RESEARCH ARTICLE

A community-science approach identifies genetic variants associated with three color morphs in ball pythons (*Python regius*)

Autumn R. Brown, Kaylee Comai, Dominic Mannino, Haily McCullough, Yamini Donekal, Hunter C. Meyers, Chiron W. Graves*, Hannah S. Seidel*, The BIO306W Consortium

Department of Biology, Eastern Michigan University, Ypsilanti, MI, United States of America

ID

The BIO306W Consortium comprises students in an undergraduate laboratory course at Eastern Michigan University. Students are listed in the Acknowledgments

* cgraves6@emich.edu (CWG); hseidel@emich.edu (HSS)

Abstract

Color morphs in ball pythons (*Python regius*) provide a unique and largely untapped resource for understanding the genetics of coloration in reptiles. Here we use a community-science approach to investigate the genetics of three color morphs affecting production of the pigment melanin. These morphs—Albino, Lavender Albino, and Ultramel—show a loss of melanin in the skin and eyes, ranging from severe (Albino) to moderate (Lavender Albino) to mild (Ultramel). To identify genetic variants causing each morph, we recruited shed skins of pet ball pythons via social media, extracted DNA from the skins, and searched for putative loss-of-function variants in homologs of genes controlling melanin production in other vertebrates. We report that the Albino morph is associated with missense and non-coding variants in the gene *TYR*. The Lavender Albino morph is associated with a deletion in the gene *OCA2*. The Ultramel morph is associated with a missense variant and a putative deletion in the gene *TYRP1*. Our study is one of the first to identify genetic variants associated with color morphs in ball pythons and shows that pet samples recruited from the community can provide a resource for genetic studies in this species.

Introduction

Color patterns are distinctive and beautiful features of many animal species. Among the many functions of color are to camouflage animals to their surroundings, protect tissues from ultraviolet radiation, warn predators of poisons, and provide signals for mating and social communication [1]. Color patterns have also been targeted by artificial selection to create novel and fanciful color patterns in domestic animals [2].

Colors are produced through a combination of chemical pigments and physical structures (structural colors). Pigments absorb light, whereas structural colors are created when light is reflected by the nanoscale geometry of a tissue. Common pigments include brown-to-black melanin, present throughout the animal kingdom, and yellow-to-red carotenoids and pteridines, common in birds, reptiles, and lower vertebrates [3]. Color-producing structures
include ordered keratin matrices in bird feathers [4], chitin layers in butterfly wings [5], and
purine crystals in the skin of fish, amphibians, and reptiles [6,7]. These color-producing struc-
tures can produce a variety of colors, including iridescent colors, depending on the wave-
lengths of light they reflect [8].

The genetics and development of color patterns in vertebrates have been studied most
extensively in mammals. Mammals largely rely on a single type of pigment (melanin), pro-
duced in the skin by a single type of cell (melanocyte) [9]. This system provides mammals with
skin and hair colors ranging from black to reddish-brown, depending on the chemical struc-
ture of the melanin [10].

Skin colors outside mammals are more diverse and include bright colors of all hues. This
increased complexity is produced through a combination of structural colors, melanin pigments,
and non-melanin pigments, and it relies on multiple types of specialized color-producing cells in
the skin [11–15]. Color cell development has been characterized to some extent in fish [16–18],
and genes controlling the use of non-melanin pigments have been identified in a few species of
fish and birds [19–25]. The genetics and development of non-mammalian color patterns are less
well understood in other vertebrates, particularly in reptiles [although see 26–29].

A unique resource for understanding the genetics of color patterns in reptiles is the ball
python (P. regius). Ball pythons are native to sub-Saharan Africa, but have become common as
pets in the United States [30]. Wild ball pythons exhibit a mottled color pattern, consisting of
brown-to-black melanin and red-to-yellow (non-melanin) pigments in the skin (Fig 1). Capti-
vite-bred ball pythons, by contrast, include many variants of the normal color pattern [26,31–
33]. These variants, referred to as ‘color morphs’, include animals having reduced melanin
(e.g. Albino), increased melanin (e.g. Cinnamon), reduced red-to-yellow coloration (e.g.
Axanthic), or complex changes in the placement of color patches on the skin (e.g. Spider,
Clown, Genetic Stripe, and Enchi). Many color morphs are heritable and show simple domi-
nant or recessive patterns of inheritance. These inheritance patterns are consistent with single-
gene causality, but only a single genetic variant associated with a ball python color morph (Pie-
bald) has been identified to date [26]. Ball pythons therefore represent a tractable yet largely
untapped resource for understanding the genetics of coloration in reptiles. An additional fea-
ture of ball pythons convenient for genetic studies is that DNA samples can be obtained non-
invasively from shed skin [34].

![Fig 1. The Albino, Lavender Albino, and Ultramel color morphs have reduced brown-to-black coloration, characteristic of a loss of melanin. Red-to-yellow coloration is unaffected in these morphs. Hypothesized causative genes represent genes in which loss-of-function variants in other vertebrates produce similar phenotypes (Table 1).](https://doi.org/10.1371/journal.pone.0276376.g001)
The goal of the current study was to perform proof-of-concept experiments to identify the genetic causes of color morphs in ball pythons, using shed skins of pet ball pythons recruited via social media. We focused on three morphs for which candidate genes could be readily identified: Albino, Lavender Albino, and Ultramel. These morphs show a loss of brown-to-black coloration in the skin and eyes, characteristic of a defect in melanin production (Fig 1). These morphs are recessive and non-allelic (i.e. crosses between these morphs yield offspring with normal coloration). Their loss of melanin ranges from severe (Albino) to moderate (Lavender Albino) to mild (Ultramel). This range of phenotypes mirrors the range of phenotypes observed for loss-of-function variants in genes required for melanin production in other vertebrates (Table 1). This similarity provided a list of candidate genes that we predicted might harbor loss-of-function variants causing the Albino, Lavender Albino, and Ultramel color morphs in ball pythons. Our study demonstrates the feasibility of using community-sourced samples for genetic studies in ball pythons and lays the groundwork for future investigations of reptile-specific color patterns in this species.

**Results**

**Obtaining a reference sequence for melanogenesis genes in a wildtype ball python**

Genes required for melanin production have been identified in humans and other vertebrates (Table 1). These genes encode enzymes that synthesize melanin (TYR, TYRP1, TYRP2) [87], a

---

| Gene | Species with identified loss-of-function variants | Role of protein | Loss-of-function phenotype |
|------|-----------------------------------------------|----------------|---------------------------|
| **TYR** | Fish | Japanese carp [35], Japanese rice fish [36] | Rate-limiting enzyme in melanin synthesis pathway | Severe loss of melanin |
| | Frogs and snakes | Japanese wild frogs (three species) [37], Japanese rat snake [38] | | |
| | Birds | Chicken [39-41] | | |
| | Ungulates | Water buffalo [42], Cattle [43], Red deer [44], Asinara donkey [45] | | |
| | Rodents | House mouse [46], Wistar rat [47], Domestic guinea pig [48] | | |
| | Primates | Crab-eating macaque [49], Capuchin monkey [50], Hamadryas baboon [51], Human [reviewed in 52] | | |
| | Other mammals | Humpback whale [53], Ferret [54], American mink [55], Domestic cat [56-58], Silver fox [59] | | |
| **OCA2** | Fish | Mexican cavefish [60], Lake Malawi cichlid [61] | Cl– channel regulating melanosome pH | Moderate loss of melanin |
| | Snakes | Corn snake [62] | | |
| | Mammals | Domestic dog [63], Bama miniature pig [64], House mouse [65], Human [reviewed in 52] | | |
| **TYRP1** | Fish | Zebrafish [66] | Enzymes contributing to melanin synthesis | Mild loss of melanin |
| | Birds | Saker falcon [67] | | |
| | Mammals | Chinese indigenous pig [68], Liangshan pig [69], Domestic dog [reviewed in 70], Domestic cat [58,71], Soay sheep [72], Valais Red sheep [73], American mink [74], Human [reviewed in 52] | | |
| **TYRP2** | Mammals | House mouse [75,76] | | |
| **SLC7A11** | Mammals | House mouse [77] | Solute transporters | |
| **SLC24A5** | Fish | Zebrafish [78] | | |
| | Mammals | Horse [79], Human [reviewed in 52] | | |
| **SLC45A2** | Fish | Japanese rice fish [80] | | |
| | Birds | Chicken and Japanese quail [81] | | |
| | Mammals | Domestic dog [82], Horse [83], House mouse [84], Bengal tiger [85], Western lowland gorilla [86], Human [reviewed in 52] | | |

[Link to Table 1: https://doi.org/10.1371/journal.pone.0276376.t001]
chloride channel required for maintaining the pH of melanosomes \((OCA2)\) [88], and transporters thought to import solutes into the cell or into organelles \((SLC7A11, SLC24A5, \text{and } SLC45A2)\) [77,89,90]. These genes are highly conserved among vertebrates and occur in single copy in most vertebrate genomes. Loss-of-function variants in these genes in humans cause a genetic disorder known as oculocutaneous albinism, which is characterized by loss of melanin in the skin, hair, and eyes [52]. This loss of melanin ranges from severe to mild, depending on the causative gene and genetic variants therein [52]. Similar phenotypes occur in other animals, where the loss of melanin extends to feathers, scales, and fur (Table 1).

To obtain a reference sequence for melanogenesis genes in ball python, we amplified and sequenced the coding regions of \(TYR, TYRP1, TYRP2, OCA2, SLC7A11, SLC24A5, \text{and } SLC45A2\) from a single ball python having normal coloration (henceforth ‘wildtype’). Primers for amplification were designed against the genome of Burmese python \((Python bivittatus)\), the closest relative of ball python for which genome sequence was available [91]. Comparison of ball python sequences to sequences from Burmese python revealed 97.2–99.2% nucleotide identity within coding regions and 96.9–98.5% nucleotide identity within flanking non-coding regions (Table 2). This analysis provided a reference sequence for genes in ball python that might harbor loss-of-function variants causing the Albino, Lavender Albino, and Ultramel color morphs.

### The Albino color morph is associated with three haplotypes of TYR

The Albino color morph in ball pythons is characterized by an absence or near absence of melanin in the skin and eyes—the brown-to-black coloration observed in wildtype is absent or severely reduced, and skin patches appear white or light beige (Fig 1D–1F). The Albino color morph is described by breeders as having three alleles \((Alb_{Albino}, Alb_{Candy}, \text{and } Alb_{Toffee})\), although prior to the current study, it remained unclear whether these alleles represented distinct molecular variants of the same gene or the same molecular variant discovered independently three times. We began our analysis of the Albino color morph by treating all animals within the color morph as a single group.

