Review

Mammalian-specific genomic functions: Newly acquired traits generated by genomic imprinting and LTR retrotransposon-derived genes in mammals

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Abstract: Mammals, including human beings, have evolved a unique viviparous reproductive system and a highly developed central nervous system. How did these unique characteristics emerge in mammalian evolution, and what kinds of changes did occur in the mammalian genomes as evolution proceeded? A key conceptual term in approaching these issues is “mammalian-specific genomic functions”, a concept covering both mammalian-specific epigenetics and genetics. Genomic imprinting and LTR retrotransposon-derived genes are reviewed as the representative, mammalian-specific genomic functions that are essential not only for the current mammalian developmental system, but also mammalian evolution itself. First, the essential roles of genomic imprinting in mammalian development, especially related to viviparous reproduction via placental function, as well as the emergence of genomic imprinting in mammalian evolution, are discussed. Second, we introduce the novel concept of “mammalian-specific traits generated by mammalian-specific genes from LTR retrotransposons”, based on the finding that LTR retrotransposons served as a critical driving force in the mammalian evolution via generating mammalian-specific genes.

Keywords: mammalian-specific genomic functions, evolution and development, genomic imprinting, newly acquired genes from LTR retrotransposon, placenta and viviparity, brain

Introduction

Mammals never develop to term without the cooperation of both of the parental genomes: embryos with either two maternally or paternally derived genomes, such as parthenogenetic or androgenetic embryos, exhibit early embryonic lethality (Fig. 1a).1–3) This is because of functional differences between the paternally and maternally derived genomes that are attributable to the presence of imprinted genes, paternally expressed genes (Peg) and maternally expressed genes (Meg), exhibiting parent-of-origin specific monoallelic expression.4–6) This mechanism is unique to mammals among the vertebrates, and is known as genomic imprinting (Fig. 1a).1–8) Therefore, it is of great interest to know how this mammalian-specific epigenetic mechanism emerged and why it has been widely conserved in mammals despite the apparent developmental disadvantage that arises because the limitation of the monoallelic expression of essential genes is quite likely to endanger the survival of individuals in both the pre- and postnatal periods. What is the advantage conferred by genomic imprinting to overcome this difficulty? Accidental or unexpected parthenogenesis may be life-threatening and undesirable in females, because the environmental conditions suit-
able for breeding pups, such as food, temperature and climate, are constrained by season. Therefore, a prohibition of parthenogenetic development might be advantageous for mammalian reproduction in terms of fitting with the seasonal conditions.4) 9) Another biological advantage for mammals may be the absence of the placental trophoblast tissues in the parthenogenetic conceptus, because they have an infiltrative nature and a means of invading the maternal uterus. Mammalian females can avoid developing the malignant ovarian teratocarcinomas that arise during parthenogenetic development.10)
These hypotheses suggest that there are certain advantages in genomic imprinting for mammalian development. However, this does not directly mean that genomic imprinting specifically evolved for the purpose of prohibiting parthenogenetic development and/or invasive trophoblast development. In general, it is difficult to prove such hypotheses by experiment. An important clue may be provided from the historical point of view. A relationship between placenta formation and genomic imprinting in mammals has also been proposed in the form of various “placenta hypotheses” [4, 9, 11]. Thus, it is expected that the continuing investigation of genomic imprinting will provide clues in the molecular mechanism connecting these phenomena, as well as in the elucidation of mammalian evolution itself as one of the mammalian-specific genomic functions.

Recent studies have clearly demonstrated the essential nature of genomic imprinting in the current form of the mammalian developmental system, where the reciprocal regulation of Pegs and Megs via DNA methylation on imprinting control regions (ICRs) plays a key role [12–14]. Pegs and Megs cannot be expressed from the individual parental chromosomes simultaneously, therefore, two parental chromosomes with different expression profiles are necessary for normal development. Several lines of evidence also indicate that this mechanism emerged as a defense against exogenous DNA, because these ICRs themselves emerged in the mammalian genome as newly integrated DNA sequences during the course of evolution [15–19].

Moreover, genomic imprinting has brought into focus a new dimension to the model of mammalian evolution through the identification of two Pegs, Peg10 and Peg11/Rtl1, which play an essential role in mammalian development via the formation and maintenance of a mammalian-specific placenta, respectively, because these two genes are mammalian-specific genes derived from an LTR-retrotransposon [20–26]. It is known that many housekeeping genes and genes related to various signaling pathways that are conserved in other organs are involved in placenta formation [27]. In addition, several placenta-specific genes generated by gene duplication in a wide range of multigene families are known to play an essential role in this process. However, Peg10 and Peg11/Rtl1 were the first examples of mammalian-specific, essential placental genes acquired from a specific LTR-retrotransposon. Thus, a new concept established which says that the evolution of mammalian-specific traits occurred by the acquisition of mammalian-specific genes from LTR retrotransposons [24, 25].

At least 30 LTR retrotransposon-derived genes exist in the human genome as therian- and eutherian-specific genes [25]. A series of knockout mouse studies on these genes has gathered persuasive evidence that at least some of these genes are essential in the current form of the developmental and reproductive systems, which suggests that they made critical contributions to mammalian evolution in a variety of ways, such as the establishment of viviparity [22–26] and presumably certain sophisticated brain functions. Thus, it is highly probable that the LTR retrotransposons served as one of the driving forces in mammalian evolution; the acquisition of novel genes from LTR retrotransposons is one of the important evolutionary mechanisms as well as being a gene/genome duplicating mechanism [28, 29]. Importantly, from the viewpoint of the LTR retrotransposon-derived genes, eutherians and marsupials are entirely different animal groups, because most genes exist as eutherian-specific genes while a few are marsupial-specific except for a single exception, Peg10, which is common to both groups [25].

In the last part of this review, we also discuss a hypothetical two-step model from LTR retrotransposons to endogenous genes during mammalian evolution in which the nearly neutral theory of molecular evolution played an essential underlying role in the first part and then was succeeded by Darwinian evolution at the critical selection step [24, 25, 30–32].

1. The investigation of “mammalian-specific genomic functions”, including our own personal research history

Mammals are considered one of the most successful animals in the Cenozoic era because they have exhibited adaptive radiation since the extinction of the dinosaurs. They have evolved an efficient viviparous reproductive system using a placenta and a highly developed cerebral neocortex. In this review, we will endeavor to address how these unique characteristics have emerged in mammalian evolution, and what kinds of changes occurred in the mammalian genomes as evolution proceeded. A key conceptual term we will employ is “mammalian-specific genomic functions”, a concept covering both mammalian-specific epigenetics and genetics. This term is necessary, because mammals have unique epigenetic mechanisms, such as X chromosome inactivation (XCI) and genomic imprinting, as well
as certain unique genes acquired from retroelements, such as retrotransposons and retroviruses.

