Adaptive Evolution in the Glucose Transporter 4 Gene \textit{Slc2a4} in Old World Fruit Bats (Family: Pteropodidae)

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Abstract

Fruitarian and nectarivorous bats are able to ingest large quantities of sugar in a short time span while avoiding the potentially adverse side-effects of elevated blood glucose. The glucose transporter 4 protein (GLUT4) encoded by the \textit{Slc2a4} gene plays a critical role in transmembrane skeletal muscle glucose uptake and thus glucose homeostasis. To test whether the \textit{Slc2a4} gene has undergone adaptive evolution in bats with carbohydrate-rich diets in relation to their insect-eating sister taxa, we sequenced the coding region of the \textit{Slc2a4} gene in a number of bat species, including four Old World fruit bats (Pteropodidae) and three New World fruit bats (Phyllostomidae). Our molecular evolutionary analyses revealed evidence that \textit{Slc2a4} has undergone a change in selection pressure in Old World fruit bats with 11 amino acid substitutions detected on the ancestral branch, whereas, no positive selection was detected in the New World fruit bats. We noted that in the former group, amino acid replacements were biased towards either Serine or Isoleucine, and, of the 11 changes, six were specific to Old World fruit bats (A133S, A164S, V377F, V386I, V441I and G459S). Our study presents preliminary evidence that the \textit{Slc2a4} gene has undergone adaptive changes in Old World fruit bats in relation to their ability to meet the demands of a high sugar diet.

Introduction

High acute blood glucose concentration causes severe physiological dysfunction and even death [1,2,3]. To maintain glucose homeostasis, and thus avoid potential toxicity, the blood glucose concentration is maintained within narrow limits by an interplay between tissue glucose uptake, hepatic glucose production and insulin production [4]. In the first of these regulatory mechanisms, it is skeletal muscle that contributes most to the removal of excess glucose from circulation [4,5], a process that is mediated by the transmembrane glucose transporter 4 protein (GLUT4) [5]. GLUT4 (encoded by the \textit{Slc2a4} gene) is a family member of sugar transporter proteins that is highly expressed in skeletal muscle, heart and adipose tissue [4]. In response to stimulation by insulin, exercise or contraction, GLUT4 is translocated from intracellular compartments to the plasma membrane of adipocytes and muscle, where it mediates glucose uptake [3,4,6]. Studies of transgenic mice revealed that GLUT4 plays a pivotal role in the regulation of skeletal muscle glucose uptake and maintenance of glucose homeostasis [7,8].

Old World fruit bats (OWFBs) (Pteropodidae) and New World fruit bats (NWFBs) (Phyllostomidae) have independently evolved a carbohydrate-rich diet comprising mostly fruit and/or nectar [9]. To meet the high energy demands of powered flight, while relying on an often highly ephemeral food source, these bats must frequently ingest large quantities of sugar in a short time span [10,11]. It has recently been shown that the majority of glucose is absorbed via a passive paracellular pathway in the intestinal tract [12,13,14]. At the same time, this has raised the issue of how fruit eating bats ingest sugars rapidly and efficiently while maintaining normal glycaemia.

To date, many efforts have been made to elucidate the means by which fruit bats regulate their blood glucose, with several studies identifying physiological and biochemical adaptations. Michelson et al. (1998) showed that the pancreas of the Old World frugivorous bat \textit{Rousettus aegyptiacus} contained an unusually large volume (9.1%) of endocrine tissue [15]. More recently, Protzek et al. (2010) found that the pancreas of the New World frugivorous bat \textit{Artibeus lituratus} contained a large population of \(\beta\)-cells, and that this species showed elevated sensitivity to glucose and insulin [16]. In addition to these assumed anatomical specializations, biochemical research on New World nectarivorous bat, \textit{Glossophaga soricina} has shown that high skeletal muscle enzymatic flux capacities in this species are similar to those of hummingbirds [17], and it has been suggested that \textit{G. soricina} has evolved a mechanism of high-activity to balance blood glucose and thus avoid hyperglycemia after excess sugar consumption [18]. Several studies have also shown that both OWFBs (e.g. \textit{R. aegyptiacus}) and NWFBs (e.g. \textit{G. soricina}) fuel their metabolic needs exclusively with exogenous sugars rather than endogenous products [19,20,21].

Physiological and biochemical adaptations for glucose metabolism in bats should not only require the effective regulation of plasma glucose levels, but in addition the efficient transportation of

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blood glucose to high energy demanding organs (e.g. flight muscles). Thus, it is reasonable to suspect that key genes involved in carbohydrate metabolism and blood glucose regulation may have been targets of molecular adaptation in the dietary switches seen in fruit and nectar eating bats.

Considering the critical role of GLUT4 in the regulation of skeletal muscle glucose uptake and maintenance of glucose homeostasis, we speculated that the Slc2a4 gene will have undergone adaptive evolution coincident with the switch in diet in the ancestral OWFBs and NWFBs. To test our hypothesis, we obtained the coding region of Slc2a4 from 16 bat species (including four OWFBs species and three NWFBs species), and assessed the molecular evolution of this gene in bats and other mammals.