We hypothesized that the Albino color morph was caused by loss of function of \(TYR\), which encodes the enzyme catalyzing the rate-limiting step of melanin production. Loss of this

| Region   | Gene     | Total nucleotide sequence aligned (bp) | Nucleotide identity (%) | Amino acid identity (%) |
|----------|----------|----------------------------------------|-------------------------|-------------------------|
| Coding   | TYR      | 1,591                                  | 98.5                    | 99.6                    |
|          | TYRP1    | 1,584                                  | 99.2                    | 99.6                    |
|          | TYRP2    | 1,473                                  | 98.3                    | 99.1                    |
|          | OCA2     | 2,577                                  | 98.9                    | 99.7                    |
|          | SLC7A11  | 1,506                                  | 98.7                    | 99.7                    |
|          | SLC24A5  | 1,379                                  | 98.2                    | 99.5                    |
|          | SLC45A2  | 1,557                                  | 97.2                    | 98.9                    |
| Non-coding | TYR     | 577                                    | 96.9                    | (not applicable)        |
|          | TYRP1    | 1,160                                  | 97.0                    |                         |
|          | TYRP2    | 1,060                                  | 97.9                    |                         |
|          | OCA2     | 2,154                                  | 98.5                    |                         |
|          | SLC7A11  | 3,407                                  | 97.9                    |                         |
|          | SLC24A5  | 1,335                                  | 97.9                    |                         |
|          | SLC45A2  | 999                                    | 97.0                    |                         |

https://doi.org/10.1371/journal.pone.0276376.t002
enzymatic activity causes a severe loss of melanin [87], which is typically more severe than the loss of melanin caused by loss of function of other melanogenesis genes (Table 1). To test whether the Albino color morph was associated with variants in TYR, we performed a small-scale association study using 50 Albinos and 56 Non-Albinos, recruited from a total of 18 states in the United States. Polymorphic sites were identified through pilot sequencing of a subset of animals (see Methods). Six polymorphic sites were selected for the association study, two near the 5’ end of TYR and four near the 3’ end (Fig 2A). Each animal was genotyped at each site, and haplotypes were reconstructed using PHASE [92,93]. Association between haplotype and color morph was tested using the case-control test of PHASE. As a negative control, we also tested for association between haplotype and color morph at TYRP1, TYRP2, and OCA2, using two polymorphic sites per gene (S1 Table). This analysis revealed a significant association between haplotype and color morph at TYR (p = 0.04, Bonferroni corrected), but no association for the other three genes (p > 0.05, Bonferroni corrected) (Fig 2A). The association for TYR was driven by two features of the haplotype distribution: (i) haplotype diversity was reduced from a total of 15 TYR haplotypes among Non-Albinos to three haplotypes among Albinos (Fig 2A and 2C), and (ii) two of the three TYR haplotypes found in Albinos were rare among Non-Albinos (Fig 2A). These results demonstrate an association between the Albino color morph and variants in TYR; further, they show that all Albino animals were homozygous (34 animals) or compound heterozygous (16 animals) for any of three haplotypes of TYR.

We hypothesized that each of the three haplotypes of TYR found among Albinos might carry a distinct loss-of-function variant in the gene. To search for such variants, we selected one Albino animal homozygous for each haplotype and sequenced the TYR coding regions and adjacent splice sites in these animals. We found that one of these animals was homozygous for a missense variant in the third coding region of TYR (MZ269492:c.A3695G), which leads to an aspartic acid-to-glycine exchange [UPQ41330.1:p.(Asp394Gly)]. This variant is termed hereafter D394G. A second animal was homozygous for a different missense variant, also in the third coding region of TYR (MZ269492:c.C3665T), which leads to a proline-to-leucine exchange [UPQ41330.1:p.(Pro384Gly)]. This variant is termed hereafter P384L. Both variants alter conserved residues (Fig 2B), and the P384L variant occurs at the same site as a similar variant (P384A) associated with oculocutaneous albinism in humans [94]. The third animal carried no coding variants and no splice-site variants compared to wildtype. These results show that two of the three TYR haplotypes found among Albinos carried missense variants that are likely disruptive for TYR protein function. We term these TYR haplotypes TYRD394G and TYRP384L. We term the third Albino haplotype, which lacked coding or splice-site variants compared to wildtype, TYRAlbino.

We hypothesized that D394G, P384L, and an unidentified variant on the TYRAlbino haplotype were causative for the Albino color morph. This hypothesis predicted that all Albinos would be homozygous or compound heterozygous for these variants, whereas Non-Albinos would be heterozygotes and non-carriers. To test this prediction, we genotyped our full panel of 50 Albinos and 56 Non-Albinos for the missense variants D394G and P384L. Consistent with our prediction, we found that the Albinos were exclusively homozygous or compound heterozygous for D394G, P384L, or the TYRAlbino haplotype (Fig 2C). The Non-Albinos were exclusively heterozygotes or non-carriers (Fig 2C). The most common haplotype among Albinos was TYRAlbino (haplotype frequency of 71%), followed by TYRP384L (16%) and TYRD394G (13%). We conclude that D394G, P384L, and an unidentified variant on the TYRAlbino haplotype are likely causative for the Albino color morph. Any combination of these variants produces the Albino phenotype.
The TYR<sub>Albino</sub> haplotype lacks an obvious loss-of-function variant

The lack of coding or splice-site variants on the TYR<sub>Albino</sub> haplotype led us to hypothesize that this haplotype carried a loss-of-function variant in a non-coding region. To search for such variants, we examined the TYR promoter. We sequenced ~2 kb immediately upstream of the
TYR start codon in a \( \text{TYR}^{\text{Albino}} \) homozygote and compared this sequence to wildtype. We found that the \( \text{TYR}^{\text{Albino}} \) haplotype differed from wildtype at a total three sites (three single-base substitutions). In all three cases, the \( \text{TYR}^{\text{Albino}} \) haplotype was homozygous for the allele shared with Burmese python (i.e. the ancestral allele). This result suggests that these variants are not causative for the Albino color morph. We conclude that the causative variant in the \( \text{TYR}^{\text{Albino}} \) haplotype does not reside within the sequenced region, \( \sim 2 \text{ kb upstream of the TYR} \) start codon.

We hypothesized that the \( \text{TYR}^{\text{Albino}} \) haplotype might contain a large insertion or deletion (indel) in an intron that disrupts gene function. Intron indels can disrupt slicing and have been found to disrupt the function of melanogenesis genes in other species [e.g. 62, 74]. To search for large intronic indels, we tiled PCR amplicons across \( \text{TYR} \) introns and compared amplicon sizes between a \( \text{TYR}^{\text{Albino}} \) homozygote and a wildtype animal. These amplicons tiled across a total of \( \sim 38 \text{ kb of intronic sequence, which included part of intron 1 and all of introns 2, 3, and 4 (Fig 2E). (Tiling across intron 1 was incomplete because this intron contains a discontinuity in the Burmese python reference genome. Amplification across this discontinuity was unsuccessful in ball python, even in wildtype.) Size differences between the \( \text{TYR}^{\text{Albino}} \) homozygote and wildtype were assessed via standard agarose gel electrophoresis, which we estimate was sensitive enough to reveal indels larger than \( \sim 50-100 \text{ bp, depending on amplicon size (which ranged from } \sim 0.7-3.5 \text{ kb). This analysis revealed one size difference (Fig 2D). Further sequencing revealed that this size difference was caused by a variable number tandem repeat (VNTR) that was \( \sim 400 \text{ bp larger in the } \text{TYR}^{\text{Albino}} \) homozygote than in wildtype. Genotyping of the 56 Non-Albinos for this indel revealed that eight of these animals were homozygous for the larger allele of the VNTR (Fig 2D). This finding suggests that the larger allele of the VNTR is not causative for the Albino color morph. We propose that the \( \text{TYR}^{\text{Albino}} \) haplotype harbors a loss-of-function variant other than a large intronic indel or outside the genomic regions analyzed here.

\( \text{TYR}^{\text{P384L}} \) is associated with reduced phenotype severity compared to \( \text{TYR}^{\text{D394G}} \) and \( \text{TYR}^{\text{Albino}} \)

Ball python breeders describe the Albino color morph as having three alleles: \( \text{Alb}^{\text{Albino}}, \text{Alb}^{\text{Candy}}, \) and \( \text{Alb}^{\text{Toffee}}. \) Evidence for this view comes in part from variation in the coloration of Albino animals. In some Albino animals, the brown-to-black coloration observed in wildtype is entirely absent, and skin patches appear white (Fig 1D). These Albinos are typically considered to be \( \text{Alb}^{\text{Albino}} \) homozygotes. Other Albinos show a less severe phenotype, where skin patches are faintly beige or lavender instead of white (Fig 1E and 1F). These Albinos are typically considered to carry one or more copies of \( \text{Alb}^{\text{Candy}} \) or \( \text{Alb}^{\text{Toffee}}. \)

We hypothesized that the Albino alleles recognized by breeders might correspond to the \( \text{TYR}^{\text{haplotypes identified through sequencing (TYR}^{\text{D394G}}, \text{TYR}^{\text{P384L}}, \text{and TYR}^{\text{Albino}} \). To test this hypothesis, we examined the breeder designations of the 50 Albinos in our panel. We found that the \( \text{Alb}^{\text{Candy}} \) or \( \text{Alb}^{\text{Toffee}} \) designations typically corresponded the \( \text{TYR}^{\text{P384L}} \) haplotype (Fig 2C). The \( \text{Alb}^{\text{Albino}} \) designation typically corresponded to the other two haplotypes (\( \text{TYR}^{\text{D394G}} \) and \( \text{TYR}^{\text{Albino}} \). This correspondence was imperfect, and exceptions existed (e.g. three animals designated as \( \text{Alb}^{\text{Albino}} \) / \( \text{Alb}^{\text{Candy}} \) compound heterozygotes did not carry the \( \text{TYR}^{\text{P384L}} \) haplotype, Fig 2C). We conclude that the \( \text{Alb}^{\text{Candy}} \) and \( \text{Alb}^{\text{Toffee}} \) designations typically (but not exclusively) represent the \( \text{TYR}^{\text{P384L}} \) haplotype. The \( \text{Alb}^{\text{Albino}} \) designation typically (but not exclusively) represents the \( \text{TYR}^{\text{D394G}} \) or \( \text{TYR}^{\text{Albino}} \) haplotype. The association between the \( \text{Alb}^{\text{Candy}} \) and \( \text{Alb}^{\text{Toffee}} \) designations and the \( \text{TYR}^{\text{P384L}} \) haplotype suggests that this haplotype may confer a slightly less severe phenotype than do \( \text{TYR}^{\text{D394G}} \) and \( \text{TYR}^{\text{Albino}} \).
The Lavender Albino color morph is associated with a deletion in OCA2

The Lavender Albino color morph is characterized by skin patches that are lavender instead of brown or black. This phenotype is thought to arise from melanin levels that are dramatically reduced but not entirely eliminated. We hypothesized that this phenotype was caused by loss of function of OCA2, which encodes a chloride channel required for maintaining the pH of melanosomes [88,95]. When the OCA2 protein is absent or non-functional, the enzymes that synthesize melanin are less active, and only small amounts of melanin are produced [96,97]. The resulting phenotype is typically intermediate in severity between the loss-of-function phenotypes of TYR versus other melanogenesis genes (Table 1). OCA2 was therefore a good candidate for the causative gene of the Lavender Albino color morph.

To search for loss-of-function variants in OCA2, we selected a single Lavender Albino animal and sequenced 23 of the 24 coding regions of OCA2 in this animal. Comparison of these sequences to wildtype revealed no coding variants and no splice-site variants. We attempted to repeat this analysis for the remaining coding region (coding region 18), but we were unable to amplify this coding region from the Lavender Albino animal. Further test amplifications revealed that the Lavender Albino animal was homozygous for a 1,514-bp deletion spanning coding region 18 (Fig 3A and 3B). This deletion removes 36 amino acids from the protein and likely introduces a frameshift into the transcript, given that the coding regions downstream of the deletion are out of frame compared to coding regions upstream of the deletion (Fig 3E). This frameshifted transcript is predicted to produce a protein lacking six of the 12 transmembrane helices present in wildtype OCA2. Truncations occurring at similar positions in the OCA2 protein have been identified in humans and are associated with oculocutaneous albinism [98]. We conclude that the deletion of OCA2 coding region 18 likely disrupts protein function and is a strong candidate for the cause of the Lavender Albino color morph.

The Lavender Albino color morph is considered by breeders to have a single allele. We therefore predicted that the OCA2 deletion would be shared by other Lavender Albino animals. We predicted that Non-Lavender Albinos would be heterozygous for the deletion or non-carriers. To test this prediction, we genotyped the OCA2 deletion in 13 additional Lavender Albino animals. We also genotyped 76 Non-Lavender Albinos and one animal described as heterozygous for the Lavender Albino color morph. We found that all 13 Lavender Albinos were homozygous for the deletion (Fig 3C and 3D). All 76 Non-Lavender Albinos were non-carriers. The animal reported to be heterozygous for Lavender Albino was heterozygous for the deletion. These findings support the conclusion that the OCA2 deletion is causative for the Lavender Albino color morph (Fig 3E).

