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Classification: Biological sciences; Genetics

Title: Genome sequence of a diabetes-prone rodent reveals a mutation hotspot around the ParaHox gene cluster

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Keywords: Desert rodent, type 2 diabetes, homeobox, Pdx1, biased gene conversion

Abstract:

The sand rat *Psammomys obesus* is a gerbil species native to deserts of North Africa and the Middle East, and is constrained in its ecology because high carbohydrate diets induce obesity and type II diabetes which, in extreme cases, can lead to pancreatic failure and death. We report the sequencing of the sand rat genome and discovery of an unusual, extensive and mutationally-biased GC-rich genomic domain. This highly divergent genomic region encompasses several functionally essential genes, and spans the ParaHox cluster which includes the insulin-regulating homeobox gene *Pdx1*. The sequence of sand rat *Pdx1* has been grossly affected by GC-biased mutation leading to the highest divergence observed for this gene across the Bilateria. In addition to genomic insights into restricted caloric intake in a desert species, the discovery of a localised chromosomal region subject to elevated mutation suggests that mutational heterogeneity within genomes could influence the course of evolution.

Significance statement:

A core question in evolutionary biology is how mutation and selection adapt and constrain species to specialized habitats. We sequenced the genome of the sand rat, a desert rodent susceptible to nutritionally-induced diabetes, and discovered an unusual chromosome region skewed towards G and C nucleotides. This region includes the *Pdx1* homeobox gene, a transcriptional activator of *insulin*, which has undergone massive sequence change likely contributing to diabetes and adaptation to low caloric intake. Our results imply that mutation rate varies within a genome and that hotspots of high mutation rate may influence ecological adaptation and constraint. In addition, we caution that divergent regions can be omitted by conventional short-read sequencing approaches, a consideration for existing and future genome sequencing projects.

Introduction

Arid environments impose extreme physiological demands on animals because of low food and water availability. The sand rat *Psammomys obesus* (Fig. 1a) is a member of the subfamily Gerbillinae, most species of which live in deserts and arid environments (Fig. 1b). *P. obesus* has emerged as a model for research into diet-induced type II diabetes because, if provided with high carbohydrate diets, the majority of individuals become obese and develop classic diabetes symptoms, in the most extreme cases leading to pancreatic failure and death (1-4).

In searching for the molecular basis of this unusual phenotype, attention has been paid to the *Pdx1* homeobox gene, also called *Ipf1*, *Idx1*, *Stf1* or *Xlox* (5-9), the central and most highly conserved member of the ParaHox gene cluster (10). *Pdx1* is the only member of the Pdx gene family in tetrapods, and encodes a homeodomain that has been invariant across their evolution. Mammalian *Pdx1* is expressed in pancreatic beta-cells and encodes a homeodomain transcription factor that acts as a transcriptional activator of *insulin* and other pancreatic hormone genes (11,12). A pivotal role in insulin regulation is also reflected in the association of heterozygous *Pdx1* mutations with maturity-onset diabetes of the young (*MODY4*) and type II diabetes mellitus in humans (13). Contrary to the usual conservation, several studies have reported inability to detect *Pdx1* in multiple gerbil species, including *P. obesus*, by immunocytochemistry, Western blotting or PCR. However, *Pdx1* is readily detectable in the closely related spiny mouse, *Acomys cahirinus* (Fig. 1b) leading to the hypothesis that the gene has been lost within the Gerbillinae subfamily, contributing to the compromised ability to regulate insulin in the sand rat (14-16). Such a conclusion would raise further questions, since in addition to its adult functions, *Pdx1* is also essential for pancreatic development in the embryo. For example, targeted deletion in mice causes loss of pancreas and anterior duodenum and is lethal (17,9). In humans, pancreatic agenesis has been reported in a patient with a homozygous frameshift mutation before the *Pdx1* homeobox, and in a compound heterozygous patient with substitution mutations in helices 1 and 2 of the homeodomain (18-20).

