



# An integrated genetic map based on EST-SNPs and QTL analysis of shell color traits in Pacific oyster *Crassostrea gigas*

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## ABSTRACT

Shell color polymorphism is common in nature for the marine mollusks. The Pacific oyster *Crassostrea gigas*, is widely cultured and one of the most economically important bivalve species with various shell color variants. Here, a total of 1061 single nucleotide polymorphism markers, developed from expressed sequence tags (EST-SNPs), were used to construct two genetic linkage maps based on reciprocal-cross between black shell and white shell oysters (MF-A and MF-B) respectively. The two separate linkage maps were combined into an integrated map spanning 947.31 cM in total length. The integrated map was composed of 351 EST-SNP markers distributing on 10 linkage groups, with an average interval of 2.78 cM between adjacent markers. For the MF-A, 11 quantitative trait loci (QTLs) for shell color traits and five QTLs for growth traits were detected, explaining 8.1%–13.8% (mean = 11.76%) and 10.4%–11.0% (mean = 10.66%) phenotypic variance respectively. For the MF-B, nine QTLs for shell color traits and eight QTLs for growth traits were detected, explaining 8.8%–21.7% (mean = 12.84%) and 8.4%–9.4% (mean = 8.88%) phenotypic variance respectively. QTLs clustering was found on the linkage group 2 of the MF-A and the linkage group 8 of the MF-B, and the QTLs showed that pleiotropism could affect at most three traits. Three shared-QTLs associated with shell color traits were identified on the integrated map, and one of them (qA-a\* 4) was significantly homologous to *C. gigas* calmodulin-like protein. The QTLs identified in the present study could be useful in finding candidate genes for the shell color and growth-related traits in future, and potentially applied to marker-assisted selection breeding programs for *C. gigas*.

## 1. Introduction

Color plays an important role in many aspects of our life. It is a key factor in the food choice by affecting taste thresholds, sweetness perception, food preference, pleasantness, and acceptability (Clydesdale, 1993). Then it could affect the price of food products to a certain extent. For the marine mollusks, shell color polymorphism is common in nature, which brings visual aesthetics and attracts great interest of naturalists and collectors for a long time (Comfort, 1951). The Pacific oyster *Crassostrea gigas* (Thunberg 1793), is natively distributed along the Pacific coast of Asia and has been introduced to many countries worldwide for aquaculture purpose (Mann, 1991; Orensanz et al., 2002). Now it is one of the most important marine economic species and its global production ranks first in the cultured aquatic animals (FAO, 2014). Just as consumers are willing to pay more for the rich red salmon fillets (Steine et al., 2005; Alfnes et al., 2006), oysters with beautiful and pure shell and mantle color are popular in market and

traded at a higher price (Nell, 2001; Kang et al., 2013). The shell pigmentation has aroused considerable interest in the oyster industry and it is considered as new potential phenotypic trait for improving commercial value (Ward et al., 2000; Brake et al., 2004; Kang et al., 2013).

In the past decades, a lot of research has been performed regarding the aspect of inheritance model (Luttikhuisen and Drent, 2008), heritability assessment (Wan et al., 2017) and transcriptome analysis (Yue et al., 2015) of shell color traits in mollusk species. It was reported that shell color may be determined by genetic factors, while environmental factors such as diet, salinity and temperature could have an influence on shell coloration in some species (Heath, 1975; Kobayashi et al., 2004; Liu et al., 2009). For *C. gigas*, Ge et al. (2015b) proposed that the shell background color was controlled by a simple pattern of inheritance, which the golden color was dominant to the white color, and it may have an epistatic effect on foreground pigmentation. Xu et al. (2017) estimated the heritabilities of growth and shell color traits and their genetic correlations in black shell strain of *C. gigas*. The results

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suggested that total weight and black shell should be taken as joint objective traits to optimize the selection strategy in selective breeding program. Through comparative transcriptome analysis, some potential shell coloration genes and related molecular mechanisms were revealed in four shell color variants of *C. gigas* (Feng et al., 2015). Despite these progresses, the knowledge of shell color formation and inheritance we have known is still limited for *C. gigas*.

Identifying quantitative trait loci (QTLs) or the genes responsible for shell pigmentation would provide insights into understanding the genetic mechanism of shell coloration. Furthermore, QTLs could be applied in marker-assisted selection (MAS) for *C. gigas* to acc

Table 1

Descriptive statistics for the growth traits and shell color parameters of the MF-A and MF-B families.

Traits	MF-A					MF-B				
	Range	Mean $\pm$ SD	CV (%)	Skewness	Kurtosis	Range	Mean $\pm$ SD	CV (%)	Skewness	Kurtosis
$L^*$	32.13–75.18	57.68 $\pm$ 9.33 <sup>a</sup>	16.182	0.080	0.080	32.13–75.18	57.68 $\pm$ 9.33 <sup>a</sup>	16.182	0.080	0.080

The dark shell pigmentation area was defined as H:255, S:255, I:120 respectively under the HSI color model.

Pearson correlations among the shell color and growth traits were estimated. The differences of phenotype values between the two families were analyzed by *t*-test. Normal tests were conducted by using the kurtosis and skewness coefficients for each measured trait. All these analyses and descriptive statistics were calculated using the SPSS 19.0 (SPSS Inc).