The Ultramel color morph is associated with a missense variant and a putative deletion in TYRP1

The Ultramel color morph is characterized by skin patches that are tan or light brown, rather than dark brown or black. This phenotype suggests a mild loss of melanin. We hypothesized that this phenotype was caused by loss of function of one of five genes: TYRP1, TYRP2, SLC7A11, SLC24A5, and SLC45A2 (Table 1). TYRP1 and TYRP2 encode enzymes involved in synthesizing melanin [87]. Their loss-of-function phenotypes are mild because of partially redundant with other enzymes in the melanin synthesis pathway. SLC7A11 encodes a transporter responsible for importing cystine into the cell [99]. Cystine is a precursor to some forms of melanin, and its reduction alters melanin levels [77]. SLC24A5 encodes a K^+-dependent Na^+-Ca^2+ exchanger [89]. SLC45A2 encodes a putative sugar transporter [90]. Loss of their encoded proteins reduces melanin through mechanisms that may involve defects in
regulation of melanosome pH [84,89,100]. These five genes were therefore good candidates for the causative gene of the Ultramel color morph.

To search for loss-of-function variants in \textit{TYRP1}, \textit{TYRP2}, \textit{SLC7A11}, \textit{SLC24A5}, and \textit{SLC45A2}, we selected a single Ultramel animal and sequenced the coding regions of each of these genes in this animal. Comparison of these sequences to wildtype revealed a single coding variant: a missense variant in the fourth coding region of \textit{TYRP1} (MZ269497:c.G1720A), which leads to an arginine-to-histidine exchange [UPQ41334.1:p.Arg305His]. The Ultramel animal was homozygous for this variant, termed hereafter \textit{R305H}. The arginine residue at this site is conserved across vertebrates (Fig 4A) and is also conserved in \textit{TYR}, which is paralogous to \textit{TYRP1} [101]. An arginine-to-histidine substitution at the homologous site in \textit{TYR} has been reported in humans and is associated with oculocutaneous albinism [102–104]; thus, histidine

Fig 3. The Lavender Albino color morph is associated with a deletion in \textit{OCA2}. (A) Schematic of the \textit{OCA2} gene. (B) PCR amplification demonstrating the deletion in Lavender Albino. (C) PCR amplification used for genotyping the deletion in the animals in panel D. (D) Genotypes of 14 Lavender Albinos, 76 Non-Lavender Albinos, and one animal described as heterozygous for Lavender Albino. This set of animals includes the original Lavender Albino animal in which the \textit{OCA2} deletion was identified. (E) Schematic of the \textit{OCA2} haplotype found in Lavender Albinos. (A, E) Hash mark, discontinuity in the Burmese python reference genome.

https://doi.org/10.1371/journal.pone.0276376.g003
at this site is likely disruptive to protein function. The R305H variant is therefore a good candidate for the cause of the Ultramel color morph.

The Ultramel color morph is considered by breeders to have a single allele. We therefore predicted that the R305H variant would be shared by other Ultramels. We predicted that Non-Ultramels would be heterozygous or non-carriers. To test this prediction, we genotyped the R305H variant in 10 additional Ultramels and 78 Non-Ultramels. We found that five of the 10 Ultramels were homozygous for the R305H variant (Fig 4B). Of the remaining Ultramels, two were heterozygous for the R305H variant, and three did not carry it. By contrast, none of the 78 Non-Ultramels carried the R305H variant. This pattern is consistent with the R305H variant causing the Ultramel phenotype in some Ultramels but not others.
We hypothesized that the R305H variant represented one of two loss-of-function alleles of TYRP1. Under this scenario, Ultramels lacking the R305H variant are predicted to be homozygous for an alternate loss-of-function allele. Ultramels heterozygous for R305H are predicted to be compound heterozygotes. To test this prediction, we amplified and sequenced coding regions 1 through 5 of TYRP1 in one of the three Ultramels lacking the R305H variant. We found no coding variants and no splice-site variants compared to wildtype. We attempted to repeat this analysis for coding regions 6 and 7 of TYRP1, but we were unable to amplify these coding regions from this animal (Fig 4C). Identical results were observed for the other two Ultramels lacking the R305H variant (S5 Table). Further test amplifications indicated that coding regions 6 and 7 were missing from the genomes of all three animals (Fig 4C, S5 Table). Loss of these coding regions was specific to Ultramels lacking the R305H variant and did not occur in any of the Non-Ultramels, nor in the Ultramels homozygous or heterozygous for R305H (Fig 4B). A simple explanation of this pattern is that the Ultramels lacking R305H were homozygous for a deletion spanning coding regions 6 and 7 of TYRP1. The two Ultramels heterozygous for R305H were presumed to be compound heterozygous for this putative deletion, although we were not able to confirm this heterozygosity because the putative deletion extended into a discontinuity in the Burmese python reference genome (and therefore could not be detected in heterozygotes). The putative deletion removes at least 2,208 bp from the genome and 117 amino acids from the TYRP1 protein, including the second of two zinc-binding domains. Truncations occurring at similar positions in the TYRP1 protein have been identified in humans and are associated with oculocutaneous albinism [105]. We conclude that the Ultramel color morph is likely caused by variants in TYRP1 and has two alleles: missense variant R305H and a putative deletion of coding regions 6 and 7 (Fig 4D).

**Discussion**

The goal of our study was to use pet samples recruited from the community to identify the genetic causes of the Albino, Lavender Albino, and Ultramel color morphs. We succeeded in recruiting 11 or more animals for each morph, along with a larger number of animals not belonging to these morphs. This sample size, albeit small, was sufficient to identify putatively causal variants for each morph (Fig 5). The Albino color morph was associated with three alleles of TYR: missense variant D394G, missense variant P384L, and haplotype TYR<sup>Albino</sup>, which lacks coding or splice-site variants compared to wildtype. The Lavender Albino color morph was associated with a single allele of OCA2, a 1,514-bp deletion that removes OCA2 coding region 18. The Ultramel color morph was associated with two alleles of TYRP1: missense variant R305H and a putative deletion that removes TYRP1 coding regions 6 and 7. Due to the small sample size of our study, we cannot exclude the possibility of additional loss-of-function alleles of these genes segregating in the ball python population. However, such alleles are expected to be at low frequency, given their absence in our sample. These findings are consistent with genetic data from other vertebrates (Table 1), indicating that the loss-of-function phenotypes of these genes range from severe (TYR) to moderate (OCA2) to mild (TYPR1). Our study demonstrates that pet ball pythons are a tractable resource for genetic analysis of coloration in reptiles, at least for color morphs having obvious candidate genes.

**Molecular functions of genetic variants associated with color morphs**

The missense variants and deletions found in TYR, TYRP1, and OCA2 are likely hypomorphic or null alleles. The deletion in OCA2 likely results in a frameshifted transcript that, if not degraded by nonsense-mediated decay, encodes a protein lacking six of the 12 transmembrane helices present in wildtype OCA2 [95]. This truncated protein is therefore unlikely to retain the Cl⁻ channel activity observed for wildtype OCA2 [88]. TYR and TYPR1 encode globular
enzymes that require copper or zinc as co-factors [87]. Variants P384L and D394G in TYR reside in the second of TYR’s two copper-binding domains and may therefore alter the ability of the TYR to bind copper. Variant R305H in TYRP1 occurs at a site that normally forms salt bridges with residues located elsewhere in the peptide chain [87]. An arginine-to-histidine substitution at this site likely disrupts these salt bridges and may therefore interfere with proper protein folding. The putative deletion in TYRP1 removes the C-terminal ~20% of the protein. This deleted region includes the second of TYRP1’s two zinc-binding domains and several alpha helices residing to the enzyme’s hydrophobic core [87]. Truncated TYRP1 protein is therefore unlikely to fold properly, nor is it likely to properly associate with zinc.

The TYR\textsuperscript{Albino} haplotype is the most common TYR haplotype found among Albinos, but it lacks an obvious loss-of-function variant. This haplotype contains no coding variants, no splice-site variants, no derived variants within 2 kb upstream of the start codon, and no large indels in three of four introns (with the exception of a longer allele of a VNTR, which is not specific to the TYR\textsuperscript{Albino} haplotype). We propose that the TYR\textsuperscript{Albino} haplotype contains a loss-of-function variant not detectable by methods used in the current study. Examples include regulatory variants farther upstream of the start codon, substitutions in introns that disrupt splicing, or large indels or rearrangements involving the first intron. Cryptic loss-of-function variants in melanogenesis genes are thought to be relatively common in humans, where ~10–20% of oculocutaneous albinism patients are heterozygous for pathogenic variants in a known melanogenesis gene but lack a coding or splice-site variant on the opposite allele [106–108].

### Multiple alleles of the Albino color morph

Prior to the current study, the Albino color morph was recognized by breeders as having three alleles: Alb\textsuperscript{Albino}, Alb\textsuperscript{Candy}, and Alb\textsuperscript{Toffee}. Alb\textsuperscript{Albino} was thought to confer a more severe
phenotype than \textit{Alb\textsubscript{Candy}} and \textit{Alb\textsubscript{Toffee}} (Fig 1D–1F). Our results confirm that the Albino color morph has three molecular alleles (\textit{TYR\textsuperscript{D394G}}, \textit{TYR\textsuperscript{P384L}}, and \textit{TYR\textsubscript{Albino}}). Yet these alleles do not perfectly correspond to the alleles recognized by breeders. The molecular allele \textit{TYR\textsuperscript{P384L}} typically corresponds to the breeder designations \textit{Alb\textsubscript{Candy}} and \textit{Alb\textsubscript{Toffee}}. The molecular alleles \textit{TYR\textsuperscript{D394G}} and \textit{TYR\textsuperscript{Albino}} typically correspond to the breeder designation \textit{Alb\textsubscript{Albino}} (Fig 2C). This correspondence is imperfect and exceptions exist (Fig 2C).

We speculate that the \textit{TYR\textsuperscript{P384L}} haplotype was discovered twice and was named \textit{Alb\textsubscript{Candy}} by one breeder and \textit{Alb\textsubscript{Toffee}} by another breeder. We speculate that \textit{TYR\textsuperscript{D394G}} and \textit{TYR\textsuperscript{Albino}} confer similar phenotypes and were not previously recognized by breeders as distinct. The less severe phenotype associated with \textit{TYR\textsuperscript{P384L}} may reflect the \textit{P384L} variant being less disruptive to gene function compared with variants on the other two alleles. Alternatively, the less severe phenotype associated with \textit{TYR\textsuperscript{P384L}} may reflect genetic linkage to modifiers in other genes. In either case, our study demonstrates that the Albino allele designations current in use by breeders do not accurately reflect molecular genotype. Renaming of the Albino alleles is warranted, although owners and breeders may be resistant to renaming due to cultural attachment to existing allele names.

**Multiple alleles of the Ultramel color morph**

Our finding of multiple \textit{TYRP1} alleles for the Ultramel color morph was unexpected. This morph was not previously described as having multiple alleles. We speculate that one of the two alleles associated with the Ultramel color morph may represent an allele originally associated with a morph known as Caramel Albino. Multiple lineages of Caramel Albinos have been described, and their coloration similar to Ultramels. Caramel Albinos have been disfavored among owners and breeders because of spinal kinking and reduced female fertility. Caramel Albinos were not included in our study because we were unable to recruit any Caramel Albino samples. We speculate that the Caramel Albino morph may be allelic with Ultramel, and that one of two \textit{TYRP1} alleles associated with Ultramel may represent an allele originally described as Caramel Albino. Alternately, Ultramel may be distinct from Caramel Albino, and the two \textit{TYRP1} alleles may represent two distinct origins of the Ultramel morph.