**Materials and Methods**

**Ethics Statement**

Our procedures involving animals were in accordance with the guidelines of Regulations for the Administration of Laboratory Animals (Decree No. 2 of the State Science and Technology Commission of the People’s Republic of China on November 14, 1988).

**Taxonomic Coverage**

We studied the diversity and evolution of the Slc2a4 gene in 16 bat species. From the suborder Yampirochiroptera, we included four Old World fruit bats: *Pteropus vampyrus*, *Gynopterus sphinx*, *Eonycteris spelaea*, and *Rousettus leschenaultii* (family Pteropodidae). From this clade we sequenced three insectivorous bat species from sister families to the Old World fruit bats: *Rhinolophus ferrumequinum* (Rhinolophidae), *Hipposideros pratti* and *H. armiger* (Hipposideridae). From the other main suborder (Yangochiroptera) we included the three New World fruit bats *Leptonycteris yerbabuenae*, *Artibeus lituratus* and *A. jamaicensis* (Phyllostomidae) and six insectivorous species representing four families: *Taphozous melanopogon* (Emballonuridae), *Tadarida brasiliensis* (Molossidae), *Scotophilus kuhlii*, *Myotis ricketti* and *M. lucifugus* (Vespertilionidae) and *Mormoops megaphylla* (Mormoopidae). Of these the latter family is sister to the New World fruit bats. All new Slc2a4 sequences were deposited to GenBank and accession numbers are JN695651–JN695665.

We also incorporated available published sequences in our analyses; from Ensembl we obtained the Slc2a4 sequence of the insectivorous bat species *M. lucifugus*, and from GenBank we obtained sequences form the seven mammal species: *Homo sapiens* (NM_001128433), *Mus musculus* (NM_001159327), *Rattus norvegicus* (NM_174604), *Sus scrofa* (NM_001159327), *Canis familiaris* (NM_101081866), *Equus caballus* (NM_001159327), *Bos taurus* (NM_174604), *Sus scrofa* (NM_001128433), *Canis familiaris* (NM_001159327), *Rhinolophus ferrumequinum* (Rhinolophidae), *Hipposideros pratti* and *H. armiger* (Hipposideridae). We also included the three New World fruit bats *Leptonycteris yerbabuenae*, *Artibeus lituratus* and *A. jamaicensis* (Phyllostomidae) and six insectivorous species representing four families: *Taphozous melanopogon* (Emballonuridae), *Tadarida brasiliensis* (Molossidae), *Scotophilus kuhlii*, *Myotis ricketti* and *M. lucifugus* (Vespertilionidae) and *Mormoops megaphylla* (Mormoopidae). Of these the latter family is sister to the New World fruit bats. All new Slc2a4 sequences were deposited to GenBank and accession numbers are JN695651–JN695665.

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**Isolation, Amplification and Sequencing**

We isolated genomic DNA using DNeasy Blood & Tissue Kit (Qiagen) from wing membrane biospies from six species (see Table S2) and designed primers to amplify four separate sections of Slc2a4 gene (first section for exons 1–2, second for exons 3–7, third for exons 8–9 and fourth for exons 10–11) (Table S2). For the remaining nine species, we isolated total RNA from pectoral muscle tissue (stored at −80°C) of euthanized bats using Trizol reagent (Invitrogen). Following the standard protocol, 5 μg total RNA was reverse transcribed to cDNA by SuperScript™ III Reverse Transcriptase kit (Invitrogen). Primers were designed to amplify the coding sequences of Slc2a4 (see Table S2). For both genomic DNA and pectoral muscle cDNA, Polymerase Chain Reaction (PCR) were conducted using Premix Ex Taq™ (TaKaRa) with the following conditions: denaturation at 95°C for 5 min, 32 amplification cycles [95°C for 0 s, annealing temperature (see Table S2) for 30 s, 72°C for 1–2 min (depending upon the target length)], and a final extension at 72°C for 10 min. All PCR products were isolated using 1% agarose gels and purified with Gel Extraction Kits (Qiagen), ligated into pGEM-T easy vector (Promega), cloned and sequenced using the Terminator kits (Applied Biosystems) on an ABI 3730 DNA sequencer.

**Phylogenetic Reconstruction**

The nucleotide sequences of 23 species were aligned using ClustalX [22] and checked for accuracy by eye, and coding sequences were translated to amino acids using MEGA4 [23]. A Bayesian phylogenetic tree was reconstructed based on aligned nucleotide sequences using MrBayes 3.1.2 [24] with the TPM2uf+Y nucleotide substitution model selected by jModelT estimator [25]. For the Bayesian analysis, we performed 10,000,000 generations of MCMC and sampled every 100 generations, with the first 2,000,000 generations discarded as burn-in. All other options and priors were default settings of MrBayes 3.1.2 software. The standard deviations of split frequencies were stably below 0.01 after 2,000,000 generations of MCMC performances.