Discoveries of XCI and genomic imprinting in mammals in the late 20th century provided an insight into how to approach these evolutionary issues from the perspective of mammalian-specific epigenetic mechanisms.1–11,33,34 Importantly, both XCI and genomic imprinting play an essential role in mammalian development. Female embryos with two active X chromosomes exhibit early embryonic lethality, so it is clearly evident that inactivation of one of the two X chromosomes in females is essential for mammalian development.11,33,34 As mentioned, parthenogenetic or androgenetic embryos also exhibit early embryonic lethality; therefore, both parental genomes are essential for mammalian development.1–3 Interestingly, in all of these cases, the developmental problems are associated with placental abnormalities. The genes involved in the mechanism underlying XCI as well as those causing a variety of imprinted phenotypes have been extensively examined using knockout mice and in human patients with diseases of genomic imprinting. It should be noted that genomic imprinting phenomena are also found in other organisms, such as certain insects (e.g. Sciara)35 and plants (e.g. Arabidopsis sp., Zea mays).36 but it is most widespread in mammals via its own, specific regulatory mechanism, and the same is true for XCI.

The completion in the early 21st century of the “human genome project” has dramatically changed the situation with regard to these evolutionary issues because this investigation has expanded to a wide range of subsequent genome projects. At present, sequencing of the whole genome of more than 100 mammalian species, including marsupial and monotreme species is available as well as that of many other vertebrate species including reptiles, birds and fish.37–42 Through comprehensive and comparative genomics, only 1.5% of the mammalian genome is comprised of protein coding genes, but it is occupied by abundant retrotransposons, such as short interspersed nuclear elements (SINEs), long interspersed nuclear elements (LINEs) and LTR retrotransposons/retroviruses. These retrotransposon-derived DNA sequences were thought to be so-called junk DNA and even potentially harmful for host organisms for a long time; however, we and others demonstrated by a series of knock-out mouse analyses that at least some of these sequence play essential roles in the current mammalian developmental system as endogenously functional genes specific to mammals.20–26,43–45 In other words, it turns out that certain LTR retrotransposon- and retrovirus-derived genes are actually responsible for a variety of mammalian-specific traits, such as the formation, maintenance and endocrinological regulation of the placenta and presumably for a variety of brain functions. The strategy of focusing on “mammalian-specific genomic functions” has thus been proven to be very successful thus far, and is expected to continue to be so in the future in terms of shedding light on the evolutionary issues related to mammals.

Two themes, genomic imprinting and the newly acquired genes from LTR retrotransposons, which serve as the representative examples of the mammalian-specific epigenetics and genetics, respectively, seem distinctly separated and independent of each other. However, the former investigation actually triggered the latter theme, because the first and second examples of LTR retrotransposon-derived genes were identified as paternally expressed imprinted genes, such as PEG10 and PEG11. Therefore, we would like to provide the behind-the-scenes story of the birth of the second theme of the acquired genes from LTR retrotransposons from the genomic imprinting work before getting into the main topic. When we started our genomic imprinting investigation in Surani’s laboratory in Cambridge in 1990, we hypothesized that functional differences between parental genomes were due to the presence of genes exhibiting parent-of-origin-specific monoallelic expression. Subsequently, we conducted a systematic screening for Pegs and Megs by developing a new method of subtracting the genes expressed in the parthenogenetic and androgenetic embryos from those in normally fertilized embryos.46–49

One of us (T. K-I.) also had the idea that genomic imprinting was somehow induced by responses against exogenous DNA integrated into the mammalian genome. Therefore, her research target has long been the imprinted genes derived from exogenous DNA, such as retrotransposons. Eventually, a similar idea on the origin of genomic imprinting was proposed by Barlow in 1993.15 Meanwhile, the other one of us (F. I.) made a plan to identify the imprinted genes that are related to embryonic lethality, because such information would provide an important clue to help elucidate the biological significance of genomic imprinting. Therefore, his ultimate target became the genes responsible for the early embryonic lethality of the parthenogenetic embryos and the late fetal/neonatal lethality caused by paternal and maternal disomies of the
mouse chromosome 12 (Chr12). Another reason to choose the latter was its medical importance, because paternal disomy of human orthologous chromosome 14 (upd(14)pat) causes a severe genomic imprinting disease associated with neonatal lethality by a respiratory problem, presumably related to a bell-shaped thorax and abdominal defects, placental overgrowth and polyhydramnios during gestation, with severe mental and postnatal growth retardation when the patients survived. Therefore, the development of a proper diagnosis and treatment of this syndrome are greatly anticipated.

In 2006 and 2008, we finally identified Peg10 and Peg11/Rtl1 as the major imprinted genes responsible for the parthenogenetic death and the uniparental disomy of mouse Chr12/human upd(14)pat, and elucidated their essential roles in placenta formation and maintenance, respectively. Interestingly, both Peg10 and Peg11/Rtl1 are derived from an LTR retrotransposon and in fact are actually newly acquired genes specific to the mammals. At that time, we realized that our targets we had been pursuing were the same genes all along. To our great pleasure, this has personal meaning for us at the same time leading to a new concept of ‘mammalian-specific traits generated by mammalian-specific genes newly acquired from LTR retrotransposons’. Since then, we have devoted a tremendous amount of effort to elucidating the biological functions of the LTR retrotransposon-derived genes, as will be described in section 3.

2. Genomic imprinting as a mammalian-specific epigenetic mechanism

2-1. Discovery of genomic imprinting and the imprinted genes, Pegs and Megs. Genomic imprinting was discovered by pronuclear exchange experiments in mice which showed that both parthenogenetic and androgenetic embryos with either two maternal or two paternal pronuclei exhibited early embryonic lethality, albeit with different morphological anomalies (Fig. 1a). In the former case, embryos were small but appeared normal yet died due to severe placental defects. The latter embryos exhibited severe growth retardation associated with abnormal placenta overgrowth. This clearly demonstrated the functional differences between the paternal and maternal genomes and critical requirement of both for normal development in mammals. As parthenogenesis is often observed naturally or experimentally among vertebrates, such as fish, amphibians, birds and reptiles, the complete absence of parthenogenesis is a unique feature of mammals.

A series of experiments using mice with Robertsonian translocations also provided strong genetic evidence for the functional differences between the paternally and maternally derived chromosomes, because mice with uniparental duplication of certain specific chromosomal regions exhibited a variety of defects in development, growth and/or behavior (Fig. 1b). As a result, approximately 20 chromosomal imprinted regions in the mouse genome have been identified. Subsequently, genomic imprinting has attracted considerable attention as a mammalian-specific epigenetic mechanism. These phenomena are an apparent exception to Mendelian genetics in mammals, so humans suffer from a variety of non-Mendelian genetic disorders by the breakdown of this mechanism.