Results

To resolve the conundrum of a putatively absent 'essential' gene, we sequenced the *P. obesus* genome using a standard shotgun strategy (Illumina), using a combination of short and long insert libraries, initially at 85.5X coverage (SI Appendix, section 1). This assembly lacked a *Pdx1* gene supporting the prevailing hypothesis of a loss of the *Pdx1* gene in gerbils. However, a synteny comparison between *P. obesus* and other mammals delineated a contiguous block of 88 genes (SI Appendix, Fig. S2) missing from the assembly including several genes essential to basic cellular functions, such as *Brca2* and *Cdk8*, in addition to *Pdx1*. This led us to suspect that standard short read sequencing may have given an incomplete genome assembly, even at high coverage. To resolve whether this represented a large-scale deletion or an unusual genomic region, we sequenced the transcriptomes of *P. obesus* liver, pancreatic islets and duodenum, which contained transcripts for many of the missed genes (SI Appendix, Tables S4-S6). Furthermore, these transcripts show unusually high GC content in most cases, indicating that a large contiguous stretch of elevated GC had either been under-represented in initial sequencing data or had failed to assemble correctly, most likely due to nucleotide compositional bias. We term such cryptic or hidden sequence 'dark DNA'. We therefore isolated GC-rich *P. obesus* genomic DNA by Caesium Chloride gradient centrifugation, sequenced this fraction after limited amplification using Illumina MiSeq overlapping paired-end reads, and re-assembled the genome incorporating this longer-read sequence data (SI Appendix, section 1.5). This gave a refined assembly with a total size of 2.38 Gb and a scaffold N50 of 10.4 Mb (Table 1; SI Appendix, sections 1,3,4,6), including much of the 'dark DNA' region in several scaffolds, and contains genes syntenic to a region of chromosome 12 in rat and a region of chromosome 5 and the subtelomeric region of chromosome 8 in mouse. Analysis indicates that the region was initially omitted by standard genome assembly methods due to lower read coverage of GC regions coupled with short sequence read lengths. Comparison of GC content between species demonstrates that sand rat genes are elevated in GC content across this chromosomal region, syntenic to 12 Mb of the rat genome (Fig. 1c; SI Appendix, section 9). This large region encompasses a 250 kb repeat-rich scaffold containing the sand rat ParaHox cluster and its well-characterised genomic neighbours. We inferred a high W (weak, A/T) to S (strong, G/C) allelic mutation rate in this region of the *P. obesus* genome when compared with randomly selected genomic regions or homologous regions in other species of rodent (Fig 1d; SI Appendix, section 12; SI Appendix, Tables S11, S12). The existence of a localised GC-biased stretch of the *P. obesus* genome is striking and of far-reaching importance, and implies the existence of elevated and biased mutational pressure, acting in one region of a mammalian genome. Gene conversion, caused by the non-reciprocal exchange of information during meiosis, is the best characterised process known to cause GC-biased mutation (21).

The full coding sequence of the *P. obesus Pdx1* gene was deduced from the refined genome and transcriptome assemblies, and the gene was found to be expressed in sand rat pancreatic islets and duodenum (SI Appendix, section 7). The 60 amino acid homeodomain of Pdx1 shows

100% conservation across other mammals for which data are available; however, in *P. obesus* there are a remarkable 15 amino acid differences in the homeodomain, making this by far the most divergent *Pdx1* gene discovered in the Bilateria (Fig. 2a). All but one of the amino acid changes are caused by A/T to G/C mutation. The N-terminal and C-terminal regions are also divergent with numerous deletions, although the hexapeptide motif used in heterodimer formation with TALE proteins is conserved (Fig. 2b). Additional RNA sequencing of Mongolian jird (*Meriones unguiculatus*) duodenum reveals that extensive sequence divergence due to GC-biased mutation in *Pdx1* is not unique to sand rat (Fig. 2a). Analysis of synonymous and non-synonymous mutations in *Pdx1* across vertebrates reveals a dN/dS ratio of 2.6 (dN=39; dS=15) in the lineage leading to *P. obesus* and *M. unguiculatus* (SI Appendix, Fig. S10). High dN/dS ratios are often taken as evidence for positive selection, but can be skewed by mutational processes such as GC-biased gene conversion (22). Despite its radical divergence, *Pdx1* is the closest homeodomain by blastp and phylogenetic analysis places it as a rodent *Pdx1* on a long branch (Fig. 2c); extensive synteny with the ParaHox region of mouse and rat confirms it is the true and single *Pdx1* ortholog (SI Appendix, Table S9). Evidence that the locus is functional includes expression in pancreas and duodenum, and the fact that extensive polymorphism is found in the 3' untranslated region but is very limited in the coding sequence (SI Appendix, Fig. S11), indicating that the coding region is under functional constraint despite extensive mutation. Extreme deviation from the expected sequence explains why antibodies and PCR failed to detect *Pdx1* in sand rat, Mongolian jird and, potentially, other gerbil species (14-16).

