild-type *japonica* and *indica* rice varieties and their *be2b* (*ae*) mutants. **Findings:** The average length of chains of A-type amylopectin was markedly shorter than

Findings: The average length of chains of A-type amylopectin was markedly shorter than 46 that of B-type amylopectin. It was also thought that A-type amylopectin had two types of 47 branches, namely the first branches were formed mainly by BEI in the basal region of the cluster and the second branches were synthesized specifically by BEIIb inside the cluster. 48 49 These differences caused the average length of double helices formed by cluster 50 constituent chains to be shorter in wild-types than that in their *be2b* mutants, and these 51 differences changed the crystalline allomorph of starch granules to B-type in the mutants 52 from A-type in their wild-types, as examined by wide angle X-ray diffraction (XRD) and 53 sum frequency generation spectroscopy (SFG).

54 **Conclusions:** Combined with these observations and calculations of chain-length of 55 hypothetical A chains in four types of amylopectin molecules, it was concluded that both 56 the average length of external segments of cluster chains and the formation of the second 57 branches inside the cluster greatly affect the crystal types of starch granules.

58 **Significance and Novelty:** The novel findings in the present study can provide new 59 insights into the structural features of amylopectin which determines the A-type or B-type 60 crystalline allomorph of starch granules of cereals.

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63 Keywords A-type starch; B-type starch; rice endosperm; starch branching enzyme

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Abbreviations *ae*, amylose extender; APTS, 8-amino-1,3,6-pyrenesulfonic acid; BE,
starch branching enzyme; DBE, starch debranching enzyme; DP, degree of
polymerization; FACE, fluorophore-assisted carbohydrate electrophoresis; Φ-LD,
phosphorylase-limit dextrins; PaISA, Isoamylase from *Pseudomonas amyloderamosa*;
SFG, sum frequency generation spectroscopy; SS, starch synthase; XRD, wide angle Xray diffraction

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74 **1 INTRODUCTION**

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76 Rice plants have been widely used as materials for studies on starch biosynthesis. Rice 77 varieties are largely classified into *japonica*-type and *indica*-type varieties. Past 78 investigations have revealed that starch is synthesized by concerted actions of starch 79 synthase (SS), BE, and starch debranching enzyme (DBE). Although three starch 80 biosynthetic enzymes have multiple isozymes, their roles and expressions in various 81 organs and tissues are greatly different in rice plants (Ohdan et al., 2005). It has been 82 established that *japonica*-type rice is deficient in SSIIa, whereas *indica*-type rice has a 83 full activity of SSIIa (Umemoto et al., 2002; Nakamura et al., 2005).

84 Starch is composed of amylopectin, a highly branched glucan, and amylose, an 85 essentially linear glucan, and the amount of amylopectin usually accounts for 65-85% of the total starch. Amylopectin is known to have a specific fine structure. This glucan has 86 87 a structural element called "cluster", which is interconnected by long chains and aligned 88 basically in tandem fashion (French, 1972; Nakamura & Kainuma, 2022). Each cluster is 89 composed of A chains (non-branched chains) and B1 chains (branched by at least one 90 chains) and each cluster is interconnected by long chains designated as B2 chains and/or 91 B3/B4 chains that span to two/three clusters and/or three clusters, respectively (Peat et 92 al., 1952; Hizukuri, 1986). Very importantly, when non-branched segments of 93 neighboring chains of the cluster exceed 10 glucosyl units or degree of polymerization 94 (DP) of 10, they form double helices (Gidley & Bulpin, 1987). The presence of double 95 helices in amylopectin molecules dramatically affects physicochemical properties of 96 starch granules including amylopectin and amylose molecules. The clusters are packed 97 by the lateral alignment of neighboring double helices (Kainuma & French, 1972; 98 Yamaguchi et al., 1979; French, 1984). It is widely known that starch granules in cereal 99 endosperm show the A-type crystalline polymorph whereas some tubers and rhizomes 100 exhibit the B-type crystalline polymorph, which can be clearly distinguished by X-ray 101 diffraction analysis (see reviews by Hizukuri, 1996; Buléon et al., 1998). The A-type 102 starch is composed of a monoclinic unit cell whereas the B-type starch has a hexagonal 103 unit cell, and thus the A-type starch is considered to be more densely packed compared 104 with the B-type starch (see review by Imberty et al., 1991).

