Introduction

Rice (*Oryza sativa* L.) is very important as a staple food of the world, especially in Southeast Asia. Rice grains sometimes show some defective phenotypes, resulting in reduced eating quality: notched grain caused by unbalanced development of the caryopsis compared to glume size (Lin et al. 2017, Takeda and Takahashi 1970), reduction of grain size caused by high night temperatures (Morita et al. 2005), and chalkiness induced by some environmental factors including high temperature, high humidity, or cultural practices at the ripening stage (Zhou et al. 2015). These grain defects have been known to be controlled by genetic factors (Kobayashi et al. 2007, Takeda and Takahashi 1970). To improve these grain qualities, the mechanisms generating these defects need to be analyzed. For this, a sectioning sample of the grain is very useful, as Morita et al. (2005) examined the endosperm cell size and number in a cross-section of a rice grain matured under high night temperatures. They also determined the endosperm cell size and number using digitalized hand-tracing images of 20-μm thick cross-sections. For this analysis, many thin cross-section samples are needed for confirming the influence of some stresses. In this regard, it is important to rapidly produce many thin cross-section samples.

Kawamoto (2003) reported the method for preparing thin frozen-sections using an adhesive film and producing almost intact sections, which were very thin and little damaged, from many organisms such as animals, plants and insects.

In this report, we tried to obtain the cross and longitudinal sections of a whole matured rice grain using cryomicrotome by the Kawamoto method using the specific adhesive film (Kawamoto and Kawamoto 2014). We were able to obtain almost perfect sections of the whole grain with high-quality cell morphology and orientation.

Materials and Methods

The rice grain used in the study was *japonica* variety, Koshihikari. The grains were dipped in water at room temperature overnight, then subjected to sectioning by cryomicrotome according to the Kawamoto method (Kawamoto and Kawamoto 2014) as follows:

1. The water absorbed rice grain was dipped into dry-ice hexane to freeze quickly.
2. SCEM medium (SECTION-LAB Co. Ltd., Japan) for embedding was poured into a stainless steel container.
3. The frozen grains were buried in the medium.
4. The container was quickly subjected to freezing in dry-ice hexane.
5. The frozen specimen block was taken out of the container.
6. The block was fixed on the cryomicrotome sample holder.
7. The sample holder was set to sample holder chuck of cryomicrotome.

Note

Easy sectioning of whole grain of rice using cryomicrotome

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To obtain a clear intact section of a ripened rice grain, which is suitable for biochemical and histological analysis, the Kawamoto method using a specific adhesive film was applied using a cryomicrotome. The longitudinal and sagittal sections were easily obtained together with the cross-section, and cell characteristics were clearly discerned in the ripened grain. It was demonstrated that the Kawamoto method is readily applicable for intact sectioning of hard tissue, including ripened grain. Intact section sampling may be useful for enzymatic analysis and transcriptomic analysis of plant tissue.

Key Words: easy sectioning, cryomicrotome, cross section, longitudinal section, matured grain, rice.

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8. The specimen block was trimmed to the required surface with a disposable blade, 30UF at –25°C.
9. The adhesive film (Cryofilm type 3C (16UF) (SECTION-LAB Co., Ltd., Japan)) was applied to the surface using a fitting tool and then the specimen was cut into 4 µm thickness.
10. The Cryofilm with the section after cutting was picked up carefully.
11. The section was left for approx. 20 seconds to dry slightly at room temperature.
12. The section was stained with hematoxylin for 60 seconds.
13. After being rinsed with running water for 4 minutes, the section was stained with 0.2 % eosin for 10 seconds.
14. After being rinsed with water for a few seconds and with 100% ethanol for 10 seconds, the section was mounted between Cryofilm and glass slide with SCMM-R2 (SECTION-LAB Co. Ltd., Japan).
15. The excessive mounting medium was removed with a filter paper.
16. The mounting medium was polymerized with UV light.

