





Role of minor splicing factors ZRSR1 and ZRSR2 in embryo genome activation

Subjects: Molecular Biology

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Minor splicing plays an important role in vertebrate development. *Zrsr1* and *Zrsr2* paralog genes have essential roles in alternative splicing, mainly participating in the recognition of minor (U12) introns. Mice embryos with mutations in both splicing factors stopped developing mainly between the 2- and 4-cell stages, just after zygotic gene activation. RNA-seq analysis of *Zrsr1/2^{mu}* 2-cell embryos showed altered gene and isoform expression of thousands of genes enriched in essential gene ontology terms and biological pathways related to ribosome, RNA transport, spliceosome, and essential zygotic gene activation steps. Alternative splicing of both U2 and U12 intron-containing genes was altered related to cell cycle and mitotic nuclear division. *Zrsr1* and *Zrsr2* were also required for the conversion of mouse-induced pluripotent stem cells into 2C-like cells. *Zrsr1* and *Zrsr2* emerge as necessary for zygotic gene activation and the conversion of induced pluripotent stem cells into 2C-like cells.

Introduction

Alternative splicing (AS) is an essential co- and post-transcriptional process through which multiple transcripts are generated from a single gene. There are two splicing machineries: the major class or U2-dependent spliceosome, which removes the majority of introns (U2-type intron); and the minor class or U12-dependent spliceosome, which removes U12-type introns. U12-type introns represent around 0.4% of all introns in the genome ^[1], and are non-randomly distributed across the genome. Despite their scarce abundance, minor introns are highly conserved across distantly related eukaryotic taxa, indicating their common evolutionary origin ^{[2][3]}. Minor splicing has been by far less studied than canonical splicing, and although at first it was described as a cytoplasmatic mechanism ^{[4][5]}, several studies have refuted this theory ^{[6][7][8]}. However, some level of cytoplasmatic splicing has been described in strongly compartmentalized cells, such as neurons or megakaryocytes ^[9], but its function remains unclear.

Further, genes with U12 introns are over-represented in functions and pathways related to development, such as RNA processing, DNA replication, or cell cycle ^{[10][11]}, and they show tissue-dependent intron retention ^[12]. Several mutations of the minor spliceosome have been associated with multiple diseases, including developmental disorders ^{[10][13][14][15][16]}, neurodegeneration ^[17], and cancer ^[18]. Additionally, the minor spliceosome is thought to play an important developmental role in plants ^{[19][20][21]}, drosophila ^[22], zebrafish ^[23], and in the mouse central nervous system ^{[24][25]}, hypothalamus ^[26], gametogenesis ^[16], and early development ^[27]. Genes involved in mRNA splicing are over-represented during early preimplantation development ^[28], before and during zygotic gene activation (ZGA) ^[29]. However, the specific role of minor splicing in preimplantation embryo development has not been fully elucidated.

The splicing factors ZRSR1 and ZRSR2 (encoded by the *Zrsr1* and *Zrsr2* genes, respectively) have been identified in all mammalian species analysed ^{[16][30]}, and are involved in the recognition of the 3'AC dinucleotide of the AT-AC class of U12-type introns or the 3'AG of U2-type introns ^[31]. Recent sequencing studies have identified frequent somatic ZRSR2 mutations in hematological malignancies, such as myelodysplastic syndrome (MDS), causing mis-splicing of U12 introns ^[18]. However, their precise role in embryo development and function in other pluripotent cell types remains unclear. *Zrsr1* is a retrotransposed copy of X-linked *Zrsr2* (located on the X chromosome in all mammalian species analysed) and is paternally expressed in the placenta and some adult tissues, while the maternal copy is methylated and silent. *Zrsr1* plays a key role in hematopoiesis, muscle strength, and spermatogenesis, leading its mutation to infertility ^[16].