**Molecular Evolution Analyses**

Before undertaking analyses of molecular evolution, we first tested for evidence of recombination in our dataset, which can lead to the erroneous detection of positive selection. We used GARD [26] implemented in the package HyPhy [27] to estimate the number of statistically supported recombination breakpoints.

For selection tests, we used a phylogenetic tree topology based on the accepted species relationships [28,29,30]. To test for positive selection in Slc2a4, we derived maximum-likelihood estimates of the rate of nonsynonymous substitutions (dN) and synonymous substitutions (dS) using PAML CODEML [31].

We first undertook branch models using two-ratio models, in which the dN/dS ratio (termed as omega or Ω) was allowed to vary between the background and foreground. Separate models were undertaken with the foreground branch set as the ancestral branch leading to OWFBs and to NWFBs. In both cases, the one-ratio model in which Ω was fixed among all branches served as the null hypothesis of no branch-wise change in selection pressure. A three-ratio model in which different dN/dS ratios were assumed for the background, the ancestral branch of the OWFBs and the ancestral branch of the NWFBs, was also conducted, and compared to the one-ratio model and the two-ratio models described. We also conducted the free ratio model, which allows the dN/dS ratios to vary among all branches, to explore the overall selection pressures of Slc2a4 in the 23 species under study [32]. For this, the model parameters were compared to those from the one-ratio model, which served as the null expectation of no variation in selection pressure among lineages.

We also applied the branch-site model Λ in combination with Bayes empirical Bayes (BEB) estimation, to detect positively selected sites along particular branches [33]. In this model, the phylogeny is divided into foreground and background branches. Four site classes of codons are assumed, of which site class 0 and 1 evolve under purifying selection (0<ΔdS<1) and neutral selection (ΔdS=1) respectively throughout the tree. The remaining site classes 2a and 2b evolve under, respectively, purifying and neutral selection on background, but are grouped together and allowed to evolve under positive selection on foreground (ΔdS>1). We applied
test 1 and test 2 of this branch-site model [34] to the ancestral branches of OWFBs and NWFBs, in which branch-site model A was the alternative hypothesis. In test 1, the null hypothesis was the M1a (Nearly Neutral) model which assumed two site classes: \( 0 < \omega_0 < 1 \) and \( \omega_1 = 1 \). While for test 2, the null hypothesis was the modified branch-site model A with \( \omega_2 \) fixed as 1. The likelihood ratio tests (LRTs) were used to compare the model fit for alternative and null hypotheses.

### Table 1. Results of branch model tests of selection pressure on the Slc2a4 gene in bats.

| Model Description | \( \text{np} \) | \( \ell \) | \( \omega_0^* \) | \( \omega_{OW}^* \) | \( \omega_{NW}^* \) | Model Compared | \( 2\Delta \ell \) | \( P \) |
|-------------------|----------------|-----------|----------------|----------------|----------------|----------------|----------------|--------|
| A. One ratio: \( \omega_0 = \omega_{OW} = \omega_{NW} \) | 45 | -7140.94 | 0.088 | - = \omega_0 \) | - = \omega_0 | | | |
| B. Two ratios: \( \omega_0 = \omega_{OW}, \omega_{NW} \) | 46 | -7140.93 | 0.088 | - = \omega_0 \) | 0.098 | B vs. A | 0.02 | 0.887 |
| C. Two ratios: \( \omega_0 = \omega_{NW}, \omega_{OW} \) | 46 | -7137.92 | 0.085 | 0.257 | = \omega_0 | C vs. A | 6.03 | 0.014 |
| D. Three ratios: \( \omega_0, \omega_{OW}, \omega_{NW} \) | 47 | -7137.90 | 0.085 | 0.257 | 0.099 | D vs. A | 6.08 | 0.048 |
| E. Free ratio | 87 | -7116.96 | — | — | — | E vs. A | 47.94 | 0.244 |

\( \omega_{OW}, \omega_{NW}, \) and \( \omega_0 \) are the \( \omega \) ratios for branches OWFBs, NWFBs, and other branches, respectively (see Figure 1b).

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Finally, we tested for positive selection among sites across the tree. For this we compared site models M7 versus M8. In this test, the M7 model (β) was the null model with an assumed β distribution of θ between 0 and 1, while the M8 model (β & θ) is the alternative model that has an additional site class of θ > 1.

In addition to the methods of PAML, we used two alternative approaches to characterize selection pressures in our dataset, both implemented using the Datamonkey web server (http://www.datamonkey.org/). First we used the GA-Branch method [35], in which branches are assigned to a range of θ classes without a priori specification of focal lineages of interest. Second, to examine site-wise variation across branches we applied the random effects branch-site model (Branch-site REL) [36], which might be more robust to errors because it does not enforce a uniform section pressure across all background branches predicted to be not under positive selection.

