

Title	Genetic signatures of evolution of the pluripotency gene regulating network across mammals
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Citation	Genome Biology and Evolution (2020)
Issue Date	2020-08-11
URL	http://hdl.handle.net/2433/254136
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Type	Journal Article
Textversion	publisher

Genetic signatures of evolution of the pluripotency gene regulating network across mammals

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Abstract

Mammalian pluripotent stem cells (PSCs) have distinct molecular and biological characteristics among species, but to date we lack a comprehensive understanding of regulatory network evolution in mammals. Here, we carried out a comparative genetic analysis of 134 genes constituting the pluripotency gene regulatory network across 48 mammalian species covering all the major taxonomic groups. We report that mammalian genes in the pluripotency regulatory network show a remarkably high degree of evolutionary stasis, suggesting the conservation of fundamental biological process of mammalian PSCs across species. Nevertheless, despite the overall conservation of the regulatory network, we discovered rapid evolution of the downstream targets of the core regulatory elements and specific amino acid residues that have undergone positive selection. Our data indicate development of lineage-specific pluripotency regulating networks that may explain observed variations in some characteristics of mammalian PSCs. We further revealed that positively selected genes could be associated with species' unique adaptive characteristics that were not dedicated to regulation of PSCs. These results provide important insight into the evolution of the pluripotency gene regulatory network underlying variations in characteristics of mammalian PSCs.

Keywords

adaptation, embryonic development, gene network, Mammalia, natural selection, pluripotent stem cells

Significance statement

Pluripotent stem cells (PSCs) have emerged as advanced cellular and genetic material for mammalian research. Despite their interspecies similarities related to their ability to differentiate into other somatic cells and to divide to make more stem cells with maintenance of the undifferentiated state, PSCs from different species show variations in some of their characteristics such as pluripotent state and gene expression. Various studies have shown molecular mechanisms underlying the maintenance of PSC characteristics that constitute a complex gene regulatory network. Unfortunately, such knowledge is largely limited to primates and rodents. The present study provides comparative analyses involving pluripotency-regulating genes across mammalian taxa. As such, it helps the understanding of the genetic basis for the evolution of the mammalian PSCs.

Introduction

Pluripotent stem cells (PSCs) are undifferentiated cells that exhibit unlimited self-renewability and pluripotency, the potential to give rise to cells from all three embryonic germ layers. Pluripotent embryonic stem cells are isolated from the inner cell mass of developing pre-implantation mouse or human blastocysts (Evans & Kaufman 1981; Martin 1981; Thomson 1998). Although PSCs of rare animal species represent powerful resources for research and conservation, little equivalent derivations have been attempted with these species for ethical and technical reasons.

Recent advances in somatic cell reprogramming into induced PSCs (iPSCs) have broadened the possibility of obtaining PSCs from variety of mammals, including endangered species (Ben-Nun et al. 2011). Derivation of iPSCs has been reported from a wide range of taxonomic groups, including Carnivora (Shimada et al. 2009; Verma et al. 2012, 2013; Menzorov et al. 2015), Cetartiodactyla (Ezashi et al. 2009; Han et al. 2011; Liu et al. 2012), Chiroptera (Mo et al. 2014), Lagomorpha (Osteil et al. 2013), Metatheria (Weeratunga et al. 2018), Monotremata (Whitworth et al. 2019), Perissodactyla (Ben-Nun et al. 2011; Breton et al. 2013), Rodentia (Takahashi & Yamanaka 2006; Liao et al. 2009; Miyawaki et al. 2016; Lee et al. 2017), and Primates (Takahashi et al. 2007; Tomioka et al. 2010; Marchetto et al.

2013; Wunderlich et al. 2014; Ramaswamy et al. 2015). However, questions remain about species variation in the properties of the PSCs, such as pluripotent state, reprogramming efficiency, and optimal culture conditions.

A highly interconnected pluripotency gene regulatory network (PGRN) controls the characteristics of PSCs (Li & Belmonte 2017). Mammals seem to possess evolutionarily conserved PGRN because iPSCs from different taxonomic groups have been derived using human genetic sequences of identical sets of transcription factors (*OCT4*, *SOX2*, *KLF4*, and *MYC*; collectively referred to as OSKM). On the other hand, various pluripotent states and configurations are observed across different species (Ezashi et al. 2016; Weinberger et al. 2016; Paterson et al. 2018), indicating diversity and specificity of the PGRN among species. However, our understandings of the PGRN are narrow, largely limited to primates and rodents (Manor et al. 2015), and to date the evolutionary history of the PGRN across different mammalian taxonomic groups has not been studied in depth.

Comparative approaches can provide insights into the evolution of gene regulatory processes by examining the conservation and divergence of networks (Thompson et al. 2015). Evolutionary conservation and adaptations can be inferred by detecting purifying and positive selection (Nielsen et al. 2007). We have recently described genetic signatures for phenotypic

adaptations in cetacean lipid metabolism (Endo et al. 2018), and numerous studies have revealed the effects of natural selection in the development of adaptive characteristics of animals and plants (Lenski 2017). Importantly, effects of changes on a regulatory process depends on the hierarchical position of the changes within the regulatory network (Erwin & Davidson 2009) so an holistic approach is recommended. Evolutionary pattern of the PGRN within the Class Mammalia and how natural selection acts on the genes involved in the PGRN, however, are yet to be described.

In this study, we aimed to investigate whether any evolutionary pattern among mammalian taxa could be the result of selection of genes involved in the PGRN. To achieve this we conducted a comparative selection analyses of the PGRN genes across mammals covering all the major taxonomic groups. (i) To assess the evolutionary conservation profiles of the mammalian PGRN, we estimated trends in the stringency of purifying selection. (ii) To identify rapidly evolving subcircuits of the PGRN, we compared the evolutionary rate based on the ratio of substitution rates at non-synonymous and synonymous sites (dN/dS). (iii) To detect phylogenetic inference of the variations in the PGRN, we identified genes under positive selection for lineages and investigated concordance in functional regions of the proteins. This paper presents the evolutionary history underlying the conservation and

variations in mammalian pluripotency gene regulatory network, PGRN, and provides the genetic basis for the evolution of the PGRN in mammals.