Results and discussion

Mutant Mice Generation and Rescue Experiments

To gain further insight into the role of *Zrsr1* and *Zrsr2* in preimplantation development, in the present study we generated *Zrsr1* and *Zrsr2* mutant mice, truncating their RNA recognition motif. Double mutants produced by crossing males

homozygous for the *Zrsr1* mutation with females homozygous for the *Zrsr2* mutation (hereafter named *Zrsr1/2^{mu}*) stopped developing just after ZGA. Rescue experiments in which *Zrsr1* mRNA was injected into single-cell *Zrsr1/2^{mu}* embryos extended the development of mutant embryos, revealing that minor splicing is essential for ZGA. A clear complementarity action between ZRSR1 and ZRSR2 during 2-cell embryo cleavage was observed, as embryos need at least one *Zrsr1* wild-type (WT) allele from the father (as it is maternally imprinted and only expressed from the paternal allele) or one *Zrsr2* WT allele from the mother (as only *Zrsr2* is expressed in the oocyte and the embryo needs the protein during early stages of development).

Transcriptomic Analysis and Conversion of Induced Pluripotent Stem Cells (iPSCs) to 2C-Like Cells

RNA-seq analysis of the transcriptome of *Zrsr1/2^{mu}* and WT embryos revealed a large number of overexpressed genes in the double mutant embryos that are supposed to be degraded during early embryo development [32][33], suggesting that the mutations cause a blockade in the degradation of the oocyte's maternal mRNAs. Intron retention was affected for both U2 and U12 introns, being exon skipping also altered. Differentially retained U2 introns located within U12-containing genes were located preferentially adjacent to U2 introns, suggesting an interplay between the major and minor spliceosomes, which goes in accordance with the results showed by Horiuchi *et al*, 2018 [16]. We also analysed the role of *Zrsr1* and *Zrsr2* in the conversion of induced pluripotent stem cells (iPSCs) to 2C-like cells (2CLC), which are a rare metastable cell population that appears at a very low frequency in mouse embryonic stem cells (mESCs) and iPSCs in culture. We discovered that both *Zrsr1* and *Zrsr2* are necessary for the generation of 2CLC, and contrary to what happens with the embryos, *Zrsr1* and *Zrsr2* are not complementary in iPSCs, as both are required for the induction of 2CLC.

The different pleiotropic phenotypes of U12 splicing factors have made it difficult to identify unifying developmental functions of U12 intron-containing genes. There are several events common to gametogenesis and preimplantation development at the precise moment *Zrsr1* and *Zrsr2* are expressed that are critical. First, *Zrsr1* and *Zrsr2* are expressed in gametogenesis during the meiotic prophase, which is equivalent to a prolonged G2 phase in the mitotic cell cycle [34]. Similarly, the G1 phase is extremely short, while G2 is very long during the second preimplantation division in mouse embryos, when the major phase of ZGA takes place [35] and *Zrsr1* is specifically expressed. The gene *Zscan4* is also expressed in oocytes and spermatocytes during the late meiotic prophase, in G2 phase ES cells and in late 2-cell stage mouse embryos [36]. Its expression is associated with heterochromatin organization [37]. Interestingly, the small proportion of genes with U12 introns in vertebrates is over-represented in specific G2 phase functions like DNA replication, cell cycle, and DNA damage repair. The disruption of multiple cell cycle and DNA damage repair genes would be expected to cause defects in cell differentiation. Another common event between meiosis, ZGA and iPSC conversion in 2CLC is the alteration in the structure of the nuclear membrane. In addition to nuclear break down that occurs in meiosis, nuclear lamina is rearranged during stem cell differentiation [38], and the nuclear envelope breaks down after the two pronuclei undergo DNA replication, and chromosomes then associate during the first mitosis [39].