105 The phenotypes of BEIIb-deficient mutants of cereals are often designated as *ae* because 106 the *ae* mutant starches have apparently the high-amylose contents in the endosperm (see 107 review by Shannon et al., 2009). Interestingly, the *ae* starches in cereal endosperm show 108 the B-type crystalline allomorph (Gérard et al., 2000; Nishi et al., 2001; Tanaka et al.,

- 109 2004). The be2b (ae) mutants in *japonica*-type rice endosperm (Yano et al., 1985) have 110 a modified amylopectin chain profile because it contains more long B chains and fewer 111 short chains of DP \leq ca. 13 (Nishi et al., 2001; Nakamura et al., 2022). This change is 112 caused by loss of the distinct role of BEIIb, playing an essential role in the synthesis of 113 short chains in the region in the crystalline lamella of amylopectin cluster (Jane et al., 114 1997; Nakamura et al., 2020). Based on these observations, recently we proposed that 115 amylopectin in endosperm of japonica-type rice and its be2b mutant is referred to as A-116 type amylopectin and B-type amylopectin, respectively (Nakamura et al., 2022). On the 117 other hand, *indica*-type amylopectin has fewer short chains of DP ≤ 10 and more 118 intermediate chains of DP 12-24 than *japonica*-type amylopectin whereas the proportion 119 of long chains is unchanged between these amylopectin (Umemoto et al., 1999; 120 Nakamura et al., 2002). However, *indica*-type starch granules exhibit A-type allomorph 121 as *japonica*-type starch granules (Hizukuri, 1996). In this sense, *indica*-type amylopectin 122 should be classified into an A-type amylopectin.
- 123 What is a criterion to distinguish the structural difference between A-type and B-type 124 amylopectin? In the present study, to analyze the structural difference between A-type 125 amylopectin and B-type amylopectin, the fine structure of endosperm amylopectin of 126 be2b (ae) mutants from both japonica rice and indica rice was analyzed in details, because 127 both *be2b* mutant starch granules would have B-type crystalline allomorph, whereas the 128 A-type amylopectin structure is known to be greatly different between *japonica* and 129 indica rice varieties. Therefore, these materials used in this study are extremely beneficial 130 to reveal the relationship between the fine structure of amylopectin and the crystalline 131 allomorph of starch granules. In addition, to analyze the internal structure of amylopectin, 132 chain-length distribution of its phosphorylase-limit dextrin (Φ -LD) was examined. To 133 reveal the relationship between the fine structure of amylopectin and the internal starch structure of starch granule, starch crystalline structures were also examined using XRD 134 135 and SFG analysis (Miyauchi et al., 2006; Kong et al., 2014).
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138 2 MATERIALS AND METHODS

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140 **2.1 Reagents**

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- 142 A fluorophore 8-amino-1,3,6-pyrenesulfonic acid (APTS) was obtained from AB SCIEX
- 143 (Tokyo, Japan). α -Glucan phosphorylase from rabbit muscle was purchased from SIGMA.