Methods

Data sampling

Target genes were selected to include all 138 genes involved with “Signaling pathways regulating pluripotency of stem cells” (hsa04550) from KEGG pathway database (Kanehisa et al. 2017). Human coding DNA sequences (CDS) of those genes were used as references and blasted against publicly available genomes of 61 mammals, covering all major taxonomic groups. *Gallus gallus*, *Xenopus tropicalis*, and *Danio rerio* were used as an outgroup. The set of genes were tested for orthologs using OMA stand alone v2.3.0 (Altenhoff et al. 2018). When a CDS was not inferred as an ortholog, another CDS was selected based on its annotation and tested with OMA again. Because the genomes were of varying quality, mammalian species with the same or lower number of orthologs than *Gallus gallus* were removed. Genes in which orthologs were found in less than 80% of the study species were also excluded from the further analyses. Orthologs were aligned with using PRANK v.140603 (Löytynoja & Goldman 2005) and the aligned sequences were further tested for their quality

with GUIDANCE2 v2.02 (Sela et al. 2015) using the default settings. For the chicken genes, *NANOG* (Gene ID: 100272166, NP_001139614.1) was included in analyses even though it was not determined as ortholog by OMA because it is well studied and known ortholog of human *NANOG* (Fuet & Pain 2017). A consensus tree was prepared based on commonly accepted mammalian phylogenies (Meredith et al. 2011; Perelman et al. 2011) and applied for later natural selection analyses. The gene ontology (GO) and the KEGG orthology (KO) were determined according to the Gene Ontology Annotation (GOA) and KEGG, respectively.

Testing the stringency of natural selection

Natural selection was tested for each gene using RELAX (Wertheim et al. 2015) in HyPhy (Pond et al. 2005). RELAX is a general hypothesis testing framework that determines whether selective strength, distinguishing purifying or positive selection, was relaxed or intensified in the subsets of interest (Wertheim et al. 2015). RELAX estimates the selection intensity parameter, k , where $k \geq 1$ indicates intensification and $k < 1$ indicates relaxation of natural selection where $\omega > 1$ represents positive selection and $\omega < 1$ represents purifying selection. Mammalian lineages were set as a test branch and non-mammalian vertebrates as a reference branch. Because RELAX requires reference branches, genes without non-mammalian

orthologs were excluded. *P* values were corrected for False Discovery Rate (FDR) using the Benjamini-Hochberg method (Benjamini & Hochberg 1995) with a cutoff of 10%.

Evolutionary rate of each gene

Evolutionary rate of each gene was estimated by using the model M0 in CodeML (PAMLv.4.8) (Yang 2007). The synonymous substitution rate (*dS*), the non-synonymous substitution rate (*dN*), and their ratio, *dN/dS* (also noted as ω) were calculated, where *dN/dS* < 1 indicates purifying selection, *dN/dS* = 1 indicates neutral, and *dN/dS* > 1 indicates positive selection. The *dN/dS* ratio was estimated for each gene using mammalian orthologs.

Comparison of conservation patterns among subcircuits

The categories of subcircuits were determined according to KEGG descriptions in the "Pathway" category, including Core, JAK-STAT, MAPK, *OCT4 (POU5F1)*, *SOX2*, and *NANOG* (OSN) activated, OSN suppressed, PI3K-Akt, TGF β , and Wnt signaling pathways.

As for the Core subcircuits, we included the genes in the core transcriptional network, OSN and the genes which interact with the OSN inside of nucleus. When one gene was tagged in multiple subcircuits, it was grouped in all the related pathways. Because subcircuits contain

unequal sample sizes, multiple paired comparisons for the means of dN/dS ratios were statistically tested using ANOVA followed by the Tukey-Kramer post-test with P value of 0.05.

Lineage-specific positive selection

Positively selected genes were identified using the branch-site model (Zhang et al. 2005) implemented in CodeML (PAMLv.4.8) (Yang 2007). The modified model A as an alternative hypothesis was compared to the model A1 as the null. P values were evaluated under the likelihood ratio test (LRT) by comparing $2\Delta l$ of the two models to the 1 : 1 mixture of 0 and χ^2 distribution. Lineage-specific positive selection was tested by setting each contemporary species and ancestral branch as *a priori* specified target separately and corrected for FDR using the Benjamini-Hochberg method (Benjamini & Hochberg 1995) with a cutoff of 10%.

Functional impact

Positively selected sites were determined using the Bayes empirical Bayes (BEB) implemented in the branch-site model (Yang et al. 2005) with 95% posterior probability. Lineage-specific amino acid substitutions were explored in positively selected genes

identified in ancestral branches as indicative of significant changes in protein function (Tian et al. 2013). The lineage-specific amino acid substitutions were identified manually by looking at a conversion of residue that occurred and fixed in more than 90% of the descendant contemporary species. Functional domain structures and regions were referred to UniPort and Pfam using human protein as a reference.

Results

Summary of sample data

To investigate the evolutionary pattern of the gene regulatory network, the PGRN, that maintains pluripotency and self-renewability of pluripotent stem cells, we performed comparative genetic analyses across mammalian species (**Figure 1**). We collected the protein coding sequences of PGRN related genes covering all the major taxonomic groups. A total of 61 mammalian species were used in this study including Classes Primates, Rodentia, Cetartiodactyla, Chiroptera, Carnivora, Insectivora, Afrotheria, and Metatheria, with at least two species from each. For each species, we searched orthologs of the 138 genes assigned to functions associated with signaling pathways regulating pluripotency of stem cells. Our filtering strategy retained 134 gene sets from 48 mammalian species for later analyses (see

Methods, Supplementary Table S1).

Evolutionarily conserved profiles across mammals

To assess evolutionary constraint and reduction of selective strength in the mammalian PGRN, we examined the level of purifying selection with each gene. Of the 127 orthologous genes analyzed, we observed intensification of purifying selection in 53 genes (41.7%) as well as relaxation in 9 genes (7.1%) at $P < 0.05$ FDR $< 10\%$, revealing that intensification of purifying selection was more common among the PGRN (**Figure 2, Supplementary Table S2**). For the commonly used reprogramming factor, OSKM, we found high degrees of genetic conservation in *SOX2*, *KLF4*, and *MYC* across mammals. We observed relaxation of purifying selection in a number of genes involved in the molecular mechanisms that determine the pluripotent states (*APC2*, *DUSP9*, *INHBE*, *LEFTY2*, and *RIF1*) and in a number of oncogenes and tumor suppressor genes across mammals (*APC2*, *FZD6*, *INHBE*, *REST*, *RIF1*, *WNT4*, and *WNT8A*). Intensification of purifying selection was observed across the PGRN, while relaxation appeared to be limited in particular subcircuits. This led to the question whether any conservation patterns among the PGRN subcircuits.