#### 2.4. EST-SNP development and genotyping

Novel EST-SNP development was conducted according to the procedure described as (Zhong et al., 2013). A total of 1794 primer pairs were designed at the flanking sequences of putative SNPs and 576 of them were successfully validated in 32 wild oysters. Information about the 576 SNPs was summarized in Supplementary Table 1.

A total of 1061 EST-SNP markers, containing previously developed 485 EST-SNPs (Zhong et al., 2013; Jin et al., 2014; Zhong et al., 2014a; Zhong et al., 2014b; Wang and Li, 2017) and the new 576 SNP markers were used for polymorphism screening within the two parents and 14 offsprings of each family. These markers were genotyped by using the high-resolution melting (HRM) genotyping technology. The PCR was carried out using the LightCycler<sup>®</sup>480 real-time PCR system (Roche) with a final volume of 10  $\mu$ L reaction mix described as Wang and Li (2017): 10  $\times$  PCR buffer, 1.5 mM MgCl<sub>2</sub>, 0.2 mM dNTP mix, 0.2  $\mu$ M forward and reverse primers, 10 ng template DNA, 0.25 U *Taq* DNA polymerase (Takara) and 5  $\mu$ M SYTO<sup>®</sup>9 (Invitrogen). Data were analyzed using the Gene Scanning and Tm Calling programs within LightCycler<sup>®</sup>480 Software 1.5 (Roche).

#### 2.5. Genetic map construction and QTL analysis

Segregation of each marker was analyzed by chi-square test for the goodness fit of Mendelian ratios. Loci exhibiting significant distorted segregation from Mendelian ratios were also used for the linkage maps construction. JoinMap 4.0 software (Van Ooijen, 2006) was used to construct the sex-average linkage maps for the two families respectively, running the cross-pollinating (CP) population type genotype codes (lm  $\times$  ll, nn  $\times$  np, hk  $\times$  hk). The grouping of linked markers was determined with the followed calculation options: independence logarithm of the odds (LOD) threshold of 6.0, regression mapping algorithm, linkages with a rec. freq. smaller than 0.4 and a LOD larger than 1.00, goodness-of-fit jump threshold for removal of loci 5.00. Recombination rate was transformed to genetic distance in centiMorgans (cM) using the Kosambi mapping function (Kosambi, 2016). Linkage groups were drawn with MapChart2.1 program (Voorrips, 2002). Two

approaches (Chakravarti et al., 1991; Fishman et al., 2001) were used for estimating the expected genome length of the genetic maps and the average of the two estimates was calculated as the estimated genome length (Ge). The observed genome length (Go) was the total length of framework map. The genome coverage was determined by Go/Ge.

QTL mapping analysis was conducted by MapQTL 6.0 software using interval mapping algorithm (Van Ooijen, 2009). The LOD score significance thresholds were estimated by 1000 permutation tests at a linkage-group-wide significance level  $\alpha < 0.05$  (Doerge and Churchill, 1996). The QTLs are named by the family name (qA or qB) followed by an abbreviation of the trait and the QTL number. Based on the position of homologous flanking markers on the integrated map, comparative QTL analysis was conducted between the two families. If a flanking marker defining a QTL was not linked on the framework map during the merging process, the next most closely linked marker on the integrated map was chosen for representation. When a QTL was mapped to a single marker, the location of this marker was used to represent for the position of the QTL (Zheng et al., 2013). Then the detected QTLs could be annotated by the tightly linked genic SNP markers. The SNP-containing sequences were aligned with the reference genome in the NCBI database using BlastX (Altschul et al., 1990).

For comparative mapping analysis, common SNP markers were identified between the integrated map and the linkage maps reported by Hedgecock et al. (2015). The SNP-containing ESTs were aligned with the draft genome of *C. gigas* (Zhang et al., 2012) for mapping corresponding genome scaffolds.

### 3. Results

#### 3.1. Phenotypic data analysis

The descriptive statistics of the shell color and growth traits for the two families were summarized in Table 1. There were significant differences between the MF-A and MF-B families in the traits of  $L^*$ ,  $a^*$ , SPC, SWi and STW ( $P < 0.01$ ). For both families, the values of variation coefficients for weight traits (TW, SWe and STW) were larger than morphological characters (SH, SL and SWi), indicating that the difference in weight traits were more significant among the individuals. All the traits except for STW and  $L^*$  were followed the normal distribution in the two families ( $P < 0.01$ ). Because of the existence of minus in the  $a^*$  dataset, we did not calculate its coefficient of variation.

Pearson correlation analysis showed that all the growth-related traits were significantly correlated in both families, with the maximum correlation coefficients 0.95 in MF-A and 0.93 in MF-B between the total weight and shell weight (Table 2). Conversely, the correlations within shell color related traits and the correlations among growth

Table 2

Correlation coefficients of shell color and growth traits of *C. gigas* (MF-A: below the diagonal; MF-B: above the diagonal).

	<i>L</i> *	<i>a</i> *	<i>b</i> *	SPC	SH	SL	SWi	TW	SWe	STW
<i>L</i> *		0.08	0.50**	0.24**	0.10	0.06	0.02	0.05	0.06	0.02
<i>a</i> *	-0.23		0.33**	0.11	0.19	0.21**	0.10	0.12	0.10	0.13
<i>b</i> *	0.23**	0.44**		0.10	0.05	0.08	0.04	0.07	0.05	0.07
SPC	0.25**	0.18*	0.15		0.12	0.05	0.06	0.13	-0.05	0.14
SH	-0.01	0.11	0.03	0.13		0.40**	0.40**	0.67**	0.61**	0.59**
SL	-0.07	0.18*								

related traits and shell color related traits were weak.