144	Isoamylase from Pseudomonas amyloderamosa (PaISA) was kindly provided from
145	Hayashibara Co., Ltd., Japan.
146	
147	2.2 Plant materials and sampling
148	
149	Two be2b mutant lines EM10 and IR36ae were generated from a japonica-type cultivar
150	Kinmaze and an <i>indica</i> -type cultivar IR36, respectively. Mature seeds were also
151	harvested from each line which was grown in the experimental field of Akita Prefectural
152	University under natural environmental conditions during summer months, and stored at
153	8°C before use.
154	
155	2.3 Preparation of starch granules from rice endosperm
156	
157	Starch granules from randomly chosen mature endosperms of each line were prepared as
158	described previously (Nakamura et al., 2020).
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160	2.4 Chain-length distribution analysis of amylopectin
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162	The chain-length distribution of amylopectin was analyzed by using the fluorophore-
163	assisted carbohydrate electrophoresis (FACE) method (O'Shea et al., 1998) after
164	treatment of amylopectin with PaISA to remove α -1, 6-glucosidic linkages, followed by
165	labelling of APTS at the reducing ends of debranched glucan chains, as described
166	previously (Nakamura et al., 2020).
167	
168	2.5 Chain-length distribution analysis of phosphorylase-limit dextrins (Φ -LD) of
169	amylopectin
170	
171	Chain-length distribution of Φ -LD was analyzed to examine the internal structure of
172	amylopectin. In this analysis, amylopectin was treated with a rabbit muscle phosphorylase
173	to form its Φ -LD, as reported previously (Sawada et al., 2014). The Φ -LD was
174	debranched and its chain-length distribution was analyzed as above.
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176	2.6 X-ray diffraction (XRD) pattern analysis of starch granules
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178	XRD measurements were performed on a Nano-viewer system (Rigaku Co., Tokyo,
179	Japan) at a wavelength of 0.154 nm (CuK α). The camera lengths were 75 mm. A Pilatus

180 1M (Dectris AG, Baden, Switzerland) detector was used, with a q range of 3.5 to 25 nm⁻¹; q is the magnitude of the scattering vector and is defined as follows:

182 $q = 4\pi \sin \theta / \lambda$ (1),

183 where 2θ and λ are the scattering angle and wavelength, respectively. The starch granule 184 samples were put into the sample cell of approximately 500 µm thickness. Data 185 processing, which included controlling the contrast of the 2D-patterns and the preparation 186 of a 1D-profile from the obtained 2D-patterns, was performed using the FIT-2D software 187 (Ver. 12.077, Andy Hammersley/ESRF, Grenoble, France).

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2.7 Optical sum frequency generation (SFG) spectroscopy of starch granules

191 The powdered starch samples of Kinmaze, IR36, EM10, and IR36ae were put in 192 transparent silica glass square cells (AS ONE Q-101; $3.5 \text{ mm} \times 12.5 \text{ mm} \times 45 \text{ mm}$). The 193 internal sizes of the cells were 1 mm in thickness and 10 mm in width. SFG of our sample 194 was observed through the glass window of the cell. The SFG spectroscopy system was 195 the same as previously described (Hieu et al., 2015; Nakamura et al., 2020). Tunable 196 infrared light pulses at wavelength of approximately 3 µm was output from an optical 197 parametric generator (EKSPLA PG401/DFG2-18P) pumped by the fundamental and third 198 harmonic output of a Nd³⁺:YAG laser (EKAPLA PL2143B) with time width 30 ps and 199 repetition rate of 10 Hz. The pulse energy of the visible light was about 10 µJ and that of 200 the infrared was about 260 µJ at the sample. The spectral width of the IR light was 6 cm⁻ 201 ¹. For more details see our previous paper (Nakamura et al., 2020).

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2043 RESULTS

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3.1 Comparison of chain-length distribution of amylopectin in mature endosperm of *be2b* mutant lines and their parent cultivars of *japonica*-type rice and *indica*-type rice

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It is widely known that starch granules including amylopectin in mature endosperm of wild-type rice from both *japonica*-type and *indica*-type varieties have the A-type crystalline allomorph. We previously proposed that the rice amylopectin structure is referred to as the A-type amylopectin or B-type amylopectin based on the criterion that its starch granules show the A-type or B-type crystalline allomorph, such as wild-type or *be2b* mutants, respectively (Nakamura et al., 2022). To characterize the structural features

