# The genome of North American beaver provides insights into the mechanisms of its longevity and cancer resistance 

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#### Abstract

The North American beaver (Castor canadensis) is an exceptionally long-lived and cancer-resistant rodent species, and thus an excellent model organism for comparative genomic studies of longevity. Here, we utilize a significantly improved beaver genome assembly to assess evolutionary changes in gene coding sequences, copy number, and expression. We found that the beaver Aldh1a1, a stem cell marker gene encoding an enzyme required for detoxification of ethanol and aldehydes, is expanded ( $\sim 10$ copies vs. two in mouse and one in human). We also show that the beaver cells are more resistant to ethanol, and beaver liver extracts show higher ability to metabolize aldehydes than the mouse samples. Furthermore, Hpgd, a tumor suppressor gene, is uniquely duplicated in the beaver among rodents. Our evolutionary analysis identified beaver genes under positive selection which are associated with tumor suppression and longevity. Genes involved in lipid metabolism show positive selection signals, changes in copy number and altered gene expression in beavers. Several genes involved in DNA repair showed a higher expression in beavers which is consistent with the trend observed in other long-lived mammals. In summary, we identified several genes that likely contribute to beaver longevity and cancer resistance, including increased ability to detoxify aldehydes, enhanced tumor suppression and DNA repair, and altered lipid metabolism.


## Introduction

Rodent species show considerable variation in their maximum lifespans. Recent studies of several long-lived rodents have provided new insights into the mechanisms of longevity. For example, studies of the naked mole rats found unique amino acid changes in proteins involved in DNA repair and cell cycle (Kim, Fang et al. 2011), insulin $\beta$-chain associated with insulin misfolding and diabetes (Fang, Seim et al. 2014), decreased expression of genes in insulin/lgf1 signaling in liver (Kim, Fang et al. 2011), and increased expression of genes involved in DNA repair signaling (MacRae, Croken et al. 2015). In blind mole rats, Tp53 protein was found with an amino acid change associated with human tumors. This mutation leads to partial inactivation of Tp53 function and promotes p53 binding to promoters of cell cycle arrest genes rather than apoptotic genes (Ashur-Fabian, Avivi et al. 2004).

North American beaver (Castor Canadensis) has a maximum lifespan over 24 years and, with adults weighing from 24 to 71 lbs , is the second largest rodent species. It also shows resistance to cancer, despite a large body size and a long lifespan. Studies of several animals with large body sizes, long lifespans, and yet low rates of cancer revealed their enhanced antitumor mechanisms. For example, elephant, the biggest land mammal, was found have enhanced Tp53 activity by Tp53 expansion ( $\sim 20$ copies) (Abegglen, Caulin et al. 2015), while bowhead whale, the possibly longest-
lived mammal, shows signals of natural selection in aging and cancer-associated genes (Keane, Semeiks et al. 2015). How beaver evolved to have cancer resistance and longevity is not clear.

The beaver genome assembly was first released in 2017 (Lok, Paton et al. 2017). Before analyzing the beaver genome, we re-sequenced and re-assembled the beaver genome to improve the assembly quality in terms of both the contig length and the assembly completeness (Zhou, Dou et al. 2020). To find potentially novel mechanisms of cancer resistance and longevity, using our improved beaver genome assembly, we explored three major types of signals of natural selection in beaver genome compared to other rodents: changes in gene copy numbers, positive selection that favored changes in amino acid residues, which further affect gene structure, interaction, and function, and changes in gene expression. Through a systematic, comparative analysis, we identified a striking multiplication of Aldh1a1, a stem cell marker gene, by around 10 copies in the beaver genome and beaver-specific expansion of several genes, including Hpgd, a tumor suppressor gene, and Cyp19a1, a gene involved in estrogen biosynthesis and linked to human longevity (Corbo, Ulizzi et al. 2011). We also found several genes associated with lipid metabolism, oxidation reduction, cancer suppression under positive selection and increased expression of DNA repair genes in beaver compared to mouse, consistent with observations in other long-lived species. In addition, we discovered that genes associated with lipid metabolism were likely under natural selection through changes in coding sequences, gene copy numbers, and gene expression levels.

## Results

## Genome annotation and phylogeny

Our beaver genome assembly (Zhou, Dou et al. 2020) consists of 739,342 scaffolds, with scaffold $\mathrm{N} 50=24.31 \mathrm{Mb}$ (the maximum was 93.06 Mb ). This beaver genome assembly has a much higher completeness, with 95\% complete BUSCOs (Benchmarking Universal Single-Copy Orthologs) (Simao, Waterhouse et al. 2015), compared to $83.1 \%$ of the beaver genome published previously (scaffold $\mathrm{N} 50=318 \mathrm{~kb}$ and the maximum was 4.2 Mb ) (Lok, Paton et al. 2017).

For gene annotation and downstream analysis, we selected scaffolds longer than 300 bp . There were 250,435 such scaffolds, and together they count for $96.6 \%$ of the assembled sequences. Using the Maker2 pipeline (Holt and Yandell 2011), we predicted 26,515 beaver genes with support from either transcriptomes or protein sequences (Figure 1A). Among these genes, 20,670 (78\%) were found with functional domains by InterProScan (v5.25) (Jones, Binns et al. 2014). A genome is considered well annotated if more than $90 \%$ genes have an annotation edit distance (AED) score, which measures the goodness of fit of each gene to the evidence supporting it, lower than 0.5 , and over $50 \%$ proteins contain a recognizable domain (Campbell, Holt et al. 2014). In our beaver
genome annotation, $\sim 90.7 \%$ gene models have $A E D$ scores lower than 0.5 (Figure 1 Supplement Figure S1) and 78\% gene products contain domains.

A total of 14 rodent species, 2 rabbit species, and human and chimpanzee (as outgroup species), are included in our analysis. With 5,087 single copy genes across them, we generated their phylogeny and estimated that beaver and its evolutionally closest species in our set, Ord's Kangroo rat, separated about 46 million years ago (Figure 1B).

## Significant expansion of beaver Aldh1a1 and its functional consequences

Using CAFE 3 (Han, Thomas et al. 2013), we identified significant (FDR<0.01) copy number increases of eight beaver genes (Figure 2 - Supplement Figure S2, see Methods for details). Most of these increases are not specific to beavers. One of them, the expansion of Aldh1a1, however, is striking in beaver, suggesting this expansion could be important for its cancer resistance and/or longevity. Therefore, we examined this gene in detail.

Expansion of beaver Aldh1a1 Compared with humans and chimpanzees, which have one copy of Aldh1a1, most rodent species have two copies of the gene (its close paralog in mouse is named as Aldh1a7). This indicates the duplication likely happened before the last common ancestor of rodents (Figure 2 - Supplement Figure S3). In contrast, 10 copies of Aldh1a1 were predicted in the beaver genome, among which 2 copies are predicted as pseudogenes because of frameshift indels (Figure 2 - Supplement Figure S4A, S4B). One copy (ID: 266.2) has relatively lower annotation quality with two repeat fragments, which may indicate a prediction with a mixture from two copies of Aldh1a1.

We next examined every predicted copy of beaver Aldh1a1 and carried out qPCR to experimentally quantify duplication events. We compared synteny of the Aldh1a1 locus among beaver, human, and mouse (Figure 2A). Missing or unordered genes in a genomic region may indicate a poor assembly quality. A perfect match of the order of corresponding genes indicates good quality of genome assembly at this locus. After duplication, different gene copies evolved independently and accumulated differences in both sequence and gene structure. Using Apollo (Lee, Helt et al. 2013) and RNA-seq data, we manually improved the gene annotation of seven functional copies of beaver Aldh1a1 (excluding two pseudogenes and the copy 266.2), which all have different gene structures (Figure 2B). Bona fide gene copies likely encode homologous proteins with different substitutions. We picked three sites in the coding sequences of exon 4 (Figure 3C), which can distinguish the seven copies from one another (Figure 2C - Supplement Figure S4C). The alignment of RNAseq reads from beaver individuals indicates that those variable sites are genuine, and the different copies were transcribed simultaneously. While the RNA-seq read coverage at these variable sites was relatively low for the copy 266.5 , we also analyzed another site, where the copy 266.5 is distinct
from all other copies and with much higher RNA-seq reads coverage (Figure 2 - Supplement Figure S4D). To further validate the duplication of Aldh1a1, we performed qPCR analysis, which showed that there are around 10 copies of Aldh1a1 in the beaver genome (Figure 2D).

Duplicated Aldh1a1 genes are transcribed and result in higher Aldh1a1 protein levels All the predicted beaver Aldh1a1 genes have regular gene structures with introns, and only two of them were identified as pseudogenes by GeneWise (Birney, Clamp et al. 2004) (see Methods and Figure 2 - Supplement Figure S5). At least seven copies of Aldh1a1 are of high annotation quality and transcribed in parallel in beaver individuals (Figure 2C). Based on their expression patterns in different tissues, beaver Aldh1a1 genes can be clustered into three groups (Figure 2E). As expected, the two pseudogenes show very low expression in all the tissues. While the highest expression of three copies in liver appears to be liver-specific, the expression of other copies is moderate in both liver and brain. To understand the overall transcriptional activity of Aldh1a1 in beaver, we compared expression levels of Aldh1a1 genes in liver and brain between beaver and mouse using RNA-seq data (see Methods for details). The overall expression of Aldh1a1 was significantly higher in beaver liver ( $\mathrm{FDR}=2.23 \mathrm{E}-28$, fold change $=10.04$ ) and brain $(\mathrm{FDR}=2.24 \mathrm{E}-$ 12 , fold change $=9.54$ ) than that in the same mouse tissue, respectively (Figure 2F). We also examined protein level of Aldh1a1 by Western blotting. Using cytosolic extracts from liver cells of beaver and mouse, we found that there is more Aldh1a1 in beaver liver tissue than mouse liver tissue (Figure 2G-H). Together these results indicate that duplicated Aldh1a1 copies are transcribed and result in higher protein levels in beaver tissue.

Potential positive selection of Aldh1a1 duplication Gene duplications are often under relaxed purifying selection, as increased gene dosage tends to provide burden on the cell. The exceptional level of duplication of Aldh1a1 in beaver, however, suggests the presence of a positive (i.e., adaptive) selection. Aldh1a1 belongs to the Aldh super family: there are 19 and 21 Aldh genes in the human and mouse genomes, respectively. To investigate whether the duplication of Aldh1a1 in beaver is a compensation for loss of other Aldh genes or a result of natural selection, we manually checked all Aldh genes in the beaver genome (Figure 2 -

Supplement Figure S6). Except Aldh1a1, beavers have the same set of Aldh genes as mice and humans with regular gene structures and open reading frames (Figure 2 - Supplement Figure S7). This conservation of other Aldh genes indicates that the significant expansion of beaver Aldh1a1 is likely a result of positive selection during evolution.

Beaver cells show enhanced tolerance of alcohol and aldehydes Aldh gene products can protect organisms against damage from oxidative stress by processing toxic aldehydes generated
as a result of lipid peroxidation (Singh, Brocker et al. 2013). Both alcohol and endogenous aldehydes can lead to DNA damage and increase mutations in stem cells (Garaycoechea, Crossan et al. 2018). We next tested the resistance of beaver cells to ethanol and aldehydes. When treated with $18 \%$ ethanol for 7 hours both beaver and mouse lung fibroblasts showed significantly decreased cell viability (Figure 3A). However, the reduction of cell viability was significantly lower for beaver lung fibroblasts than mouse cells ( $\mathrm{P}=0.002$ ). This result suggests that higher Aldh1a1 levels in beaver cells lead to higher alcohol resistance. We also tested Aldh1a1 activity for three types of endogenous aldehydes: all-trans-retinal (RET), malonaldehyde (MDA) and 4hydroxynonenal (HNE). Beaver and mouse liver extracts did not differ in their ability to process RET. However, beaver extract showed much stronger activity on MDA and HNE (Figure 3B). This result clearly shows that beaver liver possesses higher Aldh1a1 activity than mouse liver.

## Beaver-specific expansion of conserved genes

We identified 18 gene candidates for potential beaver-specific expansion and successfully validated five of them by qPCR (see Methods for details): Hpgd (15-hydroxyprostaglandin dehydrogenase), Fitm1 (Fat storage inducing transmembrane protein 1), Cyp19a1 (Cytochrome P450 family 19 subfamily A member 1), Pla2g4c (Phospholipase A2 Group IVC), and Cenpt (Centromere protein $T$ ) (Figure 4A - Supplement Figure S8-S11). There are two copies of Hpgd, a well-known tumor suppressor gene, in the beaver genome. The Hpgd loci in beaver, mouse, and human genomes show good synteny (Figure 4B), indicating a good assembly quality at this region. The beaver-specific duplication of Hpgd was validated by real-time qPCR (Figure 4A). We aligned beaver and mouse Hpgd sequences together (Figure 4C). While the two copies of beaver Hpgd have a high degree of sequence similarity, there are several amino acid residue differences. RNAseq reads from individual beavers covered variable sites in both Hpgd copies, indicating that both copies are transcribed (and likely functional) and sequence differences were not due to DNA sequencing errors but instead divergent evolution after the duplication (Figure 4D). Hpgd is a tumor suppressor of many cancer types, including cancer of liver (Lu, Han et al. 2014), colon (Myung, Rerko et al. 2006), lung (Ding, Tong et al. 2005), and breast (Wu, Liu et al. 2017). Our RNA-seq data analysis showed although Hpgd was not differentially expressed in brain between beavers and mice, it was expressed significantly higher in beaver liver (fold change $=4.39$, FDR $=4.56 \mathrm{E}-$ 40), where Hpgd is more transcriptionally active (Figure 4E). The duplication of Hpgd and its higher expression likely contribute to the cancer resistance of beavers.