In 1991, the first three imprinted genes, Insulin-like growth factor 2 (Igf2), Insulin-like growth factor 2 receptor (Igf2r) and H19, imprinted maternal expressed transcript (H19) were identified. As Igf2 and Igf2r are related functionally because the Igf2 protein promotes embryonic and placental growth as a growth factor, while the Igf2r protein inhibits embryonic and placental growth by degrading Igf2 through its ability to bind Igf2 and transport it to the lysosome. As Igf2 exhibits paternal expression (Peg) while Igf2r exhibits maternal expression (Meg), these two genes fit very well with the weak version of ‘the conflict hypothesis’ that states that paternally expressed genes promote embryonic growth, while maternally expressed genes inhibit embryonic growth as a consequence of the genetic conflict between the parental and maternal alleles during mammalian evolution. It proposes their developmental advantages were conferred on the viviparous system that females are able to accept more than two males (polygamy). Not all, but nevertheless a substantial number of imprinted genes fit with this prediction, so the concept of “parental conflict” in the genome has come to be widely accepted. On the other hand, the strong version of “the conflict hypothesis” further predicted that the genomic imprinting mechanism itself originated under continuous pressure exerted by this genetic conflict during the course of evolution. We will discuss this issue, the origin of genomic imprinting, later in section 2-4.

H19 is located next to Igf2 on the distal portion of mouse chromosome 7, therefore, these two genes are positionally related. Retrospectively, this finding can be seen to have been very important, because
this was the first report demonstrating that imprinted genes exist in clusters. It subsequently led to the concept of a critical regulatory mechanism in which Pegs (such as Igf2) and Megs (such as H19) are reciprocally regulated by the DNA methylation status in the differentially methylated regions (DMRs) in each imprinted region.\(^4\),\(^5\),\(^12\),\(^13\) Thus, DMRs are regarded as the imprinting control region (ICR). In “the insulator model” independently proposed by Tilghman’s and Felsenfeld’s groups,\(^12\),\(^13\) H19-DMR extends from the H19 promoter region to an upstream insulator sequence where insulator-binding proteins, such as CTCF, bind and block several downstream enhancers when H19-DMR is not DNA methylated, after which the repression of the upstream Igf2 occurs. In contrast, when it is DNA methylated, the CTCF proteins cannot bind to the insulator sequence, so Igf2 becomes actively expressed instead of the repression of H19 due to promoter methylation (Fig. 2). An antisense RNA model and a bipartite regulation model have been proposed in other imprinted regions, but the reciprocal regulation of Pegs and Megs by DMR methylation is the common consequence of all of the imprinted regions.\(^4\),\(^5\),\(^14\) It is of critical importance to take this aspect of genomic imprinting into account, as will be discussed in section 2-3.

Several systematic methods of screening for imprinted genes, including ours, have contributed to the further identification of novel imprinted genes and a determination of the precise location of the imprinted regions in the estimated chromosomal regions.\(^46\)–\(^49\),\(^59\)–\(^61\) By our subtraction-hybridization method using parthenogenetic and androgenetic embryos combination with genome walking from the newly identified imprinted genes, we succeeded in isolating more than 20 imprinted genes (Peg1–12 and Meg1–5, 8–10 etc. including 4 of the known imprinted genes) in 8 imprinted regions, such as the mouse sub-distal chromosome 2 (Chr2), proximal Chr6, sub-proximal Chr6, proximal Chr7, central Chr7, distal Chr7, proximal Chr11 and distal Chr12. The mapping results were ultimately in good agreement with those of the previous genetic experiments using mice with uniparental chromosomal duplications, as shown in Fig. 1b. Thus, this series of our experiments united the two original observations of genomic imprinting, the pronuclear transplantation\(^1\)–\(^3\) and the Robertsonian translocation experiments,\(^6\)–\(^8\) thereby contributing to the establishment of the current concept that genomic imprinting phenomena are attributable to the existence of Pegs and Megs.\(^4\),\(^5\),\(^14\)

2-2. Roles of Peg10 and Peg11/Rtl1 in the placenta. According to the genomic imprinting map, it appeared highly probable that the responsible gene for the parthenogenetic death should be located on mouse proximal Chr6, because this is the only imprinted region that causes early embryonic lethality upon maternal duplication, like the parthenoge-
netic embryos, with two maternally derived pronuclei (Fig. 1b)\(^{6,8}\). We then searched in the region near *Sarcoglycan epsilon* (*Sgce*), the first imprinted gene identified in this region.\(^{61}\) Using the human genome sequence information on chromosome 7q21 which is syntenic to the mouse proximal Chr6, we identified *PEG10/Peg10* next to *SGCE/Sgce* in both humans and mice in 2001 (Fig. 3a, left).\(^{20}\) In the case of the mouse distal Chr12, we extensively sequenced around *Meg3/Gtl2*\(^{49}\), the first imprinted gene in this cluster, by genome walking and determined the entire imprinted gene cluster comprising * Peg9/Dlk1, Meg3/ Gtl2, Peg11/Rtl1, Peg10/antiPeg11, Meg8/Rian* and *Meg9/Mirg* in mice (Fig. 3a, right). Almost the same result was published by Micheal Georges’ group analyzing the same imprinted region in the ovine and human genomes including *DLK1, GTL2, PEG11, antiPeg11* and *MEG8* in 2001.\(^{21}\)

Importantly, both *Peg10* (the former region) and * Peg11* (the latter region) are genes acquired from a sushi-ichi related retrotransposon because they encode proteins exhibiting a high homology with the Gag and Pol proteins of a sushi-ichi retrotransposon, respectively.\(^{20,21}\) *PEG11* was later officially renamed *Retrotransposon-like 1 (RTL1)*.\(^{62}\) They were actually TK-I’s original target of imprinted genes, obtained ten years after the project had been started. The fact that their origin was an LTR retrotransposon is very interesting, at that time nobody knew whether they were functional, and it would require a period of another five to seven years to elucidate their biological functions in mammalian development and evolution.

It was big surprise when our knockout mouse experiments on * Peg10* and * Peg11/Rtl1* clearly demonstrated that both are essential genes in development: *Peg10* is responsible for the early embryonic lethality caused by maternal duplication of the mouse proximal Chr6 (Fig. 3b)\(^{6,8,22}\) and * Peg11/Rtl1* is responsible for the late fetal/neonatal lethality caused by the maternal disomy of mouse Chr12 (Fig. 3c)\(^ {6,8,23}\). At that point, the two concepts, the retrotransposon-derived imprinted genes and the responsible genes for embryonic lethality became united and thus brought a new dimension to mammalian evolution: that LTR retrotransposon-derived genes exerted a critical impact on mammalian evolution (see section 3).\(^{24,25}\) Our first presentation on the roles of retrotransposon-derived * Peg10* and * Peg11/Rtl1* in placental formation and functions had an impact at the genomic imprinting meeting in 2004 in Montpellier and stimulated considerable work on the retrotransposon-derived genes in mammals thereafter.\(^ {63,64} \)

