Zygotic combinatorial process in plants

Evgenii Vladimirovich Levites, Svetlana Sergeevna Kirikovich

Institute of Cytology and Genetics, Siberian Branch of the Russian Academy of Sciences, Novosibirsk, Russia
Email: elevites@ngs.ru, svetak@bionet.nsc.ru

Received 2 May 2013; revised 3 June 2013; accepted 15 June 2013

ABSTRACT

Experimental data that prove the existence of the zygotic combinatorial process occurring in an embryogenesis-entering zygote are presented in the paper. The zygotic combinatorial process is found when analyzing F1 hybrid plants obtained from crossing homozygous forms different, minimum, in two marker enzymes, and it is found in that hybrid plant which, with one marker enzyme heterozygous spectrum, has a homozygous spectrum of the other. The zygotic combinatorial process leads to F1 hybrids uniformity aberration. The zygotic combinatory process revealed in the study is supposed to be conditioned by chromosome polyteny in mother plant cells and diminution of chromatin excess from the embryogenesis-entering zygote. An obligatory condition for combinatory process is the presence of free exchange of chromatides among homological chromosomes in an embryogenesis-entering cell, i.e. the presence of crossing-over analogous to the one proceeding at meiosis.

Keywords: Isozymes; Polyteny; Diminution; Zygotic Combinatorial Process; Sugar Beet

1. INTRODUCTION

Manifestation of the laws of inheritance discovered by G. Mendel is provided by two combinatorial processes, the first of which consists in a decrease of inherited factors number during sexual cells (gametes) formation, the second one—in inherited factors integration at a random contact and fusion of male and female gametes [1]. A decrease of inherited factors number, which proceeds due to a decrease of chromosome number during meiosis, is realized randomly and equi-probably, which may be designated as meiotic combinatorial process. The second combinatorial process, provided by a gametes random meeting and fusion, may be designated as pre-zygotic.

The meiotic combinatorial process takes place also in the agamospermous (apomictic, subsexual) way of plant seed reproduction realized without gametes genomes integration. The agamospermous progeny developing from egg cells is polymorphous if a mother plant was heterozygous on genes controlling marker traits [2,3]. However, the detection of polymorphism in agamospermous progenies formed from somatic cells by means of mitotic agamospermy allowed us to conclude on the existence of one more combinatorial process [4]. It was hypothesized that this combinatorial process is provided by chromosome polytenization in mother plant cells and a random equi-probable diminution (loss) of an excess of chromatin by an embryogenesis-entering cell [5-7]. The frequency of this or that allele in such an agamospermous progeny is determined by a relative number of chromatids carrying this allele in mother plant cells.

It is known that polyteny is possible not only in somatic cells, but in various embryo sac cells [8-12]. Moreover, it was shown that the ratio of phenotypic classes in agamospermous progenies can be determined by a level of polyteny that occurs in a matured egg cell at the absence of pollination and before its entering embryogenesis [13]. Such egg cells entering embryogenesis are connected with the diminution (loss) of chromatin excess, which proceeds randomly, equi-probably, i.e. combinatorially. We designated this combinatorial process as post-meiotic apozygotic [13]. Finding the phenomenon allowed us to raise a question about the possibility regarding the existence of an analogous combinatorial process also in the sexual progeny formation. To solve this point, reliable and convenient marker traits, among which isozymes are the most available and simple, are necessary. Being different in their electrophoretic mobility, isozymes have a co-dominant manifestation character in hybrids, which allows one, on isozyme spectra, to differentiate clearly both crossed forms and a hybrid from each of parent forms [14,15]. To reveal the combinatorial process in sexual reproduction, a cross of one red table beet (RT) plant and...
one green sugar beet (SB) plant was made [16]. These 
plants were homozygous on the genes controlling glu-
cosephosphate isomerase (GPI2) and malic enzyme 
(ME1), but they were different in the electrophoretic 
 mobility of these enzymes.