Ancestral sequences were reconstructed using PAML CO-DEMEL package [37], and the amino acid substitutions on each branch were then inferred. In order to determine the lineage-specific amino acid changes, we repeated the ancestral states tests across the bat phylogeny. The alignment of Slc2a4 coding sequences spanned 1530 nucleotides, equating to 510 amino acids, of which 104 were variable in eutherian mammals (Figure S1). The fact that the open reading frame was intact with no detectable stop codons or indels suggested that the Slc2a4 gene is functional in the species studied.

Bayesian phylogenetic reconstruction recovered a tree in which the main groupings agreed with the accepted species topology (Figure 1b). Specifically, the Old World fruit bats grouped with the horseshoe bats and their allies to form a well-supported monophyletic clade (suborder Yinpterochiroptera). The remaining lineages were contained within the second major recognised clade, the Yangochiroptera, which included the NWFs. These bats (all members of the family Phyllostomidae) formed a correct and well-supported monophyletic clade (Bayesian posterior probability = 1.00) that showed a sister relationship with the single representative of the Mormoopidae (Figure 1a). We found no evidence of recombination breakpoints in our dataset, adding

| Table 2. Results of branch-site model A test for detection of positively selected sites in ancestral branches of Old World fruit bats and New World fruit bats. |
|---------------------------------------------------------------|
| **Branch-site model**                                         | **np** | **Parameters** | **LRT** | **P** | **Sites with elevated omega values** |
|---------------------------------------------------------------|
| Model A (alternative hypothesis) for Old World fruit bats      | 48     | P1 = 0.09, P2 = 0.07, P3 = 0.01, P4 = 0.008 | Test 1 | 0.033 | 1335(0.751), 1645(0.760), 377F(0.612), 386(0.929), 441L(0.652), 459Y(0.701) |
| Model A (null hypothesis) for Old World fruit bats             | 47     | P1 = 0.026, P2 = 0.006, P3 = 0.008, P4 = 0.000 | Test 2 | 0.175 | Not allowed |
| Model A (alternative hypothesis) for New World fruit bats      | 48     | P1 = 0.093, P2 = 0.077, P3 = 0.000, P4 = 0.000 | Test 1 | 0.033 | 1335(0.751), 1645(0.760), 377F(0.612), 386(0.929), 441L(0.652), 459Y(0.701) |
| Model A (null hypothesis) for New World fruit bats             | 47     | P1 = 0.026, P2 = 0.006, P3 = 0.008, P4 = 0.000 | Test 2 | 0.175 | Not allowed |

* np, number of parameters.
* LRT, likelihood ratio test.
* Sites with elevated omega values detected by branch-site model A test are referred to Pteropus vampyrus. BEB posterior probabilities are shown in parentheses. doi:10.1371/journal.pone.0033197.0102
further support that the inferred gene tree was valid for molecular genetic analyses.

To assess whether an evolutionary switch to a high carbohydrate diet in OWFBs and NWFBs was associated with positive selection acting on the \textit{Slc2a4} coding sequence, we derived maximum-likelihood estimates of the rates of nonsynonymous substitution (\(d_N\)) and synonymous substitution (\(d_S\)) and undertook a number of branch-wise tests of selection. First, for each of these groups of bats, we constructed a two-ratio branch model (labeled branches in Figure 1b) and, additionally, we used a three-ratio model to estimate simultaneously the ratios for both OWFBs and NWFBs and the background. Results of these branch models are shown in Table 1. Briefly, model comparisons indicated that the two-ratio model with the OWFBs branch designated as the foreground was a significantly better fit than the null (one-ratio) model in which the \(d_N/d_S\) ratio was invariant across all branches [likelihood ratio test (LRT) statistic (\(2\Delta\ell = 6.03, df = 1, P = 0.014\)) (Table 1). The estimated value of \(d_N/d_S\) on the ancestral OWFBs branch was an order of magnitude greater than that of background (0.257 versus 0.085, respectively) suggesting a change in selection pressure in the evolutionary history of this lineage. However, no such change was found on the ancestral branch of NWFBs, with the one-ratio model not rejected by the two-ratio model (\(2\Delta\ell = 0.02, df = 1, P = 0.887\)) (Table 1). Moreover, a three-ratio model did not fit the data better than the two-ratio model with OWFBs as the foreground (\(2\Delta\ell = 0.04, df = 1, P = 0.839\)). Finally, the free-ratio model test showed that when \(d_N/d_S\) was allowed to vary among branches, the fit was no better than that of a one-ratio model (\(2\Delta\ell = 47.94, df = 42, P = 0.244\)) (Table 1).