## Beaver genes under positive selection are associated with tumor suppression and longevity

We identified 21 beaver genes putatively under positive selection (FDR < 0.01, Table 1, Supplementary file 1), using 'branch site' model (Zhang, Nielsen et al. 2005) conducted in

PosiGene (Sahm, Bens et al. 2017) pipeline followed by manual curation (see Methods for details). Although not enriched at the pathway level, several top genes under positive selection are known to be associated with lipid metabolism (e.g., Erlin2, Fabp3 and Cilp2), tumor suppression (e.g., Vwa5a and Fabp3), and oxidation reduction process (e.g., Hsd17b1, Fabp3, Cox15, Cyb5a, and Aoc1). Especially, several genes may be associated with aging/longevity. Different alleles of Hsd17b1 were found significantly associated with human longevity in females (Scarabino, Scacchi et al. 2015). A single knockout of Mtbp in mice led to an extension of lifespan (Grieb, Boyd et al. 2016). Knockdown of Mrpl37 increased lifespan of C. elegans by $41 \%$ on average (Houtkooper, Mouchiroud et al. 2013). Long-lived bats have greater abundance of Fabp3 in muscle mitochondria than that of short-lived mice, and thus its regulation of lipid may influence mitochondrial function (Pollard, Ingram et al. 2019). FABP3 also acts as a tumor suppressor in human embryonic cancer cells and breast cancer (Song, Shen et al. 2012). In addition, Ptx3 is an inflammatory protein, which protects the organism against pathogens and controls autoimmunity, and is involved in tissue remodeling and cancer development (Doni, Stravalaci et al. 2019).

Mtbp (Mdm2 Binding Protein) interacts with oncoprotein mouse double minute 2 (Mdm2) and enhances its stability, which promotes degradation of Tp53 (Figure 5 - Supplement Figure S12). Sites with positive selection signals were identified in exons encoding the "mid-domain" and the "Cdomain" of the Mtbp protein (Figure 5A). In addition to codon changes (i.e., selection signals identified by PosiGene (Sahm, Bens et al. 2017)), there are also indels in beaver Mtbp, compared with the ortholog from other rodent species. Especially, there is a 15-bp insertion in exon 18 of beaver Mtbp, which resembles an insertion at a similar location in Mtbp of the long-lived naked mole rat. We explore potential functional effects of changes in codons and indels using PROVEAN (Choi and Chan 2015), PolyPehn2 (Adzhubei, Jordan et al. 2013) and CADD (Rentzsch, Witten et al. 2019) by considering changes from those in human genome (human sequences are the consensus sequence at those loci) to those in beaver genome (Figure 5B). Two sites under positive selection are consistently predicted to be deleterious, suggesting they could result in decreased Mtbp function and hence decrease the stability of Mdm2, leading to higher Tp53 activity. We also checked the top three sites under positive selection identified in the multiple sequence alignment of Mtbp from 62 mammals (Figure 5 - Supplement Figure S12B-D). We checked a few other sequence changes at the positive selection sites in other mammals and found only changes in beaver Mtbp were predicted as deleterious (Figure 5 - Supplement Figure S12D), which may contribute to its cancer resistance. Haploinsufficiency of Mtbp in mice delays spontaneous cancer development and extend lifespan by enhancing Tp53 function (Grieb, Boyd et al. 2016).

## Beaver displays enhanced expression of DNA repair genes and changes in genes involved in lipid metabolism

Among 12,090 genes with one-to-one orthology between beaver and mouse, we detected 2,892 (1,652 up-regulated and 1,240 down-regulated) and 2,765 (1,534 up-regulated and 1,231 downregulated) differentially expressed genes (fold change $>2.5$ and FDR $<0.001$ ) in liver and brain, respectively, between beavers and mice (see Methods for details). Among REACTOM pathways, both up-regulated and down-regulated differentially expressed genes were enriched in lipid metabolism pathway in both liver and brain (Figure 6). Such result indicates significantly divergent evolution of lipid metabolism between beaver and mouse. Up-regulated genes in both livers and brains of beavers were also enriched in hemostasis, DNA repair, and cell cycle pathways. Especially, "nucleotide excision repair" and "base excision repair" pathways are enriched with upregulated genes in beaver brains. Up-regulated genes in beaver livers are enriched in the Tp53 signaling pathway among 50 hallmark gene sets from the Molecular Signatures Database (Subramanian, Tamayo et al. 2005), which represent well-defined biological states or processes with coherent expression. Linking to human longevity and age-related diseases (Wolfson, Budovsky et al. 2009), overexpression of genes in "focal adhesion" (Figure 6A, 6B) may also contribute to beaver's longevity.

Among differentially expressed genes, lgf2 (insulin-like growth factor 2) showed much higher expression in beaver liver than in mouse liver (FDR $=3.06 \mathrm{E}-135$, top 20th up-regulated gene in beaver liver), and its binding protein Igf2bp2 also shows significant up-regulation in beaver liver (FDR = 7.69E-07) (Figure 6 - Supplement Figure S13). Expression of Igf2 significantly decreases after birth in liver for most mammals, including mouse and rat. Interestingly, Igf2 and Igf2bp2 show relatively higher expression levels in livers of naked mole rats and Damaraland mole rats (Fang, Seim et al. 2014, Ma and Gladyshev 2017), both long-lived rodents.

## Discussion

Aldh gene products process aldehydes generated as a result of lipid peroxidation, thus protecting the organism from the consequence of oxidative stress (Singh, Brocker et al. 2013). A recent study showed that both alcohol and endogenous aldehydes damage chromosomes and increase the mutation rate of stem cells (Garaycoechea, Crossan et al. 2018). We found that beaver cells exhibit better tolerance to ethanol than mouse cells and show strikingly enhanced capabilities for metabolizing malonaldehyde (MDA) and 4-hydroxynonenal (HNE) (Figure 3). While MDA is the most mutagenic aldehyde product of lipid peroxidation, HNE is the most toxic (Ayala, Munoz et al. 2014). The Aldh upregulation occurs in mammals in response to lipid peroxidation (Vassalli 2019), which is a major source of endogenous aldehydes. Polyunsaturated fatty acids (PUFAs) are more susceptible to oxidation than monounsaturated fatty acids, and beavers have a high proportion of

PUFAs, which is unusual among mammals (Martysiak-Zurowska, Zalewski et al. 2009, Zalewski, Martysiak-Zurowska et al. 2009, Domaradzki, Florek et al. 2019). For example, beaver's tail fat contains over 80\% unsaturated fatty acids (Zalewski, Martysiak-Zurowska et al. 2009). Studies of diving mammals indicate potential function of PUFAs in oxygen conservation, through reducing heart rate, to enhance their diving ability (Trumble and Kanatous 2012). Consistently, the heart rate of beavers is 100 beats/minute during rest and 50 beats/minute while diving (Muller-Schwarze 2003), which is much lower than that ( $\sim 250$ beats/mins) of other rodents (Carpenter and Marion 2013). However, high level of PUFAs will result in increased lipid peroxidation, which is reversely correlated with lifespan of diverse species, including mammals (Hulbert, Kelly et al. 2014). Longevity of beavers indicates the presence of a potential protection mechanism against lipid peroxidation. Compared to Aldh2 and Aldh3a1, Aldh1a1 is the most important enzyme to oxidize aldehydes formed by lipid peroxidation in murine hepatocytes (Makia, Bojang et al. 2011). The higher susceptibility to lipid peroxidation and the associated oxidative stress in beavers may exert selective pressure for increased expression of Aldh1a1. Gene duplication is one way to increase gene dosage rapidly. From this point of view, Aldh1a1 duplication in beaver is very likely a result of natural selection, which significantly improves its tolerance against oxidative stress, and so contributes to its longevity. In addition to gene duplication, our data also showed different expression patterns of the copies across tissues (Figure 2E), which warrants future studies to explore whether diverse selections have shaped regulation of Aldh1a1 copies. While no difference in Aldh1a1 activity on RET was observed between beavers and mice, Aldh1a1 in beavers showed much higher activities on MDA and HNE than that in mice.

Increased Aldh1 activity is expected to increase organism's resistance to oxidative stress. Enhanced tolerance for the oxidative stress has been found in long-lived fruit flies comparing those with normal lifespan (Deepashree, Niveditha et al. 2019). In human, centenarians have been found with less oxidative stress damages comparing to controls (Belenguer-Varea, TarazonaSantabalbina et al. 2019). Linked to multiple age-related diseases, the oxidative stress is likely one of major contributors to aging (Liguori, Russo et al. 2018). Lipoxidation increases with age (Mitchell, Buffenstein et al. 2007), while reactive aldehyde, a known carcinogen, interferes with DNA replication, causes DNA damage, and induces formation of DNA adducts (Langevin, Crossan et al. 2011). Aldh1a1 duplication can increase cellular protection against these toxins. Our finding of better toleration of ethanol by beaver cells is consistent with a previous human study, which showed low Aldh1a1 activity might account for alcohol sensitivity in some Caucasian populations (Marchitti, Brocker et al. 2008). Stem cell exhaustion is one of hallmarks of the aging process. Aldh1a1 is also a marker gene associated with stemness of cells and expressed higher in stem cells (Li, Condello et al. 2017). Aldh1a1 duplication may also contribute to stem cell maintenance in beavers. This hypothesis is consistent with results from other studies, which found unsaturated fatty acids can
maintain cancer cell stemness (Mukherjee, Kenny et al. 2017) and positive association between Aldh1a1 expression and lipid unsaturation level (Li, Condello et al. 2017). Thus, Aldh1a1 duplication may affect beavers' aging process by increasing both resilience to oxidative stress and the stemness of beaver cells. Mice deficient in both Aldh1a1 and Aldh3a1 had fewer hematopoietic stem cells, more reactive oxygen species, and increased sensitivity to DNA damage (Gasparetto, Sekulovic et al. 2012), all of which are early aging phenotypes.

Several studies have shown the association of Aldh1a1 expression with the aging process. The expression of Aldh1a1 dramatically increases with age in mouse hematopoietic stem cells (Levi, Yilmaz et al. 2009). It is significantly lower in CD4+ T cells from aged mice than those from young mice, and its low expression may affect migration of T cells in aged mice (Park, Miyakawa et al. 2014). However, how and in which tissues the expression of Aldh1a1 is significantly changed with age is not clear. To explore this, we used human gene expression data in diverse tissues from GTEx (Carithers, Ardlie et al. 2015) and studied Aldh1a1 expression pattern during aging in the presence of several potential covariates (see Methods). In several tissues, the expression of Aldh1a1 significantly changes with age (FDR < 0.01) (Figure 7). For example, it significantly increases with age in adipose tissue $(r=0.17, \mathrm{FDR}=0.0003$ ), which may be associated with an increase in the percentage of body fat among the old (St-Onge and Gallagher 2010). It also significantly decreases with age in several human brain regions. This may be associated with an increase in some brain disorders among the old, as patients with Parkinson's disease tend to have a decreased expression of Aldh1a1 in corresponding brain regions (Galter, Buervenich et al. 2003). It is believed that ALDHs play a significant role in neuroprotection (Marchitti, Deitrich et al. 2007). For example, ALDH1A1 has been found as a marker of astrocytic differentiation during brain development (Adam, Schnell et al. 2012). Thus, Aldh1a1 duplication may also protect beaver from age-related brain impairment.

Compared to other mammals, the adipose tissue and muscles of beavers show different lipid composition, with the highest proportion of PUFAs (Martysiak-Zurowska, Zalewski et al. 2009, Zalewski, Martysiak-Zurowska et al. 2009, Domaradzki, Florek et al. 2019), which may indicate an unusual way of lipid metabolism and anti-lipoxidation in beavers. Consistent with the unusual lipid composition in beavers, we found genes differentially expressed between mice and beavers are most enriched in lipid metabolism (Figure 6). Besides, several genes expanded in the beaver genome, including Aldh1a1, Hpgd, Fitm1, Cyp19a1, and Pla2g4c, are also associated with lipid metabolism. Aldh1a1 knockout mice show decreased accumulation of both subcutaneous and visceral fat pads (Ziouzenkova, Orasanu et al. 2007), indicating that Aldh1a1 duplication may contribute to accumulation of protective layers of fat which beavers evolved to provide insulation in cold water. Furthermore, Fabp3, one of the top positively selected genes in beaver, has also been
found to be associated with intramuscular fat deposition and body weight in chicken (Ye, Chen et al. 2010) and pigs (Gerbens, van Erp et al. 1999). In mice, Fabp3 is essential for fatty acid oxidation in brown adipose tissue and plays a central role in cold tolerance (Vergnes, Chin et al. 2011).