- 216 of amylopectin, its chain-length distribution from a *japonica*-type line Kinmaze and an 217 indica-type line IR36 was measured and compared by the FACE method. Both chain 218 profiles have two major peaks; a large peak consisting of short chains of having DP 6 to 219 approximately 34 (A and B1 chains) and a small peak consisting of long chains having 220 DP > approximately 37 (cluster-interconnecting B2-B4 chains) (Figs. 1A and 1B). Figs. 221 1C and 1D show that although both of their *be2b* mutant (*ae* mutant) lines EM10 and 222 IR36ae also had two major peaks, the proportion of long chains (B2-B4 chains) to short 223 chains (A and B1 chains) was markedly increased by the loss of BEIIb activity 224 (Supplementary Table S1).
- 225 Fig. 2 shows the differences in endosperm amylopectin chain-length distribution 226 between *japonica*-type and *indica*-type lines and between wild-type and *ae* type lines. 227 The amylopectin in *japonica*-type *ae* mutant line EM10 had fewer very short chains of 228 DP 6-12 with a peak of DP approximately 8-10 and more long chains of DP \geq about 37 and intermediate chains of DP 14-30 compared with Kinmaze amylopectin (Fig. 2A). 229 230 Notably, the similar pattern of differences in chain-length distribution of amylopectin 231 between *indica*-type mutant line IR36ae and its wild-type line IR36 was detected, 232 although the extent of differences between *indica*-type lines was significantly lower than 233 that between *japonica*-type lines (Figs. 2A and 2B). The pattern of difference in chain-234 length distribution of amylopectin between the SSIIa-deficient *japonica*-type rice and the 235 SSIIa-active indica-type rice was clearly different from that between the ae-mutant and 236 its wild-type (Compare Fig. 2B with Figs. 2A and 2C). The proportion of very short 237 chains of DP 6-10 was higher but that of intermediate chains of DP 13-24 was lower in a 238 japonica-type line Kinmaze than an indica-type line IR36, whereas there was no 239 significant difference of B2-B4 chains between two lines (Fig. 2B and Supplementary 240 Table S1), consistent with our previous studies (Umemoto et al., 1999; Nakamura et al., 241 2002, 2005).
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243**3.2 Comparison of chain-length distribution of phosphorylase-limit dextrins (\Phi-LD)244of amylopectin in mature endosperm of** *be2b* **mutant lines and their parent cultivars245of** *japonica***-type rice and** *indica***-type rice**

246

To examine the internal structure of amylopectin more in details, the external segments of chains were digested by a rabbit muscle phosphorylase a (SIGMA) and then the chainlength distribution of the resulting Φ -LD was determined by the FACE method. In this analysis, almost all of the chains of DP 4 are considered to be derived from A chains. According to this criterion, the amounts of A chains (DP 4-chains) were approximately 252 52%, 57%, 52%, and 55% in Kinmaze, EM10, IR36, and IR36ae, respectively (Fig. 3), 253 indicating that the proportion of A chains in amylopectin was significantly higher in either 254 *be2b* mutant than that in its wild-type. Fig. 4 compares the chain profiles of Φ -LD. 255 Compared with wild-type Kinmaze, Φ -LD of its *be2b* mutant EM10 had fewer chains of 256 DP 6-18 and enriched chains of DP 24-37 (Fig. 4A). The result suggests that the wild-257 type amylopectin had more branches inside the cluster, and therefore some chains carried 258 multiple branched chains. This resulted in longer internal chain-length in Kinmaze amylopectin than EM10 amylopectin. Fig. 4B shows that Φ -LD from *japonica*-type 259 260 Kinmaze had more A chains but slightly less short internal segments of DP 6-9 than that 261 from *indica*-type IR36. This suggests that the IR36 amylopectin had slightly more B1 262 chains with short internal segments of DP 6-9 to some extent than the Kinmaze 263 amylopectin. Figure 5 illustrates the structural features of amylopectins of Kinmaze and 264 IR36 and their *ae* mutants, although the difference of amylopectin between EM10 and 265 IR36ae was too small to present it in the figure.