### 3.2. Segregation of SNP markers

Of the 1061 EST-SNP markers, 457 and 394 loci were polymorphic in the two-family parents and their progeny respectively. There were 174 (38.1%) loci in MF-A and 141 (35.8%) loci in MF-B deviated significantly from Mendelian segregation at  $P < 0.05$ . Most of the departed markers in both families (122 and 106, respectively) were deficient for homozygotes.

### 3.3. Genetic linkage maps

Based on the SNPs segregation data of the MF-A and MF-B families, two separate sex-average genetic maps were constructed (Table 3). For the genetic map of MF-A (Fig. 2), it was composed of 274 SNP markers which were mapped to 10 linkage groups. The map was 897.71 cM in length, with an average interval of 3.40 cM and a maximum interval of 23.04 cM. The number of SNPs distributed to each linkage group varied from 6 SNPs on LG10 to 54 SNPs on LG1, with an average of 27.4. The longest linkage group was LG2, which was composed of 40 markers with 117.29 cM in length, and the shortest linkage group was LG10, which contained 6 markers with 39.87 cM in length. The average of linkage groups length was 89.78 cM.

For the genetic map of MF-B (Fig. 3), it was composed of 241 SNP markers which were mapped to 10 linkage groups. The map was 805.00 cM in length, with an average interval of 3.48 cM and a maximum interval of 25.85 cM. The number of SNPs distributed to each linkage group varied from 4 SNPs on LG10 to 47 SNPs on LG1, with an average of 24.1. The longest linkage group was LG2, which was composed of 45 markers with 131.09 cM in length, and the shortest linkage

group was LG7, which contained 7 markers with 39.87 cM in length. The average length of linkage groups was 80.50 cM.

The estimated genome length of the MF-A and MF-B maps based on the two methods were 976.45 cM and 892.08 cM, respectively. Thus, the framework map coverage was 91.94% for the MF-A and 90.24% for the MF-B.

### 3.4. Maps integration

According to the anchor markers on the homologous linkages, the two separate linkage maps of MF-A and MF-B were combined into an integrated map (Fig. 4). The integrated map comprised 351 markers and spanned 947.31 cM (Gof) in length, with an average interval of 2.78 cM and a maximum interval of 12.85 cM. The number of SNPs distributed to each linkage group varied from 9 SNPs on LG10 to 71 SNPs on LG1, with an average of 35.1. The longest linkage group was LG6, which was composed of 32 markers with 149.23 cM in length, and the shortest linkage group was LG10, which contained 9 markers with 45.26 cM in length. The average length of linkage groups was 94.73 cM.

The expected genome length of the integration map calculated by the two methods was 1002.31 cM and 1024.41 cM, respectively, with an average of 1013.36 cM (Ge). Then the genome coverage of the integration map was 93.48% (Cof). As the genome size of *C. gigas* was estimated about 557.7 Mb (Zhang et al., 2012), the average recombination rate was  $\sim 1.84$  cM/Mb across the genome.

### 3.5. QTL mapping for shell color and growth traits

On the basis of linkage map of MF-A, 11 significant QTLs associated with shell color were detected on LG2, LG4, LG5 and LG10 (Table 4), accounting for a percentage of phenotypic variance explained (PVE) of

