Semi-permeable layer formation during seed development in *Elymus nutans* and *Elymus sibiricus*

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**Abstract**

The semi-permeable layer is a layer in the seeds of certain plants that restricts or impedes the exchange of the solute while allowing the permeability of internal and external water and gas, which is valuable protection to sustain the health and secure the growth, development and germination. In this study, the formation time and location of the semi-permeable layer in seed coats of *Elymus nutans* (Griseb.) and *Elymus sibiricus* (L.) were investigated. The experimental seed materials were gathered in the field from the flowering to seed maturation. The light microscopy and transmission electron microscopy for lanthanum nitrate identification were used to examine the characteristics of pericarp, seed coat and nucellus. The results showed that the semi-permeable layer was identified as the position, which can inhibit the penetration of the lanthanum, and it was checked as an amorphous membrane located at the outermost layer of the seed coat that is firmly attached to the seed coat. With seed development, the cells had differentiated and some parts of the ovary and the outer integument had disappeared. The semi-permeable layer originated from the outer layer of the inner integument, which was the original form of the seed coat. It can be stained by the Sudan III and clearly distinguished from other parts of the seed. The formation time of the semi-permeable layer in both species was nearly at 10 to 12 days post-anthesis (dpa), whereas seed physiological maturity was 24 to 26 dpa.

**Keywords**: semi-permeable layer, seed coat, seed development, TEM, *Elymus nutans*, *Elymus sibiricus*

**Introduction**

The seed coat or testa is the protective outer covering surrounding the plant embryo [1]. The seed coat has several functions including the protection of the embryo in the ripe seed from mechanical damage and pathogen attack and the supply of nutrients during seed development [2]. In some species the seed coat may have a semi-permeable layer allowing water uptake and gas exchange, while restricting or preventing solute transport [3–6]. The semi-permeable layer is an important structure for restricting the penetration of toxic solutes into embryos during imbibition from the soil [7] and may also play a role in water storage by holding a sheet of water adjacent to the embryo and thereby protecting the mature embryo against desiccation [8]. At the same time, this layer surrounding the embryo acts as a barrier to apoplastic permeability and radicle emergence [9]. In addition, seed quality testing studies have shown that the existence of semi-permeable layer could decrease the feasibility of testing method, based on the tetrazolium salt or conductivity [10–13]. Therefore, the semi-permeable layer is valuable protection to sustain the health and secure the growth, development, germination, and quality testing of seeds [14].

The presence of a semi-permeable layer can inhibit the infiltration of solutes into the internal seed. For example, tetrazolium chloride, which is used for vital staining, did not penetrate into the inner seed coat of watermelon (*Citrullus vulgaris* Schrad. ex Eckl. & Zeyh.) [10], and also was not able to infiltrate the seed coat of several vegetable seeds [4]. In addition, the water-soluble heavy metal lanthanum ion accumulated in the suberin-rich inner seed coat adjacent to the endosperm in tomato (*Solanum lycopersicum* L.) and pepper (*Capsicum annuum* L.) [15,16]. Castor (*Ricinus communis* L.) and switchgrass (*Panicum virgatum* L.) were impermeable to fluorescent tracers, indicating the presence of a semi-permeable barrier surrounding the embryo [17]. Also, cucumber (*Cucumis sativus* L.) and muskmelon (*Cucumis melo* L.) seeds had lipids and callose in their semi-permeable layers [18,19]. In many grass species this layer embedded with cutin or suberin in the caryopsis integuments restricts solute diffusion [20] such as barley (*Hordeum vulgare* L.) [21], Lolium perenne (L.) [22] and Johnsongrass (*Sorghum halepensis* L.) [23].

While the location and chemical composition of semi-permeable membranes in the seeds of many species have been examined, information on the anatomy and timing of the layer formation is limited. In barley, the integumentary system was
the seat of semi-permeable properties [21], while in corn (Zea
mays L.), the suberized semi-permeable layer was also derived
from the inner integument [24]. Yim and Bradford [19] had
demonstrated that the semi-permeability of the muskmelon
desperm envelope was caused by the outer walls of the
desperm cells at 40 days post-anthesis (dpa). In cucumber
seeds at 45 dpa, the epidermis of the multilayered nucellus
formed the semi-permeable layer [18]. In Sudan grass (Sorghum
dudanens Piper.) layer formation was observed at 16 dpa [25].