* Peg10* KO mice exhibited early embryonic lethality due to poor placental growth associated with complete lack of labyrinth and spongiotrophoblast layers (Fig. 3b).\(^ {22}\) The labyrinth layer is an essential part of the placenta, where nutrient and gas exchange occurs between fetal and maternal blood cells, therefore, * Peg10* KO embryos stopped growth on day 9.5 and could not grow any further. Importantly, the phenotypes of the * Peg10* KO embryos and placentas resemble those of parthenogenetic embryos and placenta closely, strongly suggesting that * Peg10* is a major gene responsible for parthenogenetic death.\(^ 1,3,22\)

One half of the * Peg11/Rtl1* KO mice exhibited late fetal lethality while the other half exhibited neonatal lethality associated with late fetal growth retardation (Figs. 3c and d).\(^ {23}\) This corresponds very well with what is observed in mice with maternal duplication of the mouse distal Chr12.\(^ {8,50}\) In the labirynth layer of the * Peg11/Rtl1* KO placenta, severe abnormality of the fetal capillaries was observed: they were clogged at many sites because endothelial cells had been phagocytosed by surrounding endothelial cells, indicating that * Peg11/Rtl1* plays an essential role in the maintenance of the fetomaternal interface of the placenta during gestation.

* Peg11/Rtl1* is overlapped with the maternally expressed non-coding RNA *antiPeg11/antiRtl1* that comprises seven micro RNAs (miRNAs) that target * Peg11/Rtl1* mRNA by an RNAi mechanism\(^ {23,62,65}\) (Fig. 3e). Our * Peg11/Rtl1* KO mice lacked most portions of both * Peg11/Rtl1* and * antiPeg11/antiRtl1*, therefore, upon maternal transmission, * antiPeg11/antiRtl1* KO mice were generated exhibiting neonatal lethality associated with placental overgrowth due to a 2–3 fold accumulation of * Peg11/Rtl1* mRNA, resulting in an overproduction of the * Peg11/Rtl1* protein. In the labyrinth layer of * antiPeg11/antiRtl1* KO mice, fetal capillary size expansion associated with severely damaged surrounding trophoblast cell layers was observed. Thus, both the lack and overexpression of * Peg11/Rtl1* cause lethality around birth, just by different mechanisms (Figs. 3c and d).

The * AntiPeg11/antiRtl1* KO mouse is a good model for paternal duplication (or paternal disomy) of mouse Chr12 as well as human Chr 14.\(^ {23,50–52}\) It should be noted that one important difference is that a double dose of * Peg11/Rtl1* without maternally expressed * antiPeg11/antiRtl1* leads to a 4–6 fold
(a) (b) (c) (d) (e)

Fig. 3
increment of Peg11/Rtl1 mRNA, causing more severe abnormal phenotypes in the case of paternal duplication (or paternal disomy) than antiPeg11/antiRtl1 KO with only one normal Peg11/Rtl1 allele. In patients with upd(14)pat-like phenotypes due to deletion of this locus that contains an imprinting control region (IG-DMR), symptom severity correlated well with the degree of PEG11/RTL1 overexpression, indicating that the overexpression of PEG11/RTL1 is attributable to the major phenotypes of upd(14)pat. In the case of upd(14)mat, both PEG11/RTL1 and PEG9/DLK1 are responsible for growth deficiency in an additive manner.

2-3. The Essential nature of genomic imprinting. Why is genomic imprinting widely conserved in mammals? An important key to answering this question lies in the reciprocal regulation mechanism of Pegs and Megs, as mentioned earlier, because the two parental chromosomes have different epigenotypes (different DNA methylation status in the DMR) that are necessary for the expression of both Pegs and Megs. As shown in the insulator model, it is apparent that Pegs and Megs cannot be expressed from the individual parental chromosomes simultaneously (Fig. 2). In this model, the essential components comprising the reciprocal regulation mechanism are four kinds of functional genomic units, such as promoters, an insulator, enhancer(s) and a DMR, as well as insulator binding proteins, such as CTCF, which interact with the insulator sequence. All of the components except the DMR are ubiquitous in eukaryotes, so the DMR may be a key factor in this mechanism, with the important constraint that the DMR overlaps with the insulator sequence as well as the neighboring promoter in this model. However, it is apparent that the reciprocal expression of Pegs and Megs is intrinsically controlled by their own genomic DNA; in other words, the mechanism is already integrated into the mammalian genome.

This provides a good explanation for why genomic imprinting, a mechanism which promotes the monoallelic expression of Pegs and Megs, is highly conserved in mammals so as to ensure the current form of the mammalian development system, even though it seems disadvantageous upon first consideration (complementation hypothesis) (Fig. 4). In other words, monoallelic expression of Pegs and Megs in the imprinted gene cluster would appear to be a good means for avoiding a situation in which one half (Pegs or Megs) is completely repressed throughout the course of life, even though the other half (Megs or Pegs, respectively) exhibits normal biallelic expression. There are two kinds of imprinted regions, paternally imprinted regions (defined as the regions where paternal allele is methylated in sperm: D in Fig. 4) and maternally imprinted regions (where maternal allele is methylated in oocyte: E in Fig. 4). In either case, the situation is the same and the Pegs and Megs are reciprocally regulated by the DMR DNA methylation status.

This Conclusion is deduced from studies which generated embryos without genomic imprinting memories, such as cloned mice produced from primordial germ cells (PGCs) that give rise to eggs and sperm in females and males, respectively. Imprinted memories (maternal and paternal imprints) are already established in both eggs and sperm as DNA methylation marks on the DMRs, respectively, and are stably maintained after fertilization in the somatic cell lineages. However, these parental memories are erased in the PGCs from embryonic day 9.5 to 12.5 (B in Fig. 4) and reestablished again in oocytes after birth (during oocyte maturation) in females (E in Fig. 4) and in spermatogonia cells just around the time of birth in males (D in Fig. 4). When using d10.5 PGCs with almost complete parental imprinted memories remained, like somatic cells, as donors, the clones (d10.5 PGC clones) developed to term. In contrast,
d12.5 PGC clones exhibited early embryonic lethality because they had completely erased imprinted memories (B and C in Fig. 4b). In this case, only half of the Pegs and half of the Megs are active, while the other halves are repressed. In explicit detail, only Pegs are active in the maternally imprinted regions where Megs are repressed, and only Megs are active in the paternally imprinted regions where Pegs are repressed in the default state of genomic imprinting (C in Fig. 4). It should be noted that the repressed Pegs in the latter regions as well as the repressed Megs in the former regions need paternal and maternal imprints (DNA methylation) for their induction, respectively (D and E in Fig. 4). In addition, it is also reported that embryos without any maternally imprinted memory (combination of normal sperm (C in Fig. 4) and oocyte in the default status (B in Fig. 4)) cannot develop to term and exhibit early embryonic lethality in both humans and mice, indicating the genomic imprinting mechanism is essential in the current form of the mammalian developmental system. It also should be noted that the imprinted gene expression pattern in Fig. 4C is illustrated based on the result of d12.5 PGC cloned embryos, and not that of PGCs themselves. However, it seems consistent with the previous results showing that migrating PGCs (d9.5–10.5) basically exhibit monoallelic expression, while d11.5 PGCs exhibit biallelic expression. Similarly, Figs. 4D and E represent the imprinted gene expression pattern of androgenetic and parthenogenetic embryos, respectively, and not those of the sperm or oocyte themselves.