These findings indicate that GC-biased mutation has driven radical changes in an otherwise highly conserved homeobox gene; these changes could be maladaptive and constrain the physiological capability of the sand rat, or adaptive enhancing ability to live in arid regions. To test if the extent of sequence divergence is unusual for sand rat proteins, we calculated a 'protein deviation index' (PDI) (SI Appendix, section 5) for all 1:1 mammalian orthologs by dividing mouse-human protein sequence identity by mouse-sand rat sequence identity (Fig. 2d). This is distinct from identifying the fastest evolving proteins, and specifically identifies proteins that have undergone uncharacteristic divergence in sand rat. We find the majority of sand rat proteins are highly similar to mouse or human (mode PDI = 1.0); in contrast, *Pdx1* is unusually divergent (mouse-sand rat 54.82%, mouse-human 91.37%; PDI = 1.67). To test if other genes implicated in glucose metabolism or pancreatic function are also divergent, we compiled a list of 45 candidates from human studies including all genes implicated in monogenic diabetes (23) and genes for which coding sequence variants have been strongly associated with T2D (24). Of the 33 genes with clear 1:1:1 orthologs between human, mouse and sand rat, 32 lie between position 225 and 10,195 in our PDI ranking, indicating that they are not unusually divergent in sand rat. *Pdx1* is ranked 1st and is the most unusually divergent protein identified in the sand rat predicted proteome (SI Appendix, section 8; SI Appendix, Tables S8, S10). Strikingly, 7 of the top 10 highest PDI results correspond to genes located within the mutational hotspot (SI Appendix, Table S8), indicating that GC-biased mutation is contributing to coding sequence divergence across this region.

The mutations fixed in sand rat *Pdx1* gene do not cause frameshifts or truncations in known domains, and molecular modelling reveals that the sand rat Pdx1 homeodomain has the ability to form all three helices required for DNA binding (Fig. 3a). To examine if these mutations have resulted in subtle effects on the stability of DNA binding we deployed molecular dynamics simulations with atomistic representation of Pdx1 homeodomains, DNA target and solvent. From the post-processing of the molecular dynamics simulations we estimated the enthalpy of binding between sand rat and mouse (or other mammal) Pdx1 and monomer DNA binding sites using the MM-PBSA (Molecular Mechanics Poisson Boltzmann Surface Area) method (SI Appendix, section 10). Target DNA sequences used were core Pdx1-binding sites of the mouse *insulin* A1 promoter and its sand rat ortholog. From 200 ns molecular dynamics simulations the enthalpy of binding for protein-DNA interaction was calculated to be lower for sand rat than for mouse Pdx1 (mean -140 kcal/mol vs. mean -122 kcal/mol), indicative of sand rat Pdx1 binding DNA more 'tightly' than is normal for the mammalian Pdx1 protein (Fig. 3b). One amino acid change was responsible for much of the difference: a Leu to Arg substitution in alpha helix 1 (homeodomain position 13), leading to the positive side chain of Arg making a new indirect contact with the phosphate backbone of DNA. A second substitution, Val to Arg in alpha helix 2 (homeodomain position 36), makes a smaller contribution (Fig. 3c). We also detect modifications to specific base interactions, with sand rat residues Met54 and Arg58 making new contacts to A and T bases within the TAAT core. Hence, stronger DNA binding is most likely driven by increased contacts with the backbone of DNA, coupled with decreased sequence-specificity of DNA interaction. These results suggest that sand rat Pdx1 is divergent in DNA-binding affinity and specificity. Conserved Pdx1-binding sites in well-characterised promoters of three downstream target genes encoding pancreatic hormones (*insulin*, *somatostatin* and *glucokinase*) show negligible divergence in sand rat compared to mouse, rat and human (SI Appendix, section 11), indicating that Pdx1 divergence alone is likely to be responsible for altered DNA binding affinity and specificity.

