Research Article

# DNA Methylation Meta-Analysis Confirms the Division of the Genome into Two Functional Groups

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## **ABSTRACT**

Based on a meta-analysis of human genome methylation data, we tested a theoretical model in which aging is explained by the redistribution of limited resources in cells between two main tasks of the organism: its self-sustenance based on the function of the Housekeeping Gene Group (HG) and functional differentiation, provided by the (IntG) integrative gene group. A meta-analysis of methylation of 100 genes, 50 in the HG group and 50 in IntG, showed significant differences (p<0.0001) between our groups in the level of absolute methylation values of genes bodies and its promoters. We showed a reliable decrease of absolute methylation values in IntG with rising age in contrast to HG, where this level remained constant. The one-sided decrease in methylation in the IntG group is indirectly confirmed by the dispersion data analysis, which also decreased in the genes of this group. The imbalance between HG and IntG in methylation levels suggests that this IntG-shift is a side effect of the ontogenesis grownup program and the main cause of aging. The theoretical model of functional genome division also suggests the leading role of slow dividing and post mitotic cells in triggering and implementing the aging process.

Keywords: Aging; Methylation; Epigenetic; Housekeeping genes; Integrative genes; Rejuvenation.

# INTRODUCTION

DNA methylation, which affects chromatin state and gene expression, is one of the most studied processes of genomic regulation today [1,2]. The specifics of the regulatory function of methylation continue to be studied, but there is no doubt that this process is related to ontogenetic development and aging [3,4]. The main direction of research in this direction focuses on the relationship between methylation and biological age in Mammalia [5-8]. The undoubted connection between methylation and aging processes provides an opportunity for experimental verification of various approaches to this problem. Data on the correlation between the methylation profile and biological age are also widely used as efficiency criteria in works, aimed at rejuvenation of organism cells [9-11]. Interest in DNA methylation has led to the emergence of a large number of studies devoted to genome methylation and the generation of accessible databases, allowing it to be used for testing different points of view on the aging processes. Currently, there are many theories that give their explanation of the causes of aging, but there is no unanimous opinion on the problem [12-14]. We share the viewpoint that in the case of a multicultural organism, aging is always accompanied by a decrease in the resources required by its cells for repair and tissue regeneration [15-18]. The reasons for such a decrease in resources are well explained in our theoretical model of the functional division of the metazoan genome [19,20]. We argue that the genome of all cells of the organism contains two functionally independent parts. It is the most conserved part of the genome, which provides all internal needs of any cell, or Housekeeping Genes (HG). One of the main criteria of this group was the stability of the genes included in it [21,22]. We proposed to identify into a separate functional group the genes, responsible for all specialized structures and their functions, making the organism a single Integrated whole (IntG). The main principle we propose in our theoretical model, is the evolution base property of all IntG genes, which gives them an advantage in the consumption of cellular resources. Given the limitation of intracellular resources, such an advantage always leads to a decrease in the number of proteins, required for cell repair and produced by the HG genes. This property is essentially a manifestation of the antagonistic pleiotropy, described by Williams [23]. The difference

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**Received:** 14-Apr-2022, Manuscript No. JCEST-22-16853; Editor assigned: 18-Apr-2022, PreQC No. JCEST-22-16853 (PQ); Reviewed: 02-May -2022, QC No. JCEST-22-16853; Revised: 09-May-2022, Manuscript No. JCEST-22-16853 (R); Published: 16-May-2022, DOI: 10.35248/2157-7013.22.S10.352.

Citation: Salnikov L, Goldberg S, Sukumaran P, Pinsky E (2022) DNA Methylation Meta-Analysis Confirms the Division of the Genome into Two Functional Groups. J Cell Sci Therapy. S10:352.