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3.3 X-ray diffraction (XRD) patterns of starch granules in in mature endosperm of *be2b* mutant lines and their parent cultivars of *japonica*-type rice and *indica*-type rice

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271 Fig. 6 shows the XRD profiles of various starch granules in mature endosperms of 272 Kinmaze, IR36, EM10 and IR36ae. The XRD profile of Kinmaze (Fig. 6A) and IR36 273 (Fig. 6B) had peaks at the scattering vector, q, of approximately 10.58 (a single peak), 12.03 and 12.71 (doublet peaks), and 16.18 nm⁻¹ (a single peak), which are characteristics 274275 of A-type starch granules in cereal endosperm. On the other hand, the XRD profile of 276 EM10 (Fig. 6C) and IR36ae (Fig. 6D) had peaks at q of approximately 3.724 (a single peak), 11.38 (a single peak), 15.54 and 16.92 nm⁻¹ (doublet peaks), which are 277 characteristics of B-type starch granules (Nagasaki et al., 2021; Nakamura et al., 2022). 278 279

3.4. Optical sum frequency generation (SFG) spectroscopy of starch granules in mature endosperm of *be2b* mutant lines and their parent cultivars of *japonica*-type rice and *indica*-type rice

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Figure 7 shows SFG spectra obtained for starch granules in mature endosperms of Kinmaze, EM10, IR36, and IR36*ae*. The curves represent the SFG intensity fit to Lorentzian curves in Eq. (1) below with fitting parameters in Table S1.

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Big peaks at around 2910 cm⁻¹ and 2970 cm⁻¹ were previously assigned to C-H and C-H₂ stretching vibrations, respectively (Kouyama et al., 2016), but the appearance of the separate peaks around 2870 cm⁻¹ suggests that we should reconsider the assignments. For the present we assigned the peaks at 2870 cm⁻¹, 2910 cm⁻¹, and 2960 cm⁻¹ as CH or CH₂ vibrations. The broad peak around 3100cm⁻¹ was tentatively assigned to H₂O peak in our previous paper (Nakamura et al., 2020).

294 In our previous paper (Nakamura et al., 2020) we reported that starch from the matured 295 endosperms of Kinmaze and EM10 show different SFG spectra according as they are A-296 and B-type, respectively (Kong et al., 2014). This difference was reproduced in Fig. 7A 297 (Kinmaze) and 7B (EM10). Figures 7A and 7D, SFG spectra of starch granules in mature 298 endosperms of IR35, an indica-type cultivar, and IR36ae, a be2b mutant line of IR35, had 299 similar shapes to Figs. 7A and 7B, respectively. This tendency was also reflected in the 300 list of resonant frequencies in Supplementary Table S2. Namely, the SFG spectra of Kinmaze and IR36 had three resonant oscillators between 2900 cm⁻¹ and 3000 cm⁻¹, while 301 302 those of EM10 and IR36ae have two. Thus, it was confirmed that the effect of BEIIb was 303 clearly correlated with the SFG spectral shapes.

304 305

306 4 DISCUSSION

307

4.1 The structural features of A-type and B-type amylopectin in rice endosperm 309

310 The present paper analyzed the fine structures of A-type and B-type amylopectins that 311 give rise to A-type and B-type crystals, respectively, of starch granules of rice (Figs. 1-312 4). The results confirmed that both *japonica*-type and *indica*-type amylopectin had two 313 types of branches, while branches of their be2b mutant ones were present only in the 314 amorphous lamellae (Fig. 5), consistent with our previous study (Nakamura, 2002, 2015, 315 Nakamura et al., 2005, 2020, 2022). This model well explains the reason why the average 316 length of external segments of cluster chains is longer in the B-type amylopectin in be2b 317 mutants than that in the A-type amylopectin in their wild-types.

It is known that BEIIb is specifically expressed in rice endosperm, affecting the structural features of amylopectin and crystalline properties of starch granules (Nishi et al. 2001; Nakamura 2018; Nakamura et al. 2020; Ying et al. 2022). When BEIIb activity was lost in the endosperm, chains of amylopectin became longer (Fig. 2), resulting in formation of longer double helices, and this caused starch granules properties to be resistant to thermal gelatinization and the crystalline allomorphs to change to the B-type

- from the A-type, as determined by XRD (Fig. 6) and SFG analyses (Fig. 7), consistent with our recent studies (Nakamura et al., 2020; Ying et al., 2022; Zhang et al., 2022) and past investigations by other groups worldwide (Wei et al., 2010; Butardo et al., 2011).
- 327

328 4.2 Structural boundary between the A-type amylopectin and the B-type 329 amylopectin