Among genes duplicated specifically in beavers, Hpgd is highly expressed in regulatory T cells, which prevents autoimmunity and maintains adipose tissue homeostasis (Schmidleithner, Thabet et al. 2019). Fitm1 plays an important role in lipid droplet accumulation. Cyp19a1 is involved in the synthesis of cholesterol, steroids, and other lipids. It also catalyzes estrogen biosynthesis, which enhances healthy aging and human longevity (Horstman, Dillon et al. 2012). A significant association between Cyp19a1 polymorphisms and longevity was observed in humans (Corbo, Ulizzi et al. 2011). As a member of the phospholipase A2 enzyme family, Pla2g4c plays a role in hydrolyzing glycerol-phospholipids to produce lysophospholipids and free fatty acids. Although not enriched on the pathway level, among the top 10 putative positive selection genes, three are associated with lipid metabolism. Erlin2 is a regulator of cytosolic lipid of cancer cells (Wang, Zhang et al. 2012). Fabp3 plays a role in metabolism and transport of long chain polyunsaturated fatty acid, and its role in lipid regulation may be important for mitochondrial function (Pollard, Ingram et al. 2019). Genetic variant association studies indicate Cilp2 regulates lipid species in both mice and humans (Jha, McDevitt et al. 2018). All these genetic and phenotypic features indicate a natural selection on lipid metabolism in beavers. Genes involved in lipid composition have been found under stronger selective pressure in long-lived species (Jobson, Nabholz et al. 2010). The evolution of lipid metabolism is connected to mammalian longevity (Li and de Magalhaes 2013), and the important role of lipid metabolism in aging and lifespan regulation have been observed in different species, including humans (Johnson and Stolzing 2019). We hypothesize that altered regulation of genes associated with lipid metabolism in the beaver genome likely contributes to its longevity. Although how altered lipid metabolism affects beavers' longevity needs further exploration, several unusual fatty acids, such as branched-chain (BCFA) and odd-chain (OCFA), which are typical for ruminants, were detected in beaver muscles. BCFA and OCFA have shown positive biological effects on anti-cancer activity (Domaradzki, Florek et al. 2019).

Like other long-lived rodent species, beavers are also cancer resistant. The malignant transformation of beaver cells needs both inactivation of Tp53, Rb1 and Pp2ca and constitutive activation of telomerase and HRas (Seluanov, Gladyshev et al. 2018). Such requirements for beaver cells are more stringent than for mouse cells and identical to those for human cells, which makes beavers a promising model to study human cancer. Several findings from this study may shed some light on the cancer resistance of beavers. Hpgd, a known tumor suppressor for many types of cancer, has been duplicated uniquely in beaver among rodents. The expansion of Aldh1a1 can better protect beaver cells against lipid aldehyde, which can damage DNA and proteins, by
metabolizing reactive aldehyde into harmless acetic acid. Increased expression of genes in DNA repair pathway can also protect beavers from cancer.

## Materials and Methods

## De novo genome assembly

We generated the beaver genome assembly using the third-generation sequencing technology and a new scaffolding method. A single 'Chicago' library (sequenced at a $\sim 87 \mathrm{X}$ coverage) was generated by Dovetail Genomics. Those sequence data were mapped to our initial assembly (with scaffold N50 $=55.69 \mathrm{~kb}$ ), which was generated by using AllPath-LG with default parameters on whole genome shutgun sequences from the Illumina HiSeq platform. And then HiRise scaffolding pipeline (Putnam, O'Connell et al. 2016) was used to generate the dovetail assembly (with a scaffold $\mathrm{N} 50=20.99 \mathrm{Mb}$ ). Finally, we used the published long PacBio reads to fill gaps in the dovetail assembly (Lok, Paton et al. 2017).

## BUSCO-based genome quality assessment

We used BUSCO v2(Simao, Waterhouse et al. 2015) to evaluate the completeness of the beaver genome assemblies. Briefly, BUSCO assesses the genome by searching for presence of nearuniversal single-copy genes from OthoDB v9(Zdobnov, Tegenfeldt et al. 2017). Absence of those conserved genes indicates incompleteness of the genome. In our analysis, we used the mammalian gene set consisting of 4,104 single-copy genes that are present in more than $90 \%$ mammalian species.

## Training for ab initio gene prediction

Augustus was trained by running BUSCO(Simao, Waterhouse et al. 2015) with the '--long' parameter, which performs a full optimization of training for Augustus gene finding. SNAP was trained following the previously described pipeline(Campbell, Holt et al. 2014) with three iterations.

## Gene structure and function annotation

Maker2(Holt and Yandell 2011) was used for gene structure prediction (see Figure 1A for the pipeline). Repeating elements were first masked by RepeatMasker (v4.07)(Tarailo-Graovac and Chen 2009), with RepBase repeat libraries (20170127)(Bao, Kojima et al. 2015) and beaverspecific repeating elements constructed by RepeatModeler (version 1.0.10) following the instruction given by
http://weatherby.genetics.utah.edu/MAKER/wiki/index.php/Repeat_Library_Construction--Basic. RepeatRunner was then used to further identified more divergent transposable protein elements provided by Maker2.

With the repeat-masked assembly, genes were predicted by ab initio gene predictors (i.e., SNAP(Korf 2004) and Augustus(Stanke and Waack 2003)) and evidence-based gene calling (i.e., using transcript assembly and protein sequences). For beaver gene transcripts, we used our 16,816 assembled transcripts (see above) and the 9,805 full length open reading frames from the published beaver genome(Lok, Paton et al. 2017). For protein evidence, about 66.7 thousands of reviewed mammalian protein sequences from Swiss-Prot(The UniProt 2017) were used for homolog-based gene prediction. These transcript and protein sequences were used to train the ab initio gene predictors (see above), polish the predicted gene models, and evaluate each predicted gene model. Finally, we predicted 26,515 beaver genes with evidence support from either transcript or protein sequences.

For each gene, Maker2 calculates an annotation edit distance (AED) score, which measures the goodness of fit of each gene to the evidence supporting it. A genome is usually considered well annotated if more than $90 \%$ genes have AED scores < 0.5 and over $50 \%$ proteins contain a recognizable domain(Campbell, Holt et al. 2014). In our beaver genome annotation, $\sim 90.7 \%$ gene models have AED scores lower than 0.5 (Figure 1 - Supplement Figure S1) and $78 \%$ of predicted gene products contain known protein domains by InterProScan (v5.25)(Jones, Binns et al. 2014).

## Orthology and phylogeny analyses

To identify gene families across species, we used OrthoDB (release 9)(Zdobnov, Tegenfeldt et al. 2017), which covers more than 600 eukaryotic species with functional annotation from more than 100 sources. In our analysis we selected 14 rodent and rabbit species from OrthoDB and mapped beaver and naked mole protein sequences to OrthoDB. We chose human and chimpanzee as outgroup species. We identified $\sim 21,000$ gene families, among which 5,087 gene families have single copy across all 18 species.

Single-copy gene families were used for phylogenetic analysis. Briefly, orthologs from each family were first aligned by MUSLE(Edgar 2004). Poorly aligned regions was removed by TrimAI(CapellaGutierrez, Silla-Martinez et al. 2009). Trimmed alignments were then concatenated and used to generate the phylogenetic tree by RAxML(Stamatakis 2014). The best substitution model for the full data matrix was determined by the Akaike information criterion in MrModeltest software(Nylander 2004). The best-scoring maximum likelihood (ML) tree was inferred using a novel rapid bootstrap algorithm combined with ML searches following 1000 RAxML runs (using the ' $f$-a' option)(Stamatakis 2014). The divergence times for the species analyzed were estimated by Reltime(Tamura, Battistuzzi et al. 2012). Diverged about 46 million years ago, Ords Kangroo rat is evolutionarily closest to beaver (Figure 1B).

Identification of significant gene expansion in beaver
With the phylogeny that we built and the gene counts from OrthoDB (Zdobnov, Tegenfeldt et al. 2017), we predicted gene family expansion using CAFE 3 (Han, Thomas et al. 2013). CAFE first calculated an error model for gene family size estimation as a part of genome assembly and annotation. For our gene counts, it estimated $\sim 4.8 \%$ of the gene families had incorrect gene numbers assigned to them. It corrected this error before calculating ancestral family sizes and then estimated a more accurate gene family evolution rate. Finally, it calculated the probability of observing the sizes of each gene family of those species by Monte Carlo re-sampling procedure. Families with large variance in size, especially observed in closely related species, will tend to have a lower $P$-value. For families with low $P$-values ( 0.01 as the default), a $P$-value for the transition between parent and child nodes for each branch in the phylogeny was also calculated to identify where the large change of family size takes place.

We identified 84 candidate gene families showing significant expansion in the beaver branch (FDR < 0.01). To sidestep false positives due to assembly and annotation artifact, pseudogenes, etc., we first removed genes without the support of RNA-seq reads from all the tissues (Supplement Table $\mathbf{S 1 )}$ in our analysis. We also identified pseudogenes and removed ones with low expression levels (TMP < 5 across tissues). We kept pseudogenes with relatively high expression levels, since they may be functional. We then further filtered genes based on the percentage of identities between beaver gene copies. Briefly, we iteratively removed the gene copy with the least average identity to other beaver genes in the family, if the average identity is below $70 \%$. After this procedure, we reduced our candidate expanded gene families to eight genes (Figure 2 - Supplement Figure S2).

## Beaver-specific expansion of conserved genes

To find beaver-specific expansion of conserved genes, we require the same number of copies in other species included in our study and a higher copy number only in the beaver genome. Initially, 234 gene candidates were collected. After removing copies without RNA-seq reads support, lowly expressed pseudogenes (TPM < 5), and gene copies with low identity between gene products (identity < 0.7), 83 genes remained. We further processed them to remove potential false positives by considering only genes with the following qualifications: (1) gene predictions with good synteny in their genomic neighborhood among beaver, mouse, and human; (2) genes with differences in the coding sequences of their copies; (3) genes with variable sites in their copies supported by RNA-seq reads (from the same individual beaver). 18 beaver-specific gene expansions met these stringent criteria and were further validated by qPCR, with five of them show more copy numbers comparing to reference genes.

## Pseudogene identification

GeneWise(Birney, Clamp et al. 2004) was used to identify pseudogenes (Figure 2 - Supplement Figure S5). For a predicted beaver gene, we extracted the genomic sequence from its locus with both upstream and downstream 5 -kb regions. Using its mouse (or human) ortholog as the reference, we then scanned the gene sequence by GeneWise and checked the presence of frameshift indels that can 'pseudogenize' the gene. We also checked if the predicted gene can be a processed pseudogene, which can be generated through mRNA retrotransposition. If a predicted beaver gene has no introns, we checked its orthologs in other rodents to determine whether it is a processed pseudogene or a single-exon gene.

## Positive selection

Branch-site likelihood method (Zhang, Nielsen et al. 2005) implemented in PosiGene (Sahm, Bens et al. 2017) was used to identify genes under positive selection in the beaver genome. Only 9,750 genes with alignment of ortholog coding sequence from at least 10 species were considered. After the initial prediction by PosiGene (Sahm, Bens et al. 2017), manual checking was carried out for all $\sim 150$ candidate positive selection genes ( $P<0.05$ ). Briefly, we removed false positives, and for genes with ambiguous signals we manually checked and improved their predicted gene structures based on protein sequence alignments of orthologs and supporting RNA-seq reads using Apollo (Lee, Helt et al. 2013). We then used the improved coding sequences as the input for PosiGene and ran the pipeline three times. Finally, we identified 21 beaver genes under positive selection with FDR $<0.01$ consistently in all three independent runs. The coding sequences of other species were download from NCBI as of January of 2020.

## RNA extraction and sequencing

Total RNA was isolated from frozen tissues (brain and liver with two replicates each) using the mRNA-Seq Sample Prep Kit Illumina (San Diego, CA. USA) in accordance with the manufacturer's instructions, and the mRNA integrity was checked by the agarose gel analysis. Polyadenylated RNA was then isolated using a poly-dT bead procedure and followed by reverse transcription. Short-insert 'paired-end' libraries were prepared using the Illumina TruSeq Sample Preparation Kit v 2 , and the sequencing was performed on the Illumina HiSeq2000 platform. The raw data were processed by NGS QC Toolkit (v.2.3.3)(Patel and Jain 2012) to remove low-quality reads.

## De novo transcriptome assembly

RNA-seq data from brain and liver (each with two replicates) were assembled using Trinity(Haas, Papanicolaou et al. 2013). We collected 16,816 high quality beaver transcripts, by requiring each transcript to meet the following conditions: (1) proper start and stop codons, (2) a correct reading
frame (codons in triplets), (3) a gene length similar to the mouse ortholog ( $\pm 20 \%$ ), (4) a good alignment with the mouse ortholog (in peptide sequence), (5) the best candidate among all beaver Trinity assembled sequences.

## qPCR validation

Candidate primers were designed by Primer-Blast (Ye, Coulouris et al. 2012) in the exons of selected genes. Specificity of the primers was first checked by Primer-Blast against all of the coding sequences in the beaver genome. Then genome-wide specificity was checked by MFEprimer (Qu, Zhou et al. 2012). Specificity of primers was further inspected by gel electrophoresis and the melting curve analysis. Amplification efficiency was checked by a standard curve analysis for each candidate primers with different amount of DNA input to make sure that the finally used primers have equal amplification efficiency. Quantitative PCR was used to quantify gene copy number, with three replicates for each of the three different amount of DNA input. Two genes - Hcfc1 on an autosome and Pelo on chromosome X - with no predicted expansion in beaver and no annotated expansion in other rodents (according to orthoDB (Zdobnov, Tegenfeldt et al. 2017)) were used as references. The copy number of each target gene relative to its corresponding reference genes was calculated by $\Delta C_{t}$. And the genomic DNA of a male beaver, different from ones whose samples were used for the genome sequencing and assembly and transcriptomics, was used for the experiment. Primers for target and reference genes are listed in Supplement Table S2.

## Western blot

Liver cytosolic extracts were quantitated using the BCA assay (Thermo). 30 ug of each extract was resolved through 4-20\% Criterion Tris-Glycine (TGX) Stain-Free SDS-PAGE (Biorad) and transferred to nitrocellulose. Prior to transfer, total proteins were imaged using a Biorad Gel Documentation System to control for protein loading. Rabbit polyclonal anti-Aldh1a1 (Invitrogen cat\# PA5-95937) was used as the primary to detect Aldh1a1 isoforms. Protein loading was also checked by performing Western blot using anti-beta Actin (Abcam, ab8227).