According to our results, the *Zrsr1/2^{mu}* phenotype observed in embryos and iPSC could be the outcome of one of several disrupted procedures: dysregulation of U12-type intron-containing genes, the disruption of AS, and/or the altered expression of other genes. Our gene ontology (GO) analysis of U12 intron-bearing genes with alterations in intron retention in *Zrsr1/2^{mu}* embryos revealed the strong enrichment of genes related to key cellular functions such as cell cycle, mitotic nuclear division, DNA damage repair, RNA processing, etc., highlighting the importance of these genes in the 2-cell blockade. However, *Zrsr1/2^{mu}* also indirectly modified the expression of many other genes reported to play an important role in embryo cleavage and iPSC conversion to 2C-like cells, some of them carrying U12 introns. For example, one important gene for reprogramming the conversion from mouse embryonic fibroblasts (MEF) to iPSC is the epithelia-splicing factor *Esrp1* [40], a marker of several cancers [41] that has an U12 intron. In our *Zrsr1/2^{mu}* embryos, *Esrp1* showed AS alterations in ES and IR of the U12 intron. In the same manner as *Esrp1*, the genes *Zrsr1* and *Zrsr2* were necessary for iPSC reprogramming to the 2C-like state, but were not required for maintenance of pluripotency, as we could maintain *Zrsr1* and *Zrsr2* mutant iPSC lines indefinitely without loss of viability. Another gene down-regulated in the *Zrsr1/2^{mu}* embryos was *Dppa3* (also known as *Stella*), a marker of pluripotency and cell division that ensures activation of endogenous retroviruses (ERVs) [42]. *Cnot7* (also known as *Caf-1*) controls ERVs and genes specific to 2-cell embryos, and its depletion in ESCs enhances the reprogramming of 2C-like cells [43]. This gene also showed AS alterations in ES and IR in *Zrsr1/2^{mu}* embryos. Also, the Developmental pluripotency associated genes 2 (*Dppa2*) and 4

(Dppa4), which are positive regulators of 2C-like cells and ZGA transcription by directly up-regulating Dux^[44], were down regulated in *Zrsr1/2^{mu}* embryos. Other genes showing high intron retention in mutant embryos were *Srsf10* and *Cul1*. *Srsf10* is required to implement DNA damage-induced splicing shifts in other transcripts involved in apoptosis, cell-cycle control and DNA repair, indicating that SRSF10 connects DNA damage to the alternative splicing of transcripts that determine cell fate^[45]. *Cul1* is the major scaffold of the SCF complex (Skp, Cullin, F-box), an E3 ubiquitin ligase that contributes to the regulation of many cell processes, including proliferation, differentiation and death, by targeting its substrate proteins for degradation via the ubiquitin-proteasome system^[46].

Conclusions

This comprehensive analysis of *Zrsr1/2^{mu}* mouse embryos reveals a new role of paternal *Zrsr1* and maternal *Zrsr2* as components of the ZGA mechanism in the embryo, as mutation of either *Zrsr1* or *Zrsr2* blocked the reprogramming of iPSCs towards 2CLC (Figure 1). Accordingly, *Zrsr1* and *Zrsr2* emerge as essential for stem cell differentiation during gametogenesis^[16], totipotent zygote differentiation (ZGA) and reprogramming of iPSCs towards 2CLC, suggesting a critical role of minor splicing in the stem cell reprogramming process.