Elymus nutans (Griseb.) and Elymus sibiricus (L.) are
perennial grasses, which have carpyoses as propagation units.
The carpyose consists of a single seed fused with the wall of
ripened ovary (pericarp), so the term “carpyose coat” was
usually used to define the combined pericarp, seed coat
and nucellus [26]. The two investigative grasses are widely
distributed in the Tibet and northern China [27] and are
characterized by a high tolerance to the stringent conditions
of sharp continental climate [28]. These species have been
selected as forage cultivars for their adaptation and forage
quality, and widely applied in re-vegetation champaigns [29].

Previous research has demonstrated that E. nutans and E.
sibiricus possessed a semi-permeable layer, which greatly
inhibited electrolyte leakage and tetrAzolium ion uptake [14].
However, our knowledge of the formation time and location
of semi-permeable membrane is still lacking. The aim of the
study was to investigate the main developmental process in
the seed coat after fertilization, and try to definite the specific
formation time and location of the semi-permeable layer in E.
nutans and E. sibiricus.

Material and methods

Plant material

The seed development of two forage species (E. nutans and
E. sibiricus) was conducted in Gannan prairie, Gansu province,
China (102°31′E, 35°12′N), from July to August 2009. Sample
spikelets were individually tagged at the flowering stage.

Seed collection and fixation

Tagged spikelets (n = 50) were harvested at 2-days interval
up to 30 dpa for each species. At each 2-day interval, the palea
and lemma from one seed were removed, and the “naked”
seed was then fixed at 4°C for 24 h in 4% glutaraldehyde in
phosphate buffer (25 mM, pH 6.8). The remaining seeds were
weighed and air-dried.

Light microscopy (LM) and transmission electron microscopy (TEM)

The fixed samples were rinsed with PBS (0.1 mol/L, pH 7.2)
three times for 10 min each time and dehydrated by alcohol
gradient, once with 15%, 30%, 50%, 70%, and 95% and twice
with 100%. The segment of carpyose coat with subtending
desperm tissue was carefully removed with a razor blade
to approximately 1 mm³ and embedded in Technovit 7100
according to described by Kuroiwa [30]. The sections used for
LM (1 μm) were cut using a semi-thin slices machine (KD202B,
LTIE, Shanghai, China), dried and flattened on a glass slide, and
stained with 0.2% Sudan III in 70% ethanol and 0.05% aniline
blue in 0.1 M phosphate buffer (pH 8.2) for approximately 10
min as described by Yim and Bradford [19]. The structure of the
carpyose coat was examined and photos were taken using the
compound microscope (Eclipse E 100, Nikon, Toyko, Japan).

Statistical analyses were performed using the Statistical
program SPSS 16.0 (IBM Corp. Armonk, New York, USA), Analysis of variance with LSD was performed to rank the quality
of the seed samples, and an arcsine transformation was applied
to the percentages prior to analysis.

Results

The anatomical structure of the semi-permeable layer formation

Development of the E. nutans carpyose coat from 2 dpa to
full maturity is shown in Fig. 1. At 2 dpa, a small nucellus
and integument cells were arranged tightly with large numbers of
ovary cells present (Fig. 1a). This stage included three layers
of outer integument and two layers of inner integument and
a thin-walled cuboidal outer layer of nucellus connected to
the other nucellus. With seed development, the cells had dif-
erentiated and some parts of the ovary had disappeared. The

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Fig. 1  LM images showing anatomical structure of cross-sections of *E. nutans* seed coat during seed development from 2 dpa to full maturity. 

a The whole ovary included ovary wall (Ov), outer integument (Oi), inner integument (Ii) and nucellus (Nu) at 2 dpa. 
b The section of ovary at 8 dpa with nucellus cells (Nu) and two layers of inner integument (Ii) and the pericarp (Pe). 
c The semi-permeable layer (Sp) had formed and was stained by the Sudan III at 12 dpa. Remaining nucellus (Nu) were stained by the aniline blue. 
d At 26 dpa, the seed coat (Sc) had formed and aleurone layer (Al) was evident. The main structure of the seed included the pericarp (Pe), the semi-permeable layer (Sp), the seed coat (Sc), the callose layer (Cl) and the aleurone layer (Al). Scale bars: a–c 20 μm; d 10 μm.