Heterozygous spectrum GPI2 was found in all the in-
vestigated 37 hybrid seeds (RT × SB) set in a red table 
beet plant. But the heterozygous spectrum of malic en-
zeyme was found only in 36 seeds as one seed had its ma-
lic enzyme spectrum presented only by the maternal 
isozyme having fast electrophoretic mobility (phenotype 
FF). This fact was explained by the thing that the egg 
cell introduced allele Me1-F with a high polyteny level 
into the zygote. It was supposed that the polyteny of al-
lele Me1-F led to combinatorial diminution on hyper-
geometrical probability distribution laws. As a result of 
this combinatorial process, the introduced male paternal 
allele Me1-S was substituted by mother allele Me1-F 
[16]. However, this conclusion required an additional 
proof as the phenotype, similar with the maternal one, is 
possible to appear not only as a result of the substitution 
of the paternal allele, but also in its inactivation. In case 
of inactivation of the paternal allele, the seed that re-
minds of a homozygote in its phenotype is the heterozy-
gote carrying the null allele. Allelic inactivation of the 
enzyme locus may proceed in many ways; therefore, it 
cannot be a weighty proof of the combinatorial process. 
Only plant seed homozygosity on the gene controlling 
one marker enzyme, under its heterozygosity on the sec-
ond marker, that is suggestive of its passed hybridization, 
can point out the existence of the combinatorial process. 
The present contribution is devoted to the finding out of 
this point and the choice between the two possible 
mechanisms.

2. MATERIAL AND METHODS

RT × SB hybrid seeds obtained from crossing red table 
beet (RT) N 19 and the green plant of inbred sugar beet 
line KWS-9c (SB) were involved in the study. The table 
beet plant had genotype Gpi2-F/Gpi2-F (short-FF) and 
Me1-F/Me1-F (short-FF) and, respectively, fast-migrat-
ing glucosephosphate isomerase isozyme (GPI2, E.C. 
5.3.1.9.) and malic enzyme (ME1, E.C. 1.1.1.40.). The 
sugar beet plant had genotype Gpi2-S/Gpi2-S (short-SS) 
and Me1-S/Me1-S (short-SS) and, respectively, slow-
migrating GPI2 and ME1.

These two plants hybridization process was realized in 
2008 by free cross-pollination in an isolated plot at the 
distance not less than two kilometers away from any 
other flowering beet plants. The obtained seeds of RT × 
SB hybrid had been sown in a hydroponic greenhouse in 
February, 2012, where the plants were then being grown 
during three months.

RT × SB hybrid plant phenotypes were determined by 
the analysis of GPI2 and ME1 isozymes in leaves and 
leafstalks. One of the hybrid plants flowered during 
growth. It allowed us to determine not only its phenotype, 
but the genotype by genetic analysis studying the seed 
progeny obtained from crossing it with test plants in 
summer, 2012.

Electrophoretic analysis of the vegetating plant phe-
notypes and the seed progeny obtained in analyzing 
crossings was carried out on the earlier-described meth-
ods [17,18].

3. RESULTS AND DISCUSSION

Forty plants were grown from the RT × SB hybrid seeds 
at our disposal, out of which 38 had their hybrid pheneto 
type revealed on malic enzyme (ME1) and 2 plants (No. 
18 and No. 28) had the phenotype on ME1 similar with 
the homozygous mother phenotype FF (Figure 1). The 
appearance of these two plants is aberration of F, hybrid 
uniformity law. Plant phenotypes on GPI2 were deter-
mined obscurely at the 2 - 3-month growth stage. In 3 
months of growth in the hydroponic greenhouse, the 
plants were planted in the open ground for further growth 
and vernalization. Out of two plants that had mother 
phenotype on ME1, one flowered without preliminary 
vernalization and that allowed us to analyze its genotype 
on the progeny. The phenotype on ME1 revealed in this 
plant could be conditioned either by genotype FF or F0. 
It is possible to differentiate these two states by, e.g., 
crossing this plant with the testing plant of genotype

![Figure 1](image)

**Figure 1.** Isozyme patterns of malic enzyme (ME1) in sugar 
beet hybrid plants RT × SB. Phenotypes ME1: 1 - 3-FS; 
4-FF; 5, 6-FS. Migration is toward the anode.
Suppose that the studied plant has genotype $FF$, then, in cross $SS \times FF$, a uniform progeny of heterozygous phenotype $FS$ is to form. If the studied plant has genotype $F0$, then, in testing cross $SS \times F0$, there forms the progeny that consists of two geno- and phenotypic classes: common heterozygotes $FS$ and heterozygotes $SS$, whose phenotype is similar with that of $SS$.