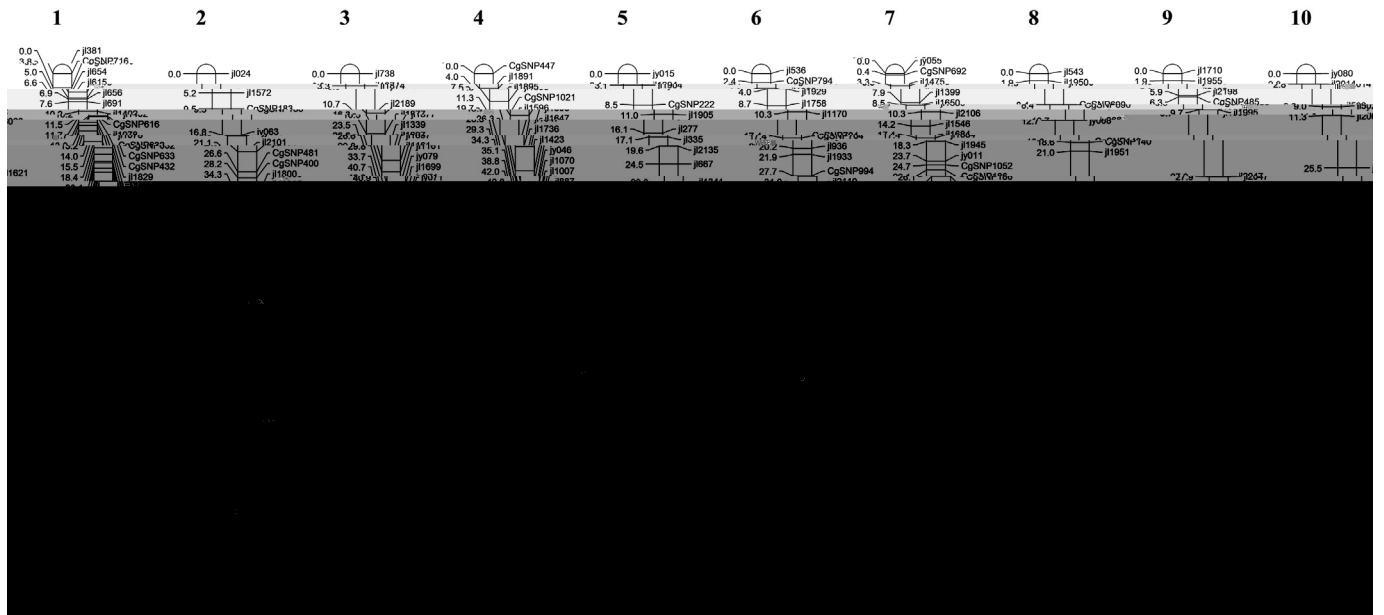


Fig. 2. The linkage map of Pacific oyster (*Crassostrea gigas*) for mapping families 1%, 10.6% and 9.7% on LG10. traits of

8.1%–13.8% with an average of 11.76%. The QTLs tended to have pleiotropism: three QTLs mapped to the same region of 71.67 cM on LG2, which located at the SNP marker j12201, were associated with  $L^*$ ,  $a^*$  and SPC simultaneously, explaining 10.4%, 11.8% and 10.8% phenotypic variation of shell color traits respectively. The QTLs associated with shell color traits of  $L^*$  and SPC were mapped to the same two locations: one was located at 71.67 cM on LG2 and one was located at the overlap interval of 33.51–44.73 cM on LG10. Three QTLs for  $b^*$  spanned discontinuously from 41.81 to 84.00 cM on LG4. For the growth-related traits, a total of five QTLs were detected on LG3 and LG5, and the average phenotypic variability explained by each QTL was 10.66% (range from 10.4% to 11.0%). Two QTLs for the shell length were at 95.26 and 99.30–102.27 cM on LG3, and one QTL were at 35.13 cM on LG5. The two QTLs (qA-TW 1 and qA-STW 1) for total weight and soft-tissue weight were detected in similar regions of 27.54–32.31 cM and 27.54–30.31 cM on LG5, with the same marker j11341 in the intervals, showing a PVE of 11.0% and 10.4%,

respectively.

For the MF-B, nine significant QTLs detected on LG5 and LG8 (Table 5) were associated with shell color traits, explaining 8.8%–21.7% phenotypic variation. Two of the nine QTLs (qB- $a^*$  1 and qB- $a^*$  2) were mapped 0.00–2.29 cM and 28.09–30.09 cM on LG5, and other seven QTLs were spanned discontinuously from 4.01 to 19.87 cM on LG8. Three QTLs controlling the  $L^*$ ,  $b^*$  and SPC were detected at the same region of 4.01 cM on LG8 within the SNP marker j1953, accounting for a percentage of 10.1%, 10.6% and 9.7% PVE respectively. Similarly, another region of overlap interval 15.30–19.87 cM of LG8 were also associated with  $L^*$ ,  $b^*$  and SPC, within the SNP marker j1819, accounting for a high percentage of PVE of 21.7%, 13.1% and 21.0%, respectively. The QTLs associated with shell color traits of  $L^*$  and SPC were mapped to the same two locations: one was located nearby 4.01 cM on LG2 and one was located at the interval of 14.30–19.87 cM on LG10. For the growth traits, one QTL associated with SL was found at 37.39 cM on LG4, showing a PVE of 9.0%. Seven QTLs related to TW,