## Ethanol treatment of cells

The effects of ethanol on cellular functions were studied with cultured primary beaver and mouse lung fibroblasts. We used two different beaver and mouse cell lines, low PD cells stabilized in culture. Cells were seed in complete medium containing $15 \%$ FBS in 96 well plate for 24 hours, then incubated in medium containing 300 mM ethanol for seven hours. Then WST-1 assay was performed by adding WST-1 reagent (cell proliferation reagent, Roche) directly to the culture wells, incubating for 4 h at $37^{\circ} \mathrm{C}$ and $5 \% \mathrm{CO}$, shaking thoroughly for 1 min on a shaker and then measuring the absorbance at 430-480 nm with TECAN spark 20M spectrophotometric reader. The stable
tetrazolium salt WST-1 was cleaved to a soluble formazan by a complex cellular mechanism that occurs primarily at the cell surface. This bio-reduction is largely dependent on the glycolytic production of $N A D(P) H$ in viable cells. Therefore, the amount of formazan dye formed directly correlates to the number of metabolically active cells in the culture.

To examine whether beaver and mouse lung fibroblast cells show significantly different rate of cell death under the ethanol treatment, relative to corresponding controls, a linear mixed effects model was used with species as the fixed effect and individuals ( $n=4$ for each species with 3 replicates each individual) as random effects ( Imer(cell death rate $\sim$ Species+(1|individual)) ), which is implemented in the R package Ime4 (Bates, Mächler et al. 2015).

## Liver aldehyde dehydrogenase activity

In addition to the cytosolic Aldh1a1, a mitochondrial enzyme, Aldh2, also plays an important role in the aldehyde metabolism. A previous study had shown that in murine liver Aldh1a1 plays a more important role in the metabolism of endogenous aldehydes than Aldh2 (Makia, Bojang et al. 2011). To exclude influence from Aldh2, we tested aldehyde dehydrogenase activity using cytosolic protein extracts from liver cells. Crude cytosolic protein extracts were prepared from wild beaver or mouse (C57/BI6) livers using phosphate buffer similar to a previously described method (Makia, Bojang et al. 2011). Specifically, tissue was resuspended in K-Phos Buffer ( 50 mM potassium phosphate $\mathrm{pH} 7.4,250 \mathrm{mM}$ sucrose, 1 mM EDTA) at $1 \mathrm{~g} / 3.0 \mathrm{ml}$. Tissues were dounce homogenized in ice using a Teflon pestle followed by ten passages through a 27 Ga needle to lyse the cells. Samples were centrifuged (Thermo/Sorvall Legend Micro21R) at $1000 \mathrm{rpm}(100 \mathrm{xg}$ ) for 10 minutes at $4^{\circ} \mathrm{C}$ to remove any remaining intact cells. Supernatant was transferred to a clean tube and centrifuged at $14,000 \mathrm{rpm}(18,800 \mathrm{x} \mathrm{g})$ for 20 minutes at $4^{\circ} \mathrm{C}$. We found that the supernatants/extracts prepared this way and rapidly aliquotted and frozen with liquid nitrogen did not lose significant activity after one freeze/thaw cycle. Just prior to assaying, a $100 \mu \mathrm{l}$ aliquot of the supernatant was thawed on ice and passaged through a 0.5 ml Zeba desalting spin column (Thermo; 7,000 mwco) equilibrated in the same K-Phos buffer to remove small molecules that might compete with added substrates. Aldehyde dehydrogenase was measured in 384-well transparent microplates in $50 \mu \mathrm{l}$ volume consisting of $10 \mu \mathrm{l}$ extract in the same K-Phos buffer supplemented with $1 \mathrm{mM} \mathrm{NAD}{ }^{+}$. Aldehyde substrates all-trans-retinal (RET), malonaldehyde tetrabutylammonium salt (MAD), and 4-hydroxynonenal (NHE) were added at indicated concentrations from DMSO stocks. Final DMSO in assay was 4\%. Dehydrogenase activity was measured as the change in absorbance at 340 nm (reduction of $\mathrm{NAD}^{+}$to NADH) using a Tecan Spark 20M plate reader preequillibrated at $37^{\circ} \mathrm{C}$. Rates were determined after a few minutes of lag during which time the plate temperature was adjusting to $37^{\circ} \mathrm{C}$. NAD+, all-trans-retinal, and malonaldehyde tetrabutylammonium salt, were purchased from Sigma. 4-hydroxynonenal was purchased from

Cayman Chemical. Aldh1a1-class of specific activity was determined by subtracting any trace amount of NAD+ to NADH conversion that occurred in the absence of added aldehyde substrate.

## Gene expression comparison between beavers and mice

For our transcriptomic analysis, we used beaver samples from two young male adults and RNAseq data from 6 weeks old young adult mice (Li, Qing et al. 2017). The RNA-Seq data of mice were generated from eight individuals ( 4 females and 4 males) for both liver and brain. Although our beaver samples were from two males, we included both male and female mice to increase the detection power with sex as a factor in the model for differential gene expression analysis. Principle component analysis showed good separation between beaver and mouse samples (Figure 6 Supplement Figure S14). We only considered protein coding genes and used Kallisto (Bray, Pimentel et al. 2016) to estimate gene expression levels, which showed better performance for genes with paralogs.

The detection of differential expressed genes between species is much more complicated than analysis within species. Because several differences between two genomes - e.g., different gene annotation quality, different number of genes, and different lengths of same genes between species - can lead to biased results, we used the following strategies together in our analysis. (1) For gene annotation quality control, we only considered 16,303 beaver genes that show high protein sequence similarity with their corresponding orthologs from other rodent species (including mouse) and could be successfully clustered into gene families by OthorDB (Zdobnov, Tegenfeldt et al. 2017). (2) We further restricted our analysis to genes $(12,089)$ that share one-to-one orthology between beaver and mouse. (3) We also controlled gene length. We used tximport (Soneson, Love et al. 2015) to obtain family-level gene expression and weighted gene lengths for each gene family. Together with estimated read counts, the weighted gene lengths were also provided to DESeq2 (Love, Huber et al. 2014), and differences in gene lengths among samples were considered for differential expression analysis.

## ALDH1A1 expression during human aging

Using data from the GTEx Project (phs000424.GTEx.v7.p2.c1) (Carithers, Ardlie et al. 2015), we studied gene expression changes of ALDH1A1 during aging. In the GTEx Project, many samples were taken from post-mortem individuals. Studies have demonstrated that expression of some genes may change in certain tissues after death and show significant association with post-mortem interval (PMI, in minutes between death and sample collection) (Ferreira, Munoz-Aguirre et al. 2018). To reduce artifacts from PMIs, we ignored tissues where the expression of ALDH1A1 shows significant association with PMI ( $\mathrm{P}<0.01$ ). For the remaining tissues, we used liner regression to
assess the association between gene expression and age with sex, body mass index, and PMI as covariates.

## Data Analyses and Availability

Genome DNA sequence data were submitted to BioProject database (https://www.ncbi.nlm.nih.gov/bioproject/) with accession numbers PRJNA505050. Genome assembly was submitted to the NCBI Assembly database (https://www.ncbi.nlm.nih.gov/assembly/) with accession number RPDE00000000. RNA-Seq data of liver and brain tissue of two beaver individuals were submitted to NCBI with accession number PRJNA627298, which will be released once the manuscript is accepted.

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## Additional files

Supplementary file 1. Coding sequences and protein sequences alignments with selection signals of beaver genes under positive selection (FDR<0.01).

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Figures and Tables
A


B


20 Million Years

Figure 1. Genome annotation and phylogeny. (A) The beaver genome annotation pipeline. As a part of the annotation, we used 9,805 full-length open reading frames (FLORFs) from the published beaver genome (Lok, Paton et al. 2017). (B) Timed phylogeny. 5,087 single copy genes were used for the analysis.


Figure 2. Expansion of Aldh1a1 in the beaver genome. (A) The synteny of the Aldh1a1 locus in the genomes of beaver, mouse, and human. We identified ten copies of Aldh1a1 in the beaver genome, including two pseudogenes (35.15 and 266.4) and a low-quality copy (266.2). (B) Gene structure of the seven functional copy of Aldh1a1. (C) RNA-seq reads coverage at genomic locations of different amino acid residues among seven Aldh1a1 gene products. Variable sites were highlighted in red, and the gray box indicates the selected sites where we checked the coverage of RNA-seq reads. We analyzed RNA-Seq reads from four beaver samples, B1-4 (Supplementary Table 1), separately. (D) Validation of the Aldh1a1 copy number by qPCR. Two single-copy genes were used as references: Hcfc 1 on an autosome and Pelo on Chromosome X. (E) Expression profiles of Aldh1a1 copies across several different beaver tissues. (F) Aldh1a1 expression in liver and brain of both beavers and mice. For each tissue, we measured the expression of Aldh1a1 in 2 beaver individuals and 8 mouse individuals.
(G) Western blot of Aldh1a1 protein from beaver and mouse liver extracts of 4 different samples of each species. (H) Quantified Aldh1a1 levels in liver extracts. It is statistically higher ( $P=0.001$ by single side Welch Two Sample $t$-test) in beaver liver than in mouse liver.

A


B


Figure 3. Functional characterization of Aldh1a1. (A) Cell viability. Beaver lung fibroblasts have lower percentage of death in the presence of high-concentration ethanol, normalized by corresponding controls ( $p=0.002$ by mixed effects model, see Methods). (B) Aldehyde metabolic activity. Enhanced Aldh1a1 activity of beaver in aldehyde metabolism. Activities are normalized by corresponding controls without any addition (see Methods). RET: all-trans-retinal; MDA: malonaldehyde; HNE: 4-hydroxynonenal. Lung fibroblasts were used in experiments of panels A. Cytosolic extracts from hepatocytes were used for the experiment of panel B. 4 different biological samples of each species were used in experiments of $\mathbf{A}$ ) and $\mathbf{B}$ ), with 3 replicates of each sample.

A


Beaver:C2678


C
82.2 MHVNGKVALVTGAAQGIGRAFAEALLHKGAKVALVDWNLEAGVKCKAALDEQFEPQKTLF 82.17 --------------------------------VALVDWNLEAGVKCKAALDEQFEPQKILF Mouse MHVNGKVALVTGAAQGIGKAFAEALLLHGAKVALVDWNLEAGVKCKAALDEQFEPQKTLF 82.2 IQCDVADQEQLRDTFRKVIDHFGRLDILVNNAGVNNEKNWEKTLQINLVSVIDATYLGLD 82.17 IQCDVADQEQLRDTFRKVVDHFGRLNILVNNAGVINEKNWEKTLQINLVSVIDATYLGLD Mouse vQCDVADQKQLRDTFRKVVDHFGRLDILVNNAGVNNEKNWEQTLQINLVSVISGTYLGLD
82.2 YMSKQNGGDGGIIINMSSLAGLLPAAQQPVYCASKHGVIGFTRSAAMAANLMNSGVRLNA 82.17 YMSKQNGGDGGIIINMSSLAGLVPVAQHPVYCASKYGVIGFTRSAAMAANLMNSGVRLNA Mouse YMSKQNGGEGGIIINMSSLAGLMPVAQQPVYCASKHGIIGFTRSAAMAANLMKSGVRLNV
82.2 ICPGFVNTPILKSIDKEENMGQYLEYTDHIKDMMKYFGILDPSMIANGLITLIEDDGLNG 82.17 ICPGFVNTPILKSFEKEKNMGQYFEYTDHIKDMMKYFGILDPSMIANGLITLIEDDGLNG Mouse ICPGFVDTPILESIEKEENMGQYIEYKDQIKAMMKFYGVLHPSTIANGLINLIEDDALNG

```
82.2 AIMKITTSKGIHFQEYDPPPSHAVITEHKHK 82.17 AIMKITTSKGIHFQEYDPPPSHAVITEHKHK
``` Mouse AIMKITASKGIHFQDYDISPLLVKAPLTS--


E


Figure 4. Beaver-specific gene expansion, including tumor suppressor Hpgd. (A) qPCR validation of the copy number of beaver-specific expanded genes. The copy numbers relative to the reference gene on chromosome X (i.e., Pelo) and on an autosome (i.e., Hcfc1) are shown for each candidate gene. (B) Synteny at the Hpgd locus. (C) Alignment of beaver and mouse Hpgd protein sequences. Variable sites are highlighted in red, and the gray box indicates the selected sites where we checked the coverage of RNA-seq reads. (D) Coverage of RNA-seq reads at the selected sites in individual beavers. (E) Expression of Hpgd in brain and liver of both beavers and mice.

A


B
\begin{tabular}{cccc}
\hline Evolutionary changes & \multicolumn{3}{c}{ Predicted deleterious score } \\
\cline { 2 - 4 }\((\) Human \(\rightarrow\) Beaver) & PROVEAN & PolyPhen2 & CADD \\
\hline S \(\rightarrow\) S (AGT \(\rightarrow\) TCG) & 0.00 & - & 10.62 \\
I \(\rightarrow\) P (ATC \(\rightarrow\) CCC) & -3.51 & 0.92 & 23.00 \\
\(S \rightarrow\) W (AGT \(\rightarrow\) TGG) & -2.94 & 0.95 & 21.70 \\
L \(\rightarrow \varnothing\) (deletion) & -3.44 & - & 13.26 \\
\(\varnothing \rightarrow\) EIRKS (insertion) & -6.12 & - & 14.50 \\
\(\mathrm{R} \rightarrow \mathrm{K}(\) AGA \(\rightarrow\) AAA) & 0.23 & 0.00 & 0.13 \\
\hline
\end{tabular}

Figure 5. Mtbp is under positive selection. (A) Positive selection signals in Mtbp gene. We observed nucleotide positions and indels under positive selection in coding sequences of three exons, corresponding to the middle and the C domains of Mtbp. (B) Nucleotide sequence changes between the beaver and the human genomes. We predicted the deleteriousness of these changes based on their human genome annotation, using the following metrics and thresholds: PROVEAN score < -2.5, 0.85 < Polyphen2 score < 1.0, and CADD score > 20.