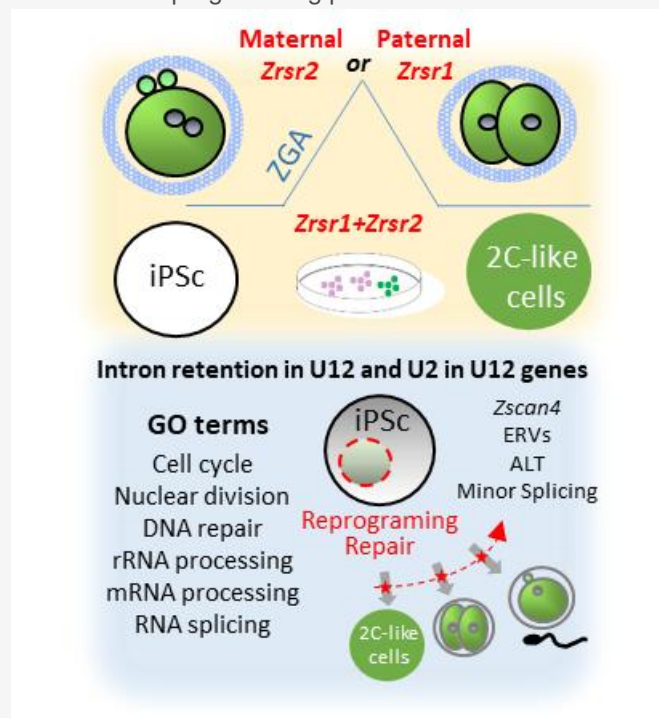


Figure 1. Synopsis of phenotypes produced by *Zrsr1* and *Zrsr2* mutations. Upper panel shows that mutation of maternal *Zrsr2* and paternal *Zrsr1* blocked early embryo development during zygotic gene activation (ZGA) and increased intron retention in U2- and U12-containing genes; and that mutation of either *Zrsr1* or *Zrsr2* led to a blockade of induced pluripotent stem cell (iPSC) conversion to 2C-like cells. The bottom panel shows gene ontology (GO) enrichment terms related to these phenotypes, indicating that disturbed U12-containing genes are critical for cell division, differentiation, and DNA damage response.

References

1. Christopher B. Burge; Richard Padgett; P A Sharp; Evolutionary Fates and Origins of U12-Type Introns. *Molecular Cell* **1998**, *2*, 773-785, 10.1016/s1097-2765(00)80292-0.
2. Anthony G. Russell; J. Michael Charette; David F. Spencer; Michael W. Gray; An early evolutionary origin for the minor spliceosome. *Nature* **2006**, *443*, 863-866, 10.1038/nature05228.
3. Sebastian Canzler; T Samuelsson; U12 type introns were lost at multiple occasions during evolution. *BMC Genomics* **2010**, *11*, 106-106, 10.1186/1471-2164-11-106.
4. Harald König; Nathalie Matter; Rüdiger Bader; Wilko Thiele; Ferenc Müller; Splicing Segregation: The Minor Spliceosome Acts outside the Nucleus and Controls Cell Proliferation. *Cell* **2007**, *131*, 718-729, 10.1016/j.cell.2007.09.043.
5. Richard Robinson; Minor spliceosome, major surprise: it's cytoplasmic. *Journal of Cell Biology* **2007**, *179*, 1086-1086, 10.1083/jcb.1796r1.
6. Kyle Friend; Nikolay G. Kolev; Mei-Di Shu; Joan A. Steitz; Minor-class splicing occurs in the nucleus of the *Xenopus* oocyte. *RNA* **2008**, *14*, 1459-1462, 10.1261/rna.1119708.