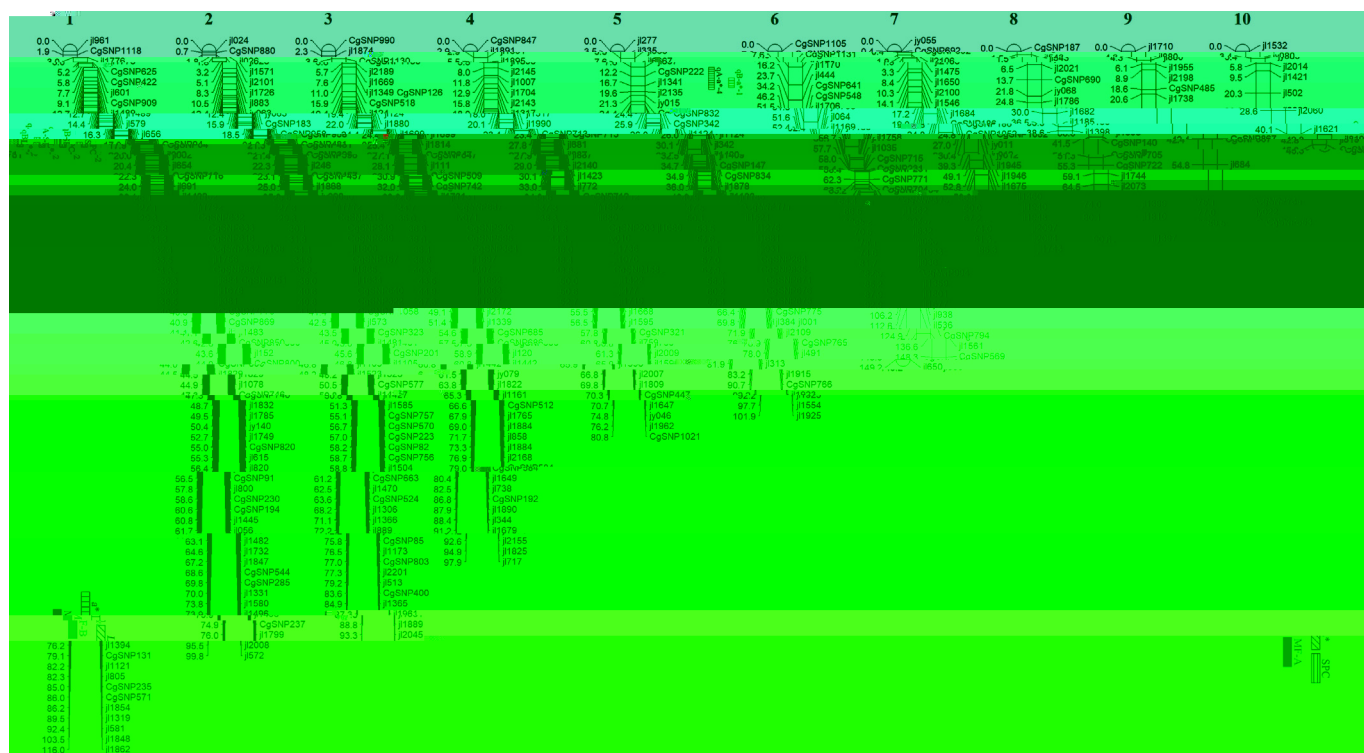


Fig. 4. The integrated linkage map of Pacific oyster (*Crassostrea gigas*) based on mapping family A and mapping family B. Marker distances in Kosambi centimorgans and marker names are indicated on the left and right of each linkage group, respectively.

SWe and STW tended to colocalize, clustering in a special region of 24.75 to 43.79 cM on LG7, and they accounted for a percentage of PVE of 8.4%–9.4%.

By comparative QTL analysis, three co-occurring significant QTLs were identified on the integrated linkage map (Fig. 4). One  $a^*$  shared-QTL ( $qA-a^* 4/qB-a^* 1$ ) was detected on the linkage group 5, while one  $L^*$  shared-QTL ( $qA-L^* 2/qB-L^* 2$ ) and one SPC shared-QTL ( $qA-SPC 2/qB-SPC 2$ ) were detected on the linkage group 10 at the same position.

### 3.6. QTL annotation and comparative mapping analysis

The functional annotation of each detected QTL was summarized in Table 4 and Table 5. The association between several annotated QTLs (like  $qA-L^* 2$  annotated as *C. gigas* furin-like protease 1 and  $qA-SL 2$  annotated as *C. gigas* cilia- and flagella-associated protein 61) and the corresponding traits still remained to be studied further. A few QTLs were annotated as hypothetical protein or their function was unknown or uncharacterized. Notably, the shared-QTL  $qA-a^* 4$  located on the linkage group 5 was highly homologous to *C. gigas* calmodulin-like protein by BlastX search.

Comparative mapping analysis identified two common SNPs (CgSNP223/502-G307268A and j11562/1492-A151764C) between the integrated map and the linkage maps constructed by Hedgecock et al. (2015). All of the 351 SNPs on the integrated map were mapped to 267 scaffolds of the draft genome (Supplementary Table 2), with 207 containing one SNP and 60 containing two or more SNPs (Supplementary Table 3). Furthermore, 34 (56.67%) of the 60 scaffolds with two or more SNPs were mapped to two or three linkage groups (Supplementary Table 4), implying that the genome scaffolds of *C. gigas* may be incorrectly assembled.

## 4. Discussion

High heterozygosity is revealed by whole-genome sequencing in marine invertebrates (Sodergren et al., 2006; Small et al., 2007). *C.*

*gigas* is a typical aquatic species with high levels of polymorphism, which means that abundant SNPs are present in the genome. It was estimated that the average SNP frequency was one per 60 bp in coding regions and one per 40 bp in non-coding regions (Sauvage et al., 2007). The level of polymorphism revealed by complete genome sequence of *C. gigas* was 2.3% (Zhang et al., 2012), higher than that in *Caernohabditis elegans* (0.06%; Hillier et al., 2008) and human (0.09%; Venter et al., 2001). The abundant variations would provide valuable resources for oyster genetic improvement in breeding practices.

Several approaches have been developed for SNP genotyping, such as single-strand conformation polymorphism (SSCP) analysis (Hayashi, 1991), denaturing gradient gel electrophoresis (DGGE) (Lerman and Silverstein, 1987), denaturing high-performance liquid chromatography (DHPLC) (Xiao and Oefner, 2001) and direct sequencing (Shattuck-Eidens et al., 1990). These mutation scanning methods require separation steps and increase the risk of contamination as the PCR products are exposed to the environment (Reed and Wittwer, 2004). Particularly, due to the rapid advance in next-generation sequencing technologies, they have been widely employed to genotype SNPs (Andrews et al., 2016). Despite the advantages in quantity, the relative high cost cannot be ignored, especially in the cases of large genomes and sequencing with high-coverage to avoid high SNP calling error rate (Qi et al., 2017). Those methods based on reduced genome sequencing, like GBS and RAD-seq, digest the genome DNA with one or two enzymes and then sequencing the selected restriction fragments with suitable length, which makes it difficult to share and transfer the data among different laboratories within different oyster populations (Elshire et al., 2011; Peterson et al., 2012). Furthermore, huge data would be obtained after sequencing using the NGS technologies, intensive and complex subsequent bioinformatics analysis steps are still needed for acquiring useful information.