2-4. Establishment of genomic imprinting regions during mammalian evolution. How did genomic imprinting emerge in the course of mammalian evolution? As mentioned previously, the differential DNA methylation of the DMRs is the most critical factor in this regulatory mechanism. Therefore, the emergence of DMRs in the genome must be a
pivotal event in the emergence of genomic imprinting in the mammals. How did these regions come to be differentially methylated in the male and female germ lines? Are there any conserved recognition sequences among them? Unfortunately, such sequence homology has never been identified in the DMRs in either the paternal nor the maternally imprinted regions.

An important clue to a way to overcome this problem came from the analysis performed on PEG10. We found that PEG10 is also present in both Australian and South American marsupial species, it is absent from the platypus, an Australian monotreme species, work done in collaboration with Renfree at the University of Melbourne and Graves at the Australian National University. This demonstrated that the putative PEG10 retrotransposon was inserted into the genome of a common therian ancestor and became an endogenous gene before the divergence of the eutherians and marsupials 160 million years ago (Ma) (Fig. 5a). An important fact is that PEG10 is also imprinted and paternally expressed in the tammar wallaby, an Australian marsupial species, and there is a PEG10-DMR overlapped with the PEG10 promoter within the 5′ non-coding region of PEG10 mRNA, suggesting that the PEG10 promoter was derived from an LTR of the ancient retrotransposon from which PEG10 originated. What is clear is that the PEG10-DMR also emerged in the therian ancestor together with PEG10, implying that the DMRs originated from inserted DNA sequences exogenous origin (Figs. 5b and 6). The PEG10 imprinted region is the first imprinted region with a common DMR in the eutherians and marsupials, demonstrating that it is at least one of the oldest imprinted regions in mammalian history.

Another imprinted region that has a common DMR in the therians is the IGF2-H19 region. Reik's group collaborated with Renfree to show that H19 and H19-DMR exist, and that IGF2 and H19 are reciprocally expressed in the tamer wallaby in the same manner as in eutherian species. Although, no information is available on the presence or absence of H19 in the monotremes, it is highly likely that there is no H19 (and H19-DMR) in the platypus, because it was reported that IGF2 is biallelically expressed. Therefore, this case further suggests that the H19-DMRs originated from inserted DNA sequences of exogenous origin together with H19 (Fig. 6). In addition, there are also several reports that the small imprinted genes that reside in the introns of other genes, such as Mcts2, Nap1l5, Inpp5f-v2, U2af1-rs1 and Nnat, are thought to have been inserted into their present positions by cDNA retrotransposition. In every case, the DMR likely emerged as novel CpG islands (CGI) at the same time as the retrotranspositioning of each gene occurred.

We examined the generality of this hypothesis by reviewing the time of novel CGI emergence for all the maternal DMR loci obtained in collaboration with Renfree. The comprehensive and comparative analyses demonstrated that the emergence of the novel CGIs occurred universally in the maternal DMR loci at different time points during mammalian evolution, presumably in correspondence with the time when the imprinting regulation started in each locus, and the same seems true for all the three paternal DMRs (Fig. 6). The evidence suggests that the acquisition of the DMRs, such as insertion events from the CpG rich exogenous DNA sequences and also by cDNA retrotransposition, is a common evolutionary pathway for generating novel imprinted regions, and that genomic imprinting originated as the defense mechanism against the insertion of exogenous DNA into the mammalian genome.

As mentioned above, imprinting arose at many different time points during mammalian evolution, presumably due to different selective pressures at different loci, and it is still continuing to evolve. It should be noted that the PEG10 and IGF2-H19 regions are the only two imprinted regions with common DMRs in the therians identified so far, implying that genomic imprinting was originally related to the prevention of parthenogenetic growth as well as the regulation of the growth of both the placenta and embryo from the beginning, that is, they may be the fundamental biological significance of this mechanism, as initially proposed by the placental hypotheses, complementary hypothesis and the weak version of the conflict hypothesis, despite the great diversity of roles that imprinted genes play at present.

3. LTR retrotransposon-derived genes in mammals

3-1. Evolution of mammalian-specific traits by acquisition of SIRH genes. As stated in section 2-2, the acquisition of novel genes took place via a conversion from the LTR retrotransposons to endogenous genes during mammalian evolution. According to the definition originally proposed by Gould and his colleagues, PEG10 and PEG11/RTL1 are very good examples of “exapted” (domes-
Fig. 5. Emergence of PEG10 in a common ancestor of marsupials and eutherians. a. Comparison of genome sequences among the higher vertebrates. The insertion of the PEG10 retrotransposon occurred after the split of the therians and monotremes, and endogenization of the PEG10 gene was completed before the eutherian-marsupial split. b. Emergence of PEG10-DMR from a newly inserted DNA. Modified from Figs. 1 and 5 in ref. 16.
ticated) genes that exhibit novel functions that are different from the original functions in the original organisms. We demonstrated that PEG10 is specific to the therian mammals,\(^{16}\) while Ferguson-Smith’s group demonstrated that PEG11/RTL1 is a eutherian-specific gene that is absent from marsupials.\(^{53}\) These facts clearly demonstrate that PEG10 was acquired after the split of the therians from the monotremes 166 or 186 Ma and before the eutherian/marsupial split 160 Ma.\(^{16},25\) Furthermore, PEG11/RTL1 was acquired after the eutherian/marsupial split and before the split of the three major eutherian lineages, boreoeutheria (including euarchontoglires and laurasiatheria), afrotheria and xenarthra 120 Ma (see Fig. 9).\(^{25},53,86\) According to the Darwinian theory of evolution, it is highly probable that PEG10 and PEG11/RTL1 were once positively selected and thus presumably contributed to the establishment of the subclasses and infraclasses of the mammals, the therians and the eutherians, respectively.\(^{14},24,25\) While this supports the previous hypothesis that retrotransposons in the genome provide genetic materials to generate novel genes,\(^{85}\) it has also led to the novel concept that at least in some cases the LTR retrotransposon-derived genes contributed to mammalian evolution, such as the emergence of viviparity.