**Prospects for genetic testing**

Many breeders of ball pythons wish to identify heterozygotes of recessive color morphs. Currently the only tool for identifying heterozygotes is test breeding, which is slow and laborious. The results of the current study will enable simple genetic testing for Albino, Lavender Albino, and Ultramel, which will allow heterozygotes to be identified prior to reproductive maturity. Testing for Lavender Albino can be performed by genotyping the \textit{OCA2} deletion. Testing for Albino can be performed by sequencing the \textit{TYR} missense variants \textit{D394G} and \textit{P384L}, and by genotyping variants that distinguish the \textit{TYR\textsubscript{Albino}} haplotype from other \textit{TYR} haplotypes. Detection of \textit{D394G} or \textit{P384L} can be considered diagnostic for Albino because these variants are likely causative for the Albino color morph. Detection of the \textit{TYR\textsubscript{Albino}} haplotype is less diagnostic because the causative variant on \textit{TYR\textsubscript{Albino}} haplotype remains unknown. Testing for one of the two Ultramel alleles can be performed by sequencing the \textit{TYRPI} missense variant \textit{R305H}. The other Ultramel allele—the putative deletion of \textit{TYRP1} coding regions 6 and 7—cannot currently be detected in heterozygotes using simple methods because the breakpoints of this putative deletion are unknown.

**Use of Burmese python reference genome**

One challenge for genetic studies in ball python is the absence of a reference genome. To fill this gap, we relied on the genome of Burmese python [91]. This genome is a scaffold-level
assembly, and some genes are fragmented across scaffolds. This fragmentation limited our analysis of the \textit{TYR}^{ Albino } haplotype and the putative deletion in \textit{TYRP1}. For \textit{TYR}^{ Albino }, a discontinuity in the first intron of \textit{TYR} in the Burmese python genome prevented us from assessing whether the \textit{TYR}^{ Albino } haplotype contained a large indel or rearrangement in this intron. For the putative deletion in \textit{TYRP1}, a discontinuity immediately downstream of \textit{TYRP1} prevented us from mapping the exact breakpoints of the deletion. As genomic resources for Burmese python and ball python expand, ideally with the use long-read technologies to improve genome assemblies and identify structural variants, we expect to identify a putatively causative variant on the \textit{TYR}^{ Albino } haplotype and to map the breakpoints of the putative deletion in \textit{TYRP1}. This information will increase the prospects for genetic testing for these alleles.

**Ball pythons as a genetic system**

The market for pet ball pythons is huge, and many owners prefer pets with novel color patterns. This demand has led breeders to propagate genetic variants affecting coloration [31,33]. This effort has fortuitously created a collection of “mutants” useful for understanding the genetics of coloration in reptiles. Examples include ball python morphs with altered red-to-yellow coloration (e.g. Axanthic) and morphs in which the normal mottled color pattern is converted to dorsal stripes (e.g. Clown, Genetic Stripe, Super Stripe). Red-to-yellow pigments are largely uncharacterized in reptiles [although see 28,29], and morphs affecting these pigments may provide insight into the metabolism and storage of these pigments. Morphs with stripes are reminiscent of evolutionary changes in color patterning across snake species [109] and may provide insights into the developmental mechanisms by which these changes evolve.

Future studies of ball python color morphs will be aided by our groundwork showing that ball python samples can be recruited effectively from pet owners, and that genetic analyses in ball python can be scaffolded using the genome of Burmese python as a reference. Similar groundwork has also been provided by a recent study characterizing the ball python morph known as Piebald [26]. We expect that a continued community-science approach will be effective in developing ball pythons into a resource for understanding the genetics of reptile coloration.

**Materials and methods**

**Recruitment of ball python sheds**

Ball python sheds were recruited from pet owners and breeders by placing announcements in Twitter, Reddit, Instagram, and Facebook, and by contacting sellers having active listings on Morph Market (www.morphmarket.com). Contributors were instructed to allow sheds to dry (if wet) and to ship sheds via standard first-class mail. Contributors sending multiple sheds were instructed to package sheds individually in plastic bags during shipping. Contributors were not provided monetary compensation for sheds, although some contributors were given pre-paid shipping envelopes to cover shipping costs. Contributors were thanked via social media whenever possible. Upon receipt, sheds were stored at -20°C to kill any insect larvae infesting the sheds.

To maximize genetic diversity within each category of morph, we limited our sample of animals of any one morph to animals contributed by different contributors. Exceptions were made if animals contributed by the same contributor had been obtained from different breeders. The goal of this strategy was to reduce the number of animals that were close relatives (e.g. siblings or parent-offspring pairs). Obtaining full pedigree information was not possible because most owners lacked this information (e.g. “I bought my animal at a pet store” or “I got my animal from a friend who was moving away”). The idea that animals in our sample were derived from multiple lineages is supported by our discovery of multiple alleles for the Albino...
and Ultramel color morphs. However, we cannot exclude the possibility that some animals in our sample may have been close relatives.

The total set of animals comprised 50 Albinos, 14 Lavender Albinos, 11 Ultramels, one animal described as heterozygous for Lavender Albino, and 46 animals described as having normal coloration (i.e. wildtype) or belonging to color morphs other than Albino, Lavender Albino, or Ultramel. The Albinos were composed of 36 animals described as Alb\textsuperscript{Albino} homozygotes, three animals described as Alb\textsuperscript{Candy} homozygotes, one animal described as an Alb\textsuperscript{Albino} / Alb\textsuperscript{Candy} compound homozygote, nine animals described as an Alb\textsuperscript{Albino} / Alb\textsuperscript{Toffee} compound homozygote, and one animal described as an Alb\textsuperscript{Albino} / Alb\textsuperscript{Toffee} compound homozygote. We use the term Non-Albino to refer to animals having normal coloration or belonging to morphs other than Albino. The Non-Albinos included some Lavender Albinos and some Ultramels. The terms Non-Lavender Albino and Non-Ultramel were used in analogous ways.

Phenotypes of Albinos, Lavender Albinos, and Ultramels were confirmed by examining shed skins for a reduction of brown-to-black coloration. The single animal described as heterozygous for Lavender Albino was included to demonstrate the three-primer assay for genotyping the OCA2 deletion. Heterozygotes for Albino or Ultramel and additional heterozygous for Lavender Albino were excluded from our study because we have found that animals described as heterozygous for recessive traits by breeders and owners are sometimes not heterozygous for these traits (e.g. due to mis-attribution of paternity).

Performing experiments in an undergraduate laboratory course

The majority of experiments and analyses described in this study were performed by undergraduate students as part of a laboratory course at Eastern Michigan University (BIO306W). This practice required that our experimental design rely on simple techniques, namely PCR and Sanger sequencing. To avoid student errors in these techniques, we implemented the following precautions. First, students never extracted DNA from more than one animal within the same laboratory period. Second, students performed negative and positive control reactions for all PCR amplifications. Data from students having incorrect controls were excluded from analysis. Third, all sequence analyses were performed independently by three or more students. When the results of all students did not all agree, sequences were re-analyzed by the instructor (HSS).

Annotation of melanogenesis genes in Burmese python

Our analyses in ball python used the genome of Burmese python as a reference. We therefore required high-confidence gene annotations in Burmese python for the genes used in our study (TYR, TYRP1, TYRP2, OCA2, SLC7A11, SLC24A5, and SLC45A2). Preliminary inspection of existing gene annotations in Burmese python suggested that many gene annotations contained errors (e.g. missing one or more coding regions). A main cause of these errors was that the Burmese python genome (Python\textit{molurus bivittatus}-5.0.2) is a scaffold-level assembly, and many genes were split across scaffolds. We therefore curated new gene annotations in Burmese python, using conservation of gene structure across species. Alignment of protein sequences for each gene from corn snake, anole lizard, chicken, and/or mouse revealed that gene structure was highly conserved for all seven genes: Genes in each species contained the same number of coding regions, and the coding-region boundaries relative to protein sequences were perfectly conserved across species, with the exception of one slight difference in mouse for one boundary of SLC45A2 (S2 Table). We therefore felt confident that conservation of coding-region boundaries could be used to curate new gene annotations in Burmese python. While it remains theoretically possible that our new gene annotations contain minor errors, our
To curate new gene annotations, we performed TBLASTN [110,111] searches of the Burmese python genome (Python_molurus_bivittatus-5.0.2) using protein sequences from anole lizard or corn snake as the query. TBLASTN is a search tool that aligns a query protein sequence all six reading frames of a nucleotide database. TBLASTN searches were performed using default parameters. Hits were examined manually. Gene annotations were built to match the coding-region boundaries conserved across species. Details for each gene annotation are given below.

**TYR.** The Burmese python genome was queried using TYR protein sequence from anole lizard (XP_003219419.1). The N-terminus of this query (coding region 1) hit Burmese python transcript XM_007438960.1 on scaffold 4418. The C-terminus of this query (coding regions 2 to 5) hit Burmese python transcript XM_007436041.2 on scaffold 3103. TYR was annotated as the union of these transcripts. The 3' boundary of coding region 1 was adjusted to match the boundary conserved across species (…TTC TCT TCA TGG CAA-3').

**TYRP1.** The Burmese python genome was queried using TYRP1 protein sequence from corn snake (XP_034266320.1). The N-terminus of this query (coding regions 1 to 5) hit Burmese python transcript XM_007426971.3 on scaffold 801. The C-terminus of this query (coding regions 6 and 7) hit unannotated regions on this same scaffold. TYRP1 was annotated as the union of transcript XM_007426971.3 and coding regions 6 and 7. The boundaries of coding regions 6 and 7 were built to match the boundaries conserved across species (coding region 6, 5'-ATA TCT CAA CAT ACC…AAG TTC AGT GGC CAT-3'; coding region 7, 5'-CAC AAG CTC TCC ATG…CAG TCA GAT GTG TGA-3').

**TYRP2.** The Burmese python genome was queried using TYRP2 protein sequence from corn snake (XP_034273310.1). The N-terminus of this query (coding regions 1 to 3) hit unannotated regions on Burmese python scaffold 4970. The C-terminus of this query (coding regions 4 to 8) hit Burmese python transcript XM_025174219.1 on this same scaffold. TYRP2 was annotated as the union of coding regions 1 to 3 and C-terminal five coding regions of transcript XM_025174219.1. Transcript XM_025174219.1 contains a sixth coding region that does not match TYRP2 and was therefore excluded from the gene annotation. The boundaries of coding regions 1 to 3 were built to match the boundaries conserved across species (coding region 1, 5'-ATG GCC TTC CTG CTG…GTT GCC AAT GCA CAG-3'; coding region 2, 5'-GAC ATT TTG CTG GCT…GAG ATA CTC TAT TAG-3'; coding region 3, 5'-GAC CAG GCC GTC CCT…GAA AGA GAT CTG CAG-3').

**OCA2.** The Burmese python genome was queried using OCA2 protein sequence from corn snake (XP_034273310.1). The N-terminus of this query (coding regions 1 to 9) hit Burmese python transcript XM_025173964.1 on scaffold 4704. The C-terminus of this query (coding regions 10 to 24) hit Burmese python transcript XM_007433276.1 on scaffold 2194. OCA2 was annotated as the union of these transcripts. The 5' boundary of coding region 10 was adjusted to match the boundary conserved across species (5'-ATT GTC CAC AGG ACA…).

**SLC7A11.** The Burmese python genome was queried using SLC7A11 protein sequence from corn snake (XP_034257397.1). All regions of the query hit Burmese python transcript XM_007430906.3 on scaffold 1543. SLC7A11 was annotated as transcript XM_007430906.3, with no further adjustments.

**SLC24A5.** The Burmese python genome was queried using SLC24A5 protein sequence from corn snake (XP_034290605.1) extended at its N-terminus to an upstream start codon present in the parent transcript (XM_034434714.1). The N-terminus of the query (coding region 1) hit an unannotated region on Burmese python scaffold 3984. The C-terminus of this query (coding regions 2 to 9) hit Burmese python transcript XM_007438007.2 on this same scaffold.
scaffold. SLC24A5 was annotated as the union of coding region 1 and transcript XM_007438007.2. The boundaries of coding region 1 were built to match the boundaries conserved across species (5'-ATG CAG CCT GCC GAG...TCC GCG AGG ATC CCG -3'). The 5' boundary of coding region 2 was adjusted to match the boundary conserved across species (5'-AGA ACG AAA CCC GCT...).