Alternatively, HRM is an economic, efficient and closed-tube SNP genotyping method. It only requires a saturation dye and a high-resolution melting instrument compare to conventional PCR to detect the differences in the sequences even with only single-base variants. After

Table 4  
Summary of quantitative trait loci for shell color and growth traits in mapping family A.

Trait	QTL	Linkage group	Interval (cM)	Peak position (cM)	Nearest marker	LOD threshold	PVE (%)	Annotation
L*	qA-L* 1	LG2	71.67	71.67	jl2201	3.4	10.4	<i>Crassostrea gigas</i> hypothetical protein (LOC105326360), transcript variant × 3, mRNA
	qA-L* 2	LG10	33.51–44.73	42.51	CgSNP751	2.5	8.1	PREDICTED: <i>Crassostrea gigas</i> furin-like protease 1 (LOC105337074), mRNA
	qA-a* 1	LG2	71.67	71.67	jl2201	3.3	11.8	<i>Crassostrea gigas</i> hypothetical protein (LOC105326360), transcript variant × 3, mRNA
	qA-a* 2	LG2	88.06–94.80	90.76	CgSNP570	3.3	9.6	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105323307 (LOC105323307), mRNA
	qA-a* 3	LG5	11.03–16.12	13.03	jl1905	3.4	11.6	PREDICTED: <i>Crassostrea gigas</i> gamma-tubulin complex component 4 (LOC105330131), mRNA
b*	qA-a* 4	LG5	19.10–25.54	24.54	jl667	3.4	11.0	PREDICTED: <i>Crassostrea gigas</i> calmodulin-like (LOC105344225), mRNA
	qA-b* 1	LG4	41.81–42.03	42.03	jl1007	3.4	10.0	PREDICTED: <i>Crassostrea gigas</i> double-stranded RNA-specific editase 1-like (LOC105329897), transcript variant × 3, mRNA
	qA-b* 2	LG4	64.69	64.69	CgSNP712	3.4	10.8	PREDICTED: <i>Crassostrea gigas</i> NAD(P) transhydrogenase, mitochondrial (LOC105343392), mRNA
SPC	qA-b* 3	LG4	83.14–84.00	84.00	jl1680	3.4	10.5	PREDICTED: <i>Crassostrea gigas</i> wybutosine-synthesizing protein 2 homolog (LOC105332610), transcript variant × 4, mRNA
	qA-SPC 1	LG2	71.67	71.67	jl2201	3.9	13.8	<i>Crassostrea gigas</i> hypothetical protein (LOC105326360), transcript variant × 3, mRNA
SL	qA-SPC 2	LG10	26.51–44.73	38.51	CgSNP751	2.7	10.8	PREDICTED: <i>Crassostrea gigas</i> furin-like protease 1 (LOC105337074), mRNA
	qA-SL 1	LG3	95.26	95.26	CgSNP647	3.5	10.6	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105342035 (LOC105342035), partial mRNA
	qA-SL 2	LG3	99.30–102.27	100.27	CgSNP364	3.5	10.7	PREDICTED: <i>Crassostrea gigas</i> cilia- and flagella-associated protein 61 (LOC105320109), transcript variant × 2, mRNA
	qA-SL 3	LG5	35.13	35.13	jl1124	3.4	10.6	PREDICTED: <i>Crassostrea gigas</i> protoadherin-like wing polarity protein stan (LOC105330278), transcript variant × 2, mRNA
TW	qA-TW 1	LG5	27.54–32.31	29.31	jl1341	3.4	11.0	PREDICTED: <i>Crassostrea gigas</i> kidney mitochondrial carrier protein 1 (LOC105333794), mRNA
	qA-STW 1	LG5	27.54–30.31	28.54	jl1341	3.6	10.4	PREDICTED: <i>Crassostrea gigas</i> kidney mitochondrial carrier protein 1 (LOC105333794), mRNA

PVE, percentage of phenotypic variance explained; SPC, shell pigmentation coverage; SL, shell length; TW, total weight; STW, soft-tissue weight.

PCR amplification, melting curves are generated by monitoring the decreasing fluorescence of saturation dye intercalated in the double-stranded DNA. Then the heterozygotes and homozygotes can be identified by a difference of melting curve shapes and a shift in melting temperature ( $T_m$ ) (Wittwer et al., 2003; Reed et al., 2007). Combining with the HRM genotyping technology, the EST-derived SNPs have been widely developed and used in animal and plant genetic studies (Distefano et al., 2013; Dong et al., 2016; Wang et al., 2015; Zhan et al., 2016). The EST-derived SNPs could be transferred and amplified well in different populations and even related species in the *Crassostrea* genus (Zhong et al., 2014b), the collinearity and synteny could be analyzed using the anchor SNP markers in comparative mapping analysis. As the EST-SNPs were genic, their location information in the genome could be identified by the specific primers, which would be highly valuable in assisting in genome assembly and identification of functional genes related with commercially and ecologically important traits.