Application of branch-site models revealed six sites on the ancestral branch of OWFBs with omega values of \(>1\) (133S, 164S, 377F, 386I, 441I and 459S) (Table 2), with branch-site model \(A\) found to fit the data better than the M1a Nearly Neutral model (\(2\Delta\ell = 6.85, df = 2, P = 0.033\); test 1). However, the explicit test of positive selection (test 2) was not significant (\(2\Delta\ell = 1.04, df = 1, P = 0.175\)) (Table 2). Thus we were unable to reject the possibility of relaxed selection, since test 1 of branch-site model \(A\) is unable to distinguish relaxed constraints from positive selection [34]. Branch-site models detected no positively selected sites on the ancestral branch of NWFBs, with the null model unable to be rejected over the alternative model in both test 1 and test 2. Other tests of positive selection (M7 versus M8 model comparison), and alternative approaches based on branch (GABranch) and branch-site comparisons (REL) also failed to detect positive selection in both OWFBs and NWFBs (data not shown).

Examination of reconstructed ancestral sequences revealed a total of 11 amino acid replacements on the ancestral branch of OWFBs, none of which were shared with the NWFBs (Figure 1b). However, of the six sites showing evidence of selection pressure change in the OWFBs (A133S, A164S, V377F, V386I, V441I and G459S), one (A133S) was also detected in \textit{Scotophilus kuhlii}, and three (V377L, V441I and G459A) were seen in the dog (Figure S2). The site with the highest probability of positive selection (386I, BPP = 0.929) was only seen in the Old World fruit bats (Figure 2). The ancestral branches of the Chiroptera, Yinpterochiroptera, Yangochiroptera and NWFBs each contained either two or three amino acid substitutions (Figure 1b). We also calculated the \(d_N/d_S\) values for each amino acid of \textit{Slc2a4} on branches leading to cow, dog, rodents and human, which all exhibited large amount of substitutions, however, no evidence of positive selection was found on these branches (Figure 3).

![Figure 2. The species tree of 23 mammals with Old World fruit bats specific amino acid replacements were highlighted by ancestral sequence reconstruction using maximum parsimony method. (a) A164S, (b) V386I. Branch lengths are not drawn to scale. The amino acids at positions 164 (A: Ala; S: Ser) and 386 (V: Val; I: Ile) of \textit{Slc2a4} for each interior and exterior node are shown. doi:10.1371/journal.pone.0033197.g002](https://www.plosone.org/doi/10.1371/journal.pone.0033197.g002)
Mapping the 11 amino acid changes in the OWFBs onto the protein secondary structure revealed that four were distributed on extracellular loops (A133S, T200I, V377F and V441I), two within intracellular regions (R19Q and A164S), and the other five within the transmembrane domains (A27S, V292I, V386I, G437S and G459S) (Figure 4). The sites showing evidence of elevated selection pressure were mainly (four of six) located in either extracellular or intracellular loops, extremely close to transmembrane domains (Figure 4).

Discussion

By comparing the Slc2a4 gene in frugivorous and nectarivorous bats to their insect-eating sister taxa, we provide some of the first data on molecular adaptation to a sugar rich diet in mammals [39]. We found evidence of a significantly higher value of $\omega$ ($d_{NS}/d_S$) on the ancestral branch of Old World fruit bats (OWFBs) compared to the rest of the tree, indicative of change in selection pressure in Slc2a4 since the divergence of this lineage from echolocating Yinpterochiroptera. Subsequent analyses revealed six sites with elevated selection pressure.

In spite of this result, based on a branch-site model test of positive selection we were unable to reject the possibility that the $\omega$ value in OWFBs had arisen due to relaxed functional constraint, and indeed alternative methods of positive selection proved negative. However, we argue that relaxed evolution at Slc2a4 in OWFBs is unlikely. This gene is highly conserved across divergent
lineages of mammals, including rodents, primates, carnivores and herbivores, a fact that is unsurprising given that the GLUT4 protein product is critical for glucose homeostasis in mammals [4]. Studies of transgenic mice revealed that targeted disruption of GLUT4 in adipocytes and muscle tissues can cause severe insulin resistance and glucose intolerance [7,8].

If we assume that relaxed selection in the Slc2a4 gene of OWFBs is indeed implausible, then the best explanation for the significantly high \( v \) in OWFBs is molecular adaptation. Previously it has been pointed out that signals of positive selection can be swamped by a history purifying selection [34], which seems possible given that OWFBs diverged from the other Yinpterochoptera nearly 60 million years ago [29]. Low sequence divergence in a dataset will also limit the power to detect positive selection [40,41,42] and in this context it is interesting to note that our free-ratio model of selection pressure did not fit the data better than a one-ratio model, emphasizing the strong sequence conservation in the Slc2a4 gene across the taxa studied.