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of our theoretical model is that we assign pleiotropic properties not to individual genes, but to the entire IntG group, functionally isolated by us. According to our hypothesis, it is the IntG advantage in competition for resources in the cell that is the primary cause of aging. The development and maturation of the organism in metazoans is under the control of the ontogenesis program. We are convinced that there is no aging program. There is a program for growing up, and aging is a side effect of it. The presence of such a program is reflected primarily in the stable lifespan in various metazoan species [24,25]. We regard the ontogenesis program as a strictly defined sequence of IntG gene expression, starting from the zygote stage, for the purpose of organism formation. The main goal of the ontogenesis program is to achieve the maximum competitive advantage by the time the organism reaches the state of fertility, since natural selection is aimed precisely at this stage of metazoan development [26-30]. Genome methylation, as a regulatory tool, is directly involved in the development and implementation of the ontogenesis program, including the processes of cell differentiation in Mammalian [31].

In addition, of interest to us is the degree of inheritability of gene expression regulation expressed in their variance, as has been shown previously [32]. We are also interested in what changes occur with the baseline, or inherited diversity of epigenetic gene tuning in the genome groups we study, based on methylation data.

To summarize, the use of a meta-analysis of genome methylation in Mammalian allows us to test our assumptions using publicly available experimental databases. In this work, we set ourselves the following goal: using mathematical statistics, to obtain results, capable of verifying our theoretical model on the basis of genome methylation data. To do this, we need to get answers to the following questions:-

- Are there statistical differences in the absolute level of methylation between the HG and IntG parts of the genome that we isolated, comparing groups in terms of both the level of methylation of gene bodies and their promoters;
- Are there any significant differences in age-related dynamics of methylation levels between these parts of the genome?
- What is the variance of the data in the highlighted groups?

#### MATERIALS AND METHODS

Genome methylation data was obtained from an open source EWAS data hub [33]. Gene methylation results from Whole blood data were used. This section not only presents the most averaged values, but also contains the largest amount of data for all ages. A sample of 100 genes from the database was used for the study. The genes were divided into two groups of fifty genes each. Data on the methylation of the gene bodies and its promoters were analysed separately. The IntG group included genes, whose activity is present in post mitotic cells and cells with low MI, which form the tissues that make up the main body mass. The TiGER open database was used to compile the gene list [34]. Genes from the Tissue-Specific Genes based on Expressed Sequence Tags (ESTs) section of the database were selected. The genes were selected using randomly generated numbers. The following permanently active genes from the listed tissues were included in the first group:

Bone-IBSP, COL2A1, ITGA11, MATN3, COL5A1, KIF23.

Kidney-UMOD, ARSF, KLK1, SALL1, RBKS, ACE2, TRPM6, VCAM1, DZIP1,TMEM12.

Liver-ALB, BAAT, FMO3, F13B, A1BG, GNMT, SPTA1, ARG1, APOA1, HBM.

Muscle-MYLK2, TNNI2, TNNC2, CKM, MYOT, TNNT3, TNNT1, MYH2, ACTN3.

Brain-MOG, GFAP, OPCML, MOBP, GALNT9, GRIN1, NGB, NLGN3.

Heart muscle-ANKRD1, NPPA, MYBPC3, MYH6, NPPA-AS1, SYPL2, SMPX.

Selection into the HG gene group was performed according to the major groups of genes, listed in the literature and categorized by the authors as "housekeeping genes" [21,22]. Genes in the first half of the list of each group, using randomly generated numbers, were included in the group. If no data were available in the EWAS database, some of the genes in both groups were replaced by the genes next in the list. The second group included the genes represented in the indicated functional groups:

Protein processing-CAPN1, CAPN7, NACA, NACA2.

Translation factors-EIF1AD, EIF1B, EIF2A.

tRNA synthesis-FARSB, GARS, HARS.

Ribosomal proteins-RPL5, RPL9, RPL10A, RPL11.

RNA polymeras-POLR1C, POLR1D, POLR1E, POLR2A, POLR2B.

Histone-HIST1H2BC, H1FX, H2AFV, H2AFFX.

Cell cycle-XRCC5, RAB14, RAB18, RAB1A.

Lysosome-CTSA, LAMP1, LAMP2.