Conservation patterns among the PGRN subcircuits

We then asked if there are differences in degree of conservation among subcircuits constituting the PGRN. In the purifying selection analysis identified above, we found relatively higher number of genes with intensified purifying selection in JAK-STAT, OSN suppressed, and PI3K-Akt signaling pathways (52.9%, 50%, and 44.8%, respectively), while more relaxed genes were found in OSN activated and Wnt signaling pathways (20% and 10%, respectively) (**Table 1**).

To gain insights into the main force that has shaped the evolution of mammalian PGRN, we estimated the evolutionary rates, dN/dS , of each gene. The estimated dN/dS values ranged from 0.0036 to 0.3614. All the genes showed dN/dS values lower than 1 and 95.5% lower than 0.2 (**Figure 3A, Supplementary Table S3**). Mean dN/dS values of each subcircuit were generally low suggesting high degree of evolutionary stasis at level of protein-coding sequence (**Supplementary Table S4**). ANOVA indicated that evolutionary rates differ among subcircuits (**Supplementary Table S5**), and the following Tukey-Kramer showed that the OSN activated subcircuits, the downstream of the core transcriptional network, had significantly higher evolutionary rate compared to other subcircuits with $P < 0.05$ (**Figure 3B, Supplementary Table S5**).

Lineage-specific positive selection

If genes in the OSN activated subdivision experience lower evolutionary constraint across mammals, some species may have developed unique characteristics through changes in these genes over evolutionary time periods. To test this, we performed positive selection analysis for the genes in the OSN activated subdivision on all the ancestral branches and contemporary species in the mammalian taxonomic tree containing our study species. Of the 15 OSN activated genes, we identified 8 positively selected genes in 4 ancestral branches and 22 contemporary species with 10% FDR correction (**Figure 4, Supplementary Table S6**). The four ancestral branches where positively selected genes were found include the common ancestor of Eutheria, Primatomorpha (Primates and the flying lemur), Megachiroptera, and Pan. The number of positive selection events was the highest in the gene *TCF3*, which is an integral component of the core regulatory circuitry of PSCs (Cole et al. 2008) (one ancestral branch and eight contemporary species). While three of the four tested epigenetic regulators (KEGG orthology, KO: 03036) have shown to be under intensified purifying selection across mammals (**Figure 2**), a relatively high number of positive selection events with *KAT6A*, *SETDB1*, and *JARID2* were observed in Primates (five events over nine total events in

Primates and over 11 events with these genes across mammals).

Functional impact of positively selected genes at nucleotide level

To assess whether the identified positively selected genes have undergone functional modification, we investigated the positively selected sites of the eight positively selected genes for each selected lineage (**Supplementary Table S7**). The positions of positively selected sites were compared to protein functional regions using humans as reference. Although the majority of the positively selected sites were found outside of known functional regions, we found sites under positive selection in functional regions of five proteins encoded by *KAT6A*, *LEFTY2*, *NANOG*, *SETDB1*, and *TCF3* (**Figure 5**). We also found lineage-specific amino acid substitutions in functional regions of the four positively selected genes at ancestral branches, including *KAT6A*, *LEFTY2*, *RIF1*, and *TCF3* (**Figure 6, Supplementary Table S8**). These lineage-specific amino acid substitutions were unique to and shared among the descendent contemporary species, indicating functional importance of these genetic mutations (Tian et al. 2013).

Discussion

Genetic conservation of the PGRN

The relatively higher number and prevalence of evolutionary conserved genes across the PGRN indicates overall comparability of the regulating mechanisms for pluripotency and self-renewability of mammalian PSCs (**Figure 2**). In context of the reprogramming somatic cells into induced pluripotent stem cells, although human sequences are effective (Ben-Nun et al. 2011), the efficient combination of reprogramming factors may differ among species (Tomioka et al. 2010; Verma et al. 2013; Mo et al. 2014; Weeratunga et al. 2018). Evidence of evolutionary conservation in the three commonly used reprogramming factors explains the efficiency of human genetic sequences for reprogramming somatic cells from variety of species (Ezashi et al. 2016). On the other hand, low success rate and inefficiency of reprogramming by human OSKM are reported in some species (Whitworth et al. 2019; Verma et al. 2013; Mo et al. 2014). For increasing reprogramming efficiency, additional expression of other pluripotency-associated genes have been shown to enhance reprogramming rate (Takahashi & Yamanaka 2016). We detected significant evidence of purifying selection across mammals for *TBX3*, which improves reprogramming efficiency in mouse (Han et al. 2010), implying that *TBX3* might be a first candidate enhancer for derivation of mammalian iPSCs with low reprogramming efficiency.

The characteristics of mammalian PSCs appear to differ among species (Ezashi et al. 2016; Weinberger et al. 2016; Paterson et al. 2018) and reduction in selective constraint highlights evolutionary flexibility and innovation (Lahti et al. 2009; Moczek 2010; Hunt et al. 2011). For example, pluripotent state of PSCs can be described with multiple types such as naive or primed states, and the amenabilities of the pluripotent state *in vitro* differ between species (Boroviak et al. 2015). Although naive state PSCs are cultured with dual inhibition (2i) of MEK and GSK3 and leukemia inhibitory factor (LIF), human and mouse embryonic stem cells show distinct transcriptional responses to the 2i/LIF (Huang et al. 2014). Our findings of relaxation of purifying selection on genes, such as *RIFI*, may explain the diversity of the amenable pluripotent state of the species. *RIFI* is involved in cellular response to LIF (GO: 1990830). *INHBE* and *LEFTY2* participate in the regulation of MAPK cascade (GO: 0043408), the target of MEK inhibitor, and the protein encoded by *DUSP9* is the essential regulator of MAPK (GO: 0000165, 0000187, 0000188). *APC2* is known to form a protein complex with the scaffold Axin and the kinases GSK3 and CK1 (Pronobis et al. 2015). Mammals lack correlation between body size or life span and cancer risk which is referred to as "Peto's paradox" (Peto et al. 1975; Caulin & Maley 2011; Peto 2015). Our findings of the relaxation of selective strength for oncogenes and tumor suppressor genes may imply

evolutional flexibility in mammalian cancer resistance.