Discussion

We show that an unusual genomic region of biased mutation arose in the evolutionary lineage of the sand rat. One consequence of this hotspot of mutation was the generation of GC-bias in the *Pdx1* gene of *P. obesus*; this forced modification of the Pdx1 protein sequence, likely affecting its ability to regulate transcription of *insulin* and other pancreatic genes. The sand rat Pdx1 hexapeptide, which mediates co-factor interactions (25), is intact, which may explain why pancreatic development proceeds permitting viable sand rat embryogenesis. We suggest mutation-driven changes have played a role in constraining or adapting the sand rat, and possibly other gerbil species, to arid environments and low caloric intake. Biased gene conversion is a known mechanism that causes GC-biased mutation (21,26); hence we suggest this mechanism, driven by elevated localised recombination, is generating a hotspot of skewed base composition. The genomic region we describe here was not detected by standard short-read sequencing approaches, known to be sensitive to nucleotide composition (27). These issues may be circumvented through the use of 3rd generation sequencing technologies offering substantially longer read lengths and reduced nucleotide bias. The possibility remains that other such dark DNA regions could be widespread features of animal genomes, thus far largely overlooked in comparative animal genomics. Indeed, GC-rich genes are also missing from the chicken genome assembly (28,29). Hotspots of mutation could drive rapid evolutionary change at the molecular level, and it will be important to decipher to what extent such hotspots have constrained and influenced evolutionary adaptation across the animal kingdom.

Materials and Methods

Sand rat genome sequencing

All animal procedures were carried out in accordance with the regulations specified under the Animals (Scientific Procedures) Act 1986, UK, or the Protection of Animals Act by the authority in Denmark, European Union and Novo Nordisk A/S. Sand rat genome sequencing libraries were constructed from a male *Psammomys obesus* obtained from Hadassah medical school, Israel. We prepared and sequenced multiple short and long-insert DNA libraries and sequenced them on an Illumina HiSeq2000. We also isolated Sand rat DNA enriched for GC content through Caesium Chloride gradient centrifugation, prepared GC-rich DNA libraries and sequenced using an Illumina MiSeq. In total we generated ~398 Gbp of sequencing data which was assembled using SOAPdenovo2 (30). Further details are provided in the SI Appendix.

Transcriptome sequencing and analysis

Total RNA was extracted and purified using either Qiagen RNeasy column-based methods (pancreatic islets and liver) or TRIreagent (duodenum). All RNA-seq libraries were prepared using Illumina chemistry. Pancreatic islet libraries were sequenced individually and as pools on the Illumina GAII. RNA-seq libraries for liver and duodenum were sequenced on the HiSeq 2000 (liver) or the HiSeq 4000 (duodenum). The pancreatic islets transcriptome was assembled using Trans-ABYSS (31) using multiple k-mer sizes (41 up to 79, in increments of 2) and the liver and duodenum transcriptomes were assembled using Trinity (32) (SI Appendix).

Gene prediction and annotation

We used multiple methods to predict genes in the sand rat genome. Repetitive elements were first masked using RepeatMasker followed by *ab initio* gene prediction with AUGUSTUS (33). Homologous proteins from mouse and human were subsequently mapped to the sand rat genome assembly using TBLASTN, with the aligned sequence being filtered and passed to GeneWise (34) to identify accurate spliced alignments. GLEAN (35) was then used to generate a consensus gene set. These were then further refined by predicting Open Reading Frames using genome-guided transcriptome assemblies assembled using TopHat (36) and Cufflinks (37).