PEG10 and PEG11/RTL1 exhibit a high homology with the same LTR retrotransposon, a sushi-ichi retrotransposon isolated from the pufferfish (Takifugu rubripes).\(^{20},21,87,88\) Using a sushi-ichi Gag amino acid sequence as a query, genes with high homology were screened from the human and mouse genomes. Eleven genes, including PEG10 and PEG11/RTL1, were identified and named sushi-ichi-related retrotransposon homologue (SIRH)1-11 (also called mammalian retrotransposon-derived (MART) or SUSHI genes.\(^{22},63,64\) Most of the SIRH genes exhibit homology with Gag, but not with Pol, except for PEG10/SIRH1, PEG11/RTL1/SIRH2 and SIRH9/ZCCHC5 (Zinc finger CCHC domain-containing 5) (Fig. 7). Interestingly, 8 out of the 11 genes, i.e. SIRH4 to SIRH11, locate on the X chromosome. Therefore, they exhibit monoallelic expression like the imprinted genes PEG10/SIRH1 and PEG11/RTL1/SIRH2. A series of knockout

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**Fig. 6.** Emergence of DMRs in imprinted regions in mammals. Paternally and maternally methylated DMRs are shown in blue and red, respectively. The locations of the mouse chromosomes are shown. Only PEG10 and H19-IGF2 DMRs are common to the eutherians and marsupials. The imprinted regulation of these two regions started in the eutherians associated with chromosome rearrangements near the DMRs. Modified from Fig. 6 in ref. 18.
mouse experiments on Sirh genes has been carried out in an effort to elucidate their biological functions in development, reproduction, growth and behavior, as well as their presumed contribution to eutherian evolution.

Female Sirh7/Ldoc1 (leucine zipper, downregulated in cancer 1, also called Mart7) KO mice had structural and related endocrinological problems in the placenta and exhibited delayed parturition along with a low pup weaning rate.26) The placenta is a major endocrine organ during gestation, and in the case of mice, a variety of placental cells, such as giant trophoblast, spongiotrophoblast and cytotrophoblast cells, produce progesterone (P4) and placental lactogen I/II (PL1 and 2) as well as several prolactin-like proteins in order to maintain pregnancy and determine the timing of parturition. Sirh7/Ldoc1 encodes a small Gag-like protein comprising only 151 amino acids, but plays an essential role in controlling the differentiation and maturation of a wide variety of placental cells, thereby regulating both the amount and timing of placental hormones (Fig. 8).26) Thus, Sirh7/Ldoc1 is another essential gene in development as well as reproduction, suggesting that it was also positively selected during the eutherian evolution because of the reproductive advantage it conferred. Together with the results on Peg10 and Peg11/Rtl1 KO mice, this additional evidence provides further support for the critical role of the LTR retrotransposon-derived genes in the evolution of the mammalian viviparous reproduction system (Fig. 8).

In addition to viviparity, another unique characteristic in mammals is higher brain function. Among the SIRH family members, Sirh11/Zcchc16 (also called Mart4) is unique in that it does not exhibit any placental expression during mouse development, but is expressed in the brain, testis, ovary and kidney. Sirh11/Zcchc16 KO mice exhibited abnormal behaviors related to cognition, including attention, impulsivity and working memory, conferring a critically important advantage in both the competition of daily life and eutherian evolution (Fig. 8).89) It also suggests that human SIRH11/ZCCHC16 may be a good candidate for X-linked intellectual disability (XLID). Another example of brain-related phenotypes was observed in Sirh3/Ldoc1 (Ldoc 1 like, also called Mart6) KO mice, such as lower activity during the night period (Irie, in preparation). As mice are nocturnal animals and the same seems likely true for the eutherian ancestors 160 Ma because they lived under the rule of dinosaurs, it is highly probable that acquisition of Sirh3/Ldoc1 in the eutherian ancestors was quite advantageous for survival in the field, thus it was positively selected in eutherian evolution (Fig. 8).
3-2. LTR retrotransposon-derived genes in eutherians and marsupials. Further comparative genomics demonstrated that all of the SIRH genes (PEG11/RTL1/SIRH2 and SIRH3-11) are eutherian-specific except for PEG10/SIRH1, strongly supporting our hypothesis that the SIRH genes exerted a major impact on therian and eutherian evolution in a variety of ways, but also in a therian- and eutherian-specific manner. We subsequently found SIRH12 only in marsupial species, such as the tammer wallaby and opossum, an Australian and a South American marsupial species, respectively, although the opossum SIRH12 is degenerated and has surely lost its function (Fig. 9). A similar result was obtained by the analysis of another LTR retrotransposon-derived gene family in mammals, the paraneoplastic Ma antigen (PNMA) family, which exhibits a high homology with gypsy DR retrotransposon isolated from zebra fish (Danio Renio). At least 19 and 15 PNMA/Pnma genes have been confirmed in humans and mice, respectively, with the difference in number due to rodent-specific deletion of PNMA6A-D. All of the PNMA genes encode only a Gag-related and not a Pol-related region, and 11 out of 19 are X-linked genes. Moreover, it is also highly likely that they are all eutherian-specific. In contrast, PNMA-MS1 (PNMA marsupial-specific 1) is only present in the marsupial species (both in the tammer wallaby and opossum), while PNMA-MS2 is specific to the opossum (Fig. 9). Thus, the number of LTR retrotransposon-derived genes exhibits a difference between the eutherians and marsupials, for example, comprising 11 and 2 genes in the case of the SIRH family, respectively, including one common gene (SIRH1/PEG10), and 19 and 2 genes in the case of the PNMA family, with no common genes. The reason for this difference remains unclear, but it should be noted that many pseudogenes are observed in the marsupial genomes and almost none in the eutherian genomes, suggesting that the eutherians have a better ability to use the LTR retrotransposons for novel purposes. An important conclusion deduced from these results is that the eutherians and marsupials are totally different animal groups in terms of the LTR retrotransposon-derived genes, suggesting that these genes contributed to the diversification of the two viviparous mammalian groups via a generation of eutherian- and marsupial-specific traits like PEG11/RTL1, SIRH7/LDOC1, SIRH3/LDOC1L and SIRH11/ZCCHC16. Therefore, analyses on the remaining Sirh KO as well as Pnma KO mice are expected to provide further insight into the impact of the LTR retrotransposons on the acquisition of a series of novel traits during the course of eutherian evolution.

3-3. How did retrotransposons become endogenous genes? According to the Darwinian theory of evolution, two essential factors in biological evolution are “mutation” and “the environment”.

Fig. 8. Biological functions of the Sirh genes in mice. The biological functions deduced from five KO mice experiments on Peg10, Peg11/RTL1, Sirh7/Ldoc1, Sirh3/Ldoc1l and Sirh11/Zcchc16 are shown. Only the major phenotypes of each KO mouse are shown.
Organisms change their traits by the mutations and the environment select the fittest by means of natural selection. In the light of alteration of the genomic information, the creation of novel genes by a gene/genome duplication mechanism\(^28\),\(^29\),\(^96\),\(^97\) is regarded as a "mutation" in addition to the known types of a simple point mutation, insertion or deletion of DNA, and chromosomal rearrangement via homologous and non-homologous recombination as well as cross-breeding between closely related species (homoploid hybrid speciation)\(^98\),\(^99\) and symbiosis between and/or among different organisms.\(^100\)–\(^103\) It is now evident that the acquisition of novel genes from exogenous DNA, such as LTR retrotransposons/retroviruses, is another important mechanism in the creation of novel genes.\(^20\)–\(^26\),\(^43\)–\(^45\),\(^63\),\(^64\),\(^88\)–\(^95\) It is an important task to understand how this process has proceeded or is still proceeding in the course of mammalian evolution.