Test plant No. 12 with genotype $Me1-S/Me1-S$, $Gpi2-F/Gpi2-S$ was used in the analysis as mother plant. In the obtained progeny, all the analyzed 45 seeds had their phenotype of heterozygotes $Me1-F/Me1-S$ (Figure 2). It was indicative of the thing that the analyzed plant No. 18 had homozygous genotype $Me1-F/Me1-F$. When studying the same progeny on enzyme GPI2, three phenotypic classes were found in it at ratio $8FF:25FS:9SS$ (Table 1). It was suggestive of the thing that plant No. 18 was produced as a result of red table beet and sugar beet hybridization. However, homozygosity of the same plant on gene $Gpi2$ is indicative of the thing that the appearance of homozygous phenotype $FF$ among $F1$ hybrid heterozygotes is not connected with the silencing of allele $Me1-S$.

Table 1. Genotypic analysis of plant No. 18.

| Genotype | ME1 | GPI2 |
|----------|-----|------|
| FF       | 0   | 45   |
| FS       | 8   | 25   |
| SS       | 9   |      |

Summing up the crossing results, it possible concluded that the analyzed plant No. 18 had genotype $Me1-F/Me1-F$, $Gpi2-F/Gpi2-S$. Heterozygosity on gene $Gpi2$ is suggestive of the thing that plant No. 18 was produced as a result of red table beet and sugar beet hybridization.

Table 1. Genotypic analysis of plant No. 18.

The obtained results confirm our earlier preliminary conclusions on the thing that aberration of $F1$ hybrids uniformity is conditioned by the combinatorial process connected with an equi-probable diminution (loss) of chromatin excess from the embryogenesis-entering cell [16].

As this process was revealed in the zygote, it is possible to designate it as zygotic combinatorial process.

The analogous facts were found earlier in the genetic analysis of agamospermous sugar beet progenies when plants, first considered as heterozygotes carrying, along with the normal enzyme locus allele, also the inactivated allele, were, actually, homozygotes on the normal allele [19,20].

The data we obtained well agree to the known results of genetic investigations being carried out in animals. Thus, for example, there were experimentally bred mice that inherited both copies of mother chromosome 11 and none of the paternal copies [21].

When studying inherited human diseases, there was found the phenomenon known as single-parent disomy: when a child has two similar chromosomes 15 but both inherited from the father or from the mother [22]. Such situations may happen both in mice and human when there is excessive chromatin generation of some chromosome being accompanied by its division into two independent identical chromosomes and by the following random release of chromatin excess from the zygotic cell proceeding so that two identical chromosomes remain in the cell and the third chromosome, not similar with them,
is lost.

This research is a logical completion of a series of contributions that began with bringing forth the hypothesis on multidimensionality of inherited information coding in plants [5-7] and were continued by obtaining genetic proofs of this hypothesis—first in agamospermous (subsexual) [13,23] and then in gamospermous (sexual) progenies [16]. This allows us to make some general conclusions regarding the peculiarities of zygotic and apozygotic combinatorial processes in plants, also to understand many biological phenomena explainable on the base of the proposed outlook on these processes.

1) Sugar beet was used as a model object to study variability in sexual (gamospermous) and subsexual (agamospermous) progenies. The agamospermous progenies of diploid and triploid sugar beet plants were diploid. The sequence of events underlying agamospermous reproduction can be briefly outlined the following way; a) separate chromosome regions or whole chromosome polytenization in egg cells, embryo sac cells or somatic nucellus and integuments cells, b) random equi-probable attachment of two out of a multitude of allelic gene copies (or chromatides) present in the nucleus to the nuclear membrane of the cell capable of agamospermous development, c) cell transition to embryogenesis and the onset of embryogenesis, d) duplication of chromosomes attached to the nuclear membrane anticipating nucleus and development, e) first embryogenetic division and diminution (loss) of chromatin, f) second embryogenetic division, g) cell transition to embryogenesis by means of agamospermy. Agamospermous progenies is determined, for homozygotes, on this enzyme. Hence, the plant share of heterozygous malic enzyme (ME1) and 3 had homozygous phenotype on this enzyme. Therefore, the plant share of heterozygous phenotype on malic enzyme (ME1) and 3 had homozygous phenotype on this enzyme. The impossibility of a precise prediction and answer to the question about the thing which parent nucleolus will determine separate chromosome regions polytenization. One can see that it is not necessary that the allele belongs to the mother genome leads to its polytenization. It is not excluded that there are differences among alleles in their predisposition to polytenization.