Evolutionary analyses

Using the gene predictions from our Sand rat genome assembly and the assembled tissue transcriptome data we carried out analyses of coding sequence GC content and GC-biased mutation within coding and intronic regions compared to other rodents. We also conducted an analysis to determine the extent of protein divergence within the Sand rat predicted proteome compared to mouse and human. Details of these analyses are described in the SI Appendix.

Molecular modelling

We used molecular dynamics simulations to calculate the enthalpy of binding of Protein-DNA complexes, namely between the sand rat or mouse *Pdx1* homeodomain and the sand rat or mouse A1 region of the *insulin* promoter, using Molecular Mechanics Poisson Boltzmann Surface Area (MMPBSA) analyses (SI Appendix).

Data availability

Raw sequencing reads have been deposited in the National Center for Biotechnology Information short-read archive (SRA) under the accessions SRA502705 (Illumina HiSeq paired-end and mate-pair reads) and SRR5084169-SRR5084170 (GC-enriched Illumina MiSeq reads). This Whole Genome Shotgun project has been deposited at DDBJ/ENA/GenBank under the accession NESX00000000. The version described in this paper is version NESX01000000. Raw transcriptomic reads have also been uploaded to the SRA database under the accessions SRR5092818-SRR5092820 (*Psammomys obesus*) and SRR5429486 (*Meriones unguiculatus*).

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Author contributions

ADH and PWHH conducted GC-rich DNA isolation, sequencing and analysis; LZ, SL, FL and GZ performed genome assembly and annotation; MTH prepared DNA samples; SVHP and ADH extracted RNA samples; KS, SB, BF and ADH performed RNA-seq and assembly; JC performed laboratory investigations underpinning subsequent work; ADH, LZ, PGJ, JFM and FM undertook bioinformatic analyses; ES and WRT ran molecular dynamic simulations; RSH, GZ and PWHH initiated and directed the research; PWHH, ADH, LZ, GZ and RSH drafted the manuscript. All authors approved the final manuscript.

Competing Interests

JC, PGJ, MTH, SVHP, SB, KS, BAF and RSH are current or former employees of Novo Nordisk.