Regulation of Growth and Ontogenesis-LHFPL1, LHFPL4.

Cytoskeletal-ANXA1, ANXA11, ARPC1A, ARPC2.

RNA splicing-BATF1, DDX39B, HNRPA1P2, PABPN1, SRSF3.

DNA repair/replication-MCM3AP, XRCC5.

Metabolism-PRKAG1, B3GALT6, GSK3B, TPI1, LDHA, TALDO1, TSTA3.

#### **RESULTS**

Data analysis was performed using SAS 9.4 (SAS Inst Inc, Cary, NC). Methylation dynamics were assessed using linear regression analysis. The T-test was used to compare the means, and the F-test was used to compare the slopes of the regression. All p-values were based on a two-tailed hypothesis test, with values less than 0.05 considered statistically significant.

Let us examine the results of the ratio of absolute values of the methylation level between gene bodies and their promoters within the groups we identified. In the HG group, the level of methylation of gene bodies was higher than that of promoters: HG (Mean) bodies (0.3560) vs. promoters was (0.2402) p<0.0001. Difference of slopes was not significant p=0.3571. In the IntG group, the level of methylation of gene bodies was also higher than that of promoters: IntG (Mean) bodies (0.6179) vs. promoters (0.5553) was p<0.0001.

Difference of slopes was significant p=0.0021.

The results of comparison of the groups by the level of methylation of genes promoters are presented in Table 1 and Figure 1.

The comparison of the studied groups, shows the average value of the absolute level of methylation in the IntG and HG groups differ significantly, both in the level of methylation of gene bodies and their promoters (p<0.0001) (Tables 1 and 2) (Figures 1 and 2). There was also a significant difference in the observed rate of methylation decrease between the IntG and HG groups (p<0.0001). Also, the level of methylation differed significantly at each metric age point. At the same time, the difference in age dynamics was more pronounced for promoter methylation, where the difference

in slope over each subsequent 5 years between IntG and HG was significant (p=0.0026). Note that similar small fluctuations in the values seen in the graphs are related to the fact that we used data from the same group of subjects in the gene groups we studied. We also examined the variation of gene promoter methylation diversity within our selected groups individually and in comparison with each other. Standard variance (STD) was considered as a measure of diversity. The results are presented in Table 2 and Figure 2.

As shown, the mean STD in the IntG group is significantly different from that in the HG group (p<0.0001) and the STD value decreases with age (p=0.0454), repeating the trajectory of the methylation process itself (Table 2) (Figure 2). In the HG group, the variance remains practically unchanged, confirming its ontogenetic stability.

**Table 1:** Absolute values of methylation levels of genes promoters.

The whole period of life		IntG			HG		Difference between IntG and HG
	N	Value ± Standard error	p-value	N	Value +/- Standard error	p-value	p-value
Mean	84985	$0.5553 \pm 0.00119$		81249	0.2402+/- 0.00103		<0.0001
Slope per year		000301 ± 0.000047	<.0001		000116+/- 0.000040	0.0037	0.0026

Table 2: Standard deviation of methylation of promoters in HG and IntG values.

The whole period of life		IntG			HG		Difference between IntG and HG
	N	Value ± Standard error	p-value	N	Value ± Standard error	p-value	p-value
Mean of STD	90	$0.3363 \pm 0.000419$		90	$0.2932 \pm 0.000674$		<0.0001
STD Slope per year		000106 ± 0.000012	< 0.0001		000049 ± 0.000026	0.0565	0.0454