SLC45A2. The Burmese python genome was queried using SLC45A2 protein sequence from corn snake (XP_034298386.1). The N-terminal region of this query (coding regions 1 and 2) hit Burmese python transcript XM_007432459.2 on scaffold 1939. The C-terminal region of this query (coding regions 5 to 7) hit Burmese python transcript XM_007437721.2 on scaffold 3858. SLC45A2 was annotated as the union of these transcripts.

DNA extraction
Sheds were rinsed in tap water to remove dirt and debris. Sheds were air dried and lysed overnight at ~60°C in ~1 ml lysis buffer (100 mM Tris-HCl pH 8.0, 100 mM EDTA, 2% sodium dodecyl sulfate, 3 mM CaCl2, 2 mg/ml Proteinase K) per ~8 cm2 piece of shed. Lysate was separated from visible fragments of undigested shed and further cleared by centrifugation at 13,000 x g for 2 min. To precipitate protein, ammonium acetate was added to supernatant to a final concentration of 1.875 M. Samples were incubated on ice for 5–10 min and centrifuged at 13,000 x g for 3–5 min at 4°C. Supernatant was mixed with an equal volume of magnetic bead mixture (10 mM Tris-HCl pH 8.0, 1 mM EDTA, 1.6 M NaCl, 0.2% Tween-20, 11% polyethylene glycol, 0.04% washed SpeedBeads [Sigma #GE45152105050250]), and samples shaken for 5–10 min. Beads were separated from supernatant using a magnet, washed twice in 0.2 ml 70% ethanol for 2 min, and air dried for ~1 min. DNA was eluted from beads in TE buffer (10 mM Tris-HCl pH 8.0, 1 mM EDTA) at 65°C for >5 min.

Primer design and PCR
Primers were designed against the genome of Burmese python or against genomic sequences from ball python obtained in an earlier step of the study. Primers were designed using Primer3 [112], using default parameters and a target annealing temperature of 60°C. Amplification was first tested at 57°C, to allow for occasional divergence between ball python and Burmese python genomic sequences. In some cases, annealing temperatures were later adjusted to 52°C, 60°C, or 61°C, to obtain stronger product or to eliminate background bands. Genomic fragments were amplified using OneTaq polymerase (NEB #M0480) or Q5 polymerase (NEB #M0491). Genotyping assays described below used OneTaq, unless otherwise specified. OneTaq reactions consisted of 1X OneTaq Standard Reaction Buffer, 200 μM dNTPs, 0.2 μM of each primer, and 0.025 U/μl OneTaq polymerase. OneTaq thermocycling conditions were as follows: 94°C for 2 min; 30–35 cycles of 94°C for 30 sec, 52–61°C for 30 sec, and 68°C for 1–4 min; and 68°C for 5 min. Q5 reactions consisted of 1X Q5 Reaction Buffer, 200 μM dNTPs, 0.5 μM of each primer, and 0.02 U/μl Q5 polymerase. Q5 thermocycling conditions were as follows: 98°C for 30 sec; 30–35 cycles of 98°C for 10 sec, 58–61°C for 15 sec, and 72°C for 1.5–3 min; and 72°C for 5 min. Reactions used 10–100 ng template DNA per 20 μl volume.

Sanger sequencing
PCR products were purified for Sanger sequencing using magnetic beads or gel extraction. For magnetic-bead purification, PCR reactions were mixed with three volumes of magnetic-bead mixture (10 mM Tris-HCl pH 8.0, 1 mM EDTA, 1.6–2.5 M NaCl, 0.2% Tween-20, 11–20% polyethylene glycol, 0.04% washed SpeedBeads [Sigma #GE45152105050250]), and agitated...
for 5 min. Beads were separated from supernatant using a magnet, washed twice in 0.2 ml 80% ethanol for >30 sec, and air-dried for 30 sec. PCR products were eluted from beads in 10 mM Tris-HCl pH 8.0 for >3 min at 65˚C. Gel extraction was performed using QIAquick Gel Extraction Kit (Qiagen #28704), according to the manufacturer guidelines. Sanger sequencing was performed by Eton Bioscience Inc (etonbio.com).

**Sequencing of coding regions and comparison to Burmese python**

Coding regions of melanogenesis genes were amplified and sequenced using primers given in S3 Table. Chromatograms were trimmed using SnapGene Viewer (snapgene.com/snapgene-viewer) and aligned to one another or to genomic sequences from Burmese python using ApE (jorgensen.biology.utah.edu/wayned/ape). Alignments were examined manually to identify divergent and polymorphic sites. Sequence identity between ball python and Burmese python was calculated across alignable sequence, excluding indels.

**Association study genotyping**

Variants for the association study were identified through amplification and pilot sequencing of genomic fragments from five animals (four Non-Albinos and one Albino). For TYR, TYRP1, and TYRP2, we identified one or more genomic fragments containing multiple polymorphic sites within the same amplicon. These genomic fragments were selected for the association study. For OCA2, we did not identify any genomic fragments containing multiple polymorphic sites; we therefore selected two genomic fragments each containing a single polymorphic site. Divergence between the Albino animal and the other four animals was not a criterion for inclusion of variants in the association study. Variants were genotyped by amplifying and sequencing genomic fragments containing the variants. Locations of variants and primers used for amplifying and sequencing variants are given in S1 Table. Genotypes are provided in S6 Table.

**Haplotype reconstruction**

Haplotypes were reconstructed using PHASE version 2.1 [92,93]. Parameters were set to 200 iterations, a thinning interval of 2, and a burn-in parameter of 100. These settings produced identical or nearly identical output for seven runs seeded with different random numbers; thus, these settings met the criteria for effective choice of parameter settings, according to the PHASE documentation. The case-control permutation test was performed by comparing Albinos to Non-Albinos.

**Genotyping assays for TYR**

TYR missense variants D394G and P384L were genotyped by amplifying and sequencing a genomic fragment containing TYR coding region 3. This fragment was amplified using primers 13F (5’-ACT TTC AGG TGG GCA GCA G-3’) and 13R (5’-GCT GAC AAC TAA AAT CTC TGC AA-3’) and an annealing temperature of 52˚C, or using primers 242F (5’-GCC ATT GTA GCT TCT TAC CAC TC-3’) and 242R (5’-TTC CAG TCC ATA TAC AAG ATA TCC AA-3’) and an annealing temperature of 57˚C. The amplicon was sequenced using primer 13F or 242F. Genotypes are provided in S6 Table.

TYR promoter and intronic regions are amplified using primers given in S4 Table. Promoter fragments were sequenced in full, and intronic fragments were sequenced from one end, to confirm that the correct region of the genome had been amplified. Sequencing primers are given in S4 Table. Size differences among intronic fragments were assessed by separating
fragments on a 1.25% agarose gel. Gels were run long enough for the shortest ladder band (100 bp) to migrate ~8 cm from its starting position.

Allele sizes of the variable number tandem repeat (VNTR) in TYR were genotyped by amplifying a genomic fragment located in TYR intron 3, using primers 214F (5'-TCT CAC CTG ATG GCA CAT TC-3') and 209R (5'-GTG CCC ACC CTG ATG TTA TT-3') and an annealing temperature of 60˚C. Amplicon sizes were analyzed as for intronic fragments, described above.

Genotyping assays for OCA2

The OCA2 deletion was initially identified by amplifying and sequencing a genomic fragment spanning OCA2 coding region 18. This fragment was amplified using primers 218F (5'-ACC CCG TAG CCT CTT CAA AT-3') and 166R (5'-TGG GTG GCA AAC AAT CAT AA-3'), an annealing temperature of 60˚C, and Q5 polymerase. Amplicons were sequenced using both primers.

The OCA2 deletion was genotyped after its initial identification using a three-primer PCR assay. This assay used one forward primer and two reverse primers. The forward primer and one of the reverse primers were located outside the deletion: 217F (5'-GGA GAG AGA ATC CAA CCC TTG -3') and 166R (5'-TGG GTG GCA AAC AAT CAT AA-3'). The second reverse primer was located within the deletion: 188R (5'-CAA AGA CCA TTG TCC ATT TCC -3'). The annealing temperature was 57˚C. This assay produces a 429-bp product for the wildtype allele and a 349-bp product for the deletion allele. Heterozygotes produce both products. Genotypes are provided in S6 Table.

Genotyping assays for TYRP1

TYRP1 missense variant R305H was genotyped by amplifying and sequencing a genomic fragment spanning TYRP1 coding region 4. This fragment was amplified using primers 18F (5'-GCT CTT TTC TCT AAG TCT GAC CTC -3') and 18R (5'-TCT TGT CCC ACA AAA GGA TTT -3') and an annealing temperature of 57˚C. The amplicon was sequenced using primer 18F.

The putative deletion of TYRP1 coding regions 6 and 7 was identified using primers given in S5 Table. The putative deletion was genotyped after its initial identification by amplifying a genomic fragment of TYRP1 spanning coding regions 6 and 7. This fragment was amplified using primers 20F (5'-GCA TTG TTT TAT CAG CCA TGA A-3') and 21R (5'-GGA ATT GAG ACA AAT CCT TGG-3') and an annealing temperature of 57˚C. These primers reside within the putative deletion and therefore cannot distinguish between animals lacking the putative deletion and animals heterozygous for it. Presence of a PCR product in this assay indicates that TYRP1 coding regions 6 and 7 are present on at least one chromosome. Absence of a PCR product in this assay indicates that TYRP1 coding regions 6 and 7 are absent on both chromosomes. We were unable to design a PCR assay to detect the putative deletion in heterozygotes because the putative deletion extends into a discontinuity in the Burmese python reference genome. Presence or absence of a PCR product in this assay is provided in S6 Table.

Protein sequence alignment

Protein sequences were aligned using Clustal Omega [113], using default parameters.

Supporting information

S1 Fig. Raw gel images. (A) Raw image of the gel displayed in Fig 2. (B) Raw image of the gel displayed in Fig 3. (C) Raw image of the gel displayed in Fig 4. Brightness and contrast settings...
have not been adjusted in these images. x, experiment unrelated to the current study.

S1 Table. Variants used in the Albino association study and primers to genotype these variants.

S2 Table. Conservation of gene structure of melanogenesis genes.

S3 Table. Primers to amplify and sequence coding regions of melanogenesis genes.

S4 Table. Primers to amplify non-coding regions of TYR.

S5 Table. Primers to investigate the putative deletion in TYRP1.

S6 Table. Genotypes.

Acknowledgments

We thank Matt Rockman and Katy Greenwald for advice on haplotype reconstruction; Bob Winning, Anne Casper, David Kass, and two anonymous reviewers for comments on the manuscript; and the Educational Course Support program of New England BioLabs for reagents used in undergraduate teaching labs. We thank the following individuals for contributing ball python sheds: Adam and Nicole Schmid; Alycia Butler; Amanda Hall; Andelyn Czajka; Brad Carter of Driftless Reptiles; Bryan Rivera; Chiron Graves; Chun Ku of Dynasty Reptiles; Dale Porcher; Daniel Ross; David Wolf of Tornado Alley Reptiles; David Burstein; Dawn and Kelsi Greene of Super Natural Balls; Dayna Plehn; Debby Brauer; Epic Vibrant Balls; Eric Chung of Chung Reptiles; Erin Burt; George Straub; Haily McCullough; Jaden Christensen; Jake Lewis; Jamie Palazzo of New Day Reptiles; Jeff Kearns; Jeff Linton; Jodi Wilkowski; Joe Myers; John Cordone of Blue Water Reptiles; Jordan Noland; Justin Kobylka of J. Kobylka Reptiles; Lindsay VanOrman; Lisa Huis; Manuel San Juan; Mark Bilger; Lynnet Melton; Maryann Barbon; Mia Hynes; Michael, Lisa, and Bodie Cole of Ballroom Pythons South; Morgan Evans and Michael Kitto of MK Pythons; Morgan Shelton; Paul and Amber Fiorito of Vivid Scales; Pets ‘n’ Things of Saline, MI; Rachel Voyt; Royal Black Balls; Ryan Boyd and Brittney Delacruz; Ryan Young of Molecular Reptile; Sergio McDole; Stephanie S. Crisp; Steve P. of Prime Pets; Zac Parpart; and several anonymous contributors. We thank the following individuals for providing images of color morphs: Beth Woodyard; Cat Church; Cormier Jason; Christine Miller; Darin Taylor; Daniel Hatcher; David C. Callahan; Donald Grinstead; Elijah Snyder; Innovative Ecototherms; Jake Lewis; Jessica Allison; Jessica Van Riper; Justin Kobylka; James Thompson; Justin Revington; Mariette van Vuuren; Mark Hopkins; Mark Smith; Matthew Lopez; Michael Freedman of The Florida Reptile Ranch; Morgan Evans; Phil Barclay, Robert Cooper; Ron Heisenberg; Ryan Young; Selectively Bred Serpents; Seb Des Légendes Celtiques; and Thananan Jivaramonaikul.