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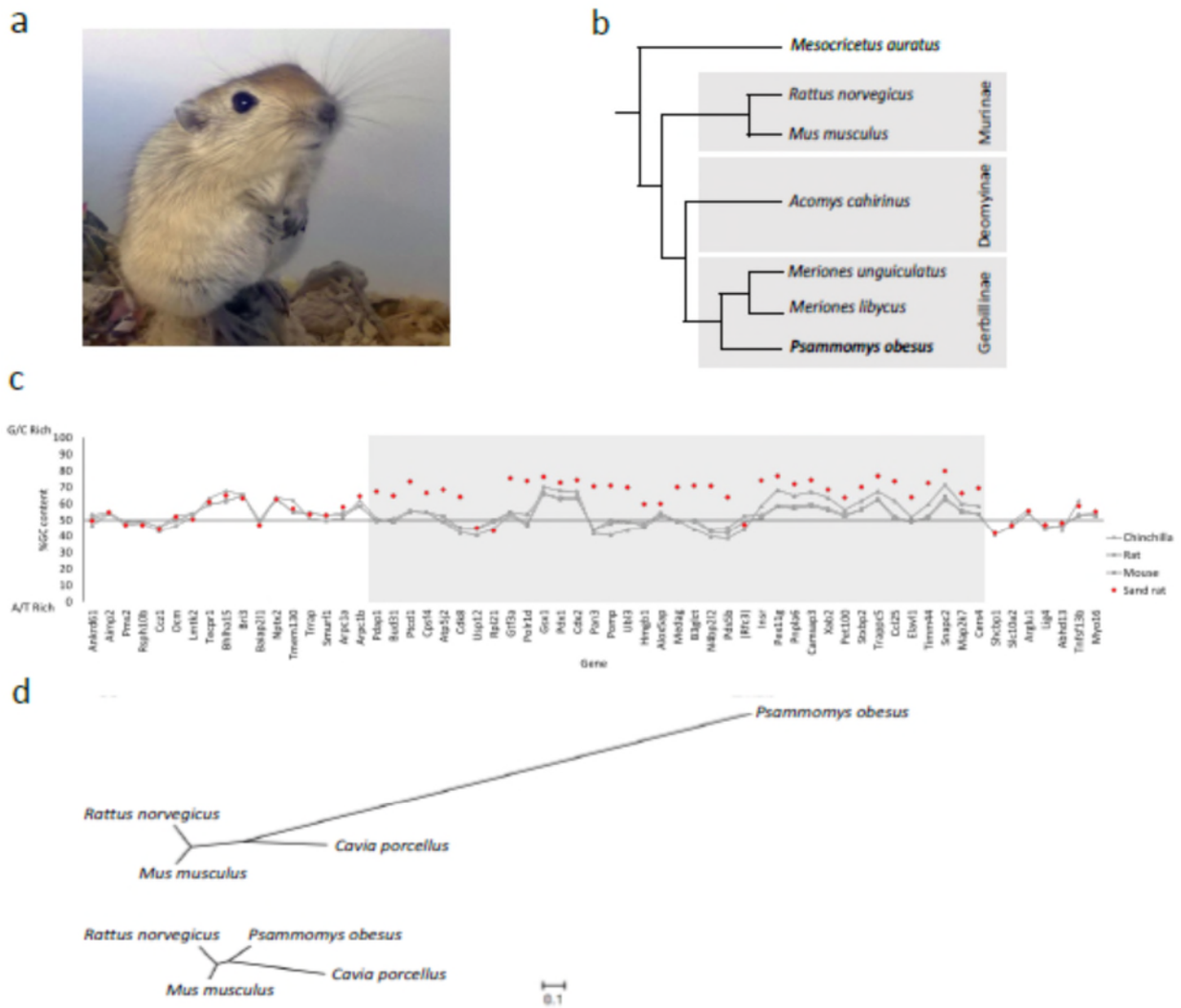


Figure 1. The sand rat and its genomic hotspot of mutation. (a) Juvenile sand rat *Psammomys obesus*. (b) Cladogram of representative murid rodents indicating the phylogenetic position of sand rat. (c) GC content of genes around the ParaHox cluster of sand rat and other rodents (*Mus musculus*, *Rattus norvegicus*, *Chinchilla lanigera*) revealing a chromosomal hotspot of GC skew in sand rat (shaded in grey). Genes shown in inferred ancestral gene order; parentheses around *Rfc3* indicate this gene has been transposed to a different genomic location in sand rat. Sand rat GC values based on transcriptome and genome sequences; when partial only alignable sequence is compared. (d) Unrooted phylogenetic trees inferred from synonymous changes (dS) only from concatenated alignments of 26 genes in the mutational hotspot (top) and 100 random genes (bottom).