7. Heli K. J. Pessa; Cindy L. Will; Xiaojuan Meng; Claudia Schneider; Nicholas Watkins; Nina Maria Perälä; Mariann Nymark; Janne Turunen; Reinhard Lührmann; Mikko J. Frilander; et al. Minor spliceosome components are predominantly localized in the nucleus. *Proceedings of the National Academy of Sciences* **2008**, *105*, 8655-8660, 10.1073/pnas.0803646105.
8. Jarnail Singh; Richard Padgett; Rates of in situ transcription and splicing in large human genes. *Nature Structural & Molecular Biology* **2009**, *16*, 1128-33, 10.1038/nsmb.1666.
9. Peter T. Buckley; Mugdha Khaladkar; Junhyong Kim; James Eberwine; Cytoplasmic intron retention, function, splicing, and the sentinel RNA hypothesis. *Wiley Interdisciplinary Reviews: RNA* **2013**, *5*, 223-230, 10.1002/wrna.1203.
10. Daniele Merico; Maian Roifman; Ulrich Braunschweig; Ryan K. C. Yuen; Roumiana Alexandrova; Andrea Bates; Brenda Reid; Thomas Nalpathamkalam; ZhuoZhi Wang; Bhooma Thiruvahindrapuram; et al. Paul E. Gray Alyson Kakakios Jane Peake Stephanie Hogarth David Manson Raymond Buncic Sergio Pereira Jo-Anne Herbrick Benjamin J. Blencowe Chaim M. Roifman Stephen W. Scherer Compound heterozygous mutations in the noncoding RNU4ATAC cause Roifman Syndrome by disrupting minor intron splicing. *Nature Communications* **2015**, *6*, 8718, 10.1038/ncomms9718.
11. Janne Turunen; Elina Niemelä; Bhupendra Verma; Mikko J. Frilander; The significant other: splicing by the minor spliceosome. *Wiley Interdisciplinary Reviews: RNA* **2012**, *4*, 61-76, 10.1002/wrna.1141.
12. Anouk M Olthof; Katery C. Hyatt; Rahul N Kanadia; Minor intron splicing revisited: identification of new minor intron-containing genes and tissue-dependent retention and alternative splicing of minor introns. *BMC Genomics* **2019**, *20*, 686-19, 10.1186/s12864-019-6046-x.
13. Huiling He; Sandya Liyanarachchi; Keiko Akagi; Rebecca Nagy; Jingfeng Li; Rosemary C. Dietrich; Wei Li; Nikhil Sebastian; Bernard Wen; Baozhong Xin; et al. Jarnail Singh Pearly Yan Hansjuerg Alder Eric Haan Dagmar Wieczorek Beate Albrecht Erik G. Puffenberger Heng Wang Judith A. Westman Richard A. Padgett David E. Symer Albert De La Chapelle Mutations in U4atac snRNA, a Component of the Minor Spliceosome, in the Developmental Disorder MOPD I. *Science* **2011**, *332*, 238-240, 10.1126/science.1200587.
14. Mahmoud Fawzi Elsaid; Nader Chalhoub; Tawfeg Ben-Omran; Pankaj Kumar; Hussein Kamel; Khalid Ibrahim; Yasmin Mohamoud; Eman Al-Dous; Iman Al-Azwani; Joel A. Malek; et al. Karsten Suhre M. Elizabeth Ross Alice Abdel Aleem Mutation in noncoding RNA RNU12 causes early onset cerebellar ataxia. *Annals of Neurology* **2017**, *81*, 68-78, 10.1002/ana.24826.
15. Patrick Edery; Charles Marcaillou; Mourad Sahbatou; Audrey Labalme; Joelle Chastang; Renaud Touraine; Emmanuel Tubacher; Faiza Senni; Michael B. Bober; Sheela Nampoothiri; et al. Pierre-Simon Jouk Elisabeth Steichen Siren Berland Annick Toutain Carol Wise Damien Sanlaville Francis Rousseau Françoise Clerget-Darpoux A. L. Leutenegger Association of TALS Developmental Disorder with Defect in Minor Splicing Component U4atac snRNA. *Science* **2011**, *332*, 240-243, 10.1126/science.1202205.