Most of the data in this study were collected by undergraduates enrolled in a laboratory course at Eastern Michigan University. These students constitute the BIO306W Consortium. These students were Alexandra Ernst, Alia Frederick, Alissa Zoltowski, Amber Northcutt, Andraya Ackerman, Anna Pathammavong, Annette Miller, Ashly Matzek, Asra Akhlaq,
Author Contributions

Conceptualization: Chiron W. Graves, Hannah S. Seidel.

Data curation: Hannah S. Seidel.

Formal analysis: Autumn R. Brown, Hannah S. Seidel.

Funding acquisition: Autumn R. Brown, Hannah S. Seidel.

Investigation: Autumn R. Brown, Kaylee Comai, Dominic Mannino, Haily McCullough, Yamini Donekal, Hunter C. Meyers, Hannah S. Seidel.

Methodology: Hannah S. Seidel.

Project administration: Hannah S. Seidel.

Resources: Chiron W. Graves, Hannah S. Seidel.

Supervision: Hannah S. Seidel.

Writing – original draft: Hannah S. Seidel.

Writing – review & editing: Autumn R. Brown, Chiron W. Graves, Hannah S. Seidel.

References

1. Cuthill IC, Allen WL, Arbuckle K, Caspers B, Chaplin G, Hauber ME, et al. The biology of color. Science. 2017; 357: eaan0221. https://doi.org/10.1126/science.aan0221 PMID: 28774901

2. Cieslak M, Reissmann M, Hofreiter M, Ludwig A. Colours of domestication. Biol Rev. 2011; 86: 885–899. https://doi.org/10.1111/j.1469-185X.2011.00177.x PMID: 21443614

3. Toews DPL, Hofmeister NR, Taylor SA. The Evolution and Genetics of Carotenoid Processing in Animals. Trends Genet. 2017; 33: 171–182. https://doi.org/10.1016/j.tig.2017.01.002 PMID: 28174022

4. Prum RO. Anatomy, physics, and evolution of avian structural colors. Bird Coloration. Cambridge, MA: Harvard University Press; 2006.

5. Structure Ghiradella H. and Development of Iridescent Lepidopteran Scales—the Papilionidae as a Showcase Family. Ann Entomol Soc Am. 1985; 78: 252–264. https://doi.org/10.1093/aesa/78.2.252

6. Bagnara JT, Taylor JD, Hadley ME. The dermal chromatophore unit. J Cell Biol. 1968; 38: 67–79. https://doi.org/10.1083/jcb.38.1.67 PMID: 5691979
7. Saenko SV, Teyssier J, van der Marel D, Milinkovitch MC. Precise colocalization of interacting structural and pigmentary elements generates extensive color pattern variation in Phelsuma lizards. BMC Biol. 2013; 11: 105. https://doi.org/10.1186/1741-7007-11-105 PMID: 24099066

8. Kinoshita S, Yoshioka S. Structural colors in nature: The role of regularity and irregularity in the structure. ChemPhysChem. 2005; 6: 1442–1459. https://doi.org/10.1002/cphc.200500007 PMID: 16015669

9. Slominski A, Tobin DJ, Shibahara S, Wortsman J. Melanin pigmentation in mammalian skin and its hormonal regulation. Physiol Rev. 2004; 84: 1155–1228. https://doi.org/10.1152/physrev.00044.2003 PMID: 15383650

10. Meredith P, Sarna T. The physical and chemical properties of eumelanin. Pigm Cell Res. 2006; 19: 572–594. https://doi.org/10.1111/j.1600-0749.2006.00345.x PMID: 17083485

11. Bagnara JT, Fernandez PJ, Fuji R. On the blue coloration of vertebrates. Pigm Cell Res. 2007; 20: 14–26. https://doi.org/10.1111/j.1600-0749.2006.00360.x PMID: 17250544

12. Olsson M, Stuart-Fox D, Ballen C. Genetics and evolution of colour patterns in reptiles. Semin Cell Dev Biol. 2013; 24: 529–541. https://doi.org/10.1016/j.semcdb.2013.04.001 PMID: 23578866

13. Shawkey MD, D’Alba L. Interactions between colour-producing mechanisms and their effects on the integumentary colour palette. Philos Trans R Soc B-Biol Sci. 2017; 372: 20160536. https://doi.org/10.1098/rstb.2016.00231 PMID: 28533449

14. Bagnara JT, Matsumoto J. Comparative Anatomy and Physiology of Pigment Cells in Nonmammalian Tissues. The Pigmentary System. John Wiley & Sons, Ltd; 2006. pp. 11–59. https://doi.org/10.1002/9780470987100.ch2

15. Singh AP, Nüsslein-Volhard C. Zebrafish Stripes as a Model for Vertebrate Colour Pattern Formation. Curr Biol. 2015; 25: R81–R82. https://doi.org/10.1016/j.cub.2014.11.013 PMID: 25602311

16. Huang D, Lewis VM, Foster TN, Toomey MB, Corbo JC, Parichy DM. Development and genetics of red coloration in the zebrafish relative Danio albolineatus. eLife. 2021; 10: e70253. https://doi.org/10.7554/eLife.70253 PMID: 34345950

17. Kimura T, Nagao Y, Hashimoto H, Yamamoto-Shiraishi Y, Yamamoto S, Yabe T, et al. Leucophores are similar to xanthophores in their specification and differentiation processes in medaka. Proc Natl Acad Sci U S A. 2014; 111: 7343–7348. https://doi.org/10.1073/pnas.1311254111 PMID: 24803434

18. Nüsslein-Volhard C, Singh AP. How fish colour their skin: A paradigm for development and evolution of adult patterns Multipotency, plasticity, and cell competition regulate proliferation and spreading of pigment cells in Zebrafish coloration. Bioessays. 2017; 39: 1600231. https://doi.org/10.1002/bies.201600231 PMID: 28176337

19. Cooke TF, Fischer CR, Wu P, Jiang T-X, Xie K, Kuo J, et al. Genetic Mapping and Biochemical Basis of Yellow Feather Pigmentation in Budgerigars. Cell. 2017; 171: 427+-. https://doi.org/10.1016/j.cell.2017.08.016 PMID: 28985565

20. Gazda MA, Araujo PM, Lopes RJ, Toomey MB, Andrade P, Afonso S, et al. A genetic mechanism for sexual dichromatism in birds. Science. 2020; 368: 1270+-. https://doi.org/10.1126/science.aba0803 PMID: 32527835

21. Kwon YM, Vranken N, Hoge C, Lichak MR, Francis KK, Camacho-Garcia J, et al. Genomic consequences of domestication of the Siamese fighting fish. 2021 Apr p. 2021.04.29.442030. https://doi.org/10.1101/2021.04.29.442030

22. Lopes RJ, Johnson JD, Toomey MB, Ferreira MS, Araujo PM, Melo-Ferreira J, et al. Genetic Basis for Red Coloration in Birds. Curr Biol. 2016; 26: 1427–1434. https://doi.org/10.1016/j.cub.2016.03.076 PMID: 27212400

23. Mundy NJ, Stapley J, Bennison C, Tucker R, Twyman H, Kim K-W, et al. Red Carotenoid Coloration in the Zebra Finch Is Controlled by a Cytochrome P450 Gene Cluster. Curr Biol. 2016; 26: 1435–1440. https://doi.org/10.1016/j.cub.2016.04.047 PMID: 27212402

24. Toomey MB, Lopes RJ, Araujo PM, Johnson JD, Gazda MA, Afonso S, et al. High-density lipoprotein receptor SCARB1 is required for carotenoid coloration in birds. Proc Natl Acad Sci U S A. 2017; 114: 5219–5224. https://doi.org/10.1073/pnas.1700751114 PMID: 28465440

25. Zhang W, Wang H, Brandt DYC, Hu B, Sheng J, Wang M, et al. The genetic architecture of phenotypic diversity in the betta fish (Betta splendens). bioRxiv. 2021; 2021.05.10.443352. https://doi.org/10.1101/2021.05.10.443352

26. Garcia-Elfring A, Roffey HL, Hendry AP, Barrett RDH. A nonsense mutation in TFEC is the likely cause of the recessive piebald phenotype in ball pythons (Python regius). 2020 Nov p. 2020.10.30.362970. https://doi.org/10.1101/2020.10.30.362970
27. Guo L, Bloom J, Sykes S, Huang E, Kashif Z, Pham E, et al. Genetics of white color and iridophoroma in “Lemon Frost” leopard geckos. PLoS Genet. 2021; 17: e1009580. https://doi.org/10.1371/journal.pgen.1009580 PMID: 3416378

28. McLean CA, Lutz A, Rankin KJ, Stuart-Fox D, Moussalli A. Revealing the Biochemical and Genetic Basis of Color Variation in a Polymorphic Lizard. Mol Biol Evol. 2017; 34: 1924–1935. https://doi.org/10.1093/molbev/msx136 PMID: 28431132

29. Ullate-Agote A, Burgelin I, Debry A, Langrez C, Montange F, Peraldi R, et al. Genome mapping of a LYST mutation in corn snakes indicates that vertebrate chromatophore vesicles are lysosome-related organelles. Proc Natl Acad Sci U S A. 2020; 117: 26307–26317. https://doi.org/10.1073/pnas.2003724117 PMID: 33020272

30. Bale R. Ball python exports raise concerns as demand for the popular pet grows. National Geographic. 2020.

31. Broghammer S. Python regius: Atlas of Colour Morphs Keeping and Breeding. NTV Natur und Tier-Verlag; 2019.

32. Irizarry KJL, Bryden RL. In Silico Analysis of Gene Expression Network Components Underlying Pigmentation Phenotypes in the Python Identified Evolutionarily Conserved Clusters of Transcription Factor Binding Sites. Adv Bioinformatics. 2016; 2016: 1286510. https://doi.org/10.1155/2016/1286510 PMID: 27698666

33. McCurley K. Complete Ball Python, A Comprehensive Guide to Care, Breeding, and Genetic Mutations: Kevin McCurley: 9780976733409: Amazon.com : Books. ECO / Serpent’s Tale NHBD; 2005.