Figure 6. Gene sets enriched with genes differentially expressed between beavers and mice. (A) KEGG signaling pathways enriched with differential expression genes in liver tissue. (B) KEGG signaling pathways enriched with differential expression genes in brain tissue. (C) REACTOM pathways enriched with differential expression genes in liver tissue. (D) REACTOM pathways enriched with differential expression genes in brain tissue. (E) Hallmark gene sets enriched with differential expression genes in liver tissue. (F) Hallmark gene sets enriched with differential expression genes in brain tissue. UP: enriched pathways of up-regulated genes in beaver tissue comparing to mouse tissue. DOWN: enriched pathways of down-regulated genes in beaver tissue comparing to mouse tissue. Top 10 enriched pathways of up- and down- regulated genes are shown for each type of gene sets. Hallmark gene sets from the Molecular Signatures Database represent specific well-defined biological states or processes and display coherent expression (Liberzon, Birger et al. 2015).


Figure 7. Age-associated expression of ALDH1A1 in human. (A) Correlation of ALDH1A1 expression with age, sex, body mass index (BMI), and post-mortem interval (PMI, minutes between death and sample collection). Tissues are ordered according to the \(P\)-values of the correlation with age. A significant correlation (FDR \(<0.01\) ) is denoted by an asterisk. (B) Significant correlation between ALDH1A1 expression and age in four human tissues. The gray shaded area around the regression line indicates \(95 \%\) confidence interval.

Table 1. Putative positive selection genes in beaver (FDR<0.01)
\begin{tabular}{lllll}
\hline Gene & Gene name & \begin{tabular}{l} 
Num. \\
seq. \({ }^{1}\)
\end{tabular} & \begin{tabular}{l} 
Num. sites \(^{2}\)
\end{tabular} & FDR \\
\hline Mtbp & MDM2 Binding Protein & 12 & 4 & \(6.58 \mathrm{E}-04\) \\
Ptx3 & Pentraxin 3 & 16 & 6 & \(1.56 \mathrm{E}-03\) \\
Tbxa2r & Thromboxane A2 Receptor & 15 & 6 & \(2.75 \mathrm{E}-03\) \\
Hsd17b1 & Hydroxysteroid 17-Beta Dehydrogenase 1 & 14 & 6 & \(3.32 \mathrm{E}-03\) \\
Erlin2 & ER Lipid Raft Associated 2 & 18 & 3 & \(3.86 \mathrm{E}-03\) \\
Fabp3 & Fatty Acid Binding Protein 3 & 17 & 7 & \(3.86 \mathrm{E}-03\) \\
Adam19 & ADAM Metallopeptidase Domain 19 & 18 & 6 & \(4.38 \mathrm{E}-03\) \\
Cilp2 & Cartilage Intermediate Layer Protein 2 & 14 & 30 & \(5.07 \mathrm{E}-03\) \\
Chst12 & Carbohydrate Sulfotransferase 12 & 18 & 5 & \(5.07 \mathrm{E}-03\) \\
Urb2 & URB2 Ribosome Biogenesis Homolog & 16 & 4 & \(5.07 \mathrm{E}-03\) \\
Mrp/37 & Mitochondrial Ribosomal Protein L37 & 18 & 6 & \(5.07 \mathrm{E}-03\) \\
Depdc7 & DEP Domain Containing 7 & 17 & 7 & \(5.61 \mathrm{E}-03\) \\
Neu2 & Neuraminidase 2 & 12 & 3 & \(6.07 \mathrm{E}-03\) \\
Vwa5a & Von Willebrand Factor A Domain Containing 5A & 10 & 16 & \(6.07 \mathrm{E}-03\) \\
Cox15 & Cytochrome C Oxidase Assembly Homolog COX15 & 18 & 4 & \(6.64 \mathrm{E}-03\) \\
Gdf2 & Growth Differentiation Factor 2 & 17 & 9 & \(7.31 \mathrm{E}-03\) \\
Sit1 & Signaling Threshold Regulating Transmembrane & 10 & 4 & \(7.31 \mathrm{E}-03\) \\
Aoc1 & Adaptor 1 & Amine Oxidase Copper Containing 1 & 17 & 9 \\
Cyb5a & Cytochrome B5 Type A & \(8.02 \mathrm{E}-03\) \\
Scn4b & odium Voltage-Gated Channel Beta Subunit 4 & 18 & 1 & \(8.52 \mathrm{E}-03\) \\
II23a & Interleukin 23 Subunit Alpha & 6 & \(8.64 \mathrm{E}-03\) \\
\hline lots & & 16 & 4 & \(1.00 \mathrm{E}-02\) \\
\hline
\end{tabular}

Notes:
1. Num. seq.: the number of available orthologous sequences from a subset of our selected 18 species to build the high-quality alignment for detecting selection signals.
2. Num. sites: the number of amino acid residues that show selection signal in beavers.


Figure S1. Related to Figure 1. Cumulative Distribution of AED score. All: 26,515 predicted genes; OrthoDB: 17,661 genes grouped into gene families by mapping beaver proteins to OrthoDB(Zdobnov, Tegenfeldt et al. 2017); PosiGene: 9,750 genes with coding sequences of at least 10 species in the alignment for identification of genes under positive selection; Expression1: 16,303 beaver genes with mouse orthologs; Expression2: 12,089 beaver genes with one-to-one mouse orthologs. An AED (Annotation Edit Distance) score measures the goodness of fit of each gene to the evidence supporting it. The dashed line shows the EAD score \(=0.5\). A genome is considered well annotated if more than \(90 \%\) genes have AED scores \(<0.5\) (Campbell, Holt et al. 2014).


Figure S2. Related to Figure 2. Gene families with significant expansion in beavers. Prl: prolactin; Aldh1a1: aldehyde dehydrogenase 1 family member A1; Prkrir: 52-KDa repressor of the inhibitor of the protein kinase; Hsfy1: heat shock transcription factor Y-linked 1; Ak6: adenylate kinase 6; Immunoglobulin-like: no gene name but with Immunoglobulin-like domain; Card16: caspase recruitment domain family member 16; Defa5: defensin alpha 5.


Figure S3. Related to Figure 2. Phylogenetic tree with Aldh1a1 copy numbers. Shown on the leaf and the ancestral nodes are the copy numbers (in black) of Aldh1a1 in species annotated by this study and OrthoDB(Zdobnov, Tegenfeldt et al. 2017) and ones (in gray) estimated by CAFE3(Han, Thomas et al. 2013), respectively.



B
Legend: ! Frame shift; [] Intron boundaries

\begin{tabular}{|c|c|c|}
\hline \multicolumn{3}{|r|}{302 CCVAASRIFV-EESVYDEFVKRSVERAKKYV-LGNPLTPGINQGPQ} \\
\hline & & \(\mathrm{C}+\mathrm{A}+\mathrm{F}\) E \(\quad \mathrm{YD}+\mathrm{FV}+\mathrm{R} \quad \mathrm{AK}+\mathrm{L} \quad \mathrm{L} \quad\) NQGPQ \\
\hline & & CVIACIQAFGWENPFYDDFVRRKC!AAKAGIFLEILLDSRNNQGPQ \\
\hline \multirow[t]{3}{*}{Beaver(266.4)} & 28058 & tgagtacgtgtgacttggtgaaat4ggaggattgaccgtaaacgcc \\
\hline & & gttcgtactggaactaaattggag ccacgtttatttacgaaagca \\
\hline & & ctatccgttgggtacttttgaggt gtaatacaattgccgttact \\
\hline
\end{tabular}
\begin{tabular}{ccrr} 
Mouse(Aldh1a1) 346 & \begin{tabular}{r} 
IDKEQHDKILDLIESGKKEGAK \\
IDKEQH+KILDLIESGKKEGA
\end{tabular} \\
Beaver(266.4) & 27919 & & IDEQHNKILDLIESGKKEGAT
\end{tabular}
\begin{tabular}{|c|c|c|}
\hline Mouse(Aldh1a1) & 368 & LECGGGRWGNKGFFVQPTVFSNVTDEMRIAKEE LECGGG WGNKG+FVQPTVFSNVTDEMRIAKEE \\
\hline & & LECGGGPWGNKGYFVQPTVFSNVTDEMRIAKEE \\
\hline Beaver(266.4) & 23326 & cgtgggctgaagttgccagttagaggacagagg \\
\hline & & taggggcggaagattaccttcatcaatgtcaaa \\
\hline
\end{tabular}

Mouse(Aldh1a1) 401
IFGPVQQIMKFKSVDDVIKRAN IFGPVQQIMKFKSVD+VIKRAN Beaver(266.4) 23227 GTAATGG Intron \(5 \quad \begin{array}{r}\text { IFGPVQQIMKFKSVDEVIKRAN } \\ \text { TAGatgcgccaaatatggggaaaga }\end{array}\) <0-----[23227 : 20278]-0>ttgctaattatactaattagca

Mouse(Aldh1a1) 423 NTTYGLAAGLFTKDLDKAITVSSALQAGVVW NT YGLAAG+FTK LD+AITVSSALQAGVVW Beaver(266.4) \(20211 \begin{aligned} & \text { NTPYGLAAGVFTKGLDRAITVSSALQAGVVW } \\ & \text { aactgtggggtaagcgagaagttgccggggt }\end{aligned}\) accagtccgttcagtagctctccctacgttg ctctcgaaactcactcacaaccttgggtagg
\begin{tabular}{ccc} 
Mouse(Aldh1a1) 285 & & \begin{tabular}{c}
-DIA-VEFAHHGVFYHQGQ \\
+A EFAH G++
\end{tabular} \\
& & +:D[gat]
\end{tabular}
\begin{tabular}{ll}
1032 & C \\
1033 & Aldh1a1 \\
1034 & 35.13 \\
1035 & 35.14 \\
1036 & 35.16 \\
1037 & 266.22 \\
1038 & 266.23 \\
1039 & 265.1 \\
1040 & 266.5 \\
1041 & Aldh1a7 (Mouse) \\
1042 & Aldh1a1 (Mouse) \\
1043 & ALDH1A1 (Human) \\
1044 & \\
1045 & 35.13 \\
1046 & 35.14 \\
1047 & 35.16 \\
1048 & 266.22 \\
1049 & 266.23 \\
1050 & 265.1 \\
1051 & 266.5 \\
1052 & Aldh1a7 (Mouse) \\
1053 & Aldh1a1 (Mouse) \\
1054 & ALDH1A1 (Human) \\
1055 & \\
1056 & 35.13 \\
1057 & 35.14 \\
1058 & 35.16 \\
1059 & 266.22 \\
1060 & 266.23 \\
1061 & 265.1 \\
1062 & 266.5 \\
1063 & Aldh1a7 (Mouse) \\
1064 & Aldh1a1 (Mouse) \\
1065 & ALDH1A1 (Human) \\
1066 & \\
1067 & 35.13 \\
1068 & 35.14 \\
1069 & 35.16 \\
1070 & 266.22 \\
1071 & 266.23 \\
1072 & 265.1 \\
1073 & 266.5 \\
1074 & Aldh1a7 (Mouse) \\
1075 & Aldh1a1 (Mouse) \\
1076 & ALDH1A1 (Human) \\
1077 & \\
1078 & 35.13 \\
1079 & 35.14 \\
1080 & 35.16 \\
1081 & 266.22 \\
1082 & 266.23 \\
1083 & 265.1 \\
1084 & 266.5 \\
1085 & Aldh1a7 (Mouse) \\
1086 & Aldh1a1 (Mouse) \\
1087 & ALDH1A1 (Human) \\
1088 & \\
1089 & 35.13 \\
1090 & 35.14 \\
1091 & 35.16 \\
1092 & 266.22 \\
1093 & 266.23 \\
1094 & 265.1 \\
1095 & 266.5 \\
1096 & Aldh1a7 (Mouse) \\
1097 & Aldh1a1 (Mouse) \\
1098 & ALDH1A1 (Human) \\
1099 & \\
&
\end{tabular}

MLSSGKPDLPTPLTNLKIQYTKIFINNEWHNSVSGKKFPVLNPATEEEICQIEEGDKADV MQFSGKPDLPTPLTNLKIQYTKIFINNEWHNSVSGKKFPVLNPATEEEICQVEEGDKADV MQSSRQPDLPAPIANLKIQYTKIFINNEWHNSVSGKKFPVLNPATEEEICQVEEGDKADV MQSSGKSDLPTPLANLKIQYTKIFINNEWHNSVSGKKFPVLNPATEEEICQVEEGDKADV MQSSGKPDLPTPLANLKIQYTKIFINNEWHNSVSGKKFSVLNPATEEEICQVEEGDKADV MQSGGKSDLPTPLANLKIQYTKIFINNEWHNSVSGKKFPVLNPATEEEICQVEEGDKADV MQSSGKSDLPTPLANLKIHYTKIFINNEWHNSVSGKKFPVLNPATEEEICQVEEGDKADV MSSPAQPAVPAPLANLKIQHTKIFINNEWHDSVSSKKFPVLNPATEEVICHVEEGDKADV MSSPAQPAVPAPLADLKIQHTKIFINNEWHNSVSGKKFPVLNPATEEVICHVEEGDKADV MSSSGTPDLPVLLTDLKIQYTKIFINNEWHDSVSGKKFPVFNPATEEELCQVEEGDKEDV