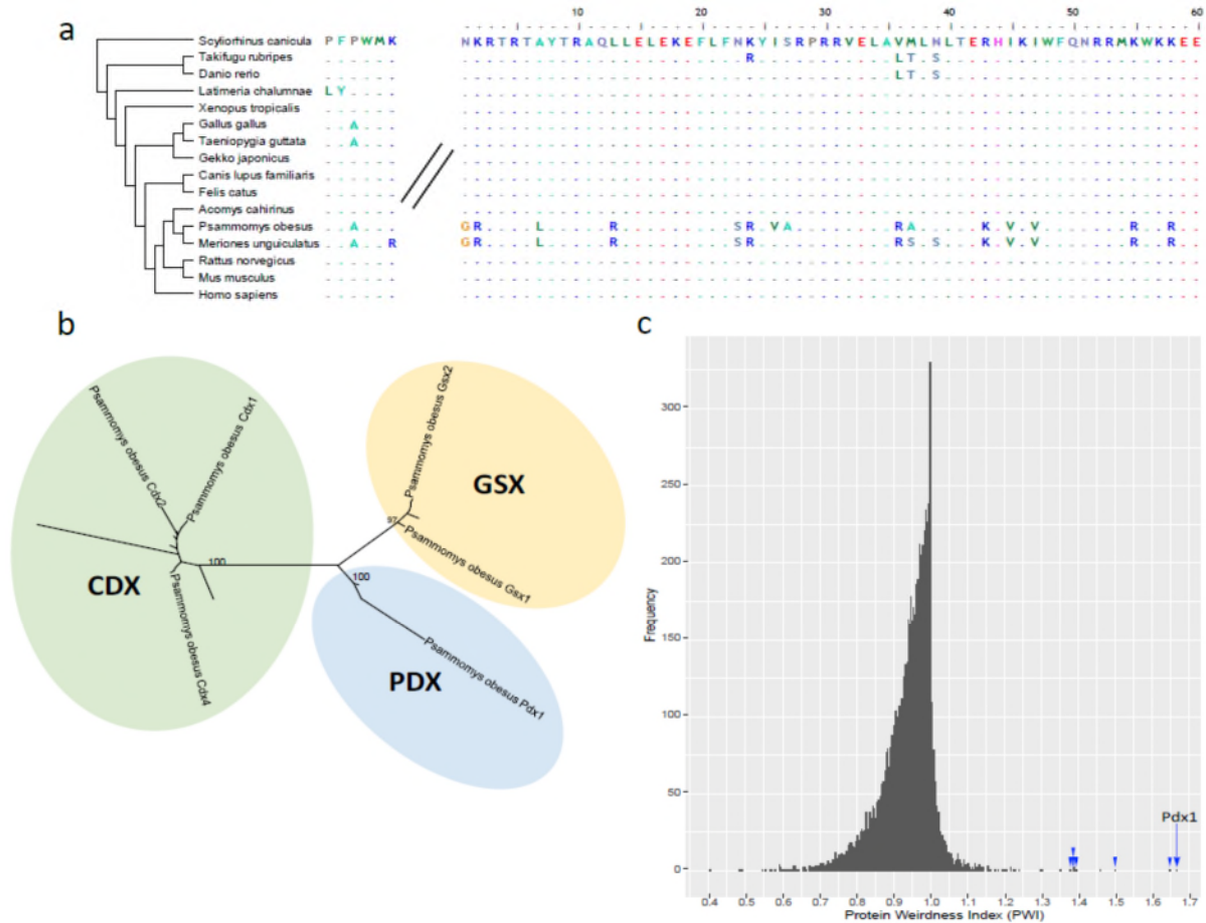


Figure 2. Molecular divergence of sand rat Pdx1. (a) Alignment of Pdx1 hexapeptide domain and homeodomain sequences across vertebrates. (b) Maximum likelihood tree of ParaHox proteins showing divergent *Psammomys obesus* Pdx1; species included are sand rat, mouse, zebra finch, spotted gar, amphioxus (full tree SI Appendix, Fig. S6). (c) Histogram of Protein Deviation Index (PDI) values for 1:1:1 mammalian orthologs of the sand rat predicted proteins: Pdx1 is marked by an arrow, other genes within the GC-rich region are marked by arrowheads.