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331 Although amylopectin chains from an *indica*-type rice were longer than those from a 332 japonica-type amylopectin in wild-type, both starches showed A-type crystalline 333 allomorph (Figs. 6 and 7). The fine structure of amylopectin was very similar between 334 the *be2b* mutants generated from the *japonica*-type line EM10 and the *indica*-type line 335 IR36ae (Fig. 2D), and in fact their starch granules exhibited the B-type allomorph (Figs. 336 6 and 7). However, it is noted that in terms of chain-length profile, the *indica*-type IR36 337 amylopectin appeared to be more similar to the be2b mutant amylopectin compared with 338 the *japonica*-type Kinmaze one (Compare Fig. 2C with Fig. 2A, Supplementary Table 339 S1). The analysis of Φ -LD of amylopectin can give us an invaluable information on the 340 features of amylopectin internal structure as well as the proportion of and chain-length 341 distribution of A-chains. The Φ -LD of the *be2b* mutant amylopectin had depleted short 342 B chains of DP approximately 6-20 and more long B chains of DP about 24-37 compared 343 to wild-type amylopectin (Fig. 4A). The result suggests that B chains of the be2b mutant 344 amylopectin had longer internal segments than those of its wild-type amylopectin. This 345 is consistent with an assumption that B1 chains of wild-type amylopectin carry branches 346 inside (the crystalline lamellae) as well as amorphous lamellae of the cluster, whereas B 347 chains of the *be2b* mutant amylopectin have branches almost in the amorphous lamellae, 348 as illustrated in Fig. 5. This idea on the structural change of the cluster in the *ae* mutant 349 is consistent with the cluster structure proposed by Waigh et al., (2000) that the average 350 length of the amylopectin helices is longer in B-type starches compared with that in A-351 type starches. The proportion of A chains was significantly higher in the be2b mutant 352 amylopectin than that in the wild-type amylopectin (Fig. 3 and 4). The results suggest 353 that new branches in the amorphous lamellae were more easily formed in B chains than 354 in A chains compared with those in the crystalline lamellae, because A chains become B 355 chains when they are used as acceptor chains.

Irrespective of phosphorolysis of amylopectin, number of total chains are unchanged. It is also considered that the number of cluster-interconnecting chains is unchanged by the treatment of amylopectin with phosphorylase. Therefore, amylopectin chain length distribution can be extrapolated by adding chain-length of DP 12 to each chain of Φ -LD. 360 In this way, the hypothetical chain profiles for A chains can be calculated. Fig. 8 compares 361 the chain-length distribution of possible A chains of amylopectin from 4 lines used in this study. If this hypothesis is correct, amylopectins of both wild-types and their be2b mutants 362 had A chains of DP approximately 6-18. Although it is known that short chains of $DP \le 9$ 363 364 are unable to form double helices (Gidley and Bulpin, 1987), the proportion of these short 365 chains in total amylopectin chains were only 15.7, 4.7, 7.8, and 4.7% in Kinmaze, EM10, IR36, and IR36ae, respectively (Fig. 8). This shows that most of A chains of amylopectin 366 367 in these lines participated in the formation of double helices, facilitating crystalline properties of starch granules. However, it is noted that the proportion of these chains of 368 369 DP 6-9 was greatly lower while the proportion of DP 10-18 chains and in particular, that 370 of DP 17 and 18 chains were significantly higher in the be2b (ae) mutants than that in 371 wild-types. The results indicate that *be2b* mutant amylopectin had longer double helices 372 than wild-type amylopectin. Therefore, it is concluded that all these structural features of 373 amylopectin determined starch granules to be the A-type or the B-type crystalline 374 allomorph. It is widely known that that the crystal type of starch is largely dependent of 375 length of external segments of amylopectin. Generally, shorter chains favor the formation 376 of A-type allomorph whereas longer chains contribute to the formation of B-type 377 allomorph (Hizukuri et al., 1983; Hizukuri, 1996). It was also shown that amylose chains 378 of DP 10-13 exhibit A-type crystals, while those of DP \geq 14 yield B- or C-type crystals 379 (Gidley and Bulpin, 1987; Pfannemüller, 1987). The ratio of number of chains DP10-380 13/number of chains of 13-18 was calculated to be approximately 1.59 and 1.30 in possible A-chains from Kinmaze and IR36, respectively, whereas that was approximately 381 382 0.97 and 0.87 in those from EM10 and IR36ae, respectively (Fig. 8), consistent with results reported previously cited above. In summary, it was confirmed that the length of 383 384 external segments of chains of amylopectin is a determinant of the crystalline allomorph 385 of starch granules.

