

## Title

Comment on “A genetic signature of the evolution of loss of flight in the Galapagos cormorant”

## Authors

Mark J. Berger<sup>1</sup>, Gill Bejerano<sup>1,2,3,4</sup>

## Affiliations

<sup>1</sup> Department of Computer Science, Stanford University, Stanford, CA, 94305-5329, USA

<sup>2</sup> Department of Developmental Biology, Stanford University, Stanford, CA, 94305-5329, USA

<sup>3</sup> Department of Pediatrics, Stanford University, Stanford, CA, 94305-5329, USA

<sup>4</sup> Correspondence – [bejerano@stanford.edu](mailto:bejerano@stanford.edu)

## Abstract

Burga et al. could not identify regulatory regions which might contribute to flightlessness in the Galapagos cormorant. Using a bird-specific alignment we discover 48 limb enhancers showing strong accelerated evolution in *P. harrisi*, including enhancers of key cilia development, hedgehog signaling, and planar cell polarity genes, such as *Prickle1*, *Twist2* and *Ndk1*, extending Burga’s proposed mechanism into the non-coding genome.

## Main text

Burga et al. (1) sequenced the genomes of four cormorant species to identify candidate mutations which may have contributed to flightlessness in the Galapagos cormorant. The authors found numerous function-altering variants in genes related to cilia development, hedgehog signaling, and the planar cell polarity (PCP) pathway, such as *Ift122*, *Gli2*, and *Fat1*. They also identified a 4-amino acid deletion in *Cux1*, a highly-conserved homeodomain transcription factor. The authors demonstrate that *Cux1* regulates the expression of key cilia and PCP genes, as well as that the 4-amino acid deletion impairs *Cux1* from activating gene expression. These results suggest that perturbations of cilia development, hedgehog signaling, and the PCP pathway may play a role in reduced keel and wings of the Galapagos cormorant.

While the authors implicated many protein-coding mutations in the disruption of the primary cilium, they found no evidence of a non-coding signature. The authors aligned phastCons (2) conserved regions from the UCSC 100-way vertebrate multiple alignment (3) to each of the four cormorant genomes using reciprocal best BLAST alignments. Using PhyloP (4), the authors tested each of the resulting 40,812 candidate regions for accelerated evolution in *P. harrisi* compared to all tetrapods. With a false-discovery rate (FDR) cutoff of 5%, Burga et al. found only 11 accelerated regions in *P. harrisi*, none of which overlapped limb enhancer marks.

Here, we show that by testing regions which are conserved among avian species, instead of those conserved among vertebrates, we discover 48 accelerated non-coding regions which overlap limb enhancer marks. Using the four cormorant species sequenced by the authors, as well as *Phalacrocorax carbo*, *Nipponia nippon*, and *Gallus gallus galGal5* assembly as reference

(Figure 1A), we assembled a genome-wide multiple alignment between the seven species using MULTIZ (5). A neutral model of evolution was constructed using fourfold degenerate sites in chicken derived from the Ensembl 89 gene set (6). Ignoring *P. harrisi*, phastCons (2) identified 894,840 highly conserved regions not overlapping a protein-coding gene in chicken. Non-coding conserved regions within 100bp of each other were merged, and regions smaller than 100bp were removed, resulting in 430,015 candidate regions (Figure 1B).

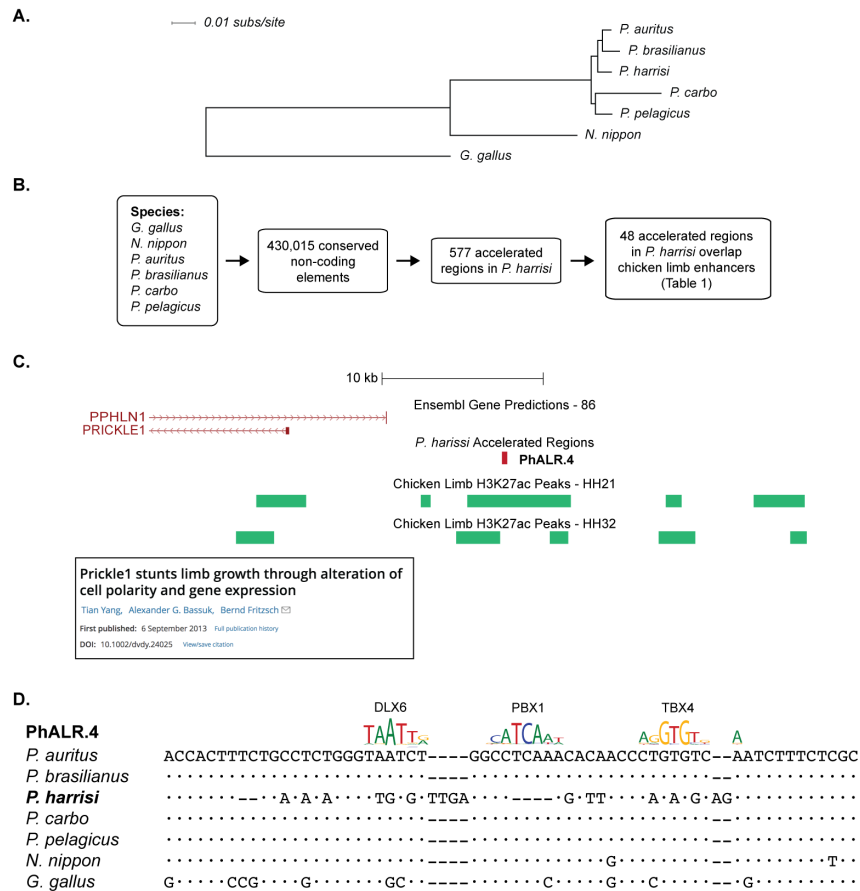
Using PhyloP (4) we asked whether any of these candidate regions have undergone accelerated molecular evolution in the Galapagos cormorant. Even with a stringent FDR cutoff of 1%, we found 577 accelerated regions in *P. harrisi* (Figure 1B). Of these, 48 elements overlap H3K27Ac enhancer marks at chicken developmental time points HH21 or HH32 by at least 50% (Table 1) (7). We refer to this set of 48 regions as *P. harrisi* Accelerated Limb Regions (PhALRs).

A number of the resulting regions corroborate Burga et al.'s proposed model. For example, PhALR.4 lies 13kb upstream of *Prickle1* (Figure 1C) and is highly diverged in the Galapagos cormorant (Figure 1D). *Prickle1* is a key member of the PCP pathway, and mice mutants with a premature stop codon in the 3<sup>rd</sup> LIM domain exhibit defective limb growth (8). Mutants also exhibit altered expression of key developmental limb genes, such as *Bmp4* and *Wnt5a* (8). Furthermore, mice with a missense mutation in *Prickle1* display primary cilium defects (9). PhALR.11 lies 31kb upstream of *Ndk1*, a highly-conserved gene which regulates Wnt signaling. Knockdown of *ndk1* in zebrafish leads to loss of cilia in Kupffer's vesicle (10).

Other regions suggest that additional mechanisms contribute to the reduced keel and wings in the Galapagos cormorant. PhALR.44 lies 11kb upstream of *Twist2*. *Twist2* is a bHLH transcription factor which represses *Grem1* to terminate the Shh/Grem1/Fgf autoregulatory loop in early limb development (11). Viral-induced misexpression of *Twist2* results in underdeveloped limbs in mice (11). PhALR.21 lies in the intron of *Dync1i1*, proximal to exon 15. Exon 15 and 17 of *DYNC1I1* have been shown to function as distal enhancers for *DLX5* and *DLX6*, two critical regulators of limb development (12). Chromosomal abnormalities of *DYNC1I1*, as well as simultaneous disruption of *Dlx5* and *Dlx6*, both lead to split hand and foot malformation in humans and mice, respectively (12, 13). Other *P. harrisi* Accelerated Limb Regions are found next to *SMAD*, *TBX*, *IRX* genes and more (Table 1).

Burga et al. identified genes in *P. harrisi* with striking loss of function mutations and demonstrated that cilia development, hedgehog signaling, and the PCP pathway may be responsible for flightlessness in the Galapagos cormorant (1). Developmental genes are highly pleiotropic and will only accumulate deleterious mutations when the organism can withstand the corresponding loss in fitness throughout all developmental contexts (14). It is plausible that additional genes contribute to flightlessness in the Galapagos cormorant, but present with no loss of function coding mutations. Instead, the oft less pleiotropic *cis*-regulatory elements for these genes may mutate more freely. Here, we discover multiple putative limb enhancers which have diverged significantly in the Galapagos cormorant, including those upstream of functionally relevant genes. Our results corroborate the model proposed by Burga et al., and extend it into the non-coding genome.

## Figures & Tables



**Figure 1. Identifying accelerated limb enhancers in the Galapagos cormorant. (A)** Phylogeny of our bird-specific multiple alignment. **(B)** Ignoring *P. harrisi*, we identified 430,015 conserved non-coding elements. Of these, 577 elements are significantly accelerated in the Galapagos cormorant. 48 of these overlap chicken limb enhancer marks. **(C)** For example, PhALR.4 lies 13kb upstream of *PRICKLE1*. Mouse *Prickle1* mutants experience stunted limb growth (8) and primary cilia defects (9). **(D)** A portion of PhALR.4 is shown highly diverged in *P. harrisi*. Bases which are identical to the sequence in *P. auritus* are represented by a dot, the absence of a basepair is denoted by a dash. Despite the very short evolutionary distance between *P. auritus* and *P. harrisi* (Figure 1A), the corresponding sequence in *P. harrisi* is even more diverged than the corresponding sequence in *G. gallus*. Using position weight matrices (15), we predicted three binding sites of important limb development regulators which appear eroded in the Galapagos cormorant.

galGal5 Chrom	Start	End	Element Name	Nearest genes (Distance to TSS)
chr1	1174874	1175514	PhALR.1	TUBAL3 (-23.3k), NET1 (-26.5k)
chr1	6994904	6995093	PhALR.2	FRMD4A (-90.4k), ENSGALG00000033226 (+357.5k)
chr1	25813481	25813762	PhALR.3	MDFIC (-555)
chr1	29721579	29721941	PhALR.4	PRICKLE1 (-13.1k), ADAMTS20 (+286.2k)
chr1	34364240	34364404	PhALR.5	LLPH (+4.9k), HMGI-C (+160.3k)
chr1	35924280	35924467	PhALR.6	ENSGALG00000040855 (+20.2k), PTPRB (-32.5k)
chr1	57866756	57866951	PhALR.7	CHRM2 (-1.1k)
chr1	62063564	62064074	PhALR.8	ENSGALG00000013042 (-24.9k), PEX26 (-50.5k)
chr1	104823969	104824070	PhALR.9	MIS18A (+57.3k), HUNK (+91.9k)
chr10	18730427	18730944	PhALR.10	SMAD6 (+31.2k), SMAD3 (-81.7k)
chr11	6499531	6499950	PhALR.11	NKD1 (-31.1k), BRD7 (-130.7k)
chr11	12075244	12075577	PhALR.12	CDH11 (-257.7k), CDH5(-372.0k)
chr11	17481889	17482561	PhALR.13	ENSGALG00000029278 (+102.0k), ENSGALG00000039008 (+209.6k)
chr12	9433627	9433759	PhALR.14	EEFSEC (+80.4k), ENSGALG00000044250 (-129.2k)
chr13	13174020	13174369	PhALR.15	TCOF1 (-5.4k), ARSI (-9.6k)
chr14	4240133	4240298	PhALR.16	RNF216 (+30.5k), FSCN1 (+33.5k)
chr15	1350352	1350509	PhALR.17	ZDHHC8 (+24.5k), PGAM5(-71.6k)
chr15	12355200	12355870	PhALR.18	TBX5 (-25.2k), TBX3 (+77.6k)
chr2	13968283	13969054	PhALR.19	ITGB1 (-77.5k), NRP1(+82.1k)
chr2	20451532	20451702	PhALR.20	FAM171A1 (-68.8k), ITGA8 (+87.4k)
chr2	24132193	24133200	PhALR.21	SLC25A13 (+110.9k), DYNC111 (+146.9k)
chr2	35320405	35320605	PhALR.22	SATB1 (+13.9k), TBC1D5 (-293.3k)
chr2	61118995	61119310	PhALR.23	JARID2 (-146.7k), CD83(+226.0k)
chr2	62397629	62398027	PhALR.24	TBC1D7 (-18.5k), GFOD1(+60.8k)
chr2	64221316	64222213	PhALR.25	SLC35B3 (-116.7k), ENSGALG00000042845 (+438.6k)
chr2	86816902	86817718	PhALR.26	IRX2 (+90.1k), IRX4(-485.9k)
chr21	3120789	3121217	PhALR.27	ENSGALG00000030881 (+27.1k), ENSGALG00000002232 (-79.0k)
chr23	2021824	2022005	PhALR.28	AHDC1 (+5.2k), WASF2 (-25.0k)
chr23	5507699	5507877	PhALR.29	HEYL (-6.9k), NT5C1A (+10.7k)
chr26	4264439	4264822	PhALR.30	ANKS1A (+7.3k), ENSGALG00000002692 (+87.0k)
chr28	4774298	4774715	PhALR.31	KDM4B (+68.6k), PTPRS(+148.4k)
chr3	14230323	14230448	PhALR.32	PLCB4(-183.5k), PLCB1(+349.7k)
chr3	54386692	54387333	PhALR.33	NHSL1 (+12.5k), ENSGALG00000035680 (+178.5k)
chr3	56803537	56804397	PhALR.34	EYA4 (-78.3k), RPS12(+161.6k)
chr33	269239	269347	PhALR.35	CERS5 (-5.0k), LIMA1 (+17.3k)
chr4	32409496	32409661	PhALR.36	NR3C2 (+161.5k), ARHGAP10 (+169.1k)
chr4	32486613	32486885	PhALR.37	NR3C2 (+84.3k), ARHGAP10(+246.3k)
chr4	39319472	39319936	PhALR.38	TACR3 (-129.2k), TET2 (-212.5k)
chr4	66532631	66532867	PhALR.39	SPATA18 (+54.3k), USP46 (+119.3k)
chr5	28082372	28082505	PhALR.40	GALNT16 (+19.7k), ERH(+56.5k)
chr6	12739800	12740116	PhALR.41	ZMIZ1 (-327.4k), RPS24(+418.0k)
chr6	33850891	33851286	PhALR.42	MGMT (-178.5k), ENSGALG00000025996 (+443.9k)
chr7	4535095	4535543	PhALR.43	MREG (+28.8k), FN1(-95.9k)
chr7	6116197	6116726	PhALR.44	TWIST2 (-11.0k), ASB1 (+142.6k)
chr7	33589086	33589506	PhALR.45	Sip1 (-892)
chr8	15800494	15800807	PhALR.46	LMO4 (+864)
chr9	20089307	20089868	PhALR.47	MECOM (+26.6k), ENSGALG00000031382 (+69.3k)

chrZ	79330495	79331788	PhALR.48	ZNF608 (-293.3k), GRAMD3 (-345.5k)
------	----------	----------	----------	------------------------------------

**Table 1. *P. harrisi* Accelerated Limb Regions (PhALRs).** Gene regulatory domains are assigned using the default rules of Genomic Regions Enrichment of Annotations Tool (GREAT) v3.0.0. The nearest transcription start sites are reported on either side of each element. Negative distances are upstream of each transcription start site and positive distances are downstream.

## References

1. A. Burga *et al.*, *Science*. **356** (2017), doi:10.1126/science.aal3345.
2. A. Siepel *et al.*, *Genome Res.* **15**, 1034–1050 (2005).
3. W. J. Kent *et al.*, *Genome Res.* **12**, 996–1006 (2002).
4. K. S. Pollard, M. J. Hubisz, K. R. Rosenbloom, A. Siepel, *Genome Res.* **20**, 110–121 (2010).
5. M. Blanchette *et al.*, *Genome Res.* **14**, 708–715 (2004).
6. A. Yates *et al.*, *Nucleic Acids Res.* **44**, D710–D716 (2016).
7. R. Seki *et al.*, *Nat. Commun.* **8**, 14229 (2017).
8. T. Yang, A. G. Bassuk, B. Fritsch, *Dev. Dyn. Off. Publ. Am. Assoc. Anat.* **242**, 1293–1306 (2013).
9. B. C. Gibbs *et al.*, *Biol. Open*, in press, doi:10.1242/bio.015750.
10. I. Schneider *et al.*, *Dev. Biol.* **348**, 22–33 (2010).
11. C. Wade, I. Brinas, M. Welfare, C. Wicking, P. G. Farlie, *Dev. Biol.* **370**, 145–153 (2012).
12. R. Y. Birnbaum *et al.*, *Genome Res.* **22**, 1059–1068 (2012).
13. R. F. Robledo, L. Rajan, X. Li, T. Lufkin, *Genes Dev.* **16**, 1089–1101 (2002).
14. S. B. Carroll, *Cell*. **134**, 25–36 (2008).
15. H. Guturu, S. Chinchali, S. L. Clarke, G. Bejerano, *PLOS Comput. Biol.* **12**, e1004711 (2016).