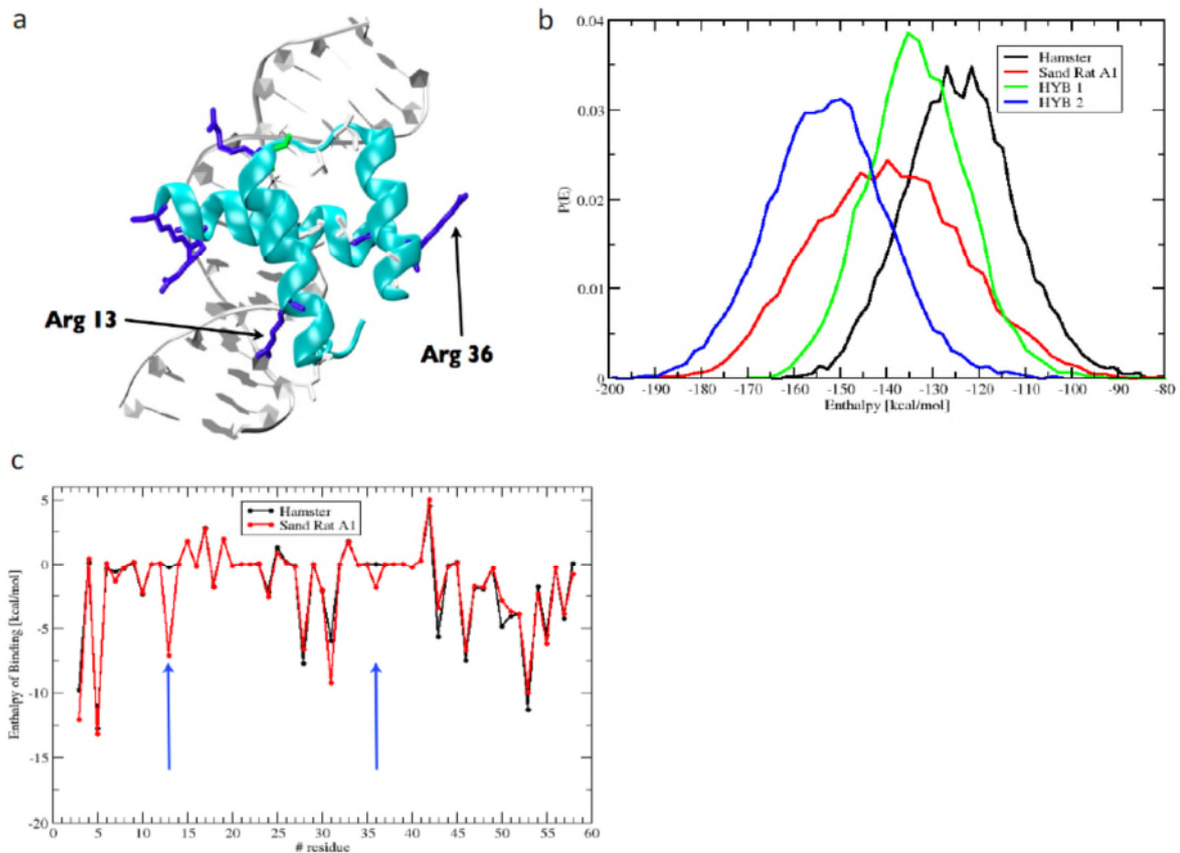


Figure 3. Molecular modelling of sand rat Pdx1 binding. (a) Molecular model of sand rat Pdx1 homeodomain bound to DNA. The two amino acid changes indicated are the largest contributors to altered enthalpy of binding. (b) Probability distributions of the enthalpy of binding of homeodomain protein-DNA interactions between hamster (normal vertebrate) Pdx1/hamster insulin A1 DNA element (black), sand rat Pdx1/sand rat A1 element (red), hamster Pdx1/sand rat A1 (green) and sand rat Pdx1/hamster A1 (blue) inferred by molecular dynamics simulations and MM-PBSA; sand rat Pdx1 homeodomain has the lowest enthalpy of binding (higher affinity) for each DNA target. (c) Per-site enthalpy of binding comparison between hamster and sand rat Pdx1 revealing contribution of amino acid changes at homeodomain positions 13 and 36 to reduced enthalpy of binding (higher affinity).

Total number of paired-end reads	724,377,486
Total number of mate-pair reads	1,780,436,140
Total bases sequenced	394,396,928,120
Estimated sequencing coverage (x)	87.6
Number of scaffolds >2 kb	1,737
Total length of assembly (bp)	2,381,209,849
Longest scaffold (bp)	54,616,910
Mean scaffold length (bp)	15,794
Scaffold N50 (bp)	10,461,538
Scaffold L50	63
Contig N50 (bp)	83,904
Percentage of assembly in scaffolds	98.6%

Table 1. Metrics of sand rat raw genomic sequencing data and final genome assembly. Coverage was calculated using an estimated genome size of 2.51 Gb based on a k-mer analysis (SI Appendix, section 1.3) and is based upon paired-end sequencing data only.