16. Keiko Horiuchi; Serafín Pérez-Cerezales; Panagiotis Papasaikas; Priscila Ramos-Ibeas; Angela Patricia Lopez-Cardona; Ricardo Laguna-Barraza; Noelia Fonseca Balvís; Eva Pericuesta; Raúl Fernández-González; Benjamin Planells; et al. Alberto Viera José Ángel Suja Pablo Juan Ross Francisco Alen Laura Orio Fernando Rodríguez De Fonseca Belen Pintado Juan Valcarcel Alfonso Gutierrez-Adan Impaired Spermatogenesis, Muscle, and Erythrocyte Function in U12 Intron Splicing-Defective Zrsr1 Mutant Mice. *Cell Reports* **2018**, *23*, 143-155, 10.1016/j.celrep.2018.03.028.
17. Thomas Doktor; Yimin Hua; Henriette Skovgaard Andersen; Sabrina Brøner; Ying Hsiu Liu; Anna Wieckowska; Maja Dembic; Gitte Hoffmann Bruun; Adrian R. Krainer; Brage Storstein Andresen; et al. RNA-sequencing of a mouse-model of spinal muscular atrophy reveals tissue-wide changes in splicing of U12-dependent introns. *Nucleic Acids Research* **2016**, *45*, 395-416, 10.1093/nar/gkw731.
18. Vikas Madan; Deepika Kanojia; Jia Li; Ryoko Okamoto; Aiko Sato-Otsubo; Alexander Kohlmann; Masashi Sanada; Vera Grossmann; Janani Sundaresan; Yuichi Shiraishi; et al. Satoru Miyano Felicitas Thol Arnold Ganser Henry Yang Torsten Haferlach Seishi Ogawa H. Phillip Koefler Aberrant splicing of U12-type introns is the hallmark of ZRSR2 mutant myelodysplastic syndrome. *Nature Communications* **2015**, *6*, 6042-6042, 10.1038/ncomms7042.
19. Tao Xu; Bo Mi Kim; Kyung Jin Kwak; Hyun Ju Jung; Hunseung Kang; The Arabidopsis homolog of human minor spliceosomal protein U11-48K plays a crucial role in U12 intron splicing and plant development. *Journal of Experimental Botany* **2016**, *67*, 3397-406, 10.1093/jxb/erw158.
20. Christine Gault; Federico Martin; Wenbin Mei; Fang Bai; Joseph B. Black; William Bradley Barbazuk; A. Mark Settles; Aberrant splicing in maize rough endosperm3 reveals a conserved role for U12 splicing in eukaryotic multicellular development. *Proceedings of the National Academy of Sciences* **2017**, *114*, E2195-E2204, 10.1073/pnas.1616173114.
21. Fang Bai; Jacob Corll; Donya N. Shodja; Ruth Davenport; Guanqiao Feng; Janaki S. Mudunkothge; Christian J. Brigolin; Federico Martin; Gertraud Spielbauer; Chi-Wah Tseung; et al. Amy E. Siebert William Bradley Barbazuk Shailesh K LaLa. Mark Settles RNA Binding Motif Protein 48 Is Required for U12 Splicing and Maize Endosperm Differentiation. *The Plant Cell* **2019**, *31*, 715-733, 10.1105/tpc.18.00754.
22. Leo R. Otake; Petra Scamborova; Carl Hashimoto; Joan A. Steitz; The divergent U12-type spliceosome is required for pre-mRNA splicing and is essential for development in Drosophila. *Molecular Cell* **2002**, *9*, 439-446, 10.1016/s1097-2765(02)00441-0.
23. Sebastian Markmiller; Nicole Cloonan; Rea M. Lardelli; Karen Doggett; M. Cristina Keightley; Yeliz Boglev; Andrew J. Trotter; Annie Y. Ng; Simon J. Wilkins; Heather Verkade; et al. Elke Ober Holly A. Field Sean Grimmond Graham J. Lieschke Didier Yr Stainier Joan Heath Minor class splicing shapes the zebrafish transcriptome during development. *Proceedings of the National Academy of Sciences* **2014**, *111*, 3062-7, 10.1073/pnas.1305536111.
24. Marybeth Baumgartner; Christopher Lemoine; Sahar Al Seesi; Devi Krishna Priya Karunakaran; Nikita Sturrock; Abdul Rouf Banday;