Most of the observed amino acid replacements in the ancestral OWFBs were seen to be concentrated in the external (i.e. intracellular and extracellular) loops of the protein, hinting at some adaptive role in efficient skeletal muscle glucose uptake. Indeed, the inferred change from valine to phenylalanine (V377F) involves the introduction of an extra aromatic ring, might enhance the binding ability of hydrophobic ligands via stacking interactions [43]. Surprisingly, nine of the 11 replacements were to either a serine or an isoleucine, a level of amino acid substitution bias that is not apparent in the other mammal branches (Figure S3 and Table S3). Of these nine, our protein comparisons showed that the residues A164S and V386I in OWFBs are homologous to residues S148 and V370 in the GLUT1 protein, in which they have been implicated both in stabilizing protein packing for facilitation of glucose transport, and in modifying hydrophobic interactions between helices 4–5 and helices 8–10 [44].

In contrast to OWFBs, we found no evidence of elevated \( v \) in their New World counterparts (NWFBs), in which Slc2a4 was under purifying selection. Moreover, none of the sites found in the OWFBs were shared by the NWFBs. Thus in spite of their convergent eating habits and similar physiological adaptations, there were no detected parallel signatures of selection in this gene, or convergent changes of the sort recently reported in the hearing genes of bats [45,46,47,48]. In this context, it is interesting to note that recent behavioural and physiological data have shown that the New World nectarivorous bat (Glossophaga soricina) regulates its blood glucose via high activity behaviour [18], perhaps reducing reliance on other mechanisms of glucose homeostasis such as insulin-stimulated transmembrane glucose uptake. We also cannot rule out the possibility that further adaptive changes underpinning glucose metabolism have occurred in promoter regions rather than in coding sequence [49]. Indeed, recent evidence has suggested that a non-coding region of Slc2a4 has been the target of natural selection in human populations [50].

Apart from Slc2a4, other genes might also have undergone adaptive change in bats (and other mammals) in relation to their diets. For example, it was recently reported that both OWFBs and NWFBs have also undergone adaptive losses of a mitochondrial targeting motif in a key dietary enzyme: AGT (alanine-glyoxylate aminotransferase 1) [39]. As a consequence of this mutation, the AGT enzyme instead accumulates in the peroxisomes, where it breaks down the accumulating plant-derived glyoxylate before it is converted to the toxic oxalate. The same study also showed that AGT has undergone positive selection in the ancestral branch of OWFBs [39]. Similarly, proteins involved in fat metabolism have also been reported to have undergone adaptive changes in species that hibernate [51,52]. We speculate that as more studies are undertaken, numerous other genes will be seen to have been targets of natural selection during the diversification of mammalian dietary habits.

Figure 4. Distribution of 11 amino acid substitutions which occurred on the ancestral branch of Old World fruit bats in the secondary structure of GLUT4 protein with 12 transmembrane domains. Protein structure is adapted from [4]. Six sites with omega values >1 (branch-site model A) are highlighted in red, and the other amino acid substitutions are indicated in blue. doi:10.1371/journal.pone.0033197.g004
Supporting Information

Figure S1 Alignment of the amino acid sequences of the Slc2a4 gene from 23 mammals (only the variable sites are shown). The species of Old World fruit bats and New World fruit bats are marked in red and blue, respectively. (TIF)

Figure S2 The species tree of 23 mammals with Old World fruit bats specific amino acid replacements were highlighted by ancestral sequence reconstruction using maximum parsimony method. (a) A133S, (b) V377F, (c) V441I and (d) G459S. Branch lengths are not drawn to scale. The amino acids for each interior and exterior node are shown. (TIF)

Figure S3 Nonsynonymous amino acid substitutions mapped onto the species topology of 23 mammals. Branch lengths are not drawn to scale. Six sites on the ancestral branch of Old World fruit bats that had estimated omega values >1 are shown in red. Ancestral branches leading to Old World fruit bats and New World fruit bats are marked with blue and green lines, respectively. (TIF)

Table S1 The information of species examined in the study. (DOC)

Table S2 The information of primers used for Slc2a4 coding sequences amplification. (DOC)

Table S3 Summarization of amino acid substitutions occurred on major branches. (DOC)

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Author Contributions

Conceived and designed the experiments: BS XH SZ. Performed the experiments: BS XH. Analyzed the data: BS XH SJR. Contributed reagents/materials/analysis tools: SZ. Wrote the paper: BS SJR SZ. Bat species sampling: JZ.