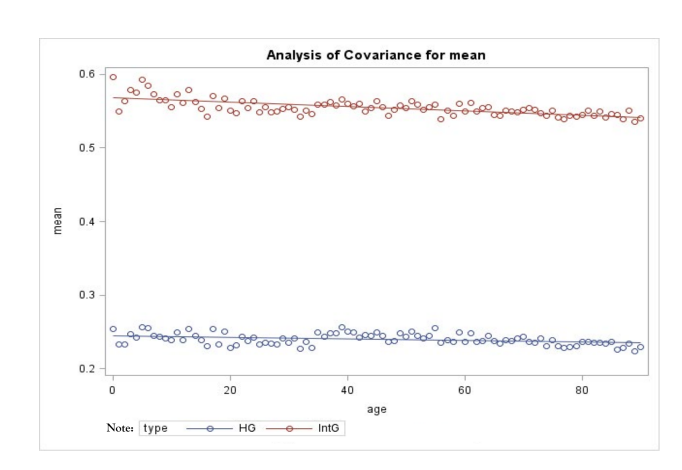


Figure 1: IntG vs. HG methylation of the genes promoters values per year.

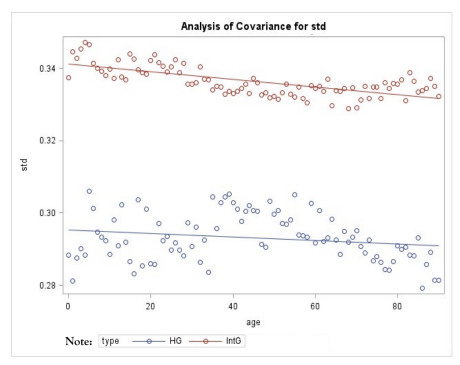


Figure 2: Standard Deviation (STD) of methylation of promoters in HG and IntG values.

## **DISCUSSION**

Discussing our results, we can conclude that our data on genome methylation supports the main statements of our theoretical aging model. For us, the approach in which we consider not individual genes but their ontogenetically determined groups is fundamental. The existence of such groups is confirmed not only by the difference between them, but also by the dynamics of their methylation with age. The data we obtained allow us to provide the following answers to the questions posed at the beginning of this work:

- Statistical differences in methylation levels between our isolated HG and IntG parts of the genome are reliable for both gene bodies and their promoters throughout the entire observation period (p<0.0001);
- In the IntG gene group there was a significant decrease in methylation levels throughout the whole period of observation, especially for promoters and proved to be statistically significant (p<0.0001);
- The variance of promoter methylation data in the IntG group is statistically significantly different from HG and decreases with age, repeating the trajectory of the methylation process itself;

A certain number of explanations should be given to the presented results. As already noted, the two functional groups of genes we identified have significant valid differences between them during the entire observation period in terms of methylation level. As the results showed, the methylation level in HG remains almost stable during the observation period. Initially, we assumed that the level of methylation in the HG genes would increase with age, but the data showed it to be unchanged with data variance remaining at the same level. In contrast, IntG methylation levels steadily decrease with age, particularly in the promoter genes. The

currently available data on the relationship between methylation level and gene biosynthesis magnitude are inconsistent and do not allow us to draw an unequivocal conclusion about an increase in IntG gene expression due to a decrease in their methylation level with age [35-39]. When discussing the role of IntG genes, which represent the basis for organismal functions, we should mention the hyperfunction theory, according to which the metabolism and energy consumption of the organism must constantly increase [40]. In contrast to this theory, for us the leading role in aging processes is played not by a direct and continuous gain of functions in the organism, but by the increase in their resource consumption, which increases after the organism reaches fertility. According to recent data, the level of metabolism reaches a stable value after puberty at the age of about 20 years and remains unchanged, gradually decreasing after 55-60 years when the deficit of self-sustaining functions begins to show [41].

By posing the question about the magnitude of the variance of methylation data in the selected groups, we wanted to find out whether there are characteristics that are unique to the IntG genes. It was found that the variance of gene promoter methylation data in the IntG group is significantly different from the HG group and decreases with age, repeating the downward trajectory of the methylation process itself. The revealed coordinated decrease in the variance of promoter methylation values with age indirectly indicates the presence of specific properties of the IntG group only.