386 Based on the present results, firstly, the average length of external segments of cluster 387 chains with DP up to approximately 24 greatly affects the type of crystalline allomorph 388 of starch granules. Secondly, it can be also pointed out that the presence of second 389 branches in the cluster synthesized by BEIIb would play an important role in the crystallization of amylopectin double helices in the A-type crystal. Currently, it is well 390 391 known that *ae* starch is highly resistant to thermal gelatinization and hydrolysis by 392 hydrolytic enzymes compared to other starches (Tsuiki et al., 2018; Wang et al., 2017; 393 Nakamura, 2018), and therefore a better understanding of the relationship between 394 structures and functional properties of starches having various structures will be 395 important for the use of starch for food and industrial applications.

396 397

398 **5 CONCLUSION**

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400 The present study aimed to examine the structural features of A-type or B-type 401 amylopectin which are responsible for the A-type or B-type crystals, respectively, of 402 starch granules by using starches in *japonica* and *indica* rice wild-type varieties and their 403 be2b (ae) mutants, because it is known that any wild-type varieties of cereals so far 404 examined have A-type starch crystals in their endosperm whereas their be2b (ae) mutants 405 produce B-type starch crystals (Hizukuri, 1996; Shannon et al., 2009). The present 406 comparative studies with starches from both *japonica*-type and *indica*-type rice wild-type 407 varieties and their be2b mutants could provide us with useful criteria to distinguish A-408 type amylopectin from B-type amylopectin, because it is widely known that the fine 409 structure of amylopectin in starch from *japonica*- and *indica*-type rice endosperm greatly 410 differs from each other (Umemoto et al., 1999; Nakamura et al., 2002, 2005). Analysis of 411 the chain-length distribution of component chains of amylopectin and their Φ -LD 412 (internal chains) indicated that the average length of cluster chains (A and B1 chains) of 413 amylopectin was clearly longer in B-type amylopectin found in both *be2b* mutants than 414 that in A-type amylopectin formed in their wild-types (Figs. 1-4 and Supplementary Table 415 S1). The results can be explained by a specific role of BEIIb in the formation of the second 416 branches inside the cluster. In the *be2b* mutants the second branches were deficient in the amylopectin cluster, and thus the average length of cluster chains became longer than that 417 in wild-types (Fig. 5). On the other hand, although chain-length of cluster chains in IR36 418 419 amylopectin was apparently longer than that in Kinmaze amylopectin (Fig. 2B), both 420 amylopectins were considered to have the second branches formed by BEIIb as well as 421 the first branches formed mainly by BEI, as presented by Figs. 5A and 5B. The present 422 study strongly suggests that the second branches synthesized by BEIIb would play an 423 important role in allying double helices of amylopectin in compact and regular manner, 424 and this results in the crystallization of amylopectin chains in the A-type allomorph. This 425 study claims that not only the average chain-length of cluster chains but also the presence 426 of the second branches in the cluster are important for the A-type crystalline allomorph 427 of starch granules.

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565 SUPPORTING INFORMATION

- 566 Additional supporting information can be found online in the Supporting Information 567 section at the end of this article.
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573 Legends for figures

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575 Fig. 1 Chain-length distribution of amylopectin in mature endosperms of a wild-type 576 japonica cultivar Kinmaze, a wild-type indica cultivar IR36, and their be2b mutant lines, 577 EM10 and IR36ae, respectively. The vertical axis presents the proportion (molar %) of 578 the amount of each chain to the total amounts of chains with degree of polymerization 579 (DP) from 6 to 90, whereas the horizontal axis shows the DP value of the chain. 580 Amylopectin of Kinmaze (A), IR36 (B), EM10 (C), and IR36ae (D). The experiments 581 were repeated at least three times until all these results were consistent, whereas each 582 figure shows one representative result. Values are the averages calculated from three 583 replicate measurements. Standard deviations were too small to be shown in the figure.