Variable evolutionary conservation patterns among PGRN subcircuits

The patterns of genetic conservation among the PGRN subcircuits provide insight into the conservation and variations of the characteristics of mammalian PSCs (**Table 1, Figure 3B**). Multiple upstream signaling pathways serve to maintain the self-renewability and pluripotency of PSCs. We found relatively high degree of conservation in JAK-STAT, OSN-suppressed, and PI3K-Akt signaling pathways, suggesting conservation of fundamental biological characteristics and homeostasis in mammalian PSCs. JAK-STAT signaling pathway is stimulated by LIF and essential for self-renewal (Niwa et al. 1998). PI3K signaling pathway plays a crucial role for proliferation in mouse embryonic stem cells (Takahashi et al. 2005) and equivalency of its function in mouse and primate ES cells has been reported (Watanabe et al. 2006). Furthermore, the role of PI3K signaling pathway in the proliferation of pluripotent stem cells has been demonstrated in the horse (Whitworth et al. 2014). Our findings support the common regulatory mechanisms observed among phylogenetically distant taxonomic groups. On the other hand, we observed the relatively higher number of relaxed genes in the Wnt signaling pathway, the downstream target of which, *ESRRB*, is

necessary and sufficient to mediate self-renewal independently of JAK-STAT signaling pathway (Martello et al. 2012), that may suggest evolutionary flexibility in the parallel pathways supporting self-renewal.

The core transcriptional network genes, OSN, orchestrate a cascade of regulatory events involving an autoregulatory loop involving the other pluripotency regulators (Cole et al. 2008; Li & Belmonte 2017). We observed relatively high evolutionary rates with the genes activated by OSN suggesting diversity of subsequent regulating mechanisms among mammals. Importantly, changes in the proximal targets of the master regulators could have profound influence on the downstream circuitry (Erwin & Davidson 2009), our observations of relatively high evolutionary rates with the OSN activated subcircuit may reflect the variations of some characteristics among mammalian PSCs. In addition, understanding mammalian diversity in the downstream targets of the core transcriptional network may help elucidate the mechanisms of reprogramming toward pluripotency and increase the reprogramming efficiency (Takahashi & Yamanaka 2016). Thus, we focused our later analyses on the genes in the OSN activated subcircuit for exploring variations in the mammalian PGRN.

Development of lineage-specific PGRN

Evidence for positive selection on PGRN genes have implications for development of lineage-specific regulatory networks (**Figure 4**). We observed evidence of frequent episodes of positive selection with *TCF3*, implying that alternation of the PGRN have occurred multiple times during mammalian evolution. *TCF3* is the integral component of the core regulatory circuitry of PSCs and encodes a member of the E protein family of helix-loop-helix transcription factors and plays a crucial role by binding to the component of the core transcription factors (Cole et al. 2008; Yi et al. 2008). We found that the common ancestor of Eutheria has been positively selected with fixed lineage-specific amino acid substitutions with *TCF3*, which targets and controls the expression of *ESRRB* (Yi et al. 2008; Martello et al. 2012). *ESRRB* has been proposed to be recruited to the Eutheria PGRN after the divergence of marsupials and Eutheria because neither Tasmanian devil nor platypus iPSCs express *ESRRB* (Weeratunga et al. 2018; Whitworth et al. 2019). Importantly, *ESRRB* not only regulates self-renewal in PSCs (Martello et al. 2012) but also plays a key role in trophoblast, which forms placenta (Luo et al. 1997; Latos et al. 2015). Our findings imply that functional modification in *TCF3* may have caused rewiring Eutherian PGRN to incorporate *ESRRB* that subsequently led to the acquisition of placenta. We also report genetic evidence of positive

selections with *TCF3* and its another target *LEFTY2* (Cole et al. 2008) in Megachiroptera. Positive selection has happened in the ancestral branch of Megachiroptera with *LEFTY2* and the two descendant species, the large flying fox (*Pteropus vampyrus*) and the rousette (*Rousettus aegyptiacus*) in *TCF3*, highlighting the evolutionary history of pre- and post-divergence of these taxa. Possible implications from this observation will be discussed in the next section. *TCF3* also regulates the expression of *NANOG* (Pereira et al. 2006; Cole et al. 2008). The reprogramming efficiency of *NANOG* varies among species; while overexpression of *NANOG* increases the reprogramming rate in felids (Verma et al. 2012, 2013), it does not increase the reprogramming rate in marmosets (Tomioka et al. 2010). We found that both the leopard and the marmoset have positively selected sites in their *TCF3*, but at different positions (**Figure 5E**), which may reflect the variations of response to *NANOG*. This could have implications in development of lineage-specific PGRNs in these taxa.

Potential drivers of mammalian PGRN variation

Our findings of lineage-specific positively selected genes among the PGRN genes provide insight into the influence of species adaptation on the PGRN and the characteristics of mammalian stem cells. The architecture of developmental gene regulatory networks, such as

the PGRN, is composed of diverse components. Certain subcircuits of regulatory networks are not dedicated to a particular biological process, but are also used for diverse functions that might have led to species adaptations. The naked mole-rat (*Heterocephalus glaber*) exhibits extraordinary longevity and cancer resistance (Buffenstein 2008; Delaney et al. 2013). The iPSCs derived from the naked mole-rat have shown to resist tumor formation through the expression of ARF (Miyawaki et al. 2016), in which the Arf/p53 pathway has a protective role from cancer and aging (Matheu et al. 2008). Consistent with this finding, we found *NANOG* and *RIFI* to be positively selected in this species. *RIFI* encodes a protein that regulates DNA replication and damage, interacting with 53BP1 (Kumar & Cheok 2014) which enhances p53 dependent transcriptional responses (Cuella-Martin et al. 2016). *NANOG* is not only one of the core transcriptional network genes, but also exhibits tumorigenic activity through interaction with p53 (Kim et al. 2016; Cuella-Martin et al. 2016). Our finding of positively selected sites on the DNA binding motifs of *NANOG* further suggests development of cancer resistance with p53 pathway in addition to ARF (**Figure 5C**).

Alternatively, the evidence of positive selection on *RIFI* may indicate convergent evolution of species adaptation for perception. We identified that *RIFI* has been positively selected in the naked mole-rat, the Cape golden mole (*Chrysochloris asiatica*), the star-nosed

mole (*Condylura cristata*), and the vampire bat (*Desmodus rotundus*). *RIFI* plays an important role in DNA replication and damage (Kumar & Cheok 2014), and these responses are important for neurogenesis, as represented in human diseases such as Meier-Grinl syndrome and Wolf-Hirschhorn syndrome (Kerzendorfer et al. 2013). Animals that live underground have developed unique sensory systems such as the somatosensory vibrissa-like body hairs on the body of the naked mole-rat (Crish et al. 2003), the middle ear structure of airborne and seismic stimuli in the Cape golden mole (Willi et al. 2006), and the "star" of the star-nosed mole (Gould et al. 1993). Among Microchiroptera, a group that uses echolocation, the vampire bat has extremely sensitive neurons that detect not only echolocation signals but also low frequency sounds, presumably for foraging prey in the dark (Schmidt et al. 1991). Our results provide insights into additional possible development of cancer resistance in mammals.

