Heterogeneous growth of the triploid Pacific oysters *Crassostrea gigas* created by chemical inhibition of polar body release

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**ABSTRACT**

Triploids have several potential advantages over diploids in aquaculture, drawing an elevated commercial reaction into the realistic application of the techniques despite we are still in the early stage of triploid industry for the Pacific oysters *Crassostrea gigas*. We traced the growth performance of the triploid *C. gigas* for over a year from hatchery spat, which was created by manipulations of chemicals (Cytochalasin B, CB or 6-Dimethylaminopurine, 6-DMAP). The growth was clearly marked by an initial longer dormancy and following a great magnitude of heterogeneity. The dormancy was almost 9 to 10-month long or even longer and was considered as a downside of the creation. The heterogeneity was magnified by appearance of extraordinarily growing oysters in part during summer season, which could be a representative upside of the triploids. Overall, however, the results were not as positive as were expected. The longer dormancy and following heterogeneity observed in our practice could be marked as an additional negative sign of the chemical use. The present study, thus, might be highly indicative in the introduction of biological cross between tetraploid and diploid to produce natural triploid embryos.

**Key words:** Heterogeneity, growth performance, chemical triploid, *Crassostrea gigas*

**INTRODUCTION**

Polyploidy referred to as a genetic state having extra sets of chromosomes beyond the normal 2, can be arisen from spontaneous somatic chromosome duplication or as a result of cytokinetic failure during meiosis. Although it is a lethal state for some higher animals, polyploidy has shown some promising results in the field of aquaculture, particularly for shellfish aquaculture (Allen and Downing, 1986; Dunham, 2004; Kong et al., 2007; Rasmussen and Morrissey, 2007). Triploid shellfish if they survive the cytokinetic manipulation, grow faster and larger than diploids, primarily because triploids are reproductively sterile in theory. As such, they put all their energy into growth instead of reproduction. Sterile status of triploids also makes them marketable year round even during reproductive season for diploids.

The techniques for creating triploid oysters were first developed in the early 1980s for *Crassostrea virginica* using cytochalasin B as a cytokinetic inhibitor of meiotic division (Stanley et al., 1981). Since then, shellfish polyploidy has been a research priority (Nell, 2002; Wang et al., 2003), creating numbers of triploid shellfish including *C. gigas*. The estimated benefits of triploids have been visualized over many shellfish, raising a great potential for aquaculture (Allen et al., 1986; Rasmussen and Morrissey, 2007). The beneficial aspects of the triploids might be bigger for *C. gigas* as the species are notable for their exceptional fecundity, accounting for up to 55% of the total dry tissue weight (Andrews, 1979; Perdue, 1983).

In spite of the potential for use of triploidy in aquaculture, several downsides have been witnessed.
Fail-safes of the sterility or stabilities of triploid state have been doubted. In other words, some of the induced triploids return back to diploids and exhibited reproductive behaviors (Allen and Downing, 1990; Guo, 1991; Wang and others 2003; Dunham 2004). More importantly, the triploids often show reduced performance for some traits and negatively affect survival rates, compared to their diploid counterparts (Dunham, 2004). A considerable part of the downsides might be attributed to chemical toxicity of the cytokinetic inhibitor such as cytochalasin B to gamete physiology (Guo et al., 1990).

For our growth performance study of strain-different C. gigas in the submerged cages, we were collecting 5 strains of seeds, two hatchery strains (a diploid and a triploid) and three habitat-different wild seeds strains. In the process, two hatchery strains came and kept in oyster bags earlier the wild strains which were still growing on seed collectors at their own habitats. In our earlier management of the two hatchery strains, the triploids stayed dormant while the diploids were growing. Their dormancy made them excluded from our study as they lost size balance by the time of the experiment. But the triploids were kept under our regular management at our study station at sea. It was about 10 months when they first exhibited a growing sign.

In the present study, we report the unique growth performance of the triploid strain. The triploid growth was also compared with longline diploids.

MATERIALS AND METHODS

1. Study site
All the oysters tested are facilitated at Shellfish Experimental Station, Southeast Sea Fisheries Research Institute located mouth of Goseong Bay, Gyeongnam, Korea.

2. Oysters
In August, 2013, about 100,000 triploid C. gigas were delivered into our study station at sea from Namhae Hatchery of NFRDI, principally created by the method of Allen and Downing (1986) using CB (cytochalasin B) or 6-DMAP (6-dimethylaminopurine) (See MOF, 2013 for detailed procedure). The control diploids were hardened wild oysters growing in the neighboring longline at our study station.

3. Oyster culture and measurement
On arrival, the triploids were contained in a commercial oyster bag (size, 45 × 90 × 10 cm; mesh, 5-13 mm; oyster density, about 3000 seeds/bag) which was suspended within 3 meters deep from longline of our study station. The oyster bags were cleaned on monthly basis, using water jets under about 100 bar pressure. Managements, otherwise mentioned, followed NFRDI Standard for Individual Oyster Culture. Shell height measurement (expressed as mean ± SD, n = 200) was monthly in our 3-replication experiment.

RESULTS AND DISCUSSION

Growth performance of the triploid C. gigas seeds was summarized in Fig. 1 and 2. It was clearly marked by an initial longer dormancy and following heterogeneity. The dormancy persisted for almost 10 months. A variety of abnormal growth performances for triploid shellfish have been reported (see Rasmussen and Morrissey, 2007). It is, however, striking to find the oysters stayed that longer...
dormancy in normal waters. It is more striking to realize that some of oysters remained in the dormant phase while others entered into the growing phase. Thus, it remains as a future research.

A representative abnormality of triploid shellfish is heterogeneous growth. Fig. 2 details the heterogeneity. The range of the heterogeneity was wider than any other range observed so far. The oysters at the left edge of the unimodal growth exhibited very slow growth or dormancy, while oysters at the other side exhibited extraordinary growth. For example, oysters at left edge of the unimodal growth were still 2 cm in shell height, but the oysters at right edge were about 11 cm in shell height (see Fig. 2). The size extremes achieved after 5-month culture are highly indicative of further study for a successful introduction of triploid oysters in aquaculture.

Principally speaking, the advantage of triploidy comes from the induced “sterility” which allows for reproductive energy to be diverted toward somatic growth, resulting in higher growth rates for triploids. This advantage has been fully realized in the shellfish aquaculture industry. At the same time, controversies are emerging in the aquaculture businesses. Representative controversies are appearance of results standing against the expected benefit. Some triploids are growing slower, at the same rate, or faster than diploids. These are much frequent in triploid fish (Dunham 2004). Shellfish, however, are not the exception (Philippe et al., 1996; Rasmussen and Morrissey, 2007). It has been marked as a downside of the triploid shellfish aquaculture.

The triploid growth was further compared with diploid growth (Fig. 3). The control diploids used as a control were hardened late seeds from Tongyeong oyster seed beds. The late hardened seeds outgrew the sizes of the test triploids by the time of the experiment. Thus, we selected the oyster seeds which exhibited retarded growth on the clutches and then contained them in the same cages for a comparable...
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Fig. 3. Growth profile of the test triploid *Crassostrea gigas* in the submerged oyster bags. No. measured: 200 for triploids, 30 for diploids. Replication: 3 times. Error bar: mean ± SD.

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REFERENCES


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