- Ashley M Kilcollins; Ion Măndoiu; Rahul N Kanadia; Minor splicing snRNAs are enriched in the developing mouse CNS and are crucial for survival of differentiating retinal neurons. *Developmental Neurobiology* **2014**, *75*, 895-907, 10.1002/dneu.22257.
25. Marybeth Baumgartner; Anouk M Olthof; Gabriela S. Aquino; Katelyn C. Hyatt; Christopher Lemoine; Kyle Drake; Nikita Sturrock; Nhut Nguyen; Sahar Al Seesi; Rahul N Kanadia; et al. Minor spliceosome inactivation causes microcephaly, owing to cell cycle defects and death of self-amplifying radial glial cells. *Development* **2018**, *145*, dev166322, 10.1242/dev.166322.
 26. Francisco Alén; Isabel Gómez-Redondo; Patricia Rivera; Juan Suarez; Priscila Ramos-Ibeas; Eva Pericuesta; Raul Fernández-González; Serafín Perez-Cerezales; Keiko Horiuchi; Laura Orío; et al. Fernando Rodríguez De Fonseca Alfonso Gutiérrez-Adán Sex-Dimorphic Behavioral Alterations and Altered Neurogenesis in U12 Intron Splicing-Defective Zrsr1 Mutant Mice. *International Journal of Molecular Sciences* **2019**, *20*, 3543, 10.3390/ijms20143543.
 27. Karen Doggett; Ben B. Williams; Sebastian Markmiller; Fan-Suo Geng; Janine Coates; Stephen Mieruszynski; Matthias Ernst; Tim Thomas; Joan Heath; Early developmental arrest and impaired gastrointestinal homeostasis in U12-dependent splicing-defective Rnpc3-deficient mice. *RNA* **2018**, *24*, 1856-1870, 10.1261/rna.068221.118.
 28. Yawei Gao; Xiaoyu Liu; Bin Tang; Chong Li; Zhaohui Kou; Lin Li; Wenqiang Liu; You Wu; Xiaochen Kou; Jing-Yi Li; et al. Yanhong ZhaoJiqing YinHong WangShe ChenLujian LiaoShaorong Gao Protein Expression Landscape of Mouse Embryos during Pre-implantation Development. *Cell Reports* **2017**, *21*, 3957-3969, 10.1016/j.celrep.2017.11.111.
 29. Lei Gao; Keliang Wu; Zhenbo Liu; Xuelong Yao; Shenli Yuan; Wenrong Tao; Lizhi Yi; Guanling Yu; Zhenzhen Hou; Dongdong Fan; et al. Yong TianJianqiao LiuZi-Jiang ChenJiang Liu Chromatin Accessibility Landscape in Human Early Embryos and Its Association with Evolution. *Cell* **2018**, *173*, 248-259.e15, 10.1016/j.cell.2018.02.028.
 30. Pablo Bermejo-Álvarez; Dimitrios Rizos; D. Rath; P. Lonergan; Alfonso Gutierrez-Adan; Sex determines the expression level of one third of the actively expressed genes in bovine blastocysts. *Proceedings of the National Academy of Sciences* **2010**, *107*, 3394-3399, 10.1073/pnas.0913843107.
 31. Haihong Shen; Xuexiu Zheng; Stephan Luecke; Michael R. Green; The U2AF35-related protein Urp contacts the 3' splice site to promote U12-type intron splicing and the second step of U2-type intron splicing. *Genes & Development* **2010**, *24*, 2389-2394, 10.1101/gad.1974810.
 32. Ken-Ichiro Abe; Satoshi Funaya; Dai Tsukioka; Machika Kawamura; Yutaka Suzuki; Masataka G. Suzuki; Richard M. Schultz; F Aoki; Minor zygotic gene activation is essential for mouse preimplantation development. *Proceedings of the National Academy of Sciences* **2018**, *115*, E6780-E6788, 10.1073/pnas.1804309115.
 33. Toshio Hamatani; Mark G. Carter; Alexei A. Sharov; M.S. Ko; Dynamics of Global Gene Expression Changes during Mouse Preimplantation Development. *Developmental Cell* **2004**, *6*, 117-131, 10.1016/s1534-5807(03)00373-3.
 34. Ursula Eichenlaub-Ritter; Oocyte ageing and its cellular basis. *The International Journal of Developmental Biology* **2012**, *56*, 841-852, 10.1387/ijdb.120141ue.