DKAVKAARQAFQIGSPWRTMDASERGRLLFKLADLIERDRLLLATMESMNAGKLFSHAYL DKAVKAARQAFQIGSPWRTMDASERGRLLFKLAKLIERDRLLLATMESINGGKIFHSTYL DKAVKAARQAFQIGSPWRTMDASQRGRLLLKLADLIERDRLLLATMESMNGGKLFPPAYL DKAVKAARQAFQIGSPWRTMDASERGRLLFKLAALIERDRLLLATMESINGGKLFPPTYL DKAVKAARQAFQIGSPWRTMDASQRGRLLFKLADLIERDRLLLATMESINGGKPFPPTYL DKAVKAARQAFQIGSPWRTMDASQRGRLLFKLADLIERDRLLLATMESINGGKLFPPTYL DKAVKAARQAFQIGSPWRTMDASQRGRLLFKLADLIERDRLLLATMESMNGGKLFPPTYL DKAVKAARQAFQIGSPWRTMDASERGRLLNKLADLMERDRLLLATMESMNAGKVFAHAYL DKAVKAARQAFQIGSPWRTMDASERGRLLNKLADLMERDRLLLATMEALNGGKVFANAYL DKAVKAARQAFQIGSPWRTMDASERGRLLYKLADLIERDRLLLATMESMNGGKLYSNAYL

MDLGGCIKTLRYCAGWADKVHGYTIPSDGDVFTYTRREPIGVCGQIIPWNFPLVMLIWKL MELGGCIETLQYFAGWADKIHGYTIPSDGDVFTYTRREPIGVCGQIIPWNFPLDMLIWKI LELGMCIKTLQYHAGWADKIHGYTIPSDGDVFTYTRREPIGVCGQIIPWNAPLLMFIWKI MELGMCIQSIQYFAGWADKVHGYTIPSDGDVFTYTRREPIGVCGQIIPWNGPLIVLLSKI MELGMCIQAIQYCAGWADKVHGYTIPSDGDVFTYTRREPIGVCGQIIPWNGPLLMFIWKI MELGMCIQVVQYCAGWADKVHGYTIPSDGDVFTYTRREPIGVCGQIIPWNGPLIILLSKI MELGTCIKALQYCAGWADKIHGYTIPSDGDVFTYTRREPIGVCGQIIPWNGPLITLIWKI LDVEISIKALQYFAGWADKIHGQTIPSDGNIFTYTRREPIGVCGQIIPWNGPLIIFTWKL SDLGGCIKALKYCAGWADKIHGQTIPSDGDIFTYTRREPIGVCGQIIPWNFPMLMFIWKI NDLAGCIKTLRYCAGWADKIQGRTIPIDGNFFTYTRHEPIGVCGQIIPWNFPLVMLIWKI

GPALSCGNTVIVKPAEQTPLTALHVASLIKEAGFPPGVVNIVPGYGPTAGAAISSHMDID APALSCGNTVIVKPAEQTPLTALHVASLIKEAGFPPGVVNIVPGYGPTAGAAISSHMDID APALSCGNTVIVKPAEQTPLTALHMASLIKEAGFPPGVVNIVPGYGPTAGAAISSHMDID APALSCGNTVIVKPAEQTPLTALHVASLIKEAGFPPGVVNIVPGYGPTAGAAISSHMDID APALTCGNTVIVKPAEQTPLTALHMASLIKEAGFPPGVVNIVPGYGPTAGAAISSHMDID APALSCGNTVIVKPAEQTPLTALHVASLIKEAGFPPGVVNIVPGYGPTAGAAISSHMDID APALSCGNTVIVKPAEQTPLTALHVASLIKEAGFPPGVVNIIPGYGPTAGAAISSHMDID GPALSCGNTVVVKPAEQTPLTALHMASLIKEAGFPPGVVNIVPGYGPTAGGAISSHMDID GPALSCGNTVVVKPAEQTPLTALHLASLIKEAGFPPGVVNIVPGYGPTAGAAISSHMDVD GPALSCGNTVVVKPAEQTPLTALHVASLIKEAGFPPGVVNIVPGYGPTAGAAISSHMDID

KVAFTGSTEVGKMIKEAAGKSNLKRVTLELGGKSPCIVFADADMDNAVELAHQGLFYHQG KVAFTGSTEVGKIIKEAAGKSNLKRVTLELGGKSPCIVFADADLEQAVEFAHQGVFFHQG KVAFTGSMEVGKMIKEAAGKSNLKRVTLELGGKSPCIVFADADLDHAVEFAHHGVFAHQG KVAFTGSTEVGKLIKEAAGKSNLKRVTLELGGKSPCIVFADADLESAVEFAHQGVFLHQG KVAFTGSTEVGKLIKEAAGKSNLKRVTLELGGKSPCIVFADADLDHAVEFAHQGVFFNQG KVAFTGSTEVGKLIQEAAGKSNLKRVSLELGGKSPCIVFADADLDHAVEFAHQGVFFHQG KVAFTGSTEVGKLIKEAAGKSNLKRVTLELGGKSPCIVFADADLDHAIEFAHQGVFFNQG KVSFTGSTEVGKLIKEAAGKSNLKRVTLELGGKSPCIVFADADLDSAVEFAHQGVFFHQG KVAFTGSTQVGKLIKEAAGKSNLKRVTLELGGKSPCIVFADADLDIAVEFAHHGVFYHQG KVAFTGSTEVGKLIKEAAGKSNLKRVTLELGGKSPCIVLADADLDNAVEFAHHGVFYHQG

QCCVAASRLFVEESIYDEFVRRSVERAKKYVLGNPLTPGITQGPQIDKEQYEKILNLIES QMCIAASRLFVEESIYDEFVRRSVERAKQYILGNPLTPGITQGPQIDKEQHNKILDLIES QICIAASRLFVDESIYDEFVRRSVERAKQYILGNPLTPGITQGPQIDKEQYNKILDLIES QICVAASRLFVEESIYDEFVRRSVERAKQYILGNPLTPGITQGPQIDKEQHNKILDLIES QICVAASRLFVEESIYDEFVRRSVERAKQYVLGNPLTPGITQGPQIDKEQHNKILDLIES QMCVAASRLFVEESIYDEFVRRSVERAKQYILGNPLTPGITQGPQIDKEQHNKILDLIES QICVAASRLFVEESIYDEFVRRSVERAKQYVLGNPLTPGITQGPQIDKEQHNKILDLIES QICVAASRLFVEESIYDEFVRRSVERAKKYILGNPLNSGINQGPQIDKEQHNKILGLIES QCCVAASRIFVEESVYDEFVKRSVERAKKYVLGNPLTPGINQGPQIDKEQHDKILDLIES QCCIAASRIFVEESIYDEFVRRSVERAKKYILGNPLTPGVTQGPQIDKEQYDKILDLIES



Figure S5. Related to Figure 2. Pseudogene identification pipeline.


1157 Figure S6. Related to Figure 2. Phylogenetic tree of beaver Aldh. The genome coordinates of 1158 each gene are given after the gene names.


Figure S7. Related to Figure 2. Phylogeny of beaver, mouse, and human ALDH. The letter prefix of the gene name denotes the species: B for beaver, M for mouse, and H for human. Gene annotation of each beaver Aldh gene was manually checked, and manually editing was done by Apollo(Lee, Helt et al. 2013), according to RNA-Seq reads alignment, when it was necessary. Seq reads coverage. C) RNA-Seq reads coverage at unique sites.

A


B
0.3 (Beaver) MVLEKLNPMDYNVTSMVLEVVPVGAMPLLLLLTGILLLIWNYENTSSIPGPAYCLGIGPII 1.25 (Beaver) MVLEKLNPMHYNVTSMVPEVVPVGAMPLLLLTGGILLLIWNYENTSSIPGPAYCLGIGPLI Cyp19a1 (Mouse) MFLEMLNPMQYNVTIMVPETVTVSAMPLLLIMGLLLLIWNCESSSSIPGPGYCLGIGPLI CYP19A1 (Human) MVLEMLNPIHYNITSIVPEAMPAATMPVLLLTGLFLLVWNYEGTSSIPGPGYCMGIGPLI
0.3 (Beaver)
1.25 (Beaver)

Cyp19a1 (Mouse)
CYP19A1 (Human)
0.3 (Beaver)
1. 25 (Beaver)

Cyp19a1 (Mouse)
CYP19A1 (Human) GLQCIGMHENGIIFNNNPSLWRTIRPFFMKALTGPGLVRMVEVCVESIKQHLDRLGEVTD GLQCIGMHEKGI IFNNNPELWKTTRPFFMKALSGPGLVRMVTVCAESLKTHLDRLEEVTN

ALGYVDVLTLMRRIMLDTSNVVFLGIPLD-----------------------------------------ALGYVDVLTLMRRIMLDTSNVLFLGIPLDESAIVNKIRGYFDAWQALLLKPNIFFKISWL TSGYVDVLTLMRHIMLDTSNMLFLGIPLDESAIVKKIQGYFNAWQALLIKPNIFFKISWL ESGYVDVLTLLRRVMLDTSNTLFLRIPLDESAIVVKIQGYFDAWQALLIKPDIFFKISWL

YKKYEKSVKDLKDAIDVLVEEKRHQVSMAEKLEDCMDFATDLIFAEKRGELTKENVNQCI YRKYERSVKDLKDEIAVLVEKKRHKVSTAEKLEDCMDFATDLIFAERRGDLTKENVNQCI YKKYEKSVKDLKDAIEVLIAEKRRRISTEEKLEECMDFATELILAEKRGDLTRENVNQCI
-----------------------------------------GERDIQMNDIQKLKVVENFI LEMLIAAPDTMSVTVYFMLSLIAKHPNVEEEIMKEIQTVVGERDIQMNDIQKLKVVENFI LEMLIAAPDTMSVTLYFMLLLVAEYPEVEAAILKEIHTVVGDRDIKIEDIQNLKVVENFI LEMLIAAPDTMSVSLFFMLFLIAKHPNVEEAIIKEIQTVIGERDIKIDDIQKLKVMENFI

YESLRYHPAVDLVMRRALEDDVIDGYPVKKGTNIILNIGRMHRLEYFPKPNEFTLENFEK YESLRYHPAVDLVMRRALEDDVIDGYPVKKGTNIILNIGRMHRLEYFPKPNEFTLENFEK NESMRYQPVVDLVMRRALEDDVIDGYPVKKGTNIILNIGRMHRLEYFPKPNEFTLENFEK YESMRYQPVVDLVMRKALEDDVIDGYPVKKGTNIILNIGRMHRLEFFPKPNEFTLENFAK

NVPYRYFQPFGFGPRGCAGKYIAMVMMKVVLVTLLRRYHVETLRGQCVEDIQKVNDLSVH NVPYRYFQPFGFGPRGCAGKYIAMVMMKVVLVTLLRRYHVETLRGQCVEDIQKVNDLSVH NVPYRYFQPFGFGPRGCAGKYIAMVMMKVVLVTLLRRFQVKTLQKRCIENIPKKNDLSLH NVPYRYFQPFGFGPRGCAGKYIAMVMMKAILVTLLRRFHVKTLQGQCVESIQKIHDLSLH

PDETSDLLEMVFIPRNSVKCLNQ-
PDETSDLLEMVFIPRNSVKCLNQPNEDRHLVEIIFSPRNSDKYLQQ* PDETKNMLEMIFTPRNSDRCLE--
\begin{tabular}{ll}
1.25 (Beaver) & PDETSDLLEMVFIPRNSVKCLNQ \\
Cyp19a1 (Mouse) & PNEDRHLVEIIFSPRNSDKYLQQ* \\
CYP19A1 (Human) & PDETKNMLEMIFTPRNSDRCLE--
\end{tabular}
cyp19A1 (Human)
SHSRFLWMGIGSACNYYNKMYGEFMRVWISGEETLIISKSSSMFHVMKHNKYISRFGSKL SHGRFLWMGIGSACNYYNKMYGEFMRVWISGEETLIISKSSSMFHVMKHNKYISRFGSKL SHGRFLWMGIGSACNYYNKMYGEFMRVWISGEETLIISKSSSMFHVMKHSHYISRFGSKR SHGRFLWMGIGSACNYYNRVYGEFMRVWISGEETLIISKSSSMFHIMKHNHYSSRFGSKL

GLQCIGMHEKGI IFNNNPDLWKEVRLFFMKALTGPGLVRMVAVCAESILRHLDKLDEVTS GLQCIGMHEKGI IFNNNPDVWKEVRPFFMKALTGPGLVRMVAVCAESILRHLDKLDEVTN
0.3 (Beaver)
1.25 (Beaver)

Cyp19a1 (Mouse)
CYP19A1 (Human)
0.3 (Beaver)
1.25 (Beaver)

Cyp19a1 (Mouse)
CYP19A1 (Human)
0.3 (Beaver)
1.25 (Beaver)

Cyp19a1 (Mouse)
CYP19A1 (Human)
0.3 (Beaver)
1.25 (Beaver)

Cyp19a1 (Mouse)
CYP19A1 (Human)
0.3 (Beaver)
1.25 (Beaver)

Cyp19a1 (Mouse)
CYP19A1 (Human)
0.3 (Beaver)
\(\qquad\)
\begin{tabular}{lllll}
\hline Gene & B1 & B2 & B3 & B4 \\
\hline 0.3 & 8 & 9 & 0 & 5 \\
1.25 & 24 & 419 & 0 & 0 \\
\hline
\end{tabular}

The amino acid letters with grey background (panel b) show where we checked the RNA-Seq reads coverage on the corresponding genome loci.
Figure S8. Related to Figure 4. Duplication of Cyp19a1. A) Gene structure of beaver Cyp19a1.