Fig. 2  LM images showing anatomical structure of cross-sections of *E. sibiricus* seed coat during seed development from 2 to 24 dpa. 

a It showed the ovary wall (Ov), outer integument (Oi), inner integument (Ii) and the nucellus (Nu) at 2 dpa. 
b The outer integument had left only one layer cell. 
c A red line located between pericarp (Pe) and inner integument (Ii) and the aleurone cell (Al) had formed at 10 dpa. The remaining nucellus (Nu) was stained by the aniline blue. 
d The main structure of the seed included the pericarp (Pe), the semi-permeable layer (Sp), the seed coat (Sc), the callose layer (Cl) and the aleurone layer (Al) at 24 dpa. Scale bars: a,b,d 20 μm; c 10 μm.
outer integument cells had degraded and had only left two layers of inner integument, which attached the inner layer of pericarp. The ovary wall parenchyma cells were fused while the inner cell had elongated to suffuse with the cytoplasm at 8 dpa (Fig. 1b). Until 12 dpa, between the pericarp and inner integument, a red line, which was only stained by the Sudan III, had appeared and this red layer was the semi-permeable layer, which was checked as an amorphous membrane under the TEM (Fig. 1c). The semi-permeable layer was visible as a continuous layer on the outer periclinal walls of the inner integument. At this developmental stage, aleurone cells also had appeared and were situated next to the remainder nucellus cell wall, which was stained by aniline blue. The inner integument lost its cytoplasm and became more and more tight to form the seed coat. By 26 dpa, the nucellus cells, which had already been reduced to one layer at 8 dpa, degraded completely leaving the apparently fully developed callose layer in the seeds at full maturity. The semi-permeable layer could be easily identified and was firmly attached to the seed coat. The pericarp left only one cell layer and the seed coat lost cell form. The periclinal and anticlinal walls of the aleurone layer continued to increase in thickness while they had increased in aleurone cytoplasmic contents. The mature caryopsis coat of *E. nutans* consisted of the pericarp, the seed coat with an associated semi-permeable layer, the callose layer and the aleurone layer (Fig. 1d).

*E. sibiricus* had the same trend with *E. nutans* with regards to the anatomical structure of the semi-permeable layer formation. At the early stages of seed development (Fig. 2a), the caryopsis also included ovary wall cells, outer integument, inner integument and nucellus. The parenchyma cells of the ovary wall and the outer integument were fused and degraded at 6 dpa (Fig. 2b). By 10 dpa, a red layer was visible, and could be

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**Fig. 3** TEM images of the semi-permeable layer formation in *E. nutans*. a,b The nucellar cells contained a large amount of lanthanum deposits, which resembled snowflakes at 2 and 4 dpa. c–e The remaining nucellus (c,d) and the aleurone cell (e) contained lanthanum deposits at 6–10 dpa. f Lanthanum was not observed between the inner integument cell walls at 12 dpa. g At 20 dpa, lanthanum was deposited at the pericarp (Pe). h A magnification of panel g. i The view of the entire seed coat (Sc) and the semi-permeable layer (Sp) at 26 dpa. The black arrows show the deposition of lanthanum, and the white arrows show the absence of lanthanum.
described as a continuous layer on the outer periclinal walls of the inner integument. The remaining nucellus cells lost their cell inclusion and appeared to develop a callose layer, which was stained by aniline blue (Fig. 2c). Until full maturity at 24 dpa, the structure of *E. sibiricus* included the pericarp, a seed coat with an associated semi-permeable layer, a callose layer and the aleurone layer (Fig. 2d).

**The TEM analysis of the semi-permeable layer formation**

The semi-permeable layer was identified as the position, which can inhibit the penetration of the lanthanum. In *E. nutans* at 2 dpa, lanthanum was deposited at the nucellus cell wall, and the electron micrographs revealed that the shape of the lanthanum resembled a snowflake (Fig. 3a). Lanthanum crystals accumulated in the inner seed at the preliminary developmental stages, which included 4 to 10 dpa. Black arrow indicated the lanthanum deposited in the nucellus cell wall (Fig. 3b) and the free endosperm cells (Fig. 3c). At 8 dpa, there were large amounts of lanthanum deposited at the remaining nucellus cell walls and around the starch granules (Fig. 3d). The lanthanum was mainly deposited at the intercellular air spaces of the aleurone cell at 10 dpa (Fig. 3e). At 12 dpa, within the pericarp surrounding the inner integument, we did not detect any lanthanum ions between the two layers of the inner integument cell wall (Fig. 3f), which is indicated by the white arrow. An abundance of lanthanum was detected at the pericarp and near the amorphous membrane, which was located outer periclinal wall of the seed coat at 20 dpa (Fig. 3g). By magnifying Fig. 3g, we observed the lanthanum deposited at the pericarp and the semi-permeable layer resembles a bright
linear membrane structure (Fig. 3h). At a higher magnification of the seeds at 26 dpa, the amorphous membrane was observed as a continuous area in the inner region of the pericarp, which was tightly connected to the seed coat (Fig. 3i).

