A Review of Epigenetic Imprints in Aquatic Animals

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Abstract

Epigenetics is one of the most rapidly expanding fields in biology. On a molecular level, covalent modifications of cytosine bases and histones, and changes in the positioning of nucleosomes are commonly regarded as the driving epigenetic mechanisms. They are fundamental to the regulation of many cellular processes, including gene and microRNA expression, DNA-protein interactions, suppression of transposable element mobility, cellular differentiation, embryogenesis, X-chromosome inactivation and genomic imprinting. Genomic imprinting is an epigenetic gene-marking phenomenon that occurs in the germ line, leads to parental-origin-specific expression of a small subset of genes in male and female gametogenesis, passed to the zygote through fertilization, maintained throughout development and adult life, and erased in primordial germ cells before the new imprints are set.

Keywords. Epigenetics; Imprinting; CpG Islands; Histone modification; Nucleosome positioning; Fish crustacean

Epigenetic and its mechanisms

Before discovering DNA as the inheritance component, scientists were concluded that all the gens in all time are not active in all organisms, although carry the same data. Then Epigenetic was introduced as a mechanism that controls the gene expression in an inheritable pathway. In a molecular level the modification of cytosine and histone alkaline modification and changes in nucleosome location are known as the typical mechanisms of epigenetic. These mechanisms are essential for regulating cellular process such as: difference, transposon movements, genomic imprinting, × deactivation and Embryogenesis. In a multi cellular organism the ability of preservation and retention of Epigenetic during genesis and transition to next generation is essential to make a lot of phenotypes that originates from the same genotype. In human, although monozygotic twins are alike in DNA sequence they are different in DNA methylation and the profiles related to histone modification. This matters even in one single cell. How stem cell can become any cell will be answered by epigenetic. The importance of epigenetic came out when their inappropriate outbreak leads to a number of diseases such as cancer [1].

Epigenetic modification and related complexes:

Epigenetic modifications divided into three branches:
1) DNA methylation
2) Histone DNA
3) Nucleosome positioning

DNA methylation

DNA methylation generally occurs in CpG dinucleotide. These nucleotides are gathered in CpG islands- which about 60% of human gene promotersare in contact with them and usually they are not methylated and have the permission to be copied and make a suitable chromatin formation for gene expression but their malapropos hypo methylation leads to deactivation of coping, but some of them will be methylated during the tissue differentiation. Generally CpG islands methylation is correlated with gene silence. DNA methylation doesn't occur just in CpG Island. The idiom of CpG Island Shores refers to the areas with low concentration of CpG which are in vicinity of CpG Islands, and methylation in these areas are completely in contact with coping suppress. It seems that most of tissue methylation occurs in CpG Island Shores instead of CpG Island. DNA methylation has a key role in genomic imprinting; it means hyper methylation in one or two maternal alleles is related to mono allele gene expression [2].

DNA methylation has several mechanisms to suppress gene expression:

1) Methylated DNA increases use of MBD protein. The family member of MBD protein calls the histone modifier complexes and chromatin remodeling’s to methylated sites [3].

2) DNA methylation suppresses transcription directly by blocking the use of joining protein (such as E2F, NF-KB, AP2) to DNA in Cis binding element areas [3].

It is so rare that DNA methylation be along with transcription activation but in plants and vertebrates, gene bodies’ methylation (the exon of some genes which is under transcription and translation) is positively related to gene expression. It is suggested that this matter be along with elongation step blockage of spurious initiation of transcription. Gene bodies tend to DE methylation in disease that leads to transcription initiation in several wrong sites.

Histone modification

Histones are key elements in epigenetic. H2B, H2A (two dimmer form) and H3, H4 (one tetra dimmer form) form nucleosome.

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Some of Germline DMRs act as Imprinting Control Region (ICR). These elements are methylated on one of paternal allele, and they are mono allele expression controller of imprinted gene and other DMRs methylation in the cluster and they are able to affect Bi directional on far distances. ICRs perform as Cis. ICR include an Insulator which blocks enhancer to join promoter in some domains and in other domains Non coding RNAs have hand in, and they makes gene silenced by absorbing chromatin modifiers complexes (just like chromosome deactivation), epigenetic correction has key role in ICR formation since ICR is just for alleles [6,7].

Elements have hand in imprinting: DNA methylation, histone correction, insulator protein such as CTCF, Non coding RNAs.

