Parallel Genetic Origin of Foot Feathering in Birds

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Abstract

Understanding the genetic basis of similar phenotypes shared between lineages is a long-lasting research interest. Even though animal evolution offers many examples of parallelism, for many phenotypes little is known about the underlying genes and mutations. We here use a combination of whole-genome sequencing, expression analyses, and comparative genomics to study the parallel genetic origin of ptilopody (Pti) in chicken. Ptilopody (or foot feathering) is a polygenic trait that can be observed in domesticated and wild avian species and is characterized by the partial or complete development of feathers on the ankle and feet. In domesticated birds, ptilopody is easily selected to fixation, though extensive variation in the type and level of feather development is often observed. By means of a genome-wide association analysis, we identified two genomic regions associated with ptilopody. At one of the loci, we identified a 17-kb deletion affecting PITX1 expression, a gene known to encode a transcription regulator of hindlimb identity and development. Similarly to pigeon, at the second loci, we observed ectopic expression of TBX5, a gene involved in forelimb identity and a key determinant of foot feather development. We also observed that the trait evolved only once as foot-feathered birds share the same haplotype upstream TBX5. Our findings indicate that in chicken and pigeon ptilopody is determined by the same set of genes that affect similar molecular pathways. Our study confirms that ptilopody has evolved through parallel evolution in chicken and pigeon.

Key words: parallel evolution, chicken, foot feathering, PITX1, TBX5, limb development.

Introduction

Parallel evolution is the independent development of similar phenotypic traits in separate but related lineages (Davis and Heywood 1963). A defining characteristic of parallel evolution is that the trait is absent in the common ancestor. Animal evolution offers many examples of parallelism. These range from seemingly simple phenotypic changes, such as similar female-limited Batesian mimicry in Papilio butterflies (Iijima et al. 2018), to more complex ones, such as the reduction of the pelvic complex in threespine and ninespine sticklebacks (Shapiro et al. 2006). Even though studies have shown that parallel evolution is not a rare event in animal evolution (Mundy et al. 2004; Shapiro et al. 2006), the genetic bases are still largely unknown for most traits. One hypothesis is that phenotypes that have evolved in parallel and that are extremely similar must have evolved by applying the same genetic mechanism due to developmental constraints. However, at the other extreme, it can be hypothesized that developmental pathways are so complex and can be perturbed in so many ways that essentially an infinite number of combinations of variations could lead to the same outcome.
that has evolved in parallel in the two species through artificial selection.

In this study, we focus on ptihopody (Pti), a trait observed in domesticated and wild avian species in which the epidermis of the ankle and foot are partially or completely covered with feathers (Boer et al. 2017; Domyan and Shapiro 2017) (fig. 1). The genetic basis of ptihopody has been extensively studied. Previous classical breeding experiments identified a small number of genetic loci (Pti-1, Pti-2, and Pti-3) of large effect in chicken (Somes 1990). However, only recently several candidate genomic regions were identified. Although Dornyshorst et al. (2010) identified one quantitative trait locus (QTL) of major effect in Silkie chickens (Dornyshorst et al. 2010), Sun et al. (2015) mapped four QTLs, two of which explain more than 20% of the phenotypic variation (Sun et al. 2015). Although recent studies have tried to associate ptihopody to a certain genomic region, it is still unclear which chromosome(s), gene(s), and mutation(s) are directly responsible for the phenotype. A better understanding of the genomic architecture underlying ptihopody comes from a recent study in domestic pigeon, in which two genomic regions containing genes responsible for a partial transformation from hindlimb to forelimb identity were implicated, mainly PITX1 and TBX5 (Domyan et al. 2016). Ptihopody in chicken and pigeon is extremely similar in appearance and this similarity is partly explained by the same genes involved (Domyan et al. 2016). Even though the same genes are involved, the question is whether a similar underlying mutation has enabled the trait to evolve in both lineages. And, even more intriguing, if indeed a similar mutation was to be involved, it becomes relevant to question how the same pathways are altered by the same regulatory mechanisms in both species. This question has never been addressed directly before, neither from a molecular nor an evolutionary perspective.

In this study, we use a combination of whole-genome sequencing (WGS), expression analyses, and comparative genomics to study the genetic basis of foot feathering in chicken. In particular, we identify the underlying causal mutations and affected molecular pathways, while investigating the parallel genetic origin of ptihopody.

Results

WGS of Scaled and Foot-Feathered Chickens

We unraveled the genetic basis and evolutionary history of ptihopody in chicken by WGS of 169 samples from a variety of domesticated chicken breeds and wild species of Gallus (i.e., G. gallus, G. sonneratii, G. lafayetii, and G. varius). On average, 148× coverage was generated for each individual after mapping to the chicken reference genome. Mapping quality was, on average, 33.4 with more than 98% of the reads successfully mapped (supplementary Additional file 1, table S3, Supplementary Material online). Since missing data comprised 12% of the total sites, we imputed missing genotypes and phased haplotypes with high accuracy using 21.0 million single-nucleotide polymorphisms (SNPs) and 1.4 million Insertion/Deletions (InDels) (supplementary Additional file 2, table S2, Supplementary Material online). Variants were also assigned to a range of functional classes, though the vast majority were located in introns (58%) and intergenic regions (30%) similarly to Lawal et al. (2018) (supplementary Additional file 2, table S3, Supplementary Material online). Of the 359,176 protein-coding variants, 217,255 were classified as synonymous, 130,147 as missense, and 11,774 as loss-of-function (supplementary Additional file 2, table S3, Supplementary Material online).