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586 Fig. 2 Difference in amylopectin between EM10 and Kinmaze, calculated from data of 587 EM10 subtracted by those of Kinmaze. A, Difference in amylopectin between EM10 and 588 Kinmaze, calculated from data of EM10 subtracted by those of Kinmaze. B, Difference 589 in amylopectin between Kinmaze and IR36, calculated from data of Kinmaze subtracted 590 by those of IR36. C, Difference in amylopectin between IR36ae and IR36, calculated 591 from data of IR36ae subtracted by those of IR36. D, Difference in amylopectin between 592 EM10 and IR36ae, calculated from data of EM10 subtracted by those of IR36ae. The 593 other conditions are the same as in Fig. 1.

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Fig. 3 Chain-length distribution of phosphorylase-limit dextrins (Φ -LD) of amylopectin in mature endosperms of a wild-type *japonica* cultivar Kinmaze, a wild-type *indica* cultivar IR36, and their *be2b* mutant lines, EM10 and IR36*ae*, respectively. Φ -LD of Kinmaze (A), IR36 (B), EM10 (C), and IR36*ae* (D). The other conditions are the same as in Fig. 1, except that data were obtained for chains with degree of polymerization (DP) from 4 to 60.

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Fig. 4 Comparison of chain-length distribution of Φ -LD of amylopectin in mature endosperms of a wild-type *japonica* cultivar Kinmaze, its *be2b* mutant line, EM10, a wild-type *indica* cultivar IR36, and its *be2b* mutant line, IR36*ae*. A, Difference in Φ -LD between EM10 and Kinmaze, calculated from data of EM10 subtracted by those of Kinmaze. B, Difference in Φ -LD between Kinmaze and IR36, calculated from data of

609	Kinmaze subtracted by those of IR36. C, Difference in Φ -LD between EM10 and IR36 <i>ae</i> ,
610	calculated from data of EM10 subtracted by those of IR36ae. The other conditions are the
611	same as in Fig. 3.
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613	
614	Fig. 5 A schematic representation of the hypothetical cluster structure of amylopectin in
615	endosperm from a wild-type <i>japonica</i> cultivar Kinmaze (A), a wild-type <i>indica</i> cultivar
616	IR36 (B), and their be2b mutant lines, EM10 (C) and IR36ae (D), respectively.
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619	Fig. 6 Wide angle X-ray scattering (WAXS) analysis of starch granules in mature
620	endosperms from a wild-type <i>japonica</i> cultivar Kinmaze (A), a wild-type <i>indica</i> cultivar
621	IR36 (B), and their be2b mutant lines, EM10 (C) and IR36ae (D), respectively. See
622	Materials and methods in details.
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625	Fig. 7 SFG spectra of starch granules in mature endosperms from a wild-type <i>japonica</i>
626	cultivar Kinmaze (A), a wild-type <i>indica</i> cultivar IR36 (B), and their <i>be2b</i> mutant lines,
627	EM10 (C) and IR36 <i>ae</i> (D), respectively. See Materials and methods in details.
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630	Fig. 8 The hypothetical chain-length distribution of A chains in a wild-type <i>japonica</i>
631	cultivar Kinmaze (A), a wild-type <i>indica</i> cultivar IR36 (B), and their <i>be2b</i> mutant lines,
632	EM10 (C) and IR36 <i>ae</i> (D), respectively. The chain-length distribution of hypothetical A
633	chains in the DP range of 6-18 were obtained by the difference of chain-length distribution
634	between native anylopectin (DP 6-60) and its Φ -LD (DP > 5), although each DP of Φ -
635	LD was added by DP12. See text in details.
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Degree of polymerization (DP)





Kinmaze (*japonica*)

IR36 (*indica*)



EM10



IR36ae