**Epigenetic in aquatic animals**

During embryonic development, epigenetic modifications of DNA occur through various processes (e.g. DNA methylation and histone acetylation) and are assumed to facilitate differentiation into specific cell types. An epigenetic alteration can be defined as a mitotically and/or meiotically heritable change in the function of a gene without alterations in the gene sequence [8]. Genes that are to be silenced from one of the parental alleles (i.e. expressed by only one allele) become methylated during the embryonic development in a process called imprinting [9]. There is only limited evidence of genomic imprinting in oviparous species, such as fish [10]. However, DNA methylation reprogramming was observed during the early embryonic development of zebrafish (Danio rerio) in a recent study using an anti-5-methylcytosine antibody in immunohistochemistry and southwestern immunoblotting [11]. This reprogramming of DNA methylation found in zebrafish is similar to the reprogramming during mammalian development [12].

Vitellogenesis is the production of yolky eggs in oviparous species, and involves transport of gene products from the liver to the ovary where proteins are deposited in the maturing oocytes. Strömqvist [13] showed that exposure to EE2 results in an alteration of DNA methylation levels in zebrafish which warrants studies of epigenetic changes in fish toxicological studies. They have for the first time shown sex and tissue differences in the level of DNA methylation in adult zebrafish [14]. Some studies show the applicability to investigate pollutant-induced alterations of methylation levels in fish [13].

Future work should include investigations into the epigenetic nature of such changes.

Genetic and epigenetic interactions between redundant genes in polyploid fish have probably influenced their evolutionary fate, leading to their current impressive biological diversity [14]. Spontaneous polyploids have been observed in several phylogenetically distant orders, including both wild and farmed fish species [15,16]. In the vertebrates, polyploid species are not exclusive to fish, since they have been reported in different groups, from amphibians [17] to occasionally even in mammals [18]. Polyploids can originate either from alterations of meiotic or mitotic processes in specimens within
a species (autopolyploidy) or by reproductive contact among species (allopolyploidy) [19].

Regulatory changes in gene expression following tetraploidisation may result in epigenetic instability, because they are more likely to be deleterious than advantageous [19].

The genetic sources of variation can be associated with the presence of the extra maternal set of chromosomes and can involve simple gene dosage (additivity) between chromosome sets or positive or negative dosage compensation effects (heterosis), epigenetic mechanisms, and transcriptional co-suppression (negative gene dosage compensation) [19].

Studies on gene dosage compensation in the allotriploid endemic Iberian minnow showed that the allelic expression patterns differ between genes and between different tissues [20]. Thus, it appears that in triploids rather than a whole haploid chromosome set (haplome) being silenced, regulatory mechanisms involve selective individual gene-copy silencing [21]. Feng et al., expressed that the live cell clusters are located at the base of the antennules and antenna, as well as the cephalic lobe, implying an epigenetic mechanism of germ cell specification in Fenneropenaeus chinensis. These cells migrate to a dorsolateral position in naupliar and zoeal stages, and gradually enter the genital ridge at the mysis 1 stage. Their findings show that the developmental expression pattern of Fc-vasa-like is different from that of other Crustaceans, and suggest an epigenetic mechanism of germ cell development in Chinese shrimp. The epigenetic mode, in which germ cell was conditionally specified and depended on inductive signals amongst embryonic cells at later embryonic stages, has been reported in the M. musculus [22,23] Strong locotrentos purpuratus [24,25] Blatta germanica (reviewed by Estavour and Aakam) [26] Platynereis dumerilii [27] and Mnemiopsis leidyi (reviewed by Estavour and Aakam) [26]. The expression pattern of Fc-vasa-like, and the origin and migratory characteristics of germ cell suggest that the specification of germ cell in F. chinensis may begin at the limb bud stage and display an epigenetic mode [21]. Characteristics of the origin and migration of the germ cell suggest that the specification of the germ cell of F. chinensis is epigenetically induced, which is different from those reported for other Crustaceans [21].

In zebrafish modulation of parental abiotic environment and nutrition confers increased resistance to the environmental stressor and alterations in cardiovascular parameters (stroke volume, heart rate, cardiac output and red blood cell concentration) to the subsequent generation [28,29].

Conclusion

One of the extremely promoting science in world is epigenetic. These science usages in different branches are spreading and the raise of studies in this field show that. Epigenetic is a process in which without affecting on main DNA succession gene expression pattern is changed. Epigenetic changes leads to malproposegene silence in tumors, better epigenetic recognition which blocks transcription, enables researchers to identify new elements for molecular treatment.

References