References

1. Kawahito S, Kishita H, Oshita S (2009) Problems associated with glucose toxicity: role of hyperglycemia-induced oxidative stress. World J Gastroenterol 15: 4137–4142.
2. Brunner Y, Schwartz D, Priego-Capote F, Couté Y, Sanchez JC (2009) Glucotoxicity and pancreatic prosecretion. J Proteomics 71: 576–591.
3. Shepherd PR, Kalun BB (1999) Glucose transporters and insulin action—implications for insulin resistance and diabetes mellitus. N Engl J Med 341: 248–257.
4. Huang S, Czech MP (2007) The GLUT4 sugar transporter. Cell Metab 5: 237–252.
5. Klip A (2009) The many ways to regulate glucose transporter 4. Appl Physiol Nutr Metab 34: 481–487.
6. Thong FSL, Dugani CB, Klip A (2005) Turning signals on and off: GLUT4 traffic in the insulin-signaling highway. Physiology 20: 271–284.
7. Abel ED, Peroni O, Kim JK, Kim VB, Boss O, et al. (2001) Adipose-selective targeting of the GLUT4 gene impairs insulin action in muscle and liver. Nature 409: 729–733.
8. Zisman A, Peroni OD, Abel ED, Michael MD, Mauruo-Jarvis F, et al. (2000) Targeted disruption of the glucose transporter 4 selectively in muscle causes insulin resistance and insulin tolerance. Nat Med 6: 924–928.
9. Ahringer JD, McOwat T, Hammond L (1996) Bats: biology and behaviour. Oxford University Press, USA.
10. Laska M (1990) Food transit times and carbohydrate use in three phyllostomid species of bats. Comp Biochem Physiol A Physiol 58: 349–352.
11. Keegan DJ (1977) Aspects of the assimilation of sugars by the Egyptian fruit bat (Rousettus aegyptiacus). Comp Biochem Physiol A Mol Physiol 58: 349–352.
12. Caviedes-Vidal E, Karasse WH, Cheddiack JG, Fassulo V, Cruz-Neto AP, et al. (2008) Paracellular absorption: a bat breaks the mammal paradigm. PLoS ONE 3: e1425.
13. Tracy CR, McWhorter TJ, Korine C, Wojciechowski MS, Pinshaw B, et al. (2007) Absorption of sugars in the Egyptian fruit bat (Rousettus aegyptiacus): a paradox explained. J Exp Biol 210: 1726–1734.
14. Caviedes-Vidal E, McWhorter TJ, Lavin SR, Cheddiack JG, Tracy CR, et al. (2007) The digestive adaptation of flying vertebrates: high intestinal paracellular absorption compensates for smaller guts. Proc Natl Acad Sci USA 104: 19132–19137.
15. Michelmore AJ, Keegan DJ, Kramer B (1998) Immunochemical identification of endocrine cells in the pancreas of the fruit bat, Rousettus aegyptiacus. Gen Comp Endocrinol 110: 319–327.
16. Proetz AOP, Rafacho A, Viscelli RA, Bosqueiro JR, Cappelli AP, et al. (2010) Insulin and glucose sensitivity, insulin secretion and β-cell distribution in endocrine pancreas of the fruit bat Heliothis humana. Comp Biochem Physiol A Mol Integr Physiol 157: 142–148.
17. Suarez RK, Welch KC, Hanna SK, Herrera MLG (2009) Flight musc enzymes and metabolic flux rates during hovering flight of the nectar bat, Glossophaga soricina: further evidence of convergence with hummingbirds. Comp Biochem Physiol A Mol Integr Physiol 153: 136–140.
18. Kelm DH, Simon R, Kuhlow D, Voigt CC, Ristow M (2011) High activity enables life on a high-sugar diet: blood glucose regulation in nectar-feeding bats. Proc R Soc B 280: 1058014;10511.0485.
19. Amitai O, Holste S, Barkan S, Amichai E, Korine C, et al. (2010) Fruit bats (Pteropodidae) fuel their metabolism rapidly and directly with exogenous sugars. J Exp Biol 213: 2693–2699.
20. Welch KC, Herrera LG, Suarez KK (2008) Dietary sugar as a direct fuel for flight in the nectarivorous bat Glossophaga soricina. J Exp Biol 211: 310–316.
21. Voigt CC, Speakman JR (2007) Nectar-feeding bats fuel their high metabolism directly with exogenous carbohydrates. Funct Ecol 21: 913–921.
22. Thompson JD, Gibson TJ, Plesnik F, Jeannousm F, Higgins DG (1997) The CLUSTAL_X windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. Nucleic Acids Res 25: 4876–4892.
23. Tamura K, Dudley J, Nei M, Kumar S (2007) MEGA4: molecular evolutionary genetics analysis (MEGA) software version 4.0. Mol Biol Evol 24: 1596–1599.
24. Ronquist F, Huelsenbeck JP (2003) MrBayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics 19: 1572–1574.
25. Posada D (2008) jModelTest: phylogenetic model averaging. Mol Biol Evol 25: 1253–1256.
26. Kosakovsky Pond SL, Posada D, Gravenor MB, Wootch KL, Frost SDW (2006) GARLI: a genetic algorithm for recombination detection. Bioinformatics 22: 3096–3098.
27. Kosakovsky Pond SL, Frost SDW, Mase SV (2005) HyPhy: hypothesis testing using phylogenies. Bioinformatics 21: 676–679.
28. Dutzmann T, von Heřenesen O, Mayer F (2010) Evolution of nectarivory in phyllostomid bats (Phyllostomidae, Gray, 1825, Chiroptera: Mammalia). BMC Evol Biol 10: 165.
29. Teeling EC, Springer MS, Madsen O, Bates P, O’Brien SJ, et al. (2005) A molecular phylogeny for bats illuminates biogeography and the fossil record. Science 307: 389–384.
30. Murphy WJ, Ezirik E, O’Brien SJ, Madsen O, Scally M, et al. (2001) Resolution of the early placental mammal radiation using Bayesian phylogenetics. Science 294: 2348–2351.
31. Yang Z (2007) PAML 4: phylogenetic analysis by maximum likelihood. Mol Biol Evol 24: 1586–1591.
32. Yang Z (1996) Likelihood ratio tests for detecting positive selection and application to primate lysozyme evolution. Mol Biol Evol 15: 568–573.
33. Yang Z, Wong WS, Nielsen R (2003) Bayes empirical Bayes inference of amino acid sites under positive selection. Mol Biol Evol 21: 1107–1118.
34. Zhang J, Nielsen R, Yang Z (2005) Evaluation of an improved branch-site likelihood method for detecting positive selection at the molecular level. Mol Biol Evol 22: 2472–2479.
35. Kosakovsky Pond SL, Frost SDW (2003) A genetic algorithm approach to detecting lineage-specific variation in selection pressure. Mol Biol Evol 20: 470–485.
36. Kosakovsky Pond SL, Murrell B, Fourmant M, Frost SDW, Delpport W, et al. (2011) A random effects branch-site model for detecting episodic diversifying selection. Mol Biol Evol 28: 3033–3043.
37. Yang Z, Kumar S, Nei M (1995) A new method of inference of ancestral nucleotide and amino acid sequences. Genetics 141: 1641–1650.
38. Maddison WP, Maddison DR (2009) Mesquite: a modular system for evolutionary analysis. Version 2.74 Available: http://mesquiteproject.org. Accessed 2010 October 3.
39. Liu Y, Xu H, Yuan X, Rossiter SJ, Zhang S (2012) Multiple adaptive losses of alanine-glyoxylate aminotransferase mitochondrial targeting in fruit eating bats. Mol Biol Evol doi: 10.1093/molbev/mss013.