To assess population stratification, we performed a principal component analysis (PCA) and Neighbor-Joining (NJ) analysis on the 19 cases and 150 controls. The PCA did not identify any distinct clustering between the two groups (supplementary Additional file 3, fig. S1, Supplementary Material online) aside from clearly separating all traditional breeds from the individuals of the four wild species of Gallus. However, the NJ tree based on the distance relationship matrix separated the 19 case individuals into three groups, one of the Breda fowl, one of the Dutch booted bantam, and one of the remaining foot-feathered samples (supplementary Additional file 3, fig. S2, Supplementary Material online). Despite that, one sample from the Marans (sample 1283), Sundheimer (sample 1769), and German Favoroles (sample 641) breed did not form any specific cluster.

Two Genomic Regions Control Foot Feathering in Chicken

We conducted a genome-wide association study (GWAS) on the 169 samples using a case/control approach through an adaptive Monte Carlo permutation test with 5,000 replications. The GWAS revealed two significant signals on chromosomes 13 and 15, respectively (fig. 2a). On chromosome 13, we identified 36 significant variants (8 intergenic, 9 upstream gene, and 19 intron variants), which are located between 16.0 and 16.1 Mb (supplementary Additional file 2, table S4, Supplementary Material online). This 57-kb region contains a protein-coding H2A histone family gene, H2AFY, a novel long noncoding RNA, ENSGALG000000048757, and one QTL, QTL127125, which was previously found to be associated with foot feathering (Sun et al. 2015) (fig. 2b). The gene H2AFY is located 145 kb upstream of PITX1, a gene that encodes a homeobox-containing transcription factor that is normally expressed in the vertebrate hindlimb but not the forelimb (Logan et al. 1998; Logan and Tabin 1999; Rodriguez-Esteban et al. 1999; Takeuchi et al. 1999). For this 57-kb region, foot-feathered breeds showed elevated levels of homozygosity relative to scaled birds, a clear signature of positive selection as indicated by the low pooled heterozygosity (ZH = −3.71 vs. ZH control = 0.80 in control individuals) (supplementary Additional file 3, fig. S4b, Supplementary Material online).

On chromosome 15, the 23 significant variants (15 intron noncoding, 3 upstream gene, and 5 intergenic) defined a 112-kb region (12.5–12.6 Mb) (supplementary Additional file 2, table S5, Supplementary Material online), in which we identified a T-box 5 protein-coding gene, TBX5, a novel IncRNA, ENSGALG00000052717, and a previously identified QTL, QTL127126 (Sun et al. 2015) (fig. 2c). Most of the variants were found in the IncRNA. Within this 112-kb region, we also identified one candidate selective sweep (15:12,560,000–
Foot Feathering Has a Single Haplotype Origin

We also performed a CNV analysis on chromosome 15 but did not observe any CNV potentially associated with the phenotype. We therefore decided to reconstruct haplotypes for each individual by taking 2 kb upstream and 2 kb downstream the intron noncoding variant with the lowest \( P \) value \((15:12,573,054)\) (supplementary Additional file 1, table S6, Supplementary Material online). The total 4-kb region \((15:12,571,054–12,575,054)\) included 44 biallelic variants, 3 of which were significantly associated with foot feathering.

RepeatMasker track to check for presence of repetitive and transposable elements at the deletion breakpoints in the chicken reference genome and identified four long interspersed nuclear elements (supplementary Additional file 2, table S6, Supplementary Material online). Even though we observed two long interspersed nuclear elements very close to each other in a genome that is not particularly enriched for them, their location (1 kb upstream and 4 kb downstream the deletion breakpoint) suggests that is unlikely that the deletion is caused by transposable elements.

Foot-Feathered Birds Share a 17-kb Deletion Upstream H2AFY

We performed a copy-number variation (CNV) analysis to test whether a CNV event is associated with foot feathering. We identified a 17-kb deletion on chromosome 13, 9 kb upstream H2AFY between 16.08 and 16.10 Mb (fig. 3b). The deletion overlapped two 100-kb bins \((13:15,964,681–16,217,433)\) estimated to have an average recombination rate of 4 cM/Mb (supplementary Additional file 3, fig. S7a, Supplementary Material online).

The breakpoints of the deletion are both within the significant peak identified by the GWAS (fig. 3c). In addition to that, the deletion fully overlaps the 44-kb deletion reported in pigeon on scaffold 79 (supplementary Additional file 3, fig. S5, Supplementary Material online). Of the 19 foot-feathered chickens, 16 were homozygous for the deletion, whereas no deletion was observed for the Sundheimer, Marans, and German Faverolles breed (fig. 3a and supplementary Additional file 1, table S4, Supplementary Material online). Of the 150 controls, 148 lacked the deletion, whereas the Phoenix and Toutenkou breed were homozygous and heterozygous, respectively (fig. 3a and supplementary Additional file 1, table S4, Supplementary Material online). Despite that, all four wild species of Gallus did not have the deletion, therefore exhibiting a normal coverage distribution (fig. 3b and supplementary Additional file 1, table S4, Supplementary Material online). The polymerase chain reaction (PCR) validated our CNV analysis, confirming the presence/absence of a deletion in our samples (supplementary Additional file 4, