Phylogeny of *Amazona barbadensis* and the Yellow-Headed Amazon Complex (Aves: Psittacidae): A New Look at South American Parrot Evolution

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### Abstract

The Yellow-shouldered Amazon (*Amazona barbadensis*) is the sole parrot of the genus *Amazona* that inhabits only dry forests. Its population has been dropping; therefore it has been the topic of many studies and conservation efforts. However, the phylogenetic relationship of this species to potential relatives classified within the Yellow-Headed Amazon (YHA) complex are still not clear. Therefore, we used more extensive data sets, including the newly sequenced mitochondrial genome of *A. barbadensis*, to conduct phylogenetic analyses. Various combinations of genes and many phylogenetic approaches showed that *A. barbadensis* clustered significantly with *A. ochrocephala ochrocephala* from Colombia and Venezuela, which created the Northern South American (NSA) lineage, clearly separated from two other lineages within the YHA complex, the Central (CA) and South American (SA). Tree topology tests and exclusion of rapidly evolving sites provided support for a NSA+SA grouping. We propose an evolutionary scenario for the YHA complex and its colonization of the American mainland. The NSA lineage likely represents the most ancestral lineage, which derived from Lesser Antillean Amazons and colonized the northern coast of Venezuela about a million years ago. Then, Central America was colonized through the Isthmus of Panama, which led to the emergence of the CA lineage. The southward expansion to South America and the origin of the SA lineage happened almost simultaneously. However, more intensive or prolonged gene flow or migrations have led to much weaker geographic differentiation of genetic markers in the SA than in the CA lineage.

### Introduction

According to the most recent classification of New World parrots, ten genera are ascribed to the Androglossini tribe [1]. One of them is the genus *Amazona* (Amazon parrots), which is represented by the largest number of species. It is widely distributed in Central and South America (from Mexico to Argentina), and throughout the Caribbean [2–4]. Amazon parrots are phenotypically characterized by their medium to large size, strong-heavy bill, short-rounded tail, prominent naked cere and the presence of a distinct notch in the upper mandible [3]. Their body plumage is predominantly green (except for a few Lesser Antillean species) with variable colorations on the head, breast, body plumage is predominantly green (except for a few Lesser Antillean species) with variable colorations on the head, breast, and lores are pale yellow or even white. The crown of this parrot resembles most closely the Blue-fronted Amazon (*A. aestiva*). Individuals of this species are found in several isolated populations in the coastal areas of Venezuela and across the Southern Caribbean region (the Venezuelan islands of Margarita and La Blanquilla as well as the Netherlands Antilles islands of Curacao and Bonaire) [6]. Its total population size is estimated at 2,500 to 10,000 individuals, and has been decreasing due to poaching, disappearance of natural habitat and the impact of non-native predators. The Yellow-shouldered Amazon is evaluated as ‘vulnerable’ by International Union for the Conservation of Nature and Natural Resources and is becoming a globally endangered parrot species. This has made the species the subject of many studies and conservation efforts.

The Yellow-shouldered Amazon is characterized by the yellow color of its shoulders, thighs, chin, upper part of the cheeks, area around the eyes, and over the ear openings (Figure S1). Forehead and lores are pale yellow or even white. The crown of this parrot sometimes has orange feathers. The lower part of cheeks and throat is usually bluish-green but sometimes also yellow. Although predominant yellow coloration of the shoulders is a characteristic feature of the species, some specimens with red patches on their shoulders were also observed.

In respect to plumage color, the Yellow-shouldered Amazon resembles most closely the Blue-fronted Amazon (*A. aestiva*). Individuals of *A. a. xanthopteryx* subspecies have also the yellow/
red coloration of shoulders as well as yellow/white/blue coloration of the head. However, this triple coloration of the head is differentially distributed in these two species. Despite the plumage similarities, a close relationship between the two species has so far never been supported by molecular research. However, Rusello and Amato [7] found that both species are nested within the Yellow-Headed Amazon (YHA) complex.

The YHA complex has been described as ‘a taxonomic headache’ [8] because the number of recognized species varies according to different taxonomic classifications. Forshaw [3] considers a single species (A. ochrocephala) to exist, with eleven subspecies in the complex; whereas other authorities recognize three separate species (A. ochrocephala, A. auropalliata, A. oratrix) also with 11 subspecies [2,4,9].

According to the latter classification [2], the species A. ochrocephala (Yellow-crowned Amazon) includes three South American subspecies (A. o. ochrocephala, A. o. xantholoma, A. o. nattereri) and one Central American subspecies (A. o. panamensis). The species A. auropalliata (Yellow-naped Amazon) contains three Central American subspecies: auropalliata, parajipes and caribaea, whereas A. oratrix (Yellow-headed Amazon) is comprised of the other four Central American subspecies: oratrix, tresmariae, belizensis and hondurensis. The split of these taxa is based on characters such as extent and position of yellow plumage on head and thighs, plumage coloration at the bend of wings and lesser wing-coverts, bill and foot pigmentation and body size [3,4]. Recently, the species A. tresmariae was distinguished from A. oratrix by the International Ornithological Committee (IOC) [10] based on results of molecular analyses [11], as was proposed by Navarro-Siguenza and Peterson [12]. In summary, according to the most recent classification [10], four different species of Amazona (ochrocephala, auropallita, oratrix and tresmariae) are currently recognized within the YHA complex.

Eberhard and Bermingham [11] presented a phylogenetic study of the YHA complex based on combined mitochondrial gene sequences with a total length of 2,515 bp (nd2, cox1, atp8, atp6). Besides ‘typical’ YHA species such as: A. tresmariae, A. oratrix (two subspecies), A. auropalliata and A. ochrocephala (one Central American and three South American subspecies), they also included the Blue-fronted Amazon (A. aestiva). Based on the four-gene data set, they found that the complex can be divided into three main lineages: the Central American (CA) lineage comprising all Central American taxa, the Northern South American (NSA) lineage represented by an individual A. ochrocephala ochrocephala from Colombia, and a South American (SA) lineage including A. ochrocephala ochrocephala (from Brazil), A. ochrocephala nattereri, A. ochrocephala xantholoma and A. aestiva. This surprising separation of the Colombian A. o. ochrocephala sample from the other South American taxa was further confirmed based on short cox1 sequence dataset (622 bp) including additional A. o. ochrocephala samples, one from Colombia and two from different localities in Venezuela [11]. Finally, the analysis of the coxl sequences demonstrated a phylogenetic separation of Northern South American A. o. ochrocephala from Central South America taxa. These results suggested strong phylogeographic structure in the Central American lineage, which was lacking in the South American lineage.

Using only the short cox1 sequence, Eberhard and Bermingham [11] reported also a putative phylogenetic position for A. barbadensis. They showed that the species represents a distinct clade, which is sister to the whole YHA complex. However, their findings disagree with the phylogenetic position of A. barbadensis that was proposed by Rusello and Amato [7].

Rusello and Amato [7] presented a phylogenetic hypothesis for the genus Amazona, which was based on 3,160 bp length concatenated alignment of mitochondrial (cox1, 12S, 16S) and nuclear (β-fibini 7, rp40, trop) loci. This apparently more detailed study of the YHA complex contained as many as nine among all eleven YHA subspecies and two subspecies of A. aestiva. However, it did not take into account the molecular data derived from A. tresmariae species. These results did not confirm the subdivision of the YHA complex into three species (oratrix, auropallita and ochrocephala) but rather suggested its paraphyly. The authors found also that A. aestiva is nested within the YHA complex. On the other hand, in contrast to the results by Eberhard and Bermingham [11], Rusello and Amato [7] showed that the species A. barbadensis is nested within the YHA complex. Despite the apparent morphological similarities between the A. barbadensis and A. aestiva xanthopteryx subspecies, the former appeared to be a distinct lineage within the YHA complex, sister to A. ochrocephala ochrocephala as well as Central American (CA) and South American (SA) clades. This single A. ochrocephala ochrocephala sample, separated from the other South American taxa, seems to correspond to the Northern South American (NSA) lineage that was proposed by Eberhard and Bermingham [11]. However, it is only a supposition because analyses by Rusello and Amato [7] included no other A. ochrocephala ochrocephala individuals whereas the geographic origin of the analyzed sample was not given. The phylogenetic position of the A. barbadensis species remains also ambiguous because trees obtained by two methods showed different topologies [7].

### Table 1. Characteristics of analysed data sets.

| Data set                          | Number of sequences | Length of alignment [bp] |
|-----------------------------------|---------------------|--------------------------|
| nd + cox + atp6 + atp8 + atp6     | 63                  | 1810                     |
| nd + cox                        | 35                  | 1663                     |
| nd + cox + atp6 + atp8 + atp6    | 31                  | 2515                     |
| 12s + 16s + cox                  | 47                  | 1485                     |
| 12s + 16s + cox + cyt           | 31                  | 2179                     |
| cox                             | 69                  | 513                      |
| cyt                             | 35                  | 694                      |

1 based on Ribas et al. [14] data set; 2 based on Ribas et al. [14] and modified data set; 3 based on Rusello and Amato [7] and modified data set; 4 similar to the set but including also cytochrome b.

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Figure 1. The Bayesian tree based on nd+cox+atp6+8 data set. Values of posterior probabilities greater than 0.50 were shown on selected branches. Three main lineages of Yellow-Headed Amazons (YHA) are marked: Central American (CA), South American (SA) and Northern South American (NSA). Species name of analyzed Amazona (A.) individuals are marked in bold. Numbers at the end of some names correspond to the labeling of individuals listed in Table S1.
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them treats *A. barbadensis* as a distinct lineage whereas the second one suggests its phylogenetic relationship with *A. ochrocephala ochrocephala* (NSA lineage) with rather low bootstrap support. Surprisingly, none of three main lineages (CA, SA, NSA) from the YHA complex was detected by Ottens-Wainright et al. [13]. Their analyses based on cytochrome *b* sequences showed the existence of closely related group including the following Amazona species: *A. aestiva*, *A. auropalliata*, *A. tresmariae*, *A. ochrocephala* (two genetically separated individuals), *A. barbadensis*, and two Lesser Antillean taxa, *A. arausiaca* and *A. versicolor*. The most basal position of this group was taken by *A. aestiva* whereas genetically diverse individuals of *A. ochrocephala* clustered together with the CA lineage taxa (*A. auropalliata*, *A. oratrix* and *A. tresmariae*). Moreover, their analyses showed that *A. barbadensis* is rather sister to both Lesser Antillean species than to the typical YHA complex.

So far, Ribas et al. [14] have presented the most informative study of the YHA complex including more samples with confirmed geographic origins. Their analysis was based on four concatenated mitochondrial gene sequences (1820 bp length): nd2, cox1, ATPase 6 and 8, which represented four *A. ochrocephala* subspecies, two *A. oratrix* subspecies, *A. auropalliata* auropalliata subspecies, *A. tresmariae* species and two *A. aestiva* subspecies. The results confirmed the existence of the three main lineages found previously by Eberhard and Bermingham [11]: the Central American (CA), South American (SA) and Northern South American (NSA) lineages. The SA lineage was further divided into two well-supported clades and occurred as a sister clade to the CA lineage. The CA lineage contained several strongly supported groups and resolved relationships between *auropalliata*, *oratrix* and *tresmariae*. Nevertheless, the results underlined the need for taxonomic revision of *Amazona ochrocephala* species. Four subspecies of *A. ochrocephala* (according to American Ornithologists' Union [9]) did not create one monophyletic clade but were distributed to the three main lineages. Even sequences ascribed to the same subspecies were separated from each other. Six individuals of *A. ochrocephala ochrocephala* from Brazil were divided into two groups within the SA clade 2, whereas the Colombian individual of this subspecies was genetically divergent from others and represented the NSA lineage. Moreover, several *A. ochrocephala nattereri* specimens were scattered within SA lineage depending on their locality.

Unfortunately, *A. barbadensis* was not sampled by Ribas et al. [14] and its taxonomic position was not determined. Therefore, it is not still clear if *A. barbadensis* is closely related to the YHA complex [7,11] or Lesser Antillean species [13]. Therefore, we used newly sequenced complete mitochondrial genome of *A. barbadensis* [15] to obtain more molecular markers useful in inferring the taxonomic position of this species. Using the most extensive data set with...
different combinations of markers and many phylogenetic approaches, we also tried to clarify relationships within the YHA complex and resolve several question concerning its phylogeography.

Materials and Methods

Phylogenetic Analyses

Phylogenetic analyses were performed on seven data sets (Table 1, Tables S1–S6) created from aligned sequences of mitochondrial genes: NADH dehydrogenase subunit 2 (nd2), cytochrome oxidase subunit I (cox1), ATPase 6 (atp6), ATPase 8 (atp8), cytochrome b (cytb), 12s rRNA (12s) and 16S rRNA (16s). Different combinations of the genetic markers, similar to those used by Ribas et al. [14] and Rusello and Amato [7], were applied. The original sets were modified by inclusion of additional sequences or exclusion of others to obtain alignments with larger number of sites. The sequences were aligned and initially analyzed in MEGA 5 [16].

To infer phylogenetic trees, we applied four approaches: Bayesian analysis in MrBayes 3.2.1 [17] as well as maximum likelihood analyses in TreeFinder [18], PAUP* 4.0b [19] and morePhyML 1.14 [20] based on PhyML 3.0 [21].

In MrBayes analyses, we assumed separate mixed+I+G(5) models for each data partition (rRNA genes and three codon positions in each of protein-coding genes) to sample appropriate models across the substitution model space in the Bayesian MCMC analysis itself avoiding the need for a priori model testing [22]. In the analyses, two independent runs starting from random trees were applied, each using eight (for cox and 12s+16s+cox+c ytb data sets) or four (for the rest sets) Markov chains. Trees were sampled every 100 generations for 10,000,000 generations. In the final analysis, we selected trees from the last 3,174,000 to 5,745,000 generations (depending on the data set) that reached the stationary phase and convergence (i.e. the standard deviation of split frequencies stabilized and was lower than the proposed threshold of 0.01).

In TreeFinder we applied search depth set to 2 and separate substitution models for each data partition (rRNA genes and three codon positions in each of protein-coding genes) as suggested by this program’s Propose Model module according to AIC, AICc, BIC and HQ criteria. Trees inferred with (more) PhyML and PAUP were based on the best-fit substitution models as found in jModeltest 2.1 [21,23] among 1624 candidate models according to all four criteria: AIC, AICc, BIC and DT. The best heuristic search algorithm, nearest neighbor interchanges (NNI) and subtree pruning and regrafting (SPR), in (more)PhyML was applied whereas in PAUP, the final tree was searched from 10 starting trees obtained by stepwise and random sequence addition followed by the tree-bisection-reconnection (TBR) branch-swapping algorithm.
To assess support for particular clades, non-parametric bootstrap analyses were performed on 1000 replicates for TreeFinder and PAUP approaches. Additionally, we applied the Local Rearrangements–Expected Likelihood Weights (LR-ELW) method in TreeFinder and the approximate likelihood ratio test (aLRT) based on a Shimodaira-Hasegawa-like procedure in morePhyML [24].

Tree topologies assuming different relationships between the three main Yellow-headed Amazona clades (Central American, South American and North-South American clades) were compared according to approximately unbiased (AU), Shimodaira-Hasegawa (SH) and weighted Shimodaira-Hasegawa (wSH) tests, which were performed in Consel v0.20 [25] assuming 10,000,000 replicates. Site-wise log-likelihoods for the analyzed trees were calculated in TreeFinder under the best fitted substitution models. In these analyses, we considered both all sites and sites with different rate categories, which were created by successive elimination of sites with the highest substitution rate in the given data set.

Divergence times of Yellow-headed Amazona complex were estimated in TreeFinder using global rate minimum deformation (GRMD) and local rate minimum deformation (LRMD) methods for five data sets (nd+cox, nd+cox+atp6+8, 12s+16s+cox, 12s+16s+cox+cytb and cytb) that included appropriate taxa for which divergence time was estimated by Schweizer et al. [26]. Based on their calculations, we used the following calibration points: Pionus menstrus – A. dufresniana/A. aestiva: 10.8 MYA, A. dufresniana – A. pretrei/A. aestiva: 6.4 MYA, A. pretrei – A. aestiva: 4.3 MYA.

Results and Discussion

Phylogenetic analyses were carried out using four approaches on seven data sets. Generally, tree topologies (Figures 1–6, Figure S2) were similar to those obtained by other authors [7,11,14]. Yellow-Headed Amazons were most closely related to Lesser Antillean taxa, A. arausiaca and A. versicolor, and separated from other Amazona sequences.

Phylogenetic Relationships of A. barbadensis

Newly obtained sequences from A. barbadensis enabled us to determine its relationship to other Yellow-Headed Amazons (YHA). All methods performed on five alignment sets consisted of at least two concatenated mitochondrial genetic markers unambiguously indicated close affiliation of A. barbadensis to A. ochrocephala ochrocephala that was represented by samples from Colombia, Guyana and Venezuela (Table 2, Figures 1–5, Figure S2).
S2). The support of this grouping was very high especially for two sets including NADH dehydrogenase, cytochrome oxidase subunit I, ATPase 6 and 8 (Figures 1 and 3). Moreover, the alignment containing rRNA and cytochrome oxidase sequences indicated lower support values of this grouping (Figure 4). Interestingly, addition of more sites from cytochrome \(b\) sequence to this set rather decreased these values (Figure S2). Phylogenies based on single markers, cytochrome oxidase subunit I (Figure 5) and cytochrome \(b\) (Figure 6) were too poorly resolved to give conclusive decision. Nevertheless, phylogenetic trees inferred in MrBayes and PAUP based on cytochrome oxidase alignment clustered \(A. \) barbadensis also with six \(A. \) ochrocephala ochrocephala sequences: two from Colombia, two from Venezuela and one from Guyana and one of unknown geographic locality (Table 2, Table S6, Figure 5). In the case of TreeFinder, this clade included also two Lesser Antillean taxa, \(A. \) arausiaca and \(A. \) versicolor but with low support (Table 2). The cluster of \(A. \) barbadensis and \(A. \) ochrocephala ochrocephala sequences correspond to the Northern South American (NSA) lineage, one of three forming the Yellow-Headed Amazon complex. The remaining two lineages, the Central American (CA) and the South American (SA) obtained also significant support in our studies.

**Phylogenetic Relationships within the Central American Lineage**

\(A. \) ochrocephala panamensis was always placed at the base of the CA lineage (Figures 1–6, Figure S2), which suggests that an expansion of this lineage to Central America started from the territory of Panama and northwest Colombia after the rise of the Isthmus of Panama (Figure 7). Assuming the gradual diversification and expansion of \(A. \) ochrocephala species to subsequent regions located in the north and west, deep into Central America, one could expect that the subsequent branching of other CA taxa in phylogenetic trees corresponds to their successive north-westward geographic distribution.

However, the successive taxon diverging within the CA lineage was \(A. \) tremariae found in Islas Marías (Mary Islands) located in the north and west limit of the CA lineage distribution, whereas the clade of \(A. \) oratrix and \(A. \) aupollaita, inhabiting more southward regions, split later (Figures 1–3). This clade obtained moderate support in our studies of three data sets: \(nd+cox+atp6+8^{1} \), \(nt+cox\) and \(nt+cox+atp6+8^{2}\) (MrBayes: 0.71, 0.80, 0.86; TreeFinder LR-ELW: 48, 54, 54; TreeFinder bootstrap: 45, 79, 84; PAUP bootstrap: 51, 78, 79; morePhyML: 0.74, 0.82, 0.78; PhyML bootstrap: 52, 83, 85, respectively). Trees for cytochrome oxidase (Figure 5) and cytochrome \(b\) (Figure 6) showed different topologies.
than described above but with very low support. The better resolved phylogenies suggest that the present localities of CA taxa may be different from the original ones and were colonized secondarily; or may be the remnants of a much wider distribution that has shrunk to the current one. It is possible that members of A. tresmariae lineage occurred originally in the south of Central America, whence they migrated northwards.

Table 2. Values obtained for different methods and analysed data sets supporting Northern South American lineage including A. barbadensis and its relative A. ochrocephala ochrocephala.

| Data set               | MrBayes | LR-ELW | TF boot | PAUP boot | aLRT | PhyML boot |
|------------------------|---------|--------|---------|-----------|------|------------|
| nd+cox+atp6+8\(^1\)    | 0.99    | 94     | 92      | 73        | 0.93 | 77         |
| nd+cox\(^2\)           | 0.74    | 50     | 70      | 52        | 0.67 | 61         |
| nd+cox+atp6+8\(^2\)    | 1       | 88     | 92      | 86        | 0.90 | 88         |
| 12s+16s+cox\(^3\)      | 0.75    | 57     | 63      | 50        | 0.76 | 59         |
| 12s+16s+cox+cytb\(^4\) | 0.38    | 43     | 48      | 41        | 0.70 | 46         |
| cox                    | 0.33    | 78*    | 40*     | 13        | -    | -          |

\(^1\)based on Ribas et al. [14] data set;  
\(^2\)based on Ribas et al. [14] and modified data set;  
\(^3\)based on Rusello and Amato [7] and modified data set;  
\(^4\)similar to the set\(^3\) but including also cytochrome b;  
\(^*\)the clade contains also A. arausiac and A. versicolor; LR-ELW - Local Rearrangements–Expected Likelihood Weights method; boot – bootstrap method; aLRT - the approximate likelihood ratio test based on a Shimodaira-Hasegawa-like procedure.

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Figure 6. The Bayesian tree for YHA complex and Lesser Antillean Amazons, LA (A) as well as all taxa (B) based on cytochrome b data set. Numbers at the end of some names correspond to the labeling of individuals listed in Table S4. See Figure 1 for other explanations.
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Figure 7. A possible scenario for YHA lineage evolution and their colonization of Central and South America. The most ancestral lineage is most likely NSA, whose distribution is restricted to the northernmost and north-west regions of South America. Their descendants colonized Central America through the isthmus of Panama as well as southward regions of South America, giving rise to CA and SA lineages, respectively. The lineages diversified further into many taxa. However, more intense migrations and introgression likely did not lead to geographic differentiation of genetic markers in the SA lineage, in contrast to the CA lineage, which was subject to greater diversification by geographic locality. doi:10.1371/journal.pone.0097228.g007

Table 3. Relationships between three main Yellow-headed Amazona lineages and their support values obtained for different methods and analysed data sets.

| Data set | MrBayes | LR-ELW | TF boot | PAUP boot | aLRT | PhyML boot |
|----------|---------|--------|---------|-----------|------|------------|
| nd+cox+atp6+8¹ | (S,NS)|0.62 | (S,NS)|0.45 | (S,NS)|0.29 | (S,NS)|0.38 |
| nd+cox² | (S,NS)|0.55 | (S,NS)|0.49 | (S,NS)|0.46 | (S,NS)|0.33 | (C,NS)|0.42 |
| nd+cox+atp6+8² | (S,NS)|0.76 | (S,NS)|0.58 | (S,NS)|0.61 | (S,NS)|0.78 | (S,NS)|0.61 |
| 12s+16s+cox³ | (S,CI)|0.79 | (S,CI)|0.59 | (S,CI)|0.65 | (S,CI)|0.79 | (S,CI)|0.79 |
| 12s+16s+cox+cystb⁴ | (S,CI)|0.54 | (S,CI)|0.16 | (S,CI)|0.47 | (S,CI)|0.42 | (C,NS)|0.15 | (C,NS)|0.41 |
| cox⁵ | (S,CI)|0.80 | (C,NS)*|0.98 | (C,NS)*|1.30 | (S,CI)|0.33 | (S,CI)|0.85 | (S,CI)|0.38 |

¹based on Ribas et al. [14] data set;
²based on Ribas et al. [14] and modified data set;
³based on Rusello and Amato [7] and modified data set;
⁴similar to the set but including also cytochrome b;
⁵the clade contains also A. arasiiaca and A. versicolor; LR-ELW - Local Rearrangements–Expected Likelihood Weights method; boot – bootstrap method; aLRT - the approximate likelihood ratio test based on a Shimodaira-Hasegawa-like procedure.

C – Central American lineage, S – South American lineage, NS – Northern South American lineage including A. barbadensis and its relative A. ochrocephala ochrocephala. doi:10.1371/journal.pone.0097228.t003
Table 4. Results of the approximately unbiased (AU), Shimodaira-Hasegawa (SH) and weighted Shimodaira-Hasegawa (wSH) tests for studied data sets and topologies assuming different relationships between three Yellow-headed *Amazona* lineages: C – Central American, S – South American, NS – Northern South American including *A. barbadensis* and its relative *A. ochrocephala ochrocephala*.

| Data set               | Selected sites | P-values from AU/SH/wSH tests for topology: |
|------------------------|----------------|---------------------------------------------|
|                        |                | (S,NS)                                      |
|                        |                | (C,NS)                                      |
|                        |                | (S,C)                                       |
| nd+cox+atp6+81         | all            | 0.775/0.763/0.755                           |
|                        |                | 0.333/0.284/0.450                           |
|                        |                | 0.212/0.293/0.482                           |
|                        | with rates <2  | 0.766/0.823/0.822                           |
|                        |                | 0.298/0.255/0.382                           |
|                        |                | 0.071/0.080/0.165                           |
|                        | with rates <1  | 1.000/1.000/1.000                           |
|                        |                | 1e-04/2e-05/2e-05                           |
|                        |                | 3e-50/0/0                                  |
|                        | with rates <0.5| 1.000/1.000/1.000                           |
|                        |                | 1e-09/0/0                                 |
|                        |                | 1e-07/0/0                                  |
|                        | with rates <0.1| 1.000/1.000/1.000                           |
|                        |                | 1e-07/0/0                                 |
|                        |                | 6e-68/0/0                                  |
| nd+cox2                | all            | 0.673/0.724/0.715                           |
|                        |                | 0.537/0.561/0.586                           |
|                        |                | 0.131/0.390/0.418                           |
|                        | with rates <2  | 0.959/0.897/0.898                           |
|                        |                | 0.064/0.113/0.197                           |
|                        |                | 0.065/0.112/0.206                           |
|                        | with rates <1  | 0.194/0.330/0.326                           |
|                        |                | 0.860/0.950/0.904                           |
|                        |                | 0.148/0.298/0.097                           |
|                        | with rates <0.5| 5e-04/0.046/3e-04                           |
|                        |                | 0.940/1.000/0.999                           |
|                        |                | 0.067/0.116/0.081                           |
|                        | with rates <0.1| 0.003/0.107/0.006                           |
|                        |                | 0.997/1.000/1.000                           |
|                        |                | 6e-55/0/0                                  |
| 12s+16s+cox3           | all            | 0.747/0.769/0.758                           |
|                        |                | 0.418/0.386/0.500                           |
|                        |                | 0.125/0.293/0.385                           |
|                        | with rates <2  | 0.996/0.999/0.998                           |
|                        |                | 0.004/0.014/0.020                           |
|                        |                | 5e-08/1e-06/1e-06                           |
|                        | with rates <1  | 0.918/0.939/0.934                           |
|                        |                | 0.095/0.097/0.166                           |
|                        |                | 0.007/0.022/0.035                           |
|                        | with rates <0.5| 0.966/0.979/0.962                           |
|                        |                | 0.034/0.040/0.070                           |
|                        |                | 5e-05/3e-04/5e-04                           |
|                        | with rates <0.1| 0.755/0.838/0.841                           |
|                        |                | 0.245/0.343/0.400                           |
|                        |                | 1e-06/1e-04/7e-06                           |
|                        | with rates <0.5| 0.578/0.725/0.715                           |
|                        |                | 0.620/0.733/0.742                           |
|                        |                | 0.154/0.175/0.176                           |
|                        | with rates <0.1| 0.896/0.935/0.972                           |
|                        |                | 0.192/0.483/0.288                           |
|                        |                | 0.028/0.047/0.044                           |
| 12s+16s+cox+cytb4      | all            | 0.147/0.394/0.451                           |
|                        |                | 0.515/0.514/0.565                           |
|                        |                | 0.670/0.722/0.712                           |
|                        | with rates <2  | 0.037/0.353/0.342                           |
|                        |                | 0.577/0.636/0.642                           |
|                        |                | 0.613/0.651/0.648                           |
|                        | with rates <1  | 0.127/0.180/0.146                           |
|                        |                | 0.835/0.922/0.922                           |
|                        |                | 0.263/0.356/0.330                           |
|                        | with rates <0.5| 0.002/0.001/0.001                           |
|                        |                | 0.971/0.994/0.994                           |
|                        |                | 0.038/0.067/0.061                           |
|                        | with rates <0.1| 4e-05/0.001/0.001                           |
|                        |                | 1.000/1.000/1.000                           |
|                        |                | 7e-93/0/0                                  |

1 based on Ribas et al. [14] data set;
2 based on Ribas et al. [14] and modified data set;
3 based on Rusello and Amato [7] and modified data set;
4 similar to the set 3 but including also cytochrome b.

The tests were performed for all sites and sites with decreasing substitution rate. P-values of the best tree are underlined whereas those less than 0.05 are in bold.
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Table 5. Results of the approximately unbiased (AU), Shimodaira-Hasegawa (SH) and weighted Shimodaira-Hasegawa (wSH) tests for the cytochrome b set and topologies assuming different relationships between three Yellow-headed *Amazona* lineages: C – Central American, S – South American, NS – Northern South American including *A. barbadensis* (Ab) and its relative *A. ochrocephala ochrocephala*.

| Selected sites | P-values from AU/SH/wSH tests for topology: |
|----------------|---------------------------------------------|
|                | (C,Ab)                                      |
|                | (C,NS)                                      |
|                | (S,NS)                                      |
|                | (S,C)                                       |
| all            | 0.771/0.816/0.852                           |
|                | 0.572/0.713/0.652                           |
|                | 0.211/0.272/0.484                           |
|                | 0.220/0.262/0.462                           |
| with rates <2  | 0.450/0.636/0.531                           |
|                | 0.867/0.897/0.943                           |
|                | 0.116/0.236/0.417                           |
|                | 0.104/0.234/0.402                           |
| with rates <1  | 0.887/0.894/0.938                           |
|                | 0.200/0.621/0.392                           |
|                | 0.033/0.091/0.197                           |
|                | 0.034/0.082/0.175                           |
| with rates <0.5| 0.096/0.519/0.061                           |
|                | 0.917/0.996/0.998                           |
|                | 0.031/0.178/0.300                           |
|                | 0.031/0.103/0.005                           |
| with rates <0.1| 0.002/0.187/0.006                           |
|                | 0.998/1.000/1.000                           |
|                | 4e-92/4e-05/7e-06                           |
|                | 1e-65/0/0                                  |

The tests were performed for all sites and sites with decreasing substitution different site rate. P-values of the best tree are underlined whereas those less than 0.05 are in bold.
doi:10.1371/journal.pone.0097228.t005
lineage. These results may indicate a misidentification of these samples or crossbreeding between *A. oratrix* subspecies as well as between *A. oratrix* and *A. auropalliata*. In agreement with that, cytochrome oxidase sequences appeared identical between these corresponding taxa (Figure 5). The same sequence showed also *A. auropalliata auropalliata* and *A. auropalliata parvipes* are compatible with findings of Lousada and Howell [28], who noted that *A. auropalliata cardiaca* from the Bay Islands off the coast of Honduras shows plumage intermediate between that of the Atlantic slope *A. oratrix hondurensis* and the Pacific slope *A. auropalliata*. It was used as one of arguments to combine *A. auropalliata* and *A. oratrix* to one taxon [28]. In agreement with that, *A. oratrix* and *A. auropalliata* were always grouped together in the phylogenetic trees.

**Phylogenetic Relationships within the South American Lineage**

The SA lineage clearly split into two significantly supported clades in our phylogenies (Figures 1–3, Figure 5) as in Ribas et al. [14]. Each of these two clades possessed a mixture of *Amazona* assigned to different species. Clade I included: *A. ochrocephala nattereri*, *A. ochrocephala xantholaema* and *A. aestica aestica*, whereas clade II included: *A. ochrocephala ochrocephala*, *A. ochrocephala nattereri* and *A. aestica xanthopteryx*. The separation of these clades corresponds neither to the course of the Amazon River nor Amazonian river dynamics in contrast to other taxa [29–31]. Their split may have resulted from other barriers, e.g. changes in type, composition and structure of vegetation that could occur during glacial cycles during the Pleistocene [32–37] and are not evident today [14]. The lack of monophyly for these species as well as the inconsistency between phylogenetic relationships and geographic distribution may result from misinterpretation of the morphological variation and hence wrong taxonomical classification especially in the case of unvouched specimens [38], incomplete lineage sorting, and migrations accompanied with introgression [14]. In support of the latter, we identified identical cytochrome oxidase sequences assigned to the different species: *A. aestica aestica* = *A. ochrocephala ochrocephala* = *A. ochrocephala nattereri*; *A. aestica xanthopteryx* = *A. ochrocephala nattereri* and *A. aestica aestica* = *A. ochrocephala xantholaema* (Figure 5). All these results, together with the clear separation of *A. ochrocephala* subspecies into the three main YHA lineages, indicate a need of revision of the whole *Amazona ochrocephala* taxon. There is some support for *A. ochrocephala panamensis* to be split from *A. ochrocephala* and form a separate species (*A. panamensis*), as it is the only representative of *A. ochrocephala* in the CA lineage. Moreover, in contrast to other *A. ochrocephala* subspecies, individuals of *A. o. panamensis* are characterized by smaller body size and pinkish-horn bill coloration [3,4].

**Phylogenetic Relationships between Three Main Lineages of the YHA Complex**

Although three lineages of the YHA complex were clearly recognized, the relationship between them was less obvious and depended on data sets rather than methods used (Table 3). Phylogenies based on nd+cox+atp6+82, nd+cox and nd+cox+atp6+82 data sets indicated the grouping of SA and NSA lineages for all methods with one exception (consensus from PhyML bootstrap trees). It suggested a closer relationship between the CA and NSA lineages than other possibilities. On the other hand, trees based on 12s+16s+cox, 12s+16s+cox+cylt and cox data sets suggested closer affiliation of SA and CA lineages by most methods. Only PhyML approaches using 12s+16s+cox+cylt and TreeFinder methods based on cytochrome oxidase preferred the CA and NSA clustering. In the latter case, the NSA lineage included also *A. aracacuara* and *A. versicolor*. Trees inferred on cytochrome *b* placed separated *A. barbadensis* and *A. ochrocephala ochrocephala*, representatives of the NSA lineage, and placed them basal to the CA lineage (Figure 6).

However, in almost all approaches support values were low or at most moderate so it was difficult to favor one of these possibilities. Therefore we performed tests on tree topologies assuming different relationships between the three YHA lineages (Table 4 and 5). As would be expected, the comparison of the different scenarios (using all sites in the alignments) preferred none of them. However, when sites with the highest substitution rate in the given data set were successively eliminated, some topologies occurred with more statistically significant than others. Two topologies that clustered NSA with CA or SA lineages were favored, whereas the grouping of CA and SA lineages was significantly worse. The NSA+CA clade was preferred for nd+cox, 12s+16s+cox+cylt and cylt data sets, while the NSA+SA clade was chosen for 12s+16s+cox and two data sets including four markers, nd+cox+atp6+82 and nd+cox+atp6+82. Interestingly, the CA+SA clade obtained in trees based on all sites of 12s+16s+cox and 12s+16s+cox+cylt alignments was rejected when rapidly evolving sited were removed.

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**Table 6. Divergence time in million years estimated by two methods for analysed data sets.**

| Data set | GRMD  | LRMD  |
|----------|-------|-------|
| nd+cox   | 0.94  | 0.99  |
| nd+cox+atp6+82 | 1.04  | 0.99  |
| 12s+16s+cox   | 1.22  | 1.16  |
| 12s+16s+cox+cylt | 1.24  | 1.23  |
| cylt     | 1.13  | 1.11  |

*based on Ribas et al. [14] and modified data set;
*based on Rusello and Amato [7] and modified data set;
*similar to the set but including also cytochrome b.

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from these data sets. Moreover, the clade grouping two members of the NSA lineage, *A. barbadensis* and *A. ochrocephala ochrocephala*, was preferred for more conserved sites than for all position of cytochrome *b* alignment (Table 5). Although these results suggest that phylogenetic signal is too low to unambiguously select one of two scenarios (NSA + CA or NSA + SA), the close relationship of NSA to SA could be considered preferentially because was based on longer alignments both for all and conserved sites. This possibility is supported by the strong affiliation of *A. barbadensis* with the taxon *A. ochrocephala ochrocephala* whose members were also assigned to the SA lineage. The ambiguity in inferring relationships between YHA lineages was probably caused by their quick evolution and divergence in a short time frame, as is also indicated by very short internal branches leading to these lineages in phylogenetic trees. Incomplete lineage sorting and introgression between newly emerging clades before their final separation could also decrease resolution of the trees and obscure the deep relationships among taxa.

**Possible Scenario for the Evolution of YHA Taxa and their Colonization of the American Mainland**

Based on these analyses, we can propose a scenario for the evolution of YHA taxa and their colonization of Central and South America (Figure 7). Since two Lesser Antillean Amazons, *A. arausiaca* and *A. versicolor*, are most closely related to YHA taxa and especially members of the NSA lineage (as found in some phylogenetic trees, Figure 5), they or their relatives seem to be good potential YHA ancestors. They probably colonized the northern coast of Venezuela, which also corresponds with the past and present distribution of *A. barbadensis* (Figure 7). In support of this, *A. barbadensis* lives or inhabited in the past also some islands in the Lesser Antilles. Then the NSA taxa represent likely descendants of the YHA ancestors. After the colonization of the mainland, rapid evolution and divergence into new lineages accompanied the occupation of new habitats and/or free ecological niches. According to our estimates (Table 6), the migration to the mainland and emergence of the YHA complex happened about million years ago and could be related to climatic changes due to glacial cycles [39,40].

A further colonization event to Central America through the already existing Isthmus of Panama [41] initiated the origin of the CA lineage (Figure 7). This is in agreement with the distribution of *A. ochrocephala panamensis*, which occupies geographic regions adjacent to those of the NSA taxa and was placed at the base of the CA lineage in phylogenetic trees (Figures 1–6, Figure S2). The southward expansion to the South America proceeded somewhat simultaneously. However, greater gene flow or migration may have occurred between NSA and SA than between more isolated CA taxa, as is suggested by tree topology tests, in which the NSA-SA clade was preferred by larger data sets (Table 4). The migrations and introgression could prevent geographic differentiation of genetic markers in the SA lineage, despite the larger occupied area, in contrast to the CA lineage [14]. It seems that changes during the Late Pleistocene in the Amazon Basin and adjacent regions inhabited by SA taxa were insufficient to cause distinct differentiation of species corresponding to their geographic distribution. Forests only in parts of eastern lowland Amazonia were at times more open and savannas may have developed only locally as a result of drier or more seasonal conditions. However, the majority of the Amazon rainforest remained intact [37,42,43]. In contrast to that, the CA taxa could undergo more intense diversification correlated with their geographic locality because of stronger influence of climatic changes on their habitat alterations and fragmentations related with glacial cycles. In fact it has been shown that high elevated regions of Central America were glaciated during the Late Pleistocene [44] with cooler conditions influencing also lowland forest vegetation and moisture pattern throughout the Central America [45–49]. This hypothesis corresponds with the results of Weir [50] who found higher speciation rates in the highland than lowland Neotropical birds during the last one million years because of climatic and glacial fluctuations. In agreement with this, CA taxa are mainly highland and upland species, whereas SA taxa favor lowlands. Nevertheless, further studies involving more samples from the whole distribution of YHA taxa represented by more genetic markers, including completely sequenced mitochondrial genomes, are necessary to verify the suggested scenario.

**Supporting Information**

Figure S1 Individual of Yellow-shouldered Amazon (*Amazona barbadensis*) whose sequences were analysed in the paper. (TIF)

Figure S2 The Bayesian tree for YHA complex and Lesser Antillean Amazons, LA (A) as well as all taxa (B) based on 12s+16s+cox+cytb data set. See Figure 1 for other explanations and Table S5 for more details. (TIF)

Table S1 List of taxa, individuals and sequences included in: nd+cox+atp6,+8', nd+cox2 and nd+cox+atp6,+8' data sets. # - number of the species individual. Length - the length of the whole combined mitochondrial sequence. (XLSX)

Table S2 List of taxa and sequences included in the 12s+16s+cox3 data set. Length - the length of the whole combined mitochondrial sequence. (XLSX)

Table S3 List of taxa, individuals and sequences included in the cytochrome oxidase data set. (XLSX)

Table S4 List of taxa, individuals and sequences included in the cytochrome *b* data set. (XLSX)

Table S5 List of taxa and sequences included in: 12s+16s+cox+cytb data set. Length - the length of the whole combined mitochondrial sequence. (XLSX)

Table S6 List of studied specimens including their voucher number, geographic location, locality code, references and institution hosted them. (XLSX)

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**Author Contributions**

Conceived and designed the experiments: AU PM. Performed the experiments: PM TS. Analyzed the data: AU PM TS. Contributed reagents/materials/analysis tools: AU PM TS. Wrote the paper: PM AU TS. Corresponded with editor and reviewers: AU PM.
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