40. Bielawski JP, Yang Z (2003) Maximum likelihood methods for detecting adaptive evolution after gene duplication. J Struct Funct Genomics 3: 201–212.

41. Anisimova M, Bielawski JP, Yang Z (2002) Accuracy and power of Bayes prediction of amino acid sites under positive selection. Mol Biol Evol 19: 950–958.

42. Yang Z, Bielawski JP (2000) Statistical methods for detecting molecular adaptation. Trends Ecol Evol 15: 496–503.

43. Betts MJ, Russell RB (2003) Amino acid properties and consequences of substitutions. In: Barnes MK, Gray IC, eds. Bioinformatics for geneticists. West Sussex: Wiley.

44. Salas-Burgos A, Iserovich P, Zuniga F, Vera JC, Fischbarg J (2004) Predicting the three-dimensional structure of the human facilitative glucose transporter Glut1 by a novel evolutionary homology strategy: insights on the molecular mechanism of substrate migration, and binding sites for glucose and inhibitory molecules. Biophys J 87: 2990–2999.

45. Liu Y, Han N, Franchini LF, Xu H, Pisciottano F, et al. (2011) The voltage-gated potassium channel subfamily KQT member 4 (KCNQ4) displays parallel evolution in echolocating bats. Mol Biol Evol. (in press).

46. Davies KT, Cotton JA, Kirwan JD, Teeling EC, Rossiter SJ (2011) Parallel signatures of sequence evolution among hearing genes in echolocating mammals: an emerging model of genetic convergence. Heredity doi:10.1038/hdy.2011.119.

47. Liu Y, Rossiter SJ, Han X, Cotton JA, Zhang S (2010) Cetaceans on a molecular fast track to ultrasonic hearing. Curr Biol 20: 1834–1839.

48. Li G, Wang J, Rossiter SJ, Jones G, Cotton JA, et al. (2008) The hearing gene Prestin reunites echolocating bats. Proc Natl Acad Sci U S A 105: 13959–13964.

49. Haygood R, Fedrigo O, Hanson B, Yokoyama KD, Wray GA (2007) Promoter regions of many neural- and nutrition-related genes have experienced positive selection during human evolution. Nat Genet 39: 1140–1144.

50. Tarazona-Santos E, Fabbri C, Yeager M, Magalhaes WC, Burdett L, et al. (2010) Diversity in the glucose transporter-4 gene (SLC2A4) in humans reflects the action of natural selection along the Old-World primates evolution. PLoS ONE 5: e9827.

51. Yuan L, Zhao X, Lin B, Rossiter SJ, He L, et al. (2011) Adaptive evolution of Leptin in heterothermic bats. PLoS ONE 6: e27189.

52. Yang J, Wang ZL, Zhao XQ, Wang DP, Qi DL, et al. (2008) Natural selection and adaptive evolution of Leptin in the Ochotona family driven by the cold environmental stress. PLoS ONE 3: e1472.