Although initiated later in linkage mapping than other agricultural animals, great progress has been made for aquatic species over the past two decades (Kocher and Kole, 2008). For *C. gigas*, several linkage maps were published mainly using amplified fragment-length polymorphism (AFLP), microsatellite or a small number of SNP markers (Hubert and Hedgecock, 2004; Li and Guo, 2004; Hubert et al., 2009; Sauvage et al., 2010; Plough and Hedgecock, 2011; Guo et al., 2012; Zhong et al., 2014a). The defect of low-density and coverage limited their potential application in QTL fine mapping. More recently, the GoldenGate microarray and GBS SNP genotyping platforms were used for constructing high density second-generation linkage maps (Hedgecock et al., 2015; Wang et al., 2016), which were not available for every laboratory in consideration of the relative high cost. In the present study, two mapping populations were established by reciprocal cross between the black shell and white shell oysters for constructing two separate linkage maps, using EST-SNP markers combined with the HRM genotyping technology, and then merging them into an integrated map. The density and coverage of the integrated map was improved compared to the two separate maps, with an average distance of 2.78 cM between markers. Using two or more reference populations, more markers would be informative in at least one of the populations and mapped into the linkage maps. This mapping strategy has been successfully applied in some aquatic species, such as the Barramundi *Lates calcarifer* (Wang et al., 2007) and grass carp *Ctenopharyngodon idella*. (Xia et al., 2010). The integrated linkage map spanned 947.31 cM in length and a total of 351 EST-SNPs were assigned to 10 LGs, which were in accordance with the haploid number of chromosomes ( $2n = 20$ ) in the species. The discordance between the linkage groups number and the haploid chromosomes number was found in previously published genetic maps of *C. gigas* (Hubert and Hedgecock, 2004; Li and Guo, 2004; Guo et al., 2012). The low density and uneven distribution of markers may lead to the result. More markers that could fill the gaps between the separate linkage groups should be added to integrate them into one linkage group. Large gaps were still observed in the present map, indicating that high recombination rates or lacking of sufficient polymorphism markers may be present in the regions. A higher density map could be obtained by adding more new SNPs and other markers in the future.

The phenomenon that significant segregation distortion from Mendelian ratios was common in linkage mapping studies and its level varied among different type markers and species (Shi et al., 2014). Of the 351 mapped markers on the present integrated map, 118 (33.62%) were departed from Mendelian segregation, higher than that of AFLP and microsatellites markers in previous studies of *C. gigas* (Li and Guo, 2004; Guo et al., 2012) and other marine mollusks species (Yu and Guo, 2003; Li et al., 2005). Several reasons may lead to the deviation from Mendelian expectations, such as zygotic selection, non-random segregation during meiosis and duplicated genes (Lyttle, 1991; Pardo-Manuel de Villena and Sapienza, 2001; Gut and Lathrop, 2004). It has been proved that a high genetic load of deleterious recessive mutations is responsible for no-Mendelian inheritance of markers in *C. gigas*

Table 5  
Summary of quantitative trait loci for shell color and growth traits in mapping family B.

Trait	QTL	Linkage group	Interval (cM)	Peak position (cM)	Nearest marker	LOD threshold	PVE (%)	Annotation
L*	qB-L* 1	LG8	4.01	4.01	j1953	2.8	10.1	Unknown
	qB-L* 2	LG8	14.30–19.87	19.87	j1819	2.8	21.7	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105346399 (LOC105346399), transcript variant × 3, mRNA
	qB-a* 1	LG5	0.00–2.29	1.29	CgSNP1077	3.3	11.6	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105341982 (LOC105341982), mRNA
	qB-a* 2	LG5	28.09–30.09	28.09	j1277	3.3	9.0	PREDICTED: <i>Crassostrea gigas</i> succinate-CoA ligase [ADP/GDP-forming] subunit alpha, mitochondrial (LOC105319941), transcript variant × 4, mRNA
b*	qB-b* 1	LG8	4.01	4.01	j1953	2.8	10.6	Unknown
	qB-b* 2	LG8	7.01–9.03	7.03	j11421	2.8	8.8	PREDICTED: <i>Crassostrea gigas</i> cell wall protein DAN4 (LOC105332709), transcript variant × 2, mRNA
	qB-b* 3	LG8	15.30–19.87	19.87	j1819	2.8	13.1	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105346399 (LOC105346399), transcript variant × 3, mRNA
SPC	qB-SPC 1	LG8	3.37–4.01	4.01	j1953	2.8	9.7	Unknown
	qB-SPC 2	LG8	14.30–19.87	19.87	j1819	2.8	21.0	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105346399 (LOC105346399), transcript variant × 3, mRNA
SL	qB-SL 1	LG4	37.39	37.39	CgSNP685	3.5	9.0	PREDICTED: <i>Crassostrea gigas</i> stabilizer of axonemal microtubules 2 (LOC105344414), transcript variant × 3, mRNA
TW	qB-TW 1	LG7	24.75–30.98	29.75	j11682	3.0	9.4	PREDICTED: <i>Crassostrea gigas</i> tRNA wybutosine-synthesizing protein 2 homolog (LOC105332610), transcript variant × 4, mRNA
Swe	qB-TW 2	LG7	34.20–37.52	35.20	j11185	3.0	9.1	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105319128 (LOC105319128), mRNA
	qB-TW 3	LG7	39.55–43.49	41.49	CgSNP140	3.0	8.9	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105336641 (LOC105336641), ncRNA
	qB-SWe 1	LG7	27.75–30.98	30.98	j11682	3.1	9.1	PREDICTED: <i>Crassostrea gigas</i> tRNA wybutosine-synthesizing protein 2 homolog (LOC105332610), transcript variant × 4, mRNA
STW	qB-SWe 2	LG7	34.20–36.20	34.20	j11185	3.1	8.5	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105319128 (LOC105319128), mRNA
	qB-STW 1	LG7	24.75–29.98	27.75	j11682	3.1	8.6	PREDICTED: <i>Crassostrea gigas</i> tRNA wybutosine-synthesizing protein 2 homolog (LOC105332610), transcript variant × 4, mRNA
	qB-STW 2	LG7	41.03–42.49	41.49	CgSNP140	3.1	8.4	PREDICTED: <i>Crassostrea gigas</i> uncharacterized LOC105336641 (LOC105336641), ncRNA