Common trends were observed with respect to the deposition of lanthanum in the two different types of seeds. At the early stages of E. sibiricus seed growth of 2 and 8 dpa, snowflake-shaped lanthanum could permeate into the nucellar cells, which was deposited in the cell walls and starch granules (Fig. 4a–c) and the inner integument cell wall (Fig. 4d). At 10 dpa (Fig. 4e), a thick periclinal wall formed that was located between the pericarp and the inner integument, and the lanthanum was blocked to penetrate into the inner integument (Fig. 4f). The arrows in Fig. 4g–i indicated the semi-permeable layer at 14, 18 and 24 dpa, which resisted the penetration of lanthanum.

In the ultrastructural investigation used for TEM, views of the entire two layers of the inner integument can be observed in Fig. 5a and Fig. 5c for E. nutans and E. sibiricus, respectively, at 12 and 10 dpa. The results showed that the outer periclinal wall of the inner integument had formed (Fig. 5b,d) to prevent the lanthanum from penetrating into the inner seed (arrow point). This layer was similar to an amorphous membrane that was firmly attached to the inner integument.

These observations showed that the timing of the formation of the semi-permeable layer was different in E. nutans and E. sibiricus but that the position of this membrane was the same, i.e., between the inner pericarp and the seed coat.

**Physiological parameters associated with seed development stages**

In E. nutans, the thousand seed weight reached a maximum of 4.0 g at 26 dpa, and no significant changes were observed from 24 to 30 dpa. The germination reached a maximum of 96% at 22 dpa. From 20 to 30 dpa, the germination exhibited no significant differences. The seed moisture content first increased from 51% to 60% between 2 and 8 dpa, and then declined rapidly during the next 22 d (Fig. 6a) to reach a minimum of 19% at 30 dpa. Based on the thousand seed weight and germination, the E. nutans seeds reached physiological maturity at 26 dpa.

Seeds at different developmental stages were immersed in distilled water for 24 h. The electrical conductivity of the seed leachate and the imbibition rate decreased and the permeability of seed was reduced with seed maturity. Between 2 and 12 dpa, the electrical conductivity decreased from 463 μS cm⁻¹ g⁻¹ to 138 μS cm⁻¹ g⁻¹ and a significant reduction in these values occurred prior to 12 dpa (Fig. 6b). After 12 dpa, the electrical conductivity exhibited only relatively steadily. The imbibition
rate decrease from 138% to 62% at 2 to 16 dpa and then had small changes that were not significant.

The seed moisture content increased slightly between 2 and 10 dpa from 53% to 64% and then declined during the next 20 days to reach a minimum of 16% at 30 dpa in *E. sibiricus* (Fig. 7a). The thousand seed weight reached a maximum of 4.1 g at 24 dpa, and no significant differences were observed from 24 to 30 dpa. The germination reached 92% at 22 dpa (Fig. 7a). Therefore, the *E. sibiricus* seeds reached physiological maturity at 24 dpa.

The electrical conductivity declined sharply from 443 μs cm⁻¹ g⁻¹ to 167 μs cm⁻¹ g⁻¹ between 2 to 10 dpa and then declined slowly. The imbibition rate decreased significantly during 2 to 14 dpa (Fig. 7b).

**Discussion**

The mature seeds of *E. nutans* and *E. sibiricus* contained a semi-permeable layer, which was similar in both species: the layer was typically amorphous, highly compact, and easily distinguished from the seed coat and pericarp under the LM and TEM observation (Fig. 1c, Fig. 2c, Fig. 3h, Fig. 4i, Fig. 5b,d). The shape of this layer was similar to the observations by Beresniewicz et al. [4] for leek (*Allium porrum* L.), onion (*Allium cepa* L.), tomato and pepper seeds, although the location of the semi-permeable layer was located at the innermost layer of the seed coat and directly adjacent to the endosperm.

However, in this present study, the semi-permeable layer was shown to be located at the outermost layer of the seed coat and was firmly attached to it. The location of this layer varied in seeds of different species. In barley, wheat (*Triticum aestivum* L.), rye (*Secale cereal* L.) [5,21,36], and corn [24] were found that the semi-permeable layer was on the seed coat, although the layer was not shown to be located in a specific site within the inner or outer part of the seed coat. Hill and Taylor [11] believe that lettuce seeds have an endospermic semi-permeable layer, which is around the embryo, forming a permeable barrier tissue. The endosperm of the castor seed prevents the penetration of fluorescent dye, indicating the presence of an endosperm semi-permeable layer [17]. In muskmelon [19] and cucumber [18], the semi-permeable layer was present in the perisperm-endosperm envelope. But in Sudan grass [13], the semi-permeable layer was located at the inner aleurone layer, connected to the undifferentiated cells.