Results and Discussion

A cross section of the Koshihikari grain was shown in Fig. 1A. It was observed that some parts of outermost exosperm, including the pericarp and seed coat of the grain, were deleted. The darkly stained outer part is the aleurone layers. A single layer was found from the ventral side to the dorsal side of the arrows, and multiple layers were developed around the dorsal peripheral side from arrows (Fig. 1B). Very clear endosperm cell shapes were observed. Especially, cell shapes at the upper side in Fig. 1A were different from those at the lower side. There were many polygonal cells at the lower side (Fig. 1C), whereas there were many relatively lengthwise cells at the upper side. The reason for the difference in cell shapes between the upper and lower sides was unknown. At the central area from ventral to the dorsal side through the central point, slender elongated cells were observed (Fig. 1D). A longitudinal section of the grain was shown in Fig. 2A. It was observed that a few aleurone layers were developed at the bottom and dorsal peripheral side (Fig. 2B). In general, lengthwise cells stretched toward the central part were found at ventral and dorsal sides of the endosperm (Fig. 2B). The embryo facing endosperm cells seemed to be collapsed (Fig. 2C). Since all sections from 16 grains showed collapse of the embryo facing endosperm cells, these collapse phenomena were considered no to be artificial during section production. The embryo facing endosperm cells slightly collapsed with cell death due to ROS generation during seed development (Nagasawa et al. 2013). Furthermore, it was reported that water absorption led to endosperm cell collapse from embryos facing the endosperm region (Hoshikawa 1993). Thus, the collapsed of embryo facing the endosperm region may increase due to water absorption. In addition, the central portion was found to consist of a population of small cells (Fig. 2D). Fig. 3A showed the sagittal section of the grain.
Cell shapes of the endosperm were similar to those in the longitudinal section. There were polygonal cells at the top part of the grain, and multiple layers of aleurone were observed at the top of the grain (Fig. 3B). Furthermore, epithelium cells and cell shape around vascular bundle of the scutellum were clearly observed as shown in Fig. 3C. Endosperm cells accumulate starch and proteins as the grain ripens. Since the starchy endosperm is hydrophilic and becomes sticky, it is difficult to obtain an intact thin section of the ripened grain. As shown in Figs. 1A, 2A and 3A, the intact cross section and longitudinal thin sections of the ripened grain were successfully and easily obtained using the Kawamoto method (Kawamoto and Kawamoto 2014).

Intact sectioning is an essential method to observe the structures of interesting organs or tissues of an organism, and the high-quality intact sections should be easily obtained. For producing thin sections of plant tissue, the paraffin section method is usually performed. This method produces uniform thin section. Ogawa et al. (2001) made paraffin sections of whole rice grain using special adhesive tape for 3D visualization of the grain. Although the section of the whole grain was observed, the quality of the section sample was poor because cell shapes of the grain were not clear. The paraffin section method has important merits for obtaining thin section samples. However, the fixation process with some fixative and several steps for paraffin infiltration in tissue are needed before sectioning. During this process, tissue components including protein, enzymes and RNA may be removed or degenerate, and some artifacts may be produced. On the other hand, fresh frozen-sections of tissue are advantageous to retain the subcellular components including protein, DNA and RNA. However, it has been difficult to readily transfer thin sections from freshly frozen samples to slide glasses. For this purpose, Kawamoto (2003) developed a frozen sectioning system using the specific adhesive film to easily obtain a very thin section without any damages and distortions because the sections are supported with an adhesive film till being mounted. The method was also applicable for obtaining the cross section of the whole leaf of rice (Maekawa and Kawamoto 2002).

Since high-quality thin sections of all tissues, including hard tissues, are easily produced by the Kawamoto method, the section can be useful in many analytical fields, e.g., histology, enzyme histochemistry, immune histochemistry, in situ hybridization and mass imaging. For example, immunohistochemical analysis using frozen section of mouse brain by Kawamoto method clearly demonstrated that some pathogenic cells accumulated at specific vessels under stress conditions, resulting in brain micro-inflammation (Arima et al. 2017). Recently, the freeze-dried section was found to be highly suitable for Laser Microdissection (LMD) for gene analysis (Kawamoto and Kawamoto 2014). Actually, fresh frozen section of mouse spine produced by Kawamoto method reveal the part at which the pathogenic cells were accumulated and was effectively utilized for LMD for concerned gene expression (Arima et al. 2012). Although any applications of LMD for gene expression using frozen-sections of plants by Kawamoto method have not been reported so far, it is expected that pinpoint gene analysis of plant tissue, including the seed or grain, could be promoted by the Kawamoto method.

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