Let us separately dwell on the analysis of methylation level diversity within the groups of genes we identified. It should be noted that methylation in the IntG group is more pronounced in tissues with high regenerative potential, such organs as liver, kidneys, connective tissue, and cartilage. As an example, the data on agerelated methylation in the genes of bone and cartilage tissue can be shown. Thus, methylation of the COL2A1 gene promoter increases in cartilage tissue, but remains almost at the same level in bone

tissue. At the same time, the level of IBSP methylation decreases with age in both tissues. Similar differences were observed in the group of HG genes. For example, the methylation level of MPPL50 remained unchanged, while that of RPL10A slightly decreased. These data illustrate the heterochronicity and heterotopicity of aging processes. However, the existing differences in methylation levels within the studied groups do not affect the obtained total result of the behavior of this index for the group as a whole.

Let's consider the prospects and directions of our future work. In the research on aging a great attention is paid to the cells and tissues with high Mitotic Index (MI). This parameter allows such cells to retain a high regenerative potential and make them a target for experiments aimed at rejuvenation [42-45]. From our point of view, only influencing the epigenetic mechanisms of cells with high MI is clearly insufficient to achieve true rejuvenation of the Mammalia organism [46]. The organs and tissues consisting of postmitotic cells and cells with low MI constitute the main body mass. The survival of this group of cells is based on intracellular repair processes. For these processes, the balance of HG activity between IntG plays a crucial role. We hypothesize that the conserved advantage of IntG genes in resource consumption, shifts the balance of consumption in their direction. This is particularly significant for postmitotic cells and cells with low MI, in contrast to cells with high MI, which compensate for this problem simply by constant division. In our opinion, it is the group of cells with low MI, constituting the basis of the body mass that plays the leading role and triggers the entire cascade of changes accompanying aging.

#### **CONCLUSION**

In conclusion, we note that our data confirm the basic statement of our theoretical model of aging - the presence of two functional parts of the genome with different behaviour during a lifetime. At this point, we can only assume the presence of common properties in a group of IntG genes aimed at achieving an advantage in their expression. A definitive answer to the question, of whether the intracellular production of the HG group of genes decreases with age, can be obtained by studying the changes in the proteome at different periods of life. This will be the main focus of our subsequent work.

### **AUTHOR CONTRIBUTIONS**

L. Salnikov has proposed the theoretical model of aging based on functional genome partition and the role of ratio of the genome two functional parts activity in ontogenesis and aging. He also proposed a way to test the hypothesis using meta-analysis of methylation data. S. Goldberg did the statistical analysis of the data. P. Sukumaran assisted with data extraction and analysis. E. Pinsky organized the work and participated in the data analysis. All authors have made a contribution to prepare the article for publication, participated in the structuring of manuscript and helped with editing. All the authors reviewed, revised and approved the final version of the manuscript.

### **FUNDING**

The authors have no relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript. This includes employment, consultancies, honoraria, stock ownership or options, expert testimony, grants or patents received or pending, or royalties. No writing assistance was utilized in the production of this manuscript.