The formation of the semi-permeable layer was accompanied by seed development. *E. nutans* formed this layer at 12 dpa (Fig. 1c), whereas in *E. sibiricus* occurred at 10 dpa (Fig. 2c). The result was different from other species. Yim and Bradford [19] had reported that in muskmelon seed, the semi-permeable layer formed at 40 dpa, whereas in cucumber seeds [18], this layer formed at 45 dpa. In Sudan grass, the semi-permeable layer emerged at 16 dpa [25]. Together with the physiological maturity parameters, the conductivity provided an indication of the permeability of caryopsis coat. After the semi-permeable layer formed, the conductivity showed no significant changes.

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**Fig. 6** Physiological parameters of *E. nutans*. a Seed moisture content (seed moisture content), viability (germination) and weight (thousand seed weight) at different development stages. b Seed conductivity and imbibition rate at different development stages. The bars denote the LSD values for each item.

**Fig. 7** Physiological parameters of *E. sibiricus*. a Seed moisture content (seed moisture content), viability (germination) and weight (thousand seed weight) at different development stages. b Seed conductivity and imbibition rate at different development stages. The bars denote the LSD values for each item.
(Fig. 6b, Fig. 7b). This layer resisted the outward diffusion of the seeds’ contents, therefore, the conductivity indicated the formation of the semi-permeable layer. This result was consistent with the anatomical results showing that the semi-permeable layer of *E. mutans* and *E. sibiricus* formed at 12 and 10 dpa, respectively.

The seed coat usually developed from one or two integuments, and the mature caryopsis coat included the pericarp, the seed coat and the nucellus [26]. At seed maturity, much of the integumentary tissue may be degenerated and absorbed by other developing tissues [26]. Therefore, it was important to identify the specific parts that had participated in the formation of the semi-permeable layer. In barley, Collins [21] thought that the integumentary system as the role of semi-permeable properties but Brown [37] indicated that the epidermis of the nucellus had formed the semi-permeable layer in barley. In the caryopsis of maize, the surface of both the inner integument and the epidermis of the nucellus became physically united and then formed the semi-permeable layer [24]. In muskmelon seeds, an extracellular layer composed primarily of callose was entirely responsible for the semi-permeable properties of the endosperm envelope [19]. However, in cucumber seeds, the semi-permeable layer appeared to differentiate in the epidermis of the multilayered nucellus [18]. But in tall wheatgrass (*Agropyron elongata*). (Host) P. Beauv., [38] and buffalograss (*Buchloe dactyloides* (Nutt.) Engelm.) [39], the outer integument had formed the semi-permeable layer. In our experiment, the cells differentiated and fused as the seeds developed, and the outer periclinal walls of the inner integument formed at 12 and 10 dpa, respectively. TEM (Fig. 3f, Fig. 4e) showed that this thickened periclinal wall prevented lanthanum penetration into the inner seeds. Therefore, this cell wall was responsible for the semi-permeability of the seed coat. The result was consistent with a report by Stiles [6] that indicated that the thickened wall might play an important role in determining the permeability.

Based on the anatomical structure of the caryopsis coat, it was evident that during the seed development, the outer integument disappeared, whereas the inner integument developed into a seed coat that was difficult to differentiate from the pericarp. The structures of these two types of seeds were similar to the caryopsis of *Triticum* [26], which included a pericarp, seed coat and aleurone layer. However, prior to the physiological maturity of the seed, the semi-permeable layer had appeared (Fig. 1, Fig. 2). This result was important for the growth of seed. As the seed develops, the cells and nutrient would continue to increase until the seed reached its maximum weight [40]. The semi-permeable layer may prevent nutrients from being lost to the environment when the plant encounters poor growth conditions. Thus, this layer can promote safe and healthy growth in seeds. Furthermore, the semi-permeability would prevent the loss of solutes to the environment until the embryo was capable of resorbing them prior to the initiation of radicle growth [19].

In conclusion, the semi-permeable layer existed in the seed coats of *E. mutans* and *E. sibiricus*. The semi-permeable layer was located at the outermost layer of the seed coat and was connected to the pericarp. As demonstrated by the anatomical observations, the semi-permeable layer was formed by the outer periclinal walls of the inner integument at 12 to 10 dpa and was similar between the two species: typically amorphous, highly compact, but easily distinguished from the pericarp and testa. Based on comparisons with physiological experiments, the formation time of the semi-permeable layer occurred prior to the seed physiological maturity.

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Authors’ contributions

The following declarations about authors’ contributions to the research have been made: designed the experiments: YW, JZ; analyzed the experimental data: JZ; wrote the paper: JZ, YW, JT.

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