Differences in sites under positive selection in the protein functional domains may reflect species unique adaptation (**Figure 5E**). As discussed previously, two Megachiroptera species have been identified under positive selection with *TCF3*, which is also associated with cancer (Patel et al. 2015) and neuronal differentiation (Kuwahara et al. 2014). We observed multiple positively selected sites on the functional motif basic helix-loop-helix (bHLH) of

TCF3 in the large flying fox (*Pteropus vampyrus*), among the largest species of bat with a wingspan of up to 1.5 meter (Kunz & Jones 2000). Potential mechanisms of reducing cancer risk through response to DNA damage has been reported in elephants (Abegglen et al. 2015; Vazquez et al. 2018). Interestingly, we also observed positively selected sites on the bHLH of *TCF3* in elephants, implying the resolution to Peto's paradox and the underlying convergent evolution between species that have developed a larger body size compared to their phylogenetic relatives. We identified positively selected sites on the transactivation domain, 9aaTAD in the Rousette bat (*Rousettus aegyptiacus*), which has been reported to exhibit enhanced infection tolerance (Pavlovich et al. 2018). Whereas the previous study revealed genetic signatures of unique signaling in NK cell receptors (Pavlovich et al. 2018), E protein encoded by *TCF3* plays a critical role in B and T lymphocyte development (Engel et al. 2001; Seet et al. 2004), suggesting multiple strategies for antiviral defense in the rousette.

The roles of transposable elements in genomic rearrangement, gene regulation, and epigenetics have been extensively studied to understand primate evolution (Lee et al. 2016). Our findings of the relatively frequent positive selection with epigenetic regulators among Primates may reflect the various impacts of transposable elements on the primate genome. Among Primates, we observed that two epigenetic regulators were under positive selection in

the gibbon (*Nomascus leucogenys*), whose chromatin interactions and epigenetic landscape has been remarkably conserved in spite of extensive genomic shuffling (Lazar et al. 2018). Interestingly, we also observed an event of positive selection at the common ancestor of Primates and the flying lemur with *LEFTY2*, with which DNA methylation plays a critical role during early embryogenesis in vertebrates (Wang et al. 2017), although the results need to be treated with caution.

Due to the habitat transition from terrestrial to aquatic environment, cetaceans have achieved a remarkable changes in their morphology (Uhen 2010). The minke whale (*Balaenoptera acutorostrata*) genome has provided support for genetic changes in *HOX* genes in this species (Yim et al. 2014), which have an important role in the body plan and embryonic development (Pearson et al. 2005). Consequently, we observed that the minke whale has been under positive selection with *KAT6A*, which regulates the expression of *HOX* gene (Voss et al. 2009), implying the morphological adaptation of the whale to the aquatic environment. However, our analyses did not indicate significant evidence of positive selection with *KAT6A* in other cetacean species. Because *KAT6A* is also associated with senescence and tumor growth (Baell et al. 2018), our findings with the minke whale may imply the adaptation of longevity and resistance to age related diseases as illustrated with the genome and

transcriptomes of the bowhead whale (Keane et al. 2015).

Overall, our data indicated that the PGRN genes positively selected in some species are also involved in their unique adaptations, which may subsequently alter their regulatory function in PSCs. Further efforts, such as in vitro genetic modification and characteristics observation, are necessary to test the functional consequences of the genetic mutations discovered in this study.

Conclusions

Our analyses illustrate the evolutionary patterns in the pluripotency gene regulatory network, PGRN, underlying the similarities and variations in characteristics among mammalian pluripotent stem cells, PSCs. This study is one of the first to compare the PGRN genes across major taxa. We showed the evolutionary conservation profiles of the mammalian PGRN and uncovered the evolutionary variable PGRN subcircuits. We identified phylogenetic inference of positive selection of genes involved in the PGRN which has enabled insights into development of lineage-specific PGRN and linkage between PGRN genes and species adaptation. These genes and the associated subcircuits will be plausible targets for future investigations exploring mammalian PSCs.

Authors' contributions

Y. E. designed the original concept and scientific objectives of the project; performed data collection and data analyses; and wrote the manuscript. M. I-M. and K. K. coordinated and participated in the design of the study, and helped draft the manuscript. All authors gave final approval for publication.

Supplementary Material

Additional results supporting this article have been uploaded as part of the online electronic supplementary material.

Acknowledgements

The authors thank Rob Ogden and Rebecca N. Johnson for constructive feedback on manuscript and English correction; Kristin Haverkamp for valuable editorial input and English correction. This work was supported by JSPS KAKENHI Grant Numbers 17H03624, 20H00420 (M. I-M.) and 17H02083 (K.K.), and Kyoto University Supporting Program for Interaction-based Initiative Team Studies (SPIRITS) to M. I-M.

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Figure Legends

Fig. 1.—Flow diagram detailing the strategy to investigate the evolution of the pluripotency regulating gene network (PGRN) across mammals. Boxes represent steps of methods in processing order. Key approaches used for each step are indicated in parenthesis.

Fig. 2.—Trends in the stringency of purifying selection on the pluripotency regulating gene network (PGRN) genes. Schematic representation of the PGRN according to KEGG is shown. Only genes with significant shifts in the stringency of purifying selection at 10% FDR or less are indicated: *Green* rectangle, intensification; *lime* circle, relaxation. Dashed lines indicate the subcircuits of the PGRN. Horizontal double line indicates the cellular envelop. Blue oval represents the nucleus. The genes under each subcircuit are simplified, see **supplementary table S4** for details.

Fig. 3.—Conservation patterns of the pluripotency regulating gene network (PGRN). (A) Histogram of gene counts for dN/dS values showing distribution of evolutionary rates for the PGRN genes. (B) Box plots of the mean dN/dS values for the PGRN subcircuits. Multiple paired comparisons were tested with the Tukey-Kramer method, where subcircuits not

assigned by the same letter are significantly different with P value of 0.05; error bars, 95% confidence interval. OSN represents *OCT4* (*POU5F1*), *SOX2*, and *NANOG*.

Fig. 4.—Mapping of positively selected genes identified in ancestral branches and contemporary species of mammals. Positively selected genes at 10% FDR or less (*red*) are indicated on a commonly accepted phylogeny of mammals. Ancestral branches with which positively selected genes were identified are shown in *red*. Colored boxes represents major taxonomic groups referred to the NCBI taxonomy.