Coding Sequence
68.67 (Beaver) 68.66 (Beaver)

> aaaaccctgcttgaaacaccttcctccaggagactgaggagccaaacagagacgactgcc AAAACCCTGCTTGAAACACCTTCCTCCAGAAGACTGAGGAGCCAAACAGAGACGACTGCC

MADFYSSDSELTTRTLLRRVLDTADSLTPRRRRSAQAGAQKTLLETPSSRRLRSQTETTA MADFYSSDSELTTRTLLRRVLDTADSLTPRRRRSAQAGAQKTLLETPSSRRLRSQTETTA MADLSFSDGDPTVRTLLRRVLETADSRTPMRRRSTRINAQRRRSQTPYSNRQGSQTKTSA MAD-HNPDSDSTPRTLLRRVLDTADPRTPRRPRSARAGARRALLETASPRKLSGQTRTIA

RQRSHGARSIGRLAHGQASGSLEEKTPRTLLRNILLTAPESSIVMPDSVVKPVSAPQVVQ RQRSHGARSIGRLAHGQASGSLEEKTPRTLLRNILLTAPESSIVMPDSVVKPVSAPQVVQ RKQSHGARSVGRSTRVQGRGRLEEQTPRTLLRNILLTAPESSTVMPDPVVKPAQVPEVAR RGRSHGARSVGRSAHIQASGHLEEQTPRTLLKNILLTAPESSILMPESVVKPVPAPQAVQ

SSRRKSSRGSLELQLPELEPPSTLAPGLPALGRRKQRLRLSVFQQEVNQELPLSQEPSG-SSRRKSSRGSLELQLPELEPPSTLAPGLPALGRRKQRLRLSVFQQEVNQELPLSQEPSGSSRRESSRGSLELHLPELEPPSTLAPGLTAPGKRKQKLRLSVFQQEVDQGLPLSQEPRRS PSRQESSCGSLELQLPELEPPTTLAPGLLAPGRRKQRLRLSVFQQGVDQGLSLSQEPQG-
-NADASALTSSLNLTFATPLQPQSVRRPGLARRPPTRRAVDVGALLQDLRDNSLAS---G -NADASALTSSLNLTFATPLQPQSVRRPGLARRPPTRRAVDVGALLQDLRDNSLAS---G RSADVSSLASSFNLTFVLPGQPETVERPGLARRRPIRQLVNAGALLQDLEDNSLASALPG -NADASSLTRSLNLTFATPLQPQSVQRPGLARRPPARRAVDVGAFLRDLRDTSLA-----

NSHRTPAAALPTDTVLEDTQPFSQPLVGCSPSLHDSLPLPTHTAVEDSERAVGHRTRSRG NSHRTPAAALPTDTVLEDTQPFSQPLVGCSPSLHDSLPLPTHTAVEDSERAVGHRTRSRG DSHRTPVAALPMDVGLEDTQPFSQSLAAFSLSGKHSLPSPSRPGVEDVERVMGPPSS--G -----------PPNIVLEDTQPFSQPMVGSP-NVYHSLPCTPHTGAEDAEQAAGRKTQSSG

PRLQNHKL--Y---P-------------------WPTTSQEVTE----------GEGFREAEVA PRLQNHKL--Y---P-------------------WPTTSQEVTE-----------GEGFREAEVA
 PGLQKNSP--G---KPAQFLAGEAEEVNAFALGFLSTSSGVSGEDEVEPLHDGVEEAEKK
----------------------------------KELEGSSGDEDTS-----DRPSKRPIAEIS -----------EAVG----------SNEAAEPKDQEGSSGYEETS-----------------ARPA MEEEGVSVSEMEATGAQGPSRVEEAEGHTEVTEAEGSQGTAEADGPGASSGDEDASGRAA
-------------------------------NLPSEPLEPMLARLPRRPRTAGPRPRQDPY CPELASSTPEFLQAKQPHEFLDPTPSLGVTILPSEPLEPMLARLPRRPRTAGPRPRQDPY SGELSSSTHDSLP------AEQPPPSPGVAVLSSEPLESVTAKCPSRTQTAGPRRRQDPH SPESASSTPESLQARRHHQFLEPAPAPGAAVLSSEPAEPLLVRHPPRPRTTGPRPRQDPH

KAGLSHYAKLFSFYAKMPMEKAAMEMVEKCLDKYFQHLCNDLEVFAAHAGRKTVRPEDLE KAGLSHYAKLFSFYAKMPMEKAAMEMVEKCLDKYFQHLCNDLEVFAAHAGRKTVRPEDLE KAGLSPYVKFFSFCTKMPVEKTALEIVEKCLDKYFQHLCNDLEVFASHAGRKIVKPEDLL KAGLSHYVKLFSFYAKMPMERKALEMVEKCLDKYFQHLCDDLEVFAAHAGRKTVKPEDLE

LLMRR--------LLHVLVERHLPLEYRQLLIPCAFSGNSVFPAQ-LLMRRQGLVTDQVSLHVLVERHLPLEYRQLLIPCAFSGNSVFPAQLLMRRQGLVTDQVSQHVLVERYLPLEYRQQLIPCAFSGNSVFPAQ* LLMRRQGLVTDQVSLHVLVERHLPLEYRQLLIPCAYSGNSVFPAQ*

\section*{Protein sequence} 68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)
68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)
68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)
68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)
68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)
68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)
68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)
68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)
68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)
68.67 (Beaver) 68.66 (Beaver) Cenpt (Mouse) CENPT (Human)


C
\begin{tabular}{lllll}
\hline Gene & B1 & B2 & B3 & B4 \\
\hline 68.67 & 18 & 21 & 5 & 16 \\
68.66 & 8 & 5 & 0 & 0 \\
\hline
\end{tabular}

The amino acid letters with grey background (panel B) show where we checked the RNA-Seq reads coverage on the corresponding genome loci. Although there are no difference on amino acid sequence, the coding sequences show differences.
Figure S9. Related to Figure 4. Duplication of Cenpt. A) Gene structure of beaver Cenpt. B) Protein sequence alignment. Sequences with grey background show where we check the RNASeq reads coverage. C) RNA-Seq reads coverage at unique sites.

1311 C
\begin{tabular}{lllll}
\hline Gene & B1 & B2 & B3 & B4 \\
\hline 10.4 .2 & 268 & 136 & 579 & 122 \\
0.2 & 70 & 31 & 265 & 34 \\
\hline
\end{tabular}

The amino acid letters with grey background (panel B) show where we checked the RNA-Seq reads coverage on the corresponding genome loci.

\section*{A}

B
10.4.2 (Beaver) CWGDQPLLCCPAEQAVSGEGREHGAGAVVGARAQIQALLGCLVKVLLWVASALLYFGSEQ
0.2 (Beaver)
------------------MERGPVVG----ARAQIQALLGCLVKVLLWVASALLYFGSEQ
Fitm1 (Mouse) ------------------MERGPTVGAGLGAGTRVRALLGCLVKVLLWVASALLYFGSEQ
FITM1 (Human) ------------------MERGPVVGAGLGAGARIQALLGCLLKVLLWVASALLYFGSEQ
10.4.2 (Beaver) AARLLGSPCLRRLYHAWLAAVVIFGSLLQFHVNSRTIFASHGNFFNIKFVNSAWGWTCTF
0.2 (Beaver) AARLLGSPCLRRLYHAWLAAVVIFGPLLQFHVNSRTIFASHGNFFNIKFVNSAWGWTCTF
Fitm1 (Mouse) AARLLGSPCLRRLYHAWLAAVVIFGPLLQFHVNSRTIFASHGNFFNIKFVNSAWGWTCTF
FITM1 (Human) AARLLGSPCLRRLYHAWLAAVVIFGPLLQFHVNPRTIFASHGNFFNIKFVNSAWGWTCTF
10.4.2 (Beaver) LGGFVLLVVFLATRRVAVTARHLSRLVVGAACGEGPA-GLPPHEDLTGSCFEPLPQGLLL
0.2 (Beaver) LGGFVLLVVFLATRRVAVTARHLSRLVVGAAVWRGAGRAFLLIEDLTGSCFEPLPQGLLL
Fitm1 (Mouse) LGGFVLLVVFLATRRVAVTARHLSRLVVGAAVWRGAGRAFLLIEDLTGSCFEPLPQGLLL
FITM1 (Human) LGGFVLLVVFLATRRVAVTARHLSRLVVGAAVWRGAGRAFLLIEDLTGSCFEPLPQGLLL
10.4.2(Beaver) HELPDRRSCLAAGHQWRGYTVSSHTFLLTFCCLLMAEEAAVFAKYLAHGLPAGTPLRLVF
0.2 (Beaver) HELPDRRSCLAAGHQWRGYTVSSHTFLLTFCCLLMAE---------------------------
Fitm1 (Mouse) HELPDRKSCLAAGHQWRGYTVSSHTFLLTFCCLLMAEEAAVFAKYLAHGLPAGAPLRLVF
FITM1 (Human) HELPDRRSCLAAGHQWRGYTVSSHTFLLTFCCLLMAEEAAVFAKYLAHGLPAGAPLRLVF
10.4.2(Beaver) LLNVLLLGLWNFLLLCTV----------VGAAVGTFAWFLTYGSWYHQPWSPGSPGH
0.2 (Beaver)
\(\begin{array}{ll}\text { Fitm1 (Mouse) } & \text { LLNVLLLGLLWNFLLLCTVIYFHQYTHKVVGAAVGTFAWYLTYGSWYHQPWSPGIPGHGLF } \\ \text { FITM1 (Human) } & \text { LLNVLLLGLWNFLLLCTVIYFHYTHKVVGAAVGTFAWYLTYGSWYHQPWSPGSPGHGLF }\end{array}\)
10.4.2 (Beaver) PHPHSSRKHN-
0.2 (Beaver) -----------
Fitm1 (Mouse) PRSRSMRKHN*
FITM1 (Human) PRPHSSRKHN*

A


\section*{B}
14.34 (Beaver)
14.18 (Beaver)
13.30 (Beaver)
13.14 (Beaver) Pla2g4c (Mouse) PLA2G4C (Human)
14.34 (Beaver)
14.18 (Beaver)
13.30 (Beaver)
13.14 (Beaver)

Pla2g4c (Mouse)
PLA2G4C (Human)
14.34 (Beaver)
14.18 (Beaver)
13.30 (Beaver)
13.14 (Beaver)

Pla2g4c (Mouse) PLA2G4C (Human)
14.34 (Beaver)
14.18 (Beaver)
13.30 (Beaver)
13.14 (Beaver)

Pla2g4c (Mouse) PLA2G4C (Human)
14.34 (Beaver)
14.18 (Beaver)
13.30 (Beaver)
13.14 (Beaver)

Pla2g4c (Mouse)
PLA2G4C (Human)
14.34 (Beaver)
14.18 (Beaver)
13.30 (Beaver)
13.14 (Beaver)

Pla2g4c (Mouse
PLA2G4C (Human)
14.34 (Beaver)
14.18 (Beaver)
13.30 (Beaver)
13.14 (Beaver)

Pla2g4c (Mouse)
PLA2G4C (Human)
14.34 (Beaver)
14.18 (Beaver)
13.30 (Beaver)

MSCEQSPKSLLSKKNRSDGVHITPELQKEEKEAVDNRKTQVVMSLNKLGIKADEAPVLAV
--------------------------------------------------------------------1
------------------HGVLITPELQKEEKEAVDNRKPQVVMSLNKLGIKADEAPVIAV MSHEQSPKSLLSKKD--TRVRITPELQKEEKEAVDNRKLQVVMSLNKLGIKADEAPVIAV MSCAESPKSLH---KRSSGVCPATRLQEAEKAAVHKRSPKVLEALRKLNIQADQAPVIAV MRTRPRPR--LRR--TENFLTAVHHGKKEEKAAVERRRLHVLKALKKLRIEADEAPVVAV

LGSGGGLRAHFACLGVLIEMKNHGLLDVITYLAGVSGSTWALSSFYTNSGNMEHIEADLE ---------------------------------------ALSSFYTNSGNVEHIEAVLE LGSGGGLRAHFACLGVLIEMKNHGLLDVITYLAGVSGSTWALSSFYTNSGNMDLIEADLE LGSGGGLRAHFACLGVLSEMKKHGLLDVITYLAGVSGSTWALSSFYTNSGNMEHIEADLE LGSGGGLRAHIACLGVLSELKELGLLDAVTYLAGVSGSTWALSSLYTKNGNMEGIEEELK LGSGGGLRAHIACLGVLSEMKEQGLLDAVTYLAGVSGSTWAISSLYTNDGDMEALEADLK

HRFELENWSIWESLQKTIEAASLENYSLTDFWAYIVVSRQTREFQGSLLSSMKKHVEKGT HRFDPENWS IWDSLQKTIEAASLENYSLTDFWAYIVVSRQTREFQDSLLSS IKKHVEKGT HRFEPENWSVRESLQKTIEVASLENYSLTDFWAYVVISRQTREFQGSLLSSMKKHVEKGT HRFDPENWSIWDSLQKTIEAASLENYSLTDFWAYIVVSRQTREFQDSLLSSIKKHVEKGT HRYEKNEWDFHESLEKAIQASKRENYSLTDFWAYLIVSRQIRELQDSNLSSLKKQVEEGV HRFTRQEWDLAKSLQKTIQAARSENYSLTDFWAYMVISKQTRELPESHLSNMKKPVEEGT