34. Burbrink FT, Castoe TA. Molecular Phylogeography of Snakes. Snakes: Ecology and Conservation. Cornell University Press; 2011. pp. 38–77. https://doi.org/10.7591/9780801459092-006

35. Liu Q, Qi Y, Liang Q, Song J, Liu J, Li W, et al. Targeted disruption of tyrosinase causes melanin reduction in Carassius auratus cuvieri and its hybrid progeny. Sci China-Life Sci. 2019; 62: 1194–1202. https://doi.org/10.1007/s11427-018-9404-7 PMID: 30593611

36. Koga A, Wakamatsu Y, Kurosawa J, Hori H. Oculocutaneous albinism in the i(6) mutant of the medaka fish is associated with a deletion in the tyrosinase gene. Pigm Cell Res. 1999; 12: 252–258. https://doi.org/10.1111/j.1600-0749.1999.tb00758.x PMID: 10454293

37. Miura I, Tagami M, Fujitani T, Ogata M. Spontaneous tyrosinase mutations identified in albinos of three wild frog species. Genes Genet Syst. 2017; 92: 189–196. https://doi.org/10.1266/ggs.16-00061 PMID: 28674275

38. Iwanishi S, Zaitsu S, Shibata H, Nitasaka E. An albinino mutant of the Japanese rat snake (Elaphe climacophora) carries a nonsense mutation in the tyrosinase gene. Genes Genet Syst. 2018; 93: 163–167. https://doi.org/10.1266/age.12386 PMID: 30158334

39. Chang C-M, Coville J-L, Coquerelle G, Gourichon D, Oulmouden A, Tixier-Boichard M. Complete association between a retroviral insertion in the tyrosinase gene and the recessive white mutation in chickens. Anim Genet. 2006; 7: 19. https://doi.org/10.1186/1471-2164-7-19 PMID: 16457736

40. Florisbal Dame MC, Xavier GM, Oliveira-Filho JP, Borges AS, Oliveira HN, Riet-Correa F, et al. A nonsense mutation in the tyrosinase gene causes albinism in water buffalo. BMC Genet. 2012; 13: 62. https://doi.org/10.1186/1471-2156-13-62 PMID: 22817390

41. Schmutz SM, Berryere TG, Ciobanu DC, Mileham AJ, Schmidt BH, Fredholm M. A form of albinism in cattle is caused by a tyrosinase frameshift mutation. Mamm Genome. 2004; 15: 62–67. https://doi.org/10.1007/s00335-002-2249-5 PMID: 14727143

42. Reiner G, Tramberend K, Niefeld F, Volmer K, Wurmsner C, Fries R, et al. A genome-wide scan study identifies a single nucleotide substitution in the tyrosinase gene associated with white coat colour in a red deer (Cervus elaphus) population. BMC Genet. 2020;21. https://doi.org/10.1186/s12863-020-0814-0 PMID: 32041521

43. Utzeri VJ, Bertolini F, Ribani A, Schiavo G, Dall'Olio S, Fontanesi L. The albinism of the feral Asinara white donkeys (Equus asinus) is determined by a missense mutation in a highly conserved position of the tyrosinase (TYR) gene deduced protein. Anim Genet. 2016; 47: 120–124. https://doi.org/10.1111/age.12386 PMID: 26763160
46. Yokoyama T, Silversides D, Waymire K, Kwon B, Takeuchi T, Overbeek P. Conserved Cysteine to Serine Mutation in Tyrosinase Is Responsible for the Classical Albino Mutation in Laboratory Mice. Nucleic Acids Res. 1990; 18: 7293–7298. https://doi.org/10.1093/nar/18.24.7293 PMID: 2124349

47. Blaszczyk WM, Arning L, Hoffmann KP, Epplen JT. A Tyrosinase missense mutation causes albinoism in the Wistar rat. Pigm Cell Res. 2005; 18: 144–145. https://doi.org/10.1111/j.1600-0749.2005.00227.x PMID: 15760344

48. Yu F, Jiao S, Lai W, Liu Z, Zhu M, Zhu W, et al. Conserved aspartate-to-glycine mutation in tyrosinase is associated with albino phenotype in domestic guinea pigs (Cavia porcellus). Anim Genet. 2018; 49: 354–355. https://doi.org/10.1111/age.12683 PMID: 29947431

49. Kim Y-H, Park S-J, Choe S-H, Lee J-R, Cho H-M, Kim S-U, et al. Identification and characterization of the tyrosinase gene (TYR) and its transcript variants (TYR_1 and TYR_2) in the crab-eating macaque (Macaca fascicularis). Gene. 2017; 630: 21–27. https://doi.org/10.1016/j.gene.2017.07.047 PMID: 28756020

50. Galante Rocha de Vasconcelos FT, Hauzman E, Henriques LD, Kilpp Goulart PR, Galvao O de F, Sano RY, et al. A novel nonsense mutation in the tyrosinase gene is related to the albinism in a capuchin monkey (Sapajus apella). BMC Genet. 2017; 18: 39. https://doi.org/10.1186/s12863-017-0504-8 PMID: 28476152

51. Koga A, Hisakawa C, Yoshizawa M. Baboon bearing resemblance in pigmentation pattern to Siamese cat carries a missense mutation in the tyrosinase gene. Genome. 2020; 63. https://doi.org/10.1139/gen-2020-0003 PMID: 32053406

52. Montoliu L, Gronskov K, Wei A-H, Martinez-Garcia M, Fernandez A, Arveiler B, et al. Increasing the complexity: new genes and new types of albinism. Pigment Cell Melanoma Res. 2014; 27. https://doi.org/10.1111/pcmr.12167 PMID: 24066960

53. Polanowski AM, Robinson-Laverick SM, Paton D, Jarman SN. Variation in the Tyrosinase Gene Associated with a White Humpback Whale (Megaptera novaeangliae). J Hered. 2012; 103: 130–133. https://doi.org/10.1038/jhered.esr106 PMID: 22140235

54. Blaszczyk WM, Distler C, Dekomen G, Arning L, Hoffmann K-P, Epplen JT. Identification of a tyrosinase (TYR) exon 4 deletion in albino ferrets (Mustela putorius furo). Anim Genet. 2007; 38: 421–423. https://doi.org/10.1111/j.1365-2052.2007.01619.x PMID: 17655555

55. Anistoroaei R, Fredholm M, Christensen K, Leeb T. Albinism in the American mink (Neovison vison) is associated with a tyrosinase nonsense mutation. Anim Genet. 2008; 39: 645–648. https://doi.org/10.1111/j.1365-2052.2008.01788.x PMID: 18822100

56. Imes DL, Geary LA, Grahn RA, Lyons LA. Albinism in the domestic cat (Felis catus) is associated with a tyrosinase (TYR) mutation. Anim Genet. 2006; 37: 175–178. https://doi.org/10.1111/j.1365-2052.2005.01409.x PMID: 16573534

57. Lyons LA, Imes DL, Rah HC, Grahn RA. Tyrosinase mutations associated with Siamese and Burmese patterns in the domestic cat (Felis catus). Anim Genet. 2006; 36: 119–126. https://doi.org/10.1111/j.1365-2052.2005.01253.x PMID: 15771720

58. Schmidt-Kuntzel A, Ezirik E, O’Brien SJ, Menotti-Raymond M. Tyrosinase and tyrosinase related protein 1 alleles specify domestic cat coat color phenotypes of the albino and brown loci. J Hered. 2005; 96: 289–301. https://doi.org/10.1093/jhered/esi086 PMID: 15858157

59. Yan S, Zhao D, Hu M, Tan X, Lai W, Kang J, et al. A single base insertion in the tyrosinase gene is associated with albino phenotype in silver foxes (Vulpes vulpes). Anim Genet. 2019; 50: 550–550. https://doi.org/10.1111/age.12816 PMID: 31246286

60. Gross JB, Wilkens H. Albinism in phylogenetically and geographically distinct populations of Astyanax cavefish arises through the same loss-of-function Oca2 allele. Heredity. 2013; 111: 122–130. https://doi.org/10.1038/hdy.2013.26 PMID: 23572122

61. Kratochwil CF, Urban S, Meyer A. Genome of the Malawi golden cichlid fish (Melanochromis auratus) reveals exon loss of oca2 in an amelanistic morph. Pigment Cell Melanoma Res. 2019; 32: 719–723. https://doi.org/10.1111/pcmr.12799 PMID: 31319885

62. Saenko SV, Lamichhane S, Barrio AM, Rafati N, Andersson L, Milinkovitch MC. Amelanism in the corn snake is associated with the insertion of an LTR-retrotransposon in the OCA2 gene. Sci Rep. 2015; 5: 17118. https://doi.org/10.1038/srep17118 PMID: 26597053

63. Caduff M, Bauer A, Jagnannathan V, Leeb T, OCA2 splice site variant in German Spitz dogs with oculo-cutaneous albinism. PLoS One. 2017; 12: e0185944. https://doi.org/10.1371/journal.pone.0185944 PMID: 28873042

64. Zhang Y, Hong Q, Cao C, Yang L, Li Y, Hai T, et al. A novel porcine model reproduces human oculo-cutaneous albinism type. Cell Discov. 2019; 5: 48. https://doi.org/10.1038/s41421-019-0117-7 PMID: 31636960
65. Shoji H, Kiniwa Y, Okuyama R, Yang M, Higuchi K, Mori M. A nonsense nucleotide substitution in the oculocutaneous albinism II gene underlies the original pink-eyed dilution allele (Oca2(p)) in mice. Exp Anim. 2015; 64: 171–179. [https://doi.org/10.1538/expanim.14-0075 PMID: 25736709]

66. Krauss J, Geiger-Rudolph S, Koch I, Nuesslein-Volhard C, Irim U. A dominant mutation in tyrp1A leads to melanophore death in zebrafish. Pigment Cell Melanoma Res. 2014; 27: 827–830. [https://doi.org/10.1111/pcmr.12272 PMID: 24891189]

67. Cortimiglia C, Castiglioni B, Pizzi F, Stella A, Capra E. Involvement of tyrosinase-related protein 1 gene in the light brown plumage phenotype of Falco cherrug. Anim Genet. 2017; 48: 125–126. [https://doi.org/10.1111/age.12506 PMID: 27611661]

68. Ren J, Mao H, Zhang Z, Xiao S, Ding N, Huang L. A 6-bp deletion in the TYRP1 gene causes the brown colouration phenotype in Chinese indigenous pigs. Heredity. 2011; 106: 862–868. [https://doi.org/10.1038/hdy.2010.129 PMID: 20978532]

69. Wu X, Zhang Y, Shen L, Du J, Luo J, Liu C, et al. A 6-bp deletion in exon 8 and two mutations in introns of TYRP1 are associated with blond coat color in Liangshan pigs. Gene. 2016; 578: 132–136. [https://doi.org/10.1016/j.gene.2015.12.011 PMID: 26680103]

70. Brancalion L, Haase B, Wade CM. Canine coat pigmentation genetics: a review. Animal Genetics. 2022; 53: 3–34. [https://doi.org/10.1111/age.13154 PMID: 34751460]

71. Lyons LA, Foe IT, Rah HC, Grahn RA. Chocolate coated cats: TYRP1 mutations for brown color domestic cats. Mamm Genome. 2005; 16: 356–366. [https://doi.org/10.1007/s00335-004-2455-4 PMID: 16104383]

72. Gratten J, Beraldi D, Lowder BV, McRae AF, Visscher PM, Pemberton JM, et al. Compelling evidence that a single nucleotide substitution in TYRP1 is responsible for coat-colour polymorphism in a free-living population of Soay sheep. Proc R Soc B-Biol Sci. 2007; 274: 619–626. [https://doi.org/10.1098/rspb.2006.3762 PMID: 17254985]

73. Paris JM, Letko A, Hafliger IM, Ammann P, Flury C, Drogemuller C. Identification of two TYRP1 loss-of-function alleles in Valais Red sheep. Anim Genet. 2019; 50: 778–782. [https://doi.org/10.1111/age.12863 PMID: 31571241]

74. Cirera S, Markakis MN, Kristiansen T, Vissenberg K, Fredholm M, Christiansen K, et al. A large insertion in intron 2 of the TYRP1 gene associated with American Palomino phenotype in American mink. Mamm Genome. 2016; 27: 135–143. [https://doi.org/10.1007/s00335-016-9620-4 PMID: 26886941]

75. Budd P, Jackson I. Structure of the Mouse Tyrosinase-Related Protein-2 Dopachrome Tautomerase (tyrp2/Dct) Gene and Sequence of 2 Novel Slaty Alleles. Genomics. 1995; 29: 35–43. [https://doi.org/10.1006/geno.1995.1212 PMID: 8530099]