Fig. 5.—Positively selected sites of the OSN activated genes: (A) *KAT6A*, (B) *LEFTY2*, (C) *NANOG*, (D) *SETDB1*, and (E) *TCF3*. Positively selected lineages and their positively selected sites with 95% posterior probability are mapped on the coded protein with the arrangements of functionally important regions. Only genes with significant positively selected sites in at least one lineage and on at least one functional region are shown. The functional region represented with bar is a region of interest that cannot be described in other subsections according to UniPort. Colored boxes represent major taxonomic groups (Afrotheria (*orange*), Carnivora (*red*), Cetartiodactyla (*fuchsia*), Chiroptera (*yellow*),

Primates/flying lemur (*lime*), Rodentia (*aqua*), and other (*grey*)).

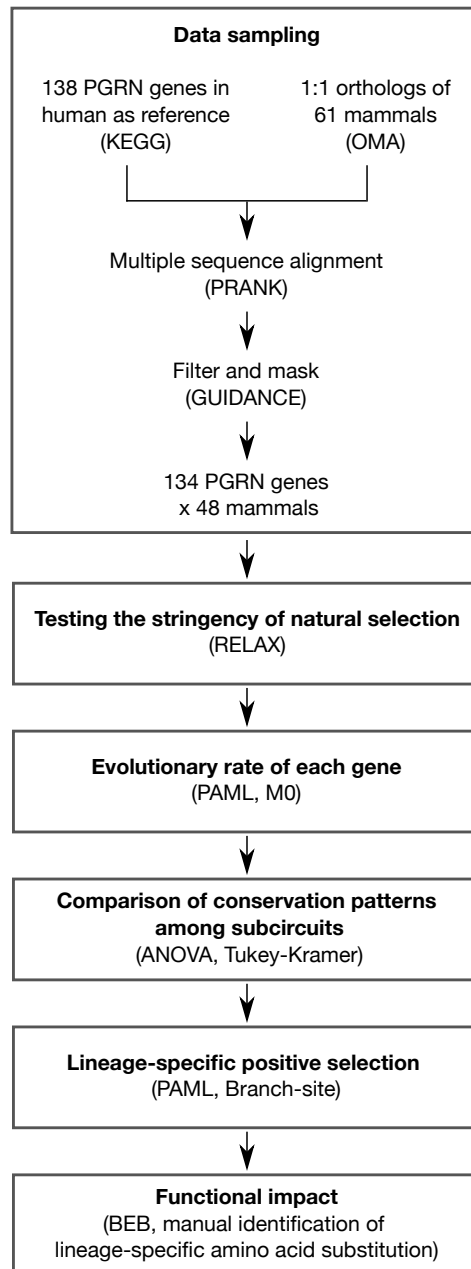
Fig. 6.—Lineage-specific amino acid substitutions in positively selected genes at ancestral branch: (A) *KAT6A*, (B) *LEFTY2*, (C) *RIF1*, and (D) *TCF3*. Positively selected lineages and their specific amino acid substitutions are mapped on the coded protein with the arrangements of functionally important regions. The functional region represented with bar is a region of interest that cannot be described in other subsections according to UniPort. Colored boxes represent major taxonomic groups (Eutheria (*purple*), Chiroptera (*yellow*), and Primates/flying lemur (*lime*)).