 35. G. Flach; Martin Johnson; P.R. Braude; R.A. Taylor; V.N. Bolton; The transition from maternal to embryonic control in the 2-cell mouse embryo. *The EMBO Journal* **1982**, *1*, 681-686, 10.1002/j.1460-2075.1982.tb01230.x.
 36. Geppino Falco; Sung-Lim Lee; Ilaria Stanghellini; Uwem C. Bassey; Toshio Hamatani; M.S. Ko; Zscan4: A novel gene expressed exclusively in late 2-cell embryos and embryonic stem cells. *Developmental Biology* **2007**, *307*, 539-550, 10.1016/j.ydbio.2007.05.003.
 37. Kei-Ichiro Ishiguro; Manuela Monti; Tomohiko Akiyama; Hiromi Kimura; Nana Chikazawa-Nohtomi; Miki Sakota; Saeko Sato; Carlo Alberto Redi; Shigeru B. H. Ko; M.S. Ko; et al. Zscan4 is expressed specifically during late meiotic prophase in both spermatogenesis and oogenesis. *In Vitro Cellular & Developmental Biology - Animal* **2016**, *53*, 167-178, 10.1007/s11626-016-0096-z.
 38. Sharon Schlesinger; Eran Meshorer; Open Chromatin, Epigenetic Plasticity, and Nuclear Organization in Pluripotency. *Developmental Cell* **2019**, *48*, 135-150, 10.1016/j.devcel.2019.01.003.
 39. Chames Kermi; Elena Lo Furno; Domenico Maiorano; Regulation of DNA Replication in Early Embryonic Cleavages. *Genes* **2017**, *8*, 42, 10.3390/genes8010042.
 40. Benjamin Cieply; Juw Won Park; Angela Nakauka-Ddamba; Thomas W. Bebee; Yang Guo; Xuequn Shang; Christopher J. Lengner; Yi Xing; Russ P. Carstens; Multiphasic and Dynamic Changes in Alternative Splicing during Induction of Pluripotency Are Coordinated by Numerous RNA-Binding Proteins. *Cell Reports* **2016**, *15*, 247-55, 10.1016/j.celrep.2016.03.025.
 41. Akira Hayakawa; Masao Saitoh; Keiji Miyazawa; Dual Roles for Epithelial Splicing Regulatory Proteins 1 (ESRP1) and 2 (ESRP2) in Cancer Progression. *Plant Promoters and Transcription Factors* **2016**, *925*, 33-40, 10.1007/5584_2016_50.
 42. Yun Huang; Jong Kyoung Kim; Dang Vinh Do; Caroline Lee; Christopher A Penfold; Jan J. Żylicz; John Marioni; Jamie A. Hackett; Azim Surani; Stella modulates transcriptional and endogenous retrovirus programs during maternal-to-zygotic transition. *eLife* **2017**, *6*, 3, 10.7554/eLife.22345.
 43. Takashi Ishiuchi; Rocío Enriquez-Gasca; Eiji Mizutani; Ana Boskovic; Céline Ziegler-Birling; Diego Rodríguez-Terrones1; Teruhiko Wakayama; Juan M. Vaquerizas; María-Elena Torres-Padilla; Early embryonic-like cells are induced by downregulating replication-dependent chromatin assembly. *Nature Structural & Molecular Biology* **2015**, *22*, 662-671, 10.1038/nsmb.3066.
 44. Melanie Eckersley-Maslin; Celia Alda-Catalinas; Marloes Blotenburg; Elisa Kreibich; Christel Krueger; Wolf Reik; Dppa2 and Dppa4 directly regulate the Dux-driven zygotic transcriptional program. *Genes & Development* **2019**, *33*, 194-208, 10.1101/gad.321174.118.
 45. Lulzim Shkreta; Johanne Toutant; Mathieu Durand; James L Manley; Benoit Chabot; SRSF10 Connects DNA Damage to the Alternative

Splicing of Transcripts Encoding Apoptosis, Cell-Cycle Control, and DNA Repair Factors.. *Cell Reports* **2016**, *17*, 1990-2003, 10.1016/j.celrep.2016.10.071.

46. Zarkower, D.; Nakayama, K.; Regulation of mitosis-meiosis transition by the ubiquitin ligase beta-TrCP in male germ cells. *Development* **2017**, *144*, 4137–4147, 10.1242/dev.158485dev.158485.

Keywords

Zrsr1;Zrsr2;U12 introns;zygotic gene activation;2C-like cell

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