### **CONFLICT OF INTEREST**

Authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

#### **REFERENCES**

- 1. Dhar GA, Saha S, Mitra P, Chaudhuri RN. DNA methylation and regulation of gene expression: Guardian of our health. Nucleus. 2021;64(3):259-270.
- 2. Lee HT, Oh S, Ro DH, Yoo H, Kwon YW. The key role of DNA methylation and histone acetylation in epigenetics of atherosclerosis. J Lipid Atheroscler. 2020;9(3):419-434.
- 3. Fei Z, Haghani A, Robeck TR, Zoller JA, Li CZ, Zhang J, et al. Universal DNA methylation age across mammalian issues. Innov Aging. 2021;5:410.
- 4. Ciccarone F, Tagliatesta S, Caiafa P, Zampieri M. DNA methylation dynamics in aging: How far are we from understanding the mechanisms? Mech Ageing Dev. 2018; 174:3-17.
- 5. Horvath S, Raj K. DNA methylation-based biomarkers and the epigenetic clock theory of aging. Nat Rev Genet. 2018;19(6):371-384
- 6. Galkin F, Mamoshina P, Aliper A, de Magalhaes JP, Gladyshev VN, Zhavoronkov A. Biohorology and biomarkers of aging: Current state-of-the-art, challenges and opportunities. Ageing Res Rev. 2020;60:101050.
- 7. Trapp A, Kerepesi C, Gladyshev VN. Profiling epigenetic age in single cells. Nat Aging. 2021;1:1189-1209.
- 8. Porter HL, Brown CA, Roopnarinesingh X, Giles CB, Georgescu C, Freeman WM, et al. Many chronological aging clocks can be found throughout the epigenome: Implications for quantifying biological aging. Aging Cell. 2021;20(11):13492.
- 9. Zhang W, Qu J, Liu GH, Belmonte JCI. The ageing epigenome and its rejuvenation. Nature reviews molecular cell biology. Nat Rev Mol Cell Biol. 2020;21(3):137-150.
- 10. Wang YX, Blau HM. Reversing aging for heart repair. Sci. 2021;373(6562):1439-1440.
- 11. Chen Y, Luttmann FF, Schoger E, Scholer HR, Zelarayan LC, Kim KP, et al. Reversible reprogramming of cardiomyocytes to a fetal state drives heart regeneration in mice. Sci. 2021;373(6562):1537-1540.
- 12. Larocca D, Lee J, West MD, Labat I, Sternberg H. No time to age: Uncoupling aging from chronological time. Genes. 2021;12(5):611.
- 13. Augsburger BN. A multidisciplinary approach to explain biological aging and longevity. BioRxiv. 2016;1:045633.

- 14. Bartke A. New directions in research on aging. Stem Cell Rev Rep. 2021;23:1-7.
- 15. Ferreira M, Francisco S, Soares AR, Nobre A, Pinheiro M, Reis A, et al. Integration of segmented regression analysis with weighted gene correlation network analysis identifies genes whose expression is remodelled throughout physiological aging in mouse tissues. Aging. 2021;13(14):18150-18190.
- 16. Voutetakis K, Chatziioannou A, Gonos ES, Trougakos IP. Comparative meta-analysis of transcriptomics data during cellular senescence and *in vivo* tissue ageing. Oxid Med Cell Longev. 2015;2015:732914.
- 17. Walther DM, Kasturi P, Zheng M, Pinkert S, Vecchi G, Ciryam P, et al. Widespread proteome remodelling and aggregation in aging C. elegans. Cell. 2015;161(4):919-932.
- 18. Lagunas-Rangel FA, Bermudez-Cruz RM. The role of DNA repairs in cellular aging process. 2019;1:1-12.
- 19. Salnikov L, Baramiya MG. The ratio of the genome two functional parts activity as the prime cause of aging. Front Aging. 2020;1:1-12.
- 20. Salnikov L, Baramiya MG. From autonomy to integration, from integration to dynamically balanced integrated co-existence: non-aging as the third stage of development. Front Aging, 2021;2:25-33.
- 21. Eisenberg E, Levanon EY. Human housekeeping genes, revisited. Trends Genet. 2013;29(10):569-574.
- 22. Hounkpe BW, Chenou F, de Lima F, De Paula EV. HRT Atlas v1.0 database: Redefining human and mouse housekeeping genes and candidate reference transcripts by mining massive RNA-seq datasets. Nucleic Acids Res. 2021;49(1):947-955.
- 23. Williams GC. Pleiotropy, natural selection, and the evolution of senescence. Evolution. 957;11(4):398–411.
- 24. Colchero F, Aburto JM, Archie EA, Boesch C, Breuer T, Campos FA, et al. The long lives of primates and the 'invariant rate of ageing' hypothesis. Nat Commun. 2021;12(1):3666.
- 25. Curtsinger JW. Retired flies, hidden plateaus, and the evolution of senescence in *Drosophila melanogaster*. Evolution. 2016;70(6):1297-1306.
- 26. Kirkwood TBL, Holliday R. The evolution of ageing and longevity. Proc R Soc London Ser B Biol Sci. 1979;205(1161):531-546.
- 27. Gems D, Kern CC, Nour J, Ezcurra M. Reproductive suicide: Similar mechanisms of aging in *C. elegans* and Pacific Salmon. Front Cell Dev Biol. 2021;9:688788.
- 28. Bilinski T, Bylak A, Kukula K, Zadrag-Tecza R. Senescence as a trade-off between successful land colonisation and longevity: Critical review and analysis of a hypothesis. Peer J. 2021;9:12286.
- 29. de Magalhaes JP, Church GM. Genomes optimize reproduction: Aging as a consequence of the developmental program. Physiol. 2005;20:252-259.
- 30. Frankel S, Rogina B. Evolution, chance, and aging. Front