76. Jackson I, Chambers D, Tsukamoto K, Fredholm M, Christensen K, et al. A 2nd Tyrosinase-Related Protein, Trp-2, Maps to and Is Mutated at the Mouse Slaty Locus. Embo J. 1992; 11: 527–535. [https://doi.org/10.1002/j.1460-2075.1992.tb05083.x PMID: 1537334]

77. Chintala S, Li W, Lamoreux ML, Ito S, Wakamatsu K, Sviderskaya EV, et al. Slc7a11 gene controls production of pheomelanin pigment and proliferation of cultured cells. Proc Natl Acad Sci U S A. 2005; 102: 10964–10969. [https://doi.org/10.1073/pnas.0502856102 PMID: 16037214]

78. Lamason RL, Mohideen M, Mest JR, Wong AC, Norton HL, Aros MC, et al. SLC24A5, a putative cation exchanger, affects pigmentation in zebrafish and humans. Science. 2005; 310: 1782–1786. [https://doi.org/10.1126/science.1116238 PMID: 15373334]

79. Mack M, Kowalski E, Grahn RA, Bras D, Penedo MCT, Bellone R. Two Variants in SLC24A5 Are Associated with “Tiger-Eye” Iris Pigmentation in Puerto Rican Paso Fino Horses. G3-Genes Genomes Genet. 2017; 7: 2799–2806. [https://doi.org/10.1534/g3.117.037866 PMID: 28655738]

80. Fukamachi S, Shimada A, Shima A. Mutations in the gene encoding B, a novel transporter protein, reduce melanin content in medaka. Nature Genet. 2001; 28: 381–385. [https://doi.org/10.1038/ng584 PMID: 11479596]

81. Gunnarsson U, Hellstrom AR, Tixier-Boichard M, Minvielle F, Bed'hom B, Ito S, et al. Mutations in SLC45A2 cause plumage color variation in chicken and Japanese quail. Genetics. 1975; 87: 867–877. [https://doi.org/10.1534/genetics.110.063107 PMID: 17151254]

82. Winkler PA, Gornik KR, Ramsey DT, Dubielzig RR, Venta PJ, Petersen-Jones SM, et al. A Partial Gene Deletion of SLC45A2 Causes Oculocutaneous Albinism in Doberman Pinscher Dogs. PLoS One. 2014; 9: e92127. [https://doi.org/10.1371/journal.pone.0092127 PMID: 24647637]

83. Mariat D, Taourit S, Guerin G. A mutation in the MATP gene causes the cream coat colour in the horse. Genet Sel Evol. 2003; 35: 119–133. [https://doi.org/10.1186/1297-9686-35-1-119 PMID: 12605854]
84. Newton JM, Cohen-Barak O, Hagiwara N, Gardner JM, Davission MT, King RA, et al. Mutations in the human orthologue of the mouse underwhite gene (uw) underlie a new form of oculocutaneous albinism. OCA4. Am J Hum Genet. 2001; 69: 981–988. https://doi.org/10.1086/324340 PMID: 11574907

85. Xu X, Dong G-X, Hu X-S, Miao L, Zhang X-L, Zhang D-L, et al. The Genetic Basis of White Tigers. Curr Biol. 2013; 23: 1031–1035. https://doi.org/10.1016/j.cub.2013.04.054 PMID: 23707431

86. Prado-Martinez J, Hernando-Herraez I, Lorente-Galdos B, Dabad M, Ramirez O, Baeza-Delgado C, et al. The genome sequencing of an albino Western lowland gorilla reveals inbreeding in the wild. BMC Genomics. 2013; 14: 363. https://doi.org/10.1186/1471-2164-14-363 PMID: 23721540

87. Lai X, Wichers HJ, Soler-Lopez M, Dijkstra BW. Structure and Function of Human Tyrosinase and Tyrosinase-Related Proteins. Chem-Eur J. 2018; 24: 47–55. https://doi.org/10.1002/chem.201704410 PMID: 29052256

88. Bellono NW, Escobar IE, Lefkowith AJ, Marks MS, Oancea E. An intracellular anion channel critical for pigmentation. eLife. 2014; 3: e04543. https://doi.org/10.7554/eLife.04543 PMID: 25513726

89. Ginger RS, Askew SE, Ogborne RM, Wilson S, Ferdinando D, Dadd T, et al. SLC24A5 encodes a trans-golgi network protein with potassium-dependent sodium-calcium exchange activity that regulates human epidermal melanogenesis. J Biol Chem. 2008; 283: 5486–5495. https://doi.org/10.1074/jbc.M707521200 PMID: 18166528

90. Vitavska O, Wieczorek H. The SLC45 gene family of putative sugar transporters. Mol Asp Med. 2013; 34: 655–660. https://doi.org/10.1016/j.mam.2012.05.014 PMID: 23506898

91. Castoe TA, de Koning APJ, Hall KT, Card DC, Schield DR, Fujita MK, et al. The Burmese python genome reveals the molecular basis for extreme adaptation in snakes. Proc Natl Acad Sci U S A. 2013; 110: 20645–20650. https://doi.org/10.1073/pnas.122315 PMID: 24297902

92. Stephens M, Scheet P. Accounting for decay of linkage disequilibrium in haplotype inference and missing-data imputation. Am J Hum Genet. 2005; 76: 449–462. https://doi.org/10.1086/428594 PMID: 15700229

93. Stephens M, Smith NJ, Donnelly P. A new statistical method for haplotype reconstruction from population data. Am J Hum Genet. 2001; 68: 978–989. https://doi.org/10.1086/319501 PMID: 11254454

94. Simeonov DR, Wang X, Wang C, Sergeev Y, Dolinska M, Bower M, et al. DNA Variations in Oculocutaneous Albinism: An Updated Mutation List and Current Outstanding Issues in Molecular Diagnostics. Hum Mutat. 2013; 34: 827–835. https://doi.org/10.1002/humu.22315 PMID: 23504663

95. Gardner J, Nakatsu Y, Gondo Y, Lee S, Lyon M, King R, et al. The Mouse Pink-Eyed Dilution Gene—Association with Human Prader-Willi and Angelman Syndromes. Science. 1992; 257: 1121–1124. https://doi.org/10.1126/science.257.5073.1121 PMID: 1509264

96. Ni-Komatsu L, Orlow SJ. Heterologous expression of tyrosinase recapitulates the misprocessing and mistrafficking in oculocutaneous albinism type 2: Effects of altering intracellular pH and pink-eyed dilution gene expression. Exp Eye Res. 2006; 82: 519–528. https://doi.org/10.1016/j.exer.2005.08.013 PMID: 16199032

97. Puri N, Gardner JM, Brilliant MH. Aberrant pH of melanosomes in pink-eyed dilution (p) mutant melanocytes. J Invest Dermatol. 2000; 115: 607–613. https://doi.org/10.1046/j.1523-1747.2000.00108.x PMID: 10998131

98. Lee S, Nicholls R, Bundey S, Laxova R, Musarella M, Spritz R. Mutations of the P Gene in Oculocutaneous Albinism, Ocular Albinism, and Prader-Willi-Syndrome Plus Albinism. N Engl J Med. 1994; 330: 529–534. https://doi.org/10.1056/NEJM199402243300803 PMID: 8302318

99. Sato H, Tamba M, Ishii T, Bannai S. Cloning and expression of a plasma membrane cystine/glutamate exchange transporter composed of two distinct proteins. J Biol Chem. 1999; 274: 11455–11458. https://doi.org/10.1074/jbc.274.17.11455 PMID: 10206947

100. Bin B-H, Bhin J, Yang SH, Shin M, Nam Y-J, Choi D-H, et al. Membrane-Associated Transporter Protein (MATP) Regulates Melanosomal pH and Influences Tyrosinase Activity. PLoS One. 2015; 10: e0129273. https://doi.org/10.1371/journal.pone.0129273 PMID: 26057890

101. Sturm R, Osulivian B, Box N, Smith A, Smit S, Puttick E, et al. Chromosomal Structure of the Human Tyr1 and Tyr2 Loci and Comparison of the Tyrosinase-Related Protein Cone Family. Genomics. 1995; 29: 24–34. https://doi.org/10.1006/geno.1995.1211 PMID: 8530077

102. Gershonibauch R, Rosenmann A, Drooto S, Holmes S, Tripathi R, Spritz R. Mutations of the Tyrosinase Gene in Patients with Oculocutaneous Albinism from Various Ethnic-Groups in Israel. Am J Hum Genet. 1994; 54: 586–594. PMID: 8128955

103. Tripathi RK, Strunk KM, Giebel LB, Weleber RG, Spritz RA. Tyrosinase gene mutations in type I (tyrosinase-deficient) oculocutaneous albinism define two clusters of missense substitutions. Am J Med Genet. 1992; 43: 865–871. https://doi.org/10.1002/ajmg.1320430523 PMID: 1642278
104. Tsai CH, Tsai FJ, Wu JY, Lin SP, Chang JG, Yang CF, et al. Insertion/deletion mutations of type I ocu-
locutaneous albinism in chinese patients from Taiwan. Hum Mutat. 1999; 14: 542. https://doi.org/10.
1002/(SICI)1098-1004(199912)14:6<542::AID-HUMU14>3.0.CO;2-3 PMID: 10571953

105. Forshew T, Khalig S, Tee L, Smith U, Johnson CA, Mehdi SQ, et al. Identification of novel TYR and 
TYRP1 mutations in oculocutaneous albinism. Clin Genet. 2005; 68: 182–184. https://doi.org/10.
1111/j.1399-0004.2005.00460.x PMID: 15996218

106. Lasseaux E, Plaisant C, Michaud V, Pennamen P, Trimouille A, Gaston L, et al. Molecular characteri-
zation of a series of 990 index patients with albinism. Pigment Cell Melanoma Res. 2018; 31: 466– 
474. https://doi.org/10.1111/pcmr.12688 PMID: 29345414

107. Rooryck C, Morice-Picard F, Elcioglu NH, Lacombe D, Taieb A, Arveiler B. Molecular diagnosis of ocu-
locutaneous albinism: new mutations in the OCA1-4 genes and practical aspects. Pigment Cell Melano-
noma Res. 2008; 21: 583–587. https://doi.org/10.1111/j.1755-148X.2008.00496.x PMID: 18821858

108. Rooryck C, Morice F, Lacombe D, Taieb A, Arveiler B. Genetic basis of oculocutaneous albinism. 
Expert Review of Dermatology. 2009; 4: 611–622. https://doi.org/10.1586/edm.09.53

109. Allen WL, Baddeley R, Scott-Samuels NE, Cuthill IC. The evolution and function of pattern diversity in 
snakes. Behav Ecol. 2013; 24: 1237–1250. https://doi.org/10.1093/beheco/art058

110. McGinnis S, Madden TL. BLAST: at the core of a powerful and diverse set of sequence analysis tools. 
Nucleic Acids Res. 2004; 32: W20–W25. https://doi.org/10.1093/nar/gkh435 PMID: 15215342

111. Gertz EM, Yu Y-K, Agarwala R, Schäffer AA, Altschul SF. Composition-based statistics and translated 
nucleotide searches: Improving the TBLASTN module of BLAST. BMC Biology. 2006; 4: 41. https:// 
doi.org/10.1186/1741-7007-4-41 PMID: 17156431

112. Untergasser A, Cutcutache I, Koressaar T, Ye J, Faircloth BC, Remm M, et al. Primer3-new capabilities 
and interfaces. Nucleic Acids Res. 2012; 40: e115. https://doi.org/10.1093/nar/gks596 PMID: 
22730293

113. Sievers F, Wilm A, Dineen D, Gibson TJ, Karplus K, Li W, et al. Fast, scalable generation of high-quality 
protein multiple sequence alignments using Clustal Omega. Mol Syst Biol. 2011; 7: 539. https://doi. 
org/10.1038/msb.2011.75 PMID: 21988835