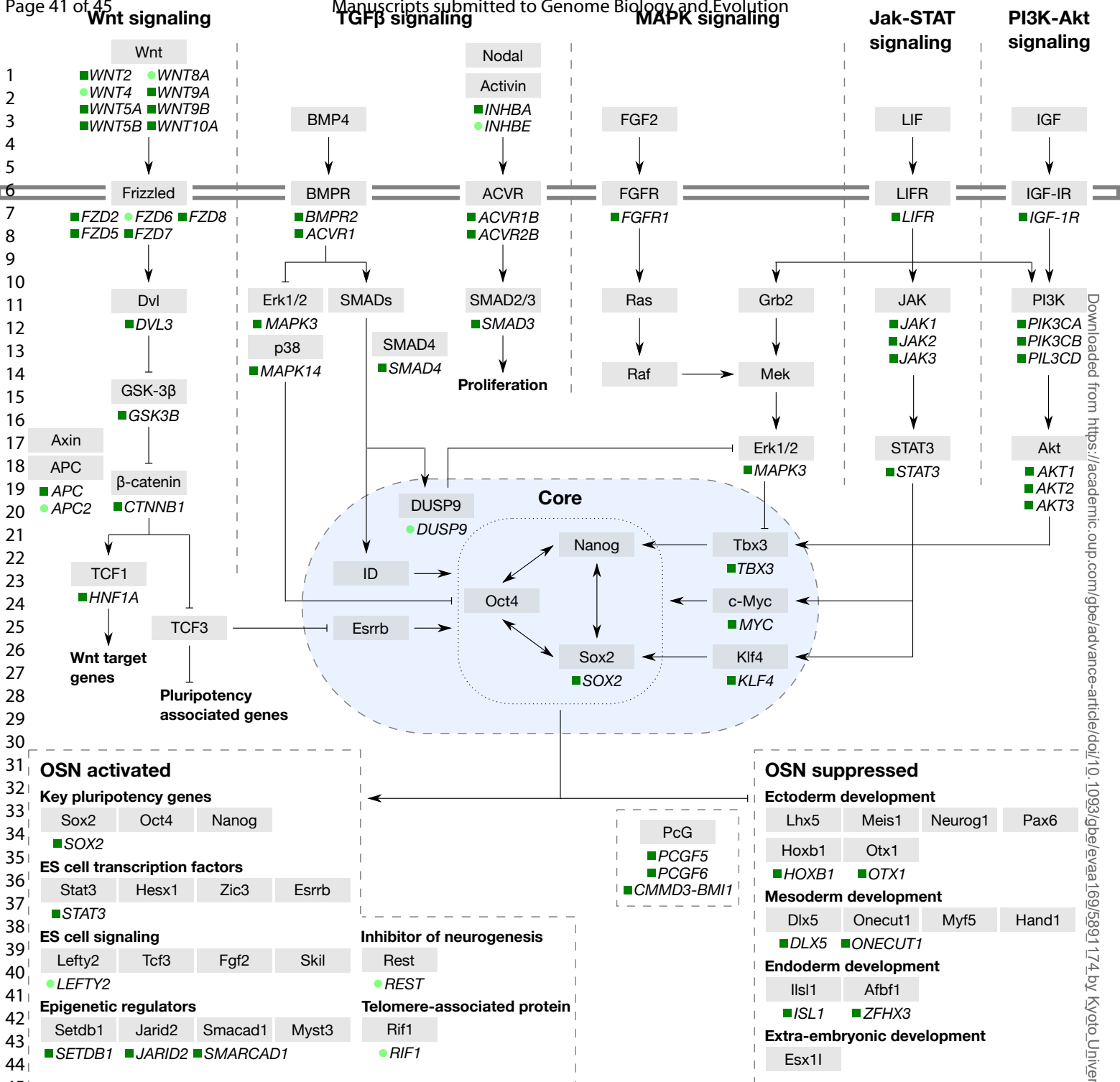
Table 1

Comparison of selective pressure among the subcircuits

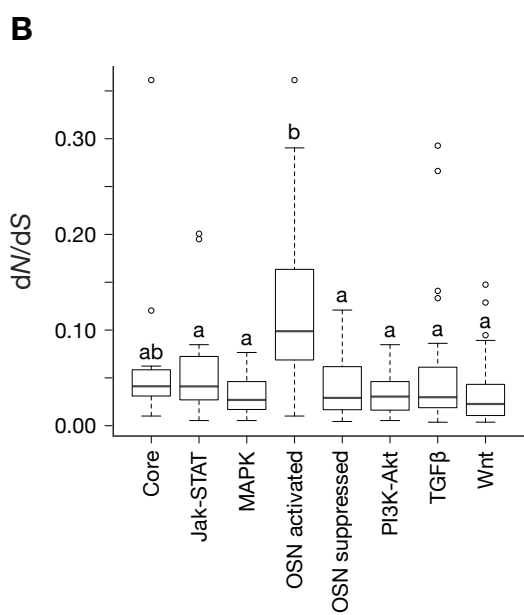
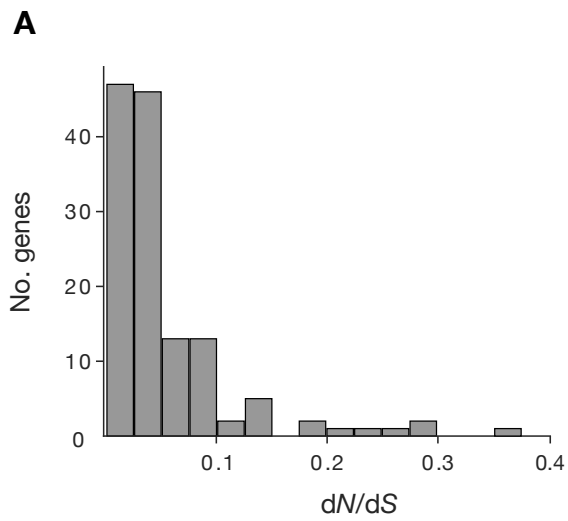
Subcircuits	No. gene	No. gene with significant shifts in strength of purifying selection.	
		Intensified	Relaxed
Core	11	4 (36.4%)	1 (9%)
JAK-STAT	17	9 (52.9%)	0
MAPK	26	8 (30.8%)	1 (3.8%)
OSN ^a activated	15	4 (26.7%)	3 (20%)
OSN ^a suppressed	18	9 (50%)	0
PI3K-Akt	29	13 (44.8%)	0
TGFβ	27	9 (33.3%)	2 (7.4%)
Wnt	40	16 (40%)	4 (10%)

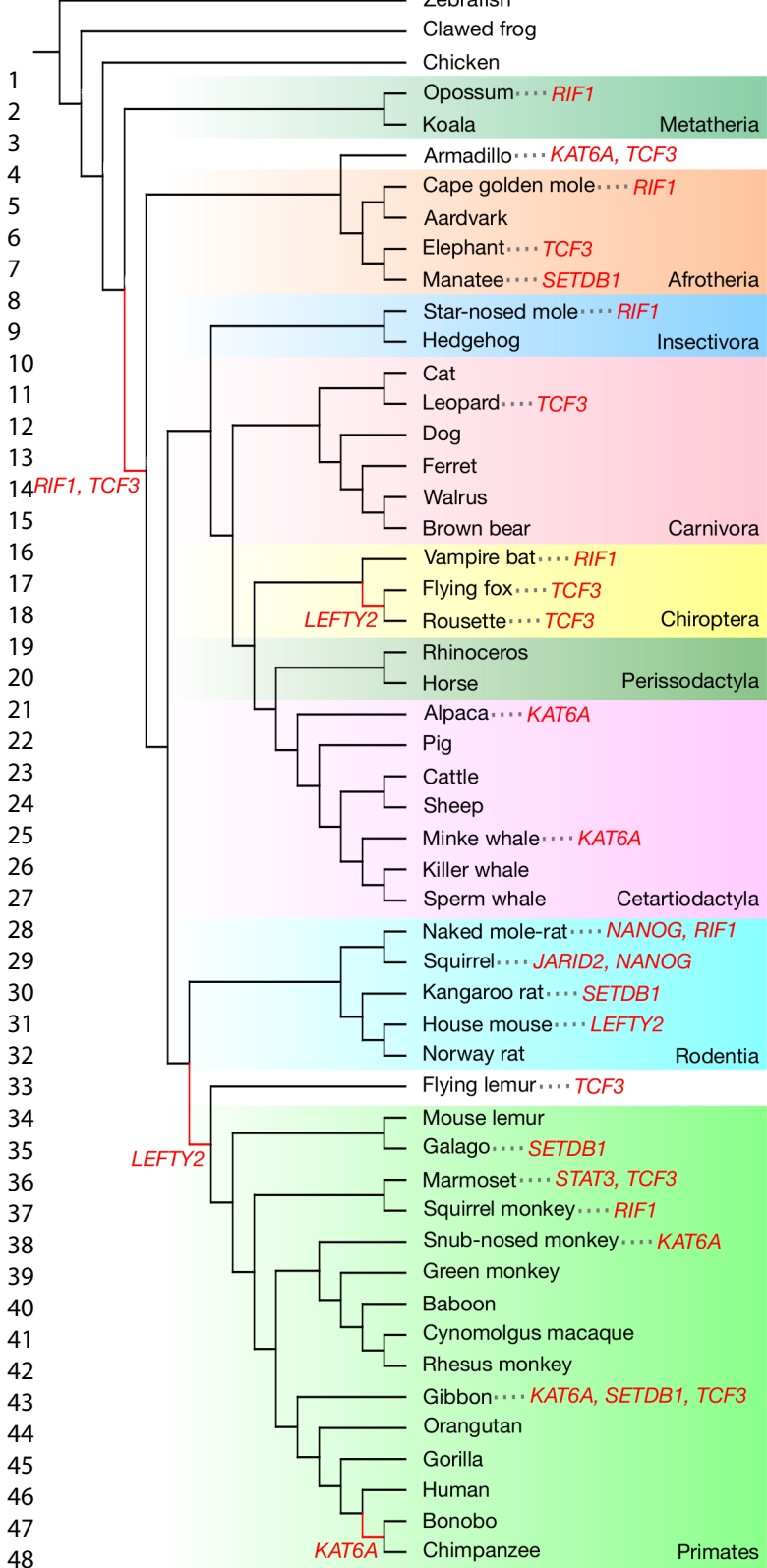
Notes.—^aOCT4 (*POU5F1*), SOX2, and NANOG.