LPYPIFAAIDNDLHPGWKDQKTQKSWFEFTPHHAGYPALKSYIPITQFGSQFENGRLVKS LPYPIFAAIDNDLHPVWKDHKTRKSWFEFTPHHAGYPALQAYIPITQFGSQFENGRLVKS LPYPIFAAIDNDLHDDWKDHKTQKSWFEFTPHHAGYPALQAYIPITQLGSQFENGRLVKS LPYPIFAAIDNDLHPVWKDHKTRKSWFEFTPHHAGYPALKAYIPITQFGSQFQNGRLVKS LPYPIFAAIDEDLLADWRERKTQNSWFEFTPHHAGYPALGAYVPITEFGSRFENGKLVKS LPYPIFAAIDNDLQPSWQEARAPETWFEFTPHHAGFSALGAFVSITHFGSKFKKGRLVRT

ALERDLSFLRGLWGSAVANTEENKKFIWG-------------------MIE----KEITVD APERDLSFLRGLWGSAIANTEENKKFIWDEFLSLKEKLLGKHQLTQGMIT----EETAVD APERDLSFLRGLWGSAVANAEENEKFIWDEFLSLKEKLLGKYQLTQGMIT----EETAVD VPERDISFLR------------------DEFLSLKDKLLGKHQLTQGMIT----EETAVD EPERDLTFLRGLWGSAFADIKEIKNYILNYFRNP----FGKLKFIEGPVTYSEAPRMNVD HPERDLTFLRGLWGSALGNTEVIREYIFDQLRNLTLKGLWRRAVA---------NAKSIG

EALLELTVDYIKDEKDPSIQKKLQALQQALDAGRDEHGEPECRKVAMMIQNWSNASQKEQ EALLELMVAYIKDEKDPSIQKKLQALQQALGARRGKQGEPEYRELAMMIRNWSKASLQER EALLELMVAYIKDEKDPSIQEKLQALQQILDARRDEHGEPEYKKLAMMIQNWSNASQEKQ EALLELMVAYIKGEKDSSIQKKLQALQQALDARKGKQGEPEYRELAMMIRNWSKASLQER AMLLDLVMAYFTDMNDPSIKDKLCALQQALGTETDEFGI----EMAEIIQNWNETSAEKK ----HLIFARLLRLQ--------ESSQGEHPPPEDEGGEPEHTWLTEMLENWTRTSLEKQ

GLILETLVCHFTGQASTMALSTALSVSK--------------------------------------GQILETLVGHFTRQASTMAVSRALSVYRVTFWDILDFLAKTVMCIWNWEWGTVHNFLLKL

 EQFLDHLLDRFKKTQEDTTTYSLMNWNTGLVWDRCVFVNETRKCVSKWQWGTVYNFLYK-EQPHEDPE------------------RKGSLSNLMDFVKKTGICASKWEWGTTHNFLYK-
-CSNVCITDEDMCSRKLLHLVDAGLAINSPYPLLLPPAREVQLILSFDFSDGDPFETVRA YGSRVGVTDDEMCSRKLLHLVDAGLAINSPYPLLLPPAREVQLILSFDFSAGDPFETVRA -CSNISITDEDMCSRKLLHLVDAGLAINSPYPLLLPPAREVQLILSFDFSDGDPFETVRA
\begin{tabular}{|c|c|}
\hline 13.14 (Beaver) & -GSRVGVTDNEMYSRKLLHLVDAGLAINSPYPLLLPPAREVQLILSFDFSAGDPFETVRA \\
\hline Pla2g4c (Mouse) & --HGKIADETMCSRELLHLVDAGFAINTPYPLVLPPVRETHLILSFDFSAGDPLETIRA \\
\hline PLA2G4C (Human) & ---HGGIRDKIMSSRKHLHLVDAGLAINTPFPLVLPPTREVHLILSFDFSAGDPFETIRA \\
\hline 14.34 (Beaver) & TADYCHHHKIPFPLVKEADLKEWAEAPTSCYILKGESGPVVMHFPLFNKDNCGDDINTWR \\
\hline 14.18 (Beaver) & TADYCYRHKIPFPLVKEADLKEWAKAPSSCYILKGESGPVVMHFPLFNKDNCGDDINTWR \\
\hline 13.30 (Beaver) & TADYCHHHKIPFPLVKEADLKEWAEAPTSCYILKGESGPMVMHFPLFNKDNCGDEINTWR \\
\hline 13.14 (Beaver) & TADYCYRHKIPFPLVKEADLKEWAKAPSSCYILKGESGPVVMHFPLFNKDNCGGDISTWR \\
\hline Pla2g4c (Mouse) & TADYCQRHEIPFPEVSEDQLKEWAKAPASCYVLRGETGPVVMHFTLFNKDNCGDDIETWR \\
\hline PLA2G4C (Human) & TTDYCRRHKIPFPQVEEAELDLWSKAPASCYILKGETGPVVMHFPLFNIDACGGDIEAWS \\
\hline 14.34 (Beaver) & DKYGTFKLSDTYSVQLVKDLLEKSKENVRKNKEKIIRTIKEVVG \\
\hline 14.18 (Beaver) & DKYATFKLSDTYSVQLVKDLLEKSKENVRKNKENIFRAIKEAVRSCPQTS \\
\hline 13.30 (Beaver) & EKYGTFKLSDTYSVQVVKDLLEKSKENVRKNKEKIIRTIKEVVG \\
\hline 13.14 (Beaver) & DKYATFKLSDTYSVQVVKDLLEKSKENVRKNKEKIIRTIKEVVG \\
\hline Pla2g4c (Mouse) & KKYGTVKLSDSYTPDLVRDLLRVSKENVKKNKINILSEMRKVAGNPGNIPRVNKEACLGD \\
\hline PLA2G4C (Human) & DTYDTFKLADTYTLDVVVLLLALAKKNVRENKKKILRELMNVAGLYYPKD-SARSCCLA* \\
\hline 14.34 (Beaver) & \\
\hline 14.18 (Beaver) & \\
\hline 13.30 (Beaver) & --------------------- \\
\hline 13.14 (Beaver) & \\
\hline Pla2g4c (Mouse) & RVKDPQGSQTVEFKKSHNISKD* \\
\hline PLA2G4C (Human) & \\
\hline
\end{tabular}

\section*{C}
\begin{tabular}{lllll}
\hline Gene & B1 & B2 & B3 & B4 \\
\hline 13.14 & 0 & 1 & 3 & 0 \\
13.30 & 0 & 0 & 6 & 2 \\
14.18 & 0 & 3 & 10 & 102 \\
14.34 & 5 & 0 & 7 & 2
\end{tabular}

The amino acid letters with grey background (panel B) show where we checked the RNA-Seq reads coverage on the corresponding genome loci.

Figure S11. Related to Figure 4. Duplication of Pla2gc. A) Gene structure of beaver Pla2gc.
B) Protein sequence alignment. Sequences with grey background show where we check the RNA-Seq reads coverage. C) RNA-Seq reads coverage at unique sites. Consistent with genome sequencing result and qPCR also showed there are duplication of Pla2gc in beaver genome (Figure 4A). However, we did not observe 4 times signals by qPCR. It may be because the primer we designed do not perfectly match sequences from each of those copies. The exact copy number of Pla2gc in beaver genome need further exploration.

A



B

\begin{tabular}{lcllllc}
\hline \(\mathbf{C}\) & & & & & \\
\hline & Codon & & Num. of mammals & AF in humans & CADD \\
\hline Position & Human & Beaver & & \\
\hline 1 & A & T & 0 & 0 & 22.8 \\
2 & G & C & 0 & 0 & 22.4 \\
3 & T & G & 0 & 0 & 20.6 \\
\hline
\end{tabular}

1411

D
\begin{tabular}{ccccc}
\hline \multicolumn{2}{c}{ Amino acid (codon) } & \multirow{2}{*}{ RROVEAN } & PolyPhen2 & \multirow{2}{*}{ CADD } \\
\hline Human & Other & & \\
\hline & P (CCC) & -3.51 & 0.916 & 23.0 \\
& L (CTC) & -0.44 & 0 & 13.32 \\
I (ATC) & V (GTC; GTG) & -0.47 & 0.005 & \(12.83 ; 14.90\) \\
& M (ATG) & -1.14 & 0.005 & 14.33 \\
& T (ACT) & -2.28 & 0.275 & 19.95 \\
\hline & W (TGG) & -2.94 & 0.948 & 21.7 \\
& G (GGC; GGT) & -1.31 & 0.000 & \(11.21 ; 12.00\) \\
& T (ACT) & -0.09 & 0.000 & 0.779 \\
S (AGT) & C (TGT) & -1.81 & 0.001 & 13.56 \\
& H (CAT) & -1.83 & 0.614 & 13.07 \\
& N (AAC; AAT) & -0.81 & 0.000 & \(6.38 ; 4.25\) \\
& M (ATG) & -1.76 & 0.020 & 9.42 \\
\hline
\end{tabular}

Figure S12. Related to Figure 5. MTBP function and positive selection. (A) Functional interaction among MTBP and other genes related to longevity or cancer. MTBP enhances MDM2 stability by inhibiting auto-ubiquitination of MDM2, which in turn promotes TP53 degradation through MDM2-mediated ubiquitination of TP53. MTBP and MYC are transcriptional co-factors. Decreased expression of either MTBP and MYC show increased longevity and enhanced healthspan(Hofmann, Zhao et al. 2015, Grieb, Boyd et al. 2016). (B) Alignment of 62 mammalian species at the three sites likely under positive selection in beavers. These three sites have selection probabilities higher than \(88 \%\). The alignment were extracted from 100-way vertebrates' alignments. (C) One codon (S:TCG) under positive selection in beavers. It is a synonymous change (S/S: TCG/AGT) between human and beaver. 'Num. of mammals' is the number of mammalian species (among the 62 in the phylogeny) with the same nucleotide as beavers. 'AF in humans' is the frequency of the allele in humans identical to the nucleotide in beavers from gnomAD (Karczewski, Francioli et al. 2019). The CADD score quantifies the deleteriousness of the sequence change between human and beaver at each nucleotide position in the codon. A CADD score greater than 20 indicates a very likely deleterious change. (D) Other two codons (P:CCC and W:TGG) under positive selection in beavers. They are both non-synonymous change (P/I: CCC/ATC and W/S: TGG/AGT) between human and beaver. We predicted the functional effects of the nonsynonymous changes between human and other mammalian species including beaver. Green letters are nucleotide changes from the human codons. PROVEAN scores lower than -2.5 indicate deleterious changes, which Polyphen2 scores between 0.85 to 1.0 indicate damaging changes.


Figure S13. Related to Figure 6. Comparison of Igf2 and Igf2bp2 expression between beavers and mice.

A


B


Figure S14. Related to Figure 6. Principle component analysis of RNA-Seq data from beavers and mice. (A) Liver. (B) Brain.

Table S1. RNA-Seq data from four beaver samples.
\begin{tabular}{llllll}
\hline Sample & Source & Sex & Age & Tissue & Num. of reads \\
\hline B1 & Wild-caught & Male & Young adult & Brain & \(23,287,737\) \\
& & & & Liver & \(22,152,280\) \\
B2 & Wild-caught & Male & Young adult & Brain & \(22,888,968\) \\
& & & & Liver & \(23,371,745\) \\
B3 & Toronto Zoo & Male & \multirow{2}{*}{10 years } & Leukocyte & \(120,041,781\) \\
& & & & NA & Moscle
\end{tabular}

B1 \& B2 are from our own study. B3 was from (Lok, Paton et al. 2017). B4 was from the beaver genome project in Oregon University who generated RNA-seq data from pooled 16 tissues: skeletal muscle, kidney, spleen, ovaries, placenta, castor gland, tail, toe webbing, whole blood, brain, lung, liver, heart, stomach, tongue, intestine.

Table S2. qPCR primers.
\begin{tabular}{lll}
\hline \begin{tabular}{l} 
Gene \\
target
\end{tabular} & Forward primer Sequence \(\left(5^{\prime} \rightarrow 3^{\prime}\right)\) & Reverse primer Sequence \(\left(5^{\prime} \rightarrow 3^{\prime}\right)\) \\
\hline Aldh1al & GACAGGCTTTCCAGATTGGTTCTC & CTAGCAGCAGACGATCTCTTTCAAT \\
Cenpt & TTCAGAGCCTTTGGAGCCTAT & GTCCAGCCTTGTAGGGATCTT \\
Cyp19al & CATGTGGAGACATTGCGAGG & ATTTCCAGCAGGTCACTCGT \\
Fitm1 & TCTGCCTTGCTGTATTTTGGA & ATGGTAGAGGCGCCGTAAG \\
Hpgd & TGAAGATGATGGCTTAAATGGTGCT & GAGATGGAGGTGGGTCATACTC \\
Pla2g4c & TCCAGGGCTCTCTTTTGTCCAG & AACCAGGGTGAAGGTCATTATCAA \\
Hcfcl & CCGTTAGTCACCATGCGACC & GTGGGCACAACCATTCGCAC \\
Pelo & TCCACTATCCGCAAGGTTCAG & GGCTTGGGAGTCAAAGTCGAT \\
\hline
\end{tabular}```