As retrotransposons are usually intrinsically harmful to host organisms, host organisms possess several defense mechanisms for regulating their transcription, thereby preventing their excessive propagation. Mammals use epigenetic mechanisms for this purpose, such as DNA methylation and histone modifications.\(^104\) the integrated retrotransposons are usually heavily DNA methylated and transcriptionally silenced in somatic cells\(^72\),\(^105\) while some are repressed in ES cells, mainly by histone H3K9 methylation.\(^106\) As a result, they behave like neutral genes and can be transmitted to the next generation (Fig. 10a).

According to the "neutral theory of molecular evolution" proposed by Kimura, neutral mutations can become fixed in a population by the mechanism of random drift.\(^30\),\(^31\) Ohta extended this theory and proposed the "nearly neutral theory of molecular evolution" that states that, even nearly neutral (less harmful) mutations can become fixed in a population when the population size is small enough.\(^32\) We think it highly probable that the neutral and nearly neutral modes of evolution played an essential background role by inactivating and neutralizing the integrated retrotransposons so as to generate the potential genetic material in the first place. A gradual conversion from being potentially harmful to slightly advantageous must have taken place as the result of multiple mutations of the genes during this period. At this point, Darwinian forces came into play, and a fraction of these genes were selected and became more advantageous by natural selection so as to generate the prototype and present type of novel

Fig. 9. Emergence of the SIRH and PNMA genes in two mammalian lineages. Most of the SIRH and PNMA genes are eutherian-specific, with a few being marsupial-specific, while the single exception of PEG10 is conserved in both the eutherians and marsupials. Interestingly, no SIRH or PNMA genes are present in egg-lying montremes. Modified from Fig. 3 in ref. 25.
Fig. 10. Acquisition of LTR retrotransposon-derived genes (a two-step model). Hypothetical scenarios for the embryonic (a) and extraembryonic tissues (b) are shown. Inserted retroelements are usually fully DNA methylated in somatic cells, while leaky expression of these elements are observed in extraembryonic tissues, such as the placenta, due to the lower DNA methylation level. As a result, they behave as neutral (a) and nearly neutral (less harmful) (b) genes and become fixed in a population according to the mechanisms described in the neutral and nearly neutral theory of molecular evolution, respectively. A key point is the rapid shift to the pressures of Darwinian evolution occurs when they become slightly advantageous.
endogenous genes for the host organisms, and subsequently have been maintained by a purifying selection process (Two step model, Fig. 10a). However, there are two apparent problems with this hypothetical scenario. First, how is the gene that confers this slightly advantage recognized in somatic cells, since it was completely repressed by DNA methylation? Second, is it possible that random mutation could have generated such a slightly advantageous gene from a potentially harmful gene without selection on multiple occasions?

In extraembryonic tissues, such as the yolk sac and placenta, a leaky expression of retroposonos and retroviruses occurs because of a lower level of DNA methylation (Fig. 10b). It should be pointed out that this two-step process may be accelerated in the placenta, because the placenta is vulnerable to whether the integrated retroposonos and its mutated forms turn out to be harmful or advantageous. In such a situation, a swift transition from the first (the state of nearly neutral evolution) to the second step (that of Darwinian evolution) would take place more readily. In this light, the placenta may be regarded as a natural laboratory for mammalian evolution. It should also be noted that the nearly neutral evolution proposed by Ohta makes a contribution to phenotypical evolution as well as Darwinian evolution, while the neutral evolution originally proposed by Kimura actively contributes to phenotypical evolution, as in the case of gene duplication.

This idea is consistent with the fact that the acquired PEG10, PEG11/RTL1 and SIRH7/LDOC1 actually play essential roles in the placenta. However, what about the cases of SIRH3/LDOC1L and SIRH11/ZCCHC16? Sirh3/Ldoc1l expression is observed in the mouse placenta although its major functional site seems to be in the brain, while Sirh11/Zcche16 does not exhibit any placental expression, as mentioned. It seems that the original SIRH genes in the eutherian ancestors might have some placental function, but subsequently shifted their central expression site to the brain after they became endogenous genes. Alternatively, such SIRH genes might be generated by the gene duplication from other SIRH genes playing a major role in the placenta, with the newly differentiated copy taking on a new role in the brain during evolution.

4. Discussion and future prospects

In this review, we have discussed two themes, genomic imprinting and the LTR retroposonos-derived genes, as mammalian-specific genomic functions. We would like to also emphasize that genomic imprinting is a very important issue, not only for genomic imprinting diseases, but also for future regenerative medicine and reproductive technology, because genomic imprinting (parental DNA methylation) memories are labile in ES, iPSC and even in somatic cells in vitro, although they are stably maintained throughout life in vivo. Recently, PGC-like cells that give rise to germ cells, such as sperm and oocytes, were reportedly induced from human and mouse iPSC cells in vitro. Morphologically, such oocytes look normal, but it remains unknown whether the imprinted memories are in fact normally erased and reestablished in these artificial processes in vitro. It is well known that both a lack of and abnormally imprinted memories lead to severe developmental failure and genomic imprinting diseases, respectively. Thus, genomic imprinting issue may be a big hurdle for the application of ES and iPSC cells to human regenerative medicine and reproductive technology.

The mechanism of imprint erasure in PGCs is a very important part of mammalian-specific epigenetics. The recent discovery of hydroxymethyl cytosine (hmC) as a key intermediate of DNA demethylation has completely changed the previous view. It is now clear that mC is first converted to hmC by the Tet enzyme. However, it is now under intense debate which plays the major role in this process, the passive (dilution) DNA demethylation mechanism via continuous cell divisions under a condition of no hemimethylation activity of DNMT1, or the active DNA demethylation mechanism proceeded without cell division. Alternatively, both may play different but essential roles in this process, respectively. It is possible that the erasing mechanism is much more complex than generally thought.