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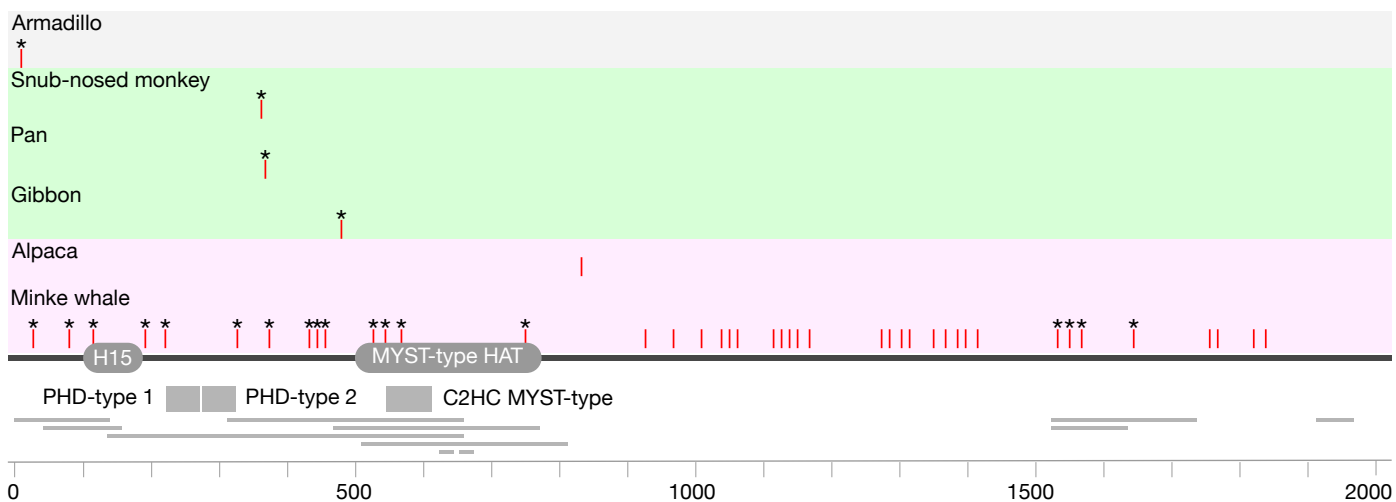




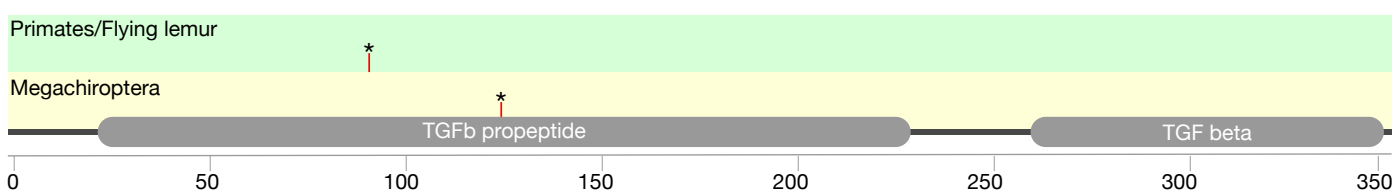
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| Positively selected site ● Domain and repeat ⬡ Motif
* Positively selected site on functional region ■ Zinc finger — Region

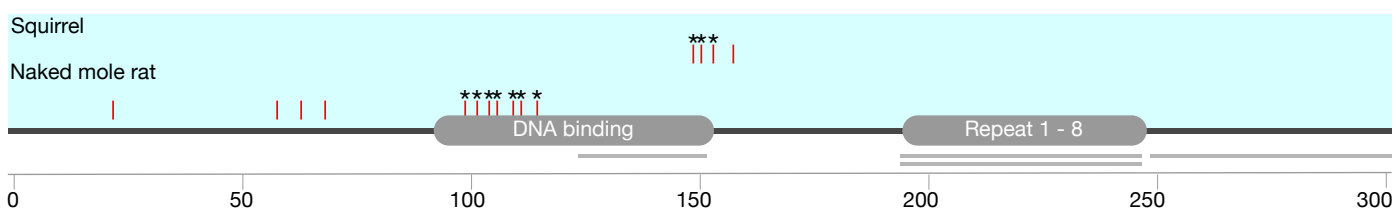
A KAT6A



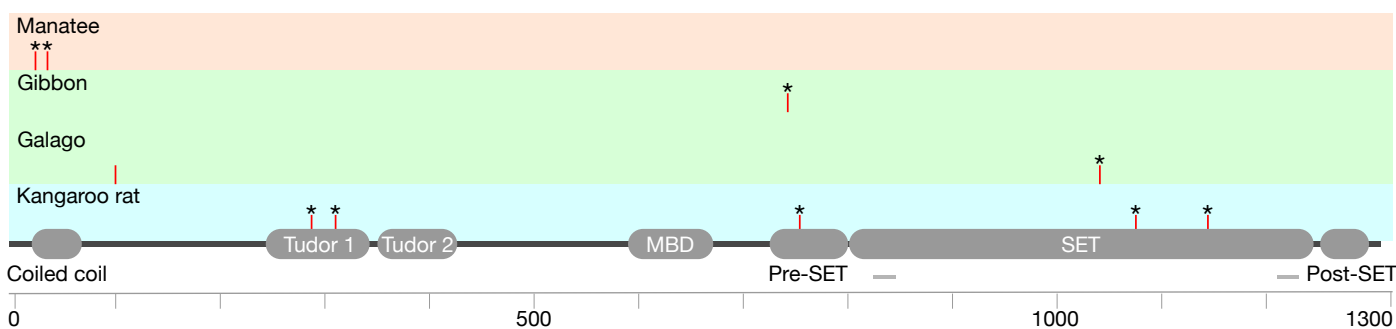
B LEFTY2



C NANOG



D SETDB1



E TCF3