2) The combinatorial process manifests itself only under mutual exchange of chromatides between chromosomes. Hence, in an embryogenesis-entering cell, there proceeds the crossing-over, analogous to the one which takes place at meiosis.

3) Manifestation of only mother allele of locus Me1 in hybrid RT × SB is indicative of the thing that its polyteny degree in the zygote, before its entering embryogenesis, was higher than that of the introduced paternal allele. This well agrees to the thing that the frequency of polyteny in cells of female generative sphere is many times higher than that of the male.

4) Polytenization of different genes alleles proceeds independently and it is conditioned not only by these genes different chromosome localization, but by the presence of a multitude of independent reduplication onset points in each plant chromosome. It is this that explains our earlier-obtained data on the differences, revealed in agamospermous progenies, in the phenotypic classes ratios of enzymes controlled by non-allelic genes [4,23].

5) Revealing the combinatorial effect on locus Mel in only one of reciprocal hybrids, namely in hybrid RT × SB and the absence of such effect in hybrid SB × RT [16] points out the non-univocality of the processes that determine separate chromosome regions polytenization. One can see that it is not necessary that the allele belonging to the mother genome leads to its polytenization. It is not excluded that there are differences among alleles in their predisposition to polytenization.

6) The viewpoints on polyteny, diminution and combinatorial process allow us to hypothesize the mechanism underlying the phenomenon of “penetrance” [25]. 37 F1 RT × SB hybrid plants were investigated in our experiments at the seed stage [16] and 40 plants, in this experiment, —at the vegetation stage. Out of all 77 F1 plants, 74 plants had their heterozygous phenotype on malic enzyme (ME1) and 3 had homozygous phenotype on this enzyme. Hence, the plant share of heterozygous phenotype, in which paternal allele Mel-S was expressed, is equal to 74/77 = 0.96. This value can be considered as the penetrance value of allele Mel-S.

Analogously, it is possible to explain also such a widely known phenomenon as nucleolar dominance [26]. The impossibility of a precise prediction and answer to the question about the thing which parent nucleolus will be expressed in the hybrid is explained, as we believe, by the combinatorial process that, as is known, is probable. One can suppose the presence of a high polyteny level of both parents nucleolus-coding chromosome regions leading to a random equi-probable diminution (loss) of chromatin excess in an embryogenesis-entering zygote.

7) The frequency of combinatorial processes in sexual progenies is considerably lower than that in the agamospermous. It points out the existence the specific mechanisms that hinder polyteny in gametes and the absence of such mechanisms in cells capable of passing on to embryogenesis by means of agamospermy. Aga-
mospermous reproduction may be accompanied by incomplete diminution of chromatin excess and an increase of certain genes dosage [23]. It is this thing that can explain some increase of DNA content in cell nuclei of agamospermous progenies estimated both cytometrically and on the number of chloroplasts in stomata guard cells [27,28].

Thus, we obtained additional proofs of the thing that aberration of the Mendel ratios in sexual progenies can be explained by polyteny of chromosome regions carrying marker genes and by excessive chromatin copies diminution from the embryogenesis-entering zygote. This well agrees to the thing that its on the base of these viewpoints that it becomes possible to explain phenotypic ratios in most of the sugar beet progenies, produced in agamospermous way, we analyzed earlier.

The evolutionary meaning of differential polyteny and combinatorial processes, and diminution connected with them consists not only in their influence on phenotypic ratios in further generations, but also in the thing that polyteny, just as the whole process of DNA synthesis, depends on a big number of internal and external factors [29-34]. Therefore, differential polyteny can be considered as a way of recording information about acquired traits.

4. ACKNOWLEDGEMENTS

The authors would like to express their gratitude to Alexander V. Zhuravlev for the English version of this article. Investigations were carried out within Integration Project #99 for 2009-2011, supported by Siberian Branch of the Russian Academy of Sciences.

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