- Genet. 2021;12:733184.
- 31. Olova N, Simpson DJ, Marioni RE, Chandra T. Partial reprogramming induces a steady decline in epigenetic age before loss of somatic identity. Aging Cell. 2019;18(1):12877.
- 32. Yamamoto R, Chung R, Vazquez JM, Sheng H, Steinberg P, Ioannidis NM, et al. Tissue-specific impacts of aging and genetics on gene expression patterns in humans. BioRxiv. 2021;11:23-43.
- 33. EWAS Data Hub. 2022.
- 34. Tissue-specific Gene Expression and Regulation. 2022.
- 35. Anastasiadi D, Esteve-Codina A, Piferrer F. Consistent inverse correlation between DNA methylation of the first intron and gene expression across tissues and species. Epigenetics Chromatin. 2018;37:1-17.
- 36. Pearce K, Cai D, Roberts AC, Glanzman DL. Role of protein synthesis and DNA methylation in the consolidation and maintenance of long-term memory in *Aplysia*. Elife. 2017;6:18299.
- 37. Spainhour JC, Lim HS, Yi SV, Qiu P. Correlation patterns between DNA methylation and gene expression in the cancer genome atlas. Cancer Inform. 2019;18:1176935119828776.
- 38. Zhang Y, Liu C, Cheng H, Tian S, Liu Y, Wang S, et al. DNA methylation and its effects on gene expression during primary to secondary growth in poplar stems. BMC Genomics. 2020;21(1):498.
- 39. Sales AJ, Maciel IS, Suavinha ACDR, Joca SRL. Modulation of DNA methylation and gene expression in rodent cortical neuroplasticity pathways exerts rapid antidepressant-like effects. Mol Neurobiol. 2021;58(2):777-794.
- 40. Gems D. The hyperfunction theory: An emerging paradigm for the biology of aging. Ageing Res Rev. 2022;74:101557.
- 41. Rhoads TW, Anderson RM. Taking the long view on metabolism. Sci. 2021;373(6556):738-739.
- 42. Lapasset L, Milhavet O, Prieur A, Besnard E, Babled A, Ait-Hamou N, et al. Rejuvenating senescent and centenarian human cells by reprogramming through the pluripotent state. Genes Dev. 2011;25(21):2248-2253.
- 43. Lemaitre JM. Reversibility of cellular aging by reprogramming through an embryonic-like state: a new paradigm for human cell rejuvenation. Cent Asian J Glob Health. 2014;2:88.
- 44. Lehmann M, Canatelli-Mallat M, Chiavellini P, Console GM, Gallardo MD, Goya RG. Partial reprogramming as an emerging strategy for safe induced cell generation and rejuvenation. Curr Gene Ther. 2019;19(4):248-254.
- 45. Cheng Z, Peng HL, Zhang R, Fu XM, Zhang GS. Rejuvenation of cardiac tissue developed from reprogrammed aged somatic cells. Rejuvenation Res. 2017;20(5):389-400.
- 46. Zhang B, Trapp A, Kerepesi C, Gladyshev VN. Emerging rejuvenation strategies-Reducing the biological age. Aging Cell. 2021;21(1):13538.