PVE, percentage of phenotypic variance explained; SPC, shell pigmentation coverage; SL, shell length; TW, total weight; Swe, shell weight; STW, soft-tissue weight.



(Launey and Hedgecock, 2001). Hubert and Hedgecock (2004) constructed consensus linkage maps for the *C. gigas* by genotyping 11-day-old larvae and the distorted segregation ratios was effectively reduced in the mapping families, indicating that a high genetic load of deleterious recessive mutations leading to strong zygotic selection during the larval stage was a cause of segregation distortion. There have been several studies demonstrated that adding distorted markers moderately would benefit linkage map construction (Fishman et al., 2001; Lanteri et al., 2006). Here, the distorted markers were utilized for constructing the genetic maps, as they were useful in mapping the deleterious recessive genes (Li and Guo, 2004; Guo et al., 2012).

QTL mapping analyses have been conducted mainly for the traits of growth and disease resistance (Yu and Guo, 2006; Sauvage et al., 2010; Guo et al., 2012; Wang et al., 2016), and seldom for inbreeding depression (Plough and Hedgecock, 2011), glycogen and shell pigmentation in oysters (Zhong et al., 2014a; Ge et al., 2015a). Color as an economic important trait, a few of color-related QTLs have been detected in aquaculture species (Qin et al., 2007; Boulding et al., 2008; Baranski et al., 2010; Petersen et al., 2012). Ge et al. (2014) identified a SCAR marker that linked to a major gene/QTL controlling the shell pigmentation for *C. gigas* and mapped it onto a reference genetic map. Seven QTLs associated with shell nacre color with a percentage of PVE of 19.7% to 22.8%, were detected in triangle sail mussel *Hyriopsis cumingii* (Bai et al., 2016). Of the seven QTLs, five QTLs were clustered in a special region on one linkage group. A similar result was found in the present two mapping families, QTLs for both shell color and growth traits showed a tendency to concentrate within narrow intervals, indicating that the genes controlling the shell color and growth traits maybe located in these regions. In the present study, we found the phenomenon of pleiotropism, that is, a same region (like 71.67 cM on LG2 of the MF-A and 27.75 cM–30.98 cM on LG7 of the MF-B) on the linkage group could be associated with diverse traits simultaneously. This conformed to the significant correlations detected among shell color and growth traits. However, the correlations among shell color and grow traits were relative low, implying that the indirect selection for shell color through growth is not feasible and we should take them as joint target traits in selective breeding practices. This conclusion has also been drawn in other studies (Wan et al., 2017; Xu et al., 2017). As markers may not segregate in two specific parents, some important linkage information would lost in QTLs mapping, and therefore the ability to detect QTLs in a single population is often limited. Conducting QTL mapping analysis with multiple populations would improve the efficiency and accuracy of detecting QTLs and provide insight into their functional allelic variation and distribution (Marcel et al., 2007; Wang et al., 2008). In this study, most of the detected QTLs (81.25% for MF-A and 82.35% for MF-B) were family-specific. Three shared-QTLs for shell color traits but no shared-QTL for growth traits were detected within the two mapping families. This may be because growth traits are controlled by a number of genes with small effects, which is consistent with the result of the relative low phenotypic variances explained by the QTLs.

The shared QTL of qA-a\* 4 for shell color traits was annotated as the *C. gigas* calmodulin-like protein, which have been identified to be involved in shell formation in bivalve species (Yan et al., 2007; Fang et al., 2008; Shi et al., 2013). In *P. yessoensis*, it is suggested that the calmodulin-like protein gene is also probably related to melanin biosynthesis of the shell (Sun et al., 2015). In future, more studies should be conducted on the gene to help us understand the relationship between biomineralization and pigmentation in molluscs.

In conclusion, we provided an extensive and valuable resource of EST-SNP markers for *C. gigas*. An integrated linkage map was constructed based on two F<sub>1</sub> families with EST-SNPs, which is potential to play a role in QTL fine mapping and comparative mapping analysis. The QTLs detected in the present study would facilitate in identifying genes controlling shell color and growth traits and provide a basis of MAS for *C. gigas*.

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