Another major remaining question with regard to genomic imprinting is how the maternal and paternal DNA methylation imprints are established in the female and male germ cells in a locus-specific manner. The de novo DNA methyltransferases DNMT3A and DNMT3B, as well as the co-factor DNMT3L, are major players in the general mechanisms underlying DNA methylation, but the locus specificity is not explained by this alone. What is the specific targeting mechanism of DNA methylation? Histone H3K4 methylation may be a key involved in this process, because histone H3K4 methylation blocks the association of DNMT3L
with the N-terminal of histone H3, thus preventing the DNMT3A-DNMT3L complexes from accessing DNA. Importantly, the deletion of KDMB1, one of histone H3K4 demethylation enzymes, causes the loss of maternal imprints at certain maternal DMRs, presumably due to a substantial increment of H3K4 methylation, but without any abnormality in any of the three paternal and the remaining maternal DMRs. These data demonstrate that both DNA methylation and histone modification play crucial roles in establishing the parental imprints and suggest that certain histone modification enzymes function as locus- and parental-specific targeting of the de novo DNA methylation machinery.

However, recent genome-wide DNA methylation analyses on oocyte and sperm have presented a completely different perspective on this issue. It is now clear that the oocyte and the sperm are at the outset poorly and highly DNA methylated, respectively, so the di
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the preimplantation stage. The loss of ZFP57, a
DNA methylation and histone modification play crucial roles in establishing the parental imprints and suggest that certain histone modification enzymes function as locus- and parental-specific targeting of the de novo DNA methylation machinery.

It should be noted that the interaction between DNA methylation and histone modification also play crucial roles in maintaining the parental imprints in the preimplantation stage. The loss of ZFP57, a KRAB domain zinc finger protein, induces somatic loss of DNA methylation at multiple DMRs. It functions together with its cofactor, KRAB-associated protein 1 (KAP1, also called TRIM28). ZFP57 brings KAP1 to chromatin, thereby promoting the recruitment of repressive chromatin modifiers, including ESET/SETDB1, which controls H3K9 trimethylation. ZFP57/KAP1 complexes also interact with DNMTs and the hemimethylated DNA binding protein UHRF1 (also called NP95), which are essential for the maintenance of DNA methylation.

It is also known that maternal KAP1 in the oocyte is essential for the maintenance of DMR methylation at several DMRs in the first round of cell division. Thus, many epigenetic factors are known to be involved in the maintenance of the DNA methylation of DMRs in the preimplantation stage, but the elucidation of the mechanisms of the locus- and parental origin-specific DMRs of genomic imprinting awaits further investigation.

In the second part of this review, we discussed the emergence of the LTR retrotransposon-derived genes in mammalian-lineages. From the viewpoint of mammalian evolution, endogenous retroviruses (ERV) may be another important factor. Retroviruses are structurally related to the LTR retrotransposons: both have LTR sequences at both ends and encode GAG and POL proteins, while the former also encodes an additional ENV protein for infection into cells. It is known that ENV genes have also become essential endogenous genes in a lineage-specific manner in the eutherians and marsupials, e.g. several syncytins and FEMATRIN-I, that is, they were acquired independently from different retroviruses. The capacity of human syncytin to fuse cells to produce multinucleated cells was first demonstrated in vitro. As it exhibits a high level of expression in syncytiotrophoblast cells in the human placenta, it was proposed that syncytin is essential for formation of these cells. Out of 18 ENV related DNA sequences in the human genome, only syncytin-1 and -2 that are derived from human-specific ERVs, HERV-W and HERV-FRD, and were integrated into a primate lineage 25 and >40 Ma, respectively, exhibited the fusogenic activity in cell fusion assays.

Mice have two syncytin genes, syncytin-A and syncytin-B, derived from Muridae family-specific integrations of HERV-F/H related ERV(s) approximately 20 Ma. Importantly, syncytin-A knockout mice exhibit mid-fetal lethality, presumably due to a functional abnormality of one of the two syncytiotrophoblast layers, and double knockout of both syncytin-A and -B was shown to lead to a more severe phenotype and early embryonic lethality, demonstrating that they are clearly essential for syncytiotrophoblast cell formation in vivo, as expected. Among the eutherians, placentation morphology have substantially diverged. Therefore, it is highly probable that other syncytins also have important roles in the placenta in an order- or family-specific manner during the time of mammalian radiation (Fig. 11).
the periods when the therian ancestors appeared, and other SIRH and PNMA genes in the eutherian and marsupial ancestors appeared. This is the time when mammalian viviparity had begun and two different types of viviparous systems using chorioallantoic and choriovitelline (yolk sac) placentas were established in the eutherians and marsupials, respectively (Figs. 9 and 11).

What were the environmental condition(s) that promoted these radical changes in the mammalian reproductive system? It is reported that in the Phanerozoic eon (from 550 Ma to the present), the highest atmospheric oxygen (30%) was observed in the Carboniferous period 300 Ma, however, it dramatically decreased to 15% around one of five major extinctions that occurred at the Permian-Triassic (P-Tr) boundary 250 Ma (Fig. 11). The period with the lowest atmospheric oxygen (12–13%) was estimated to have taken place in a period from 190 to 150 Ma after the next mass extinction of Triassic-Jurassic (Tr-J) boundary approximately 200 Ma. This scenario correlates well with the timing of the divergence of the therians from the monotremes (186 or 166 Ma), eutherians and marsupials (160 Ma) and three eutherian lineages (120 Ma), such as boreoetheria (including euarchontoglires and Laurasiatheria), afrotheria and xenarthra, is shown (see Fig. 9). The three major mass extinctions deeply related to mammalian evolution, such as Permian-Triassic (P-Tr: 250 Ma), Triassic-Jurassic (Tr-J: 200 Ma) and Kreide (Cretaceous)-Paleogene (K-Pg: 65 Ma) extinctions are shown.

It is well known that an adaptive radiation of mammals took place after the extinction of the dinosaurs at the Kreide (Cretaceous)-Paleogene...
(K-Pg) boundary 65 Ma. At that time, atmospheric oxygen concentration was gradually increased to a point near the present concentration (~20%) despite a slight lag just after K-Pg mass extinction, therefore, the low oxygen concentration was no longer a major selective pressure. It is highly probable that large vacant ecological niches on both land and sea allowed the ancestors of eutherian and marsupial lineages to become diversified by continuing the acquisition of a variety of retrovirus-derived genes, suggesting that release from powerful selective pressure also serves as a major driving force in macroevolution.\textsuperscript{30,148}

Interestingly, Koala retrovirus (KoRV) is currently undergoing endogenization in the koala genome and this process possibly started within the last 200 years.\textsuperscript{149–151} It is reported that KoRV infection has expanded in the Northern Australia Koala population to reportedly cause leukemia, lymphoma and immunosuppression, and the integrated KoRV DNAs are actually transmitted to the next generation in the manner of endogenous genes. Therefore, retrotransposon and retroviral endogenization may be a fairly ordinary process in the long course of evolution. Novel genes may continuously appear by this mechanism, both at present and in the future, although this may depend upon a mechanism of random gene drift such that it remains to be fixed in the koala population. Therefore the strategy we have introduced in this review can also be applied to elucidate human evolution by identifying human-lineage-specific acquired genes derived from the different kinds of retroviruses that have infected us and become endogenized at different stages of the ongoing process toward the primate and finally the human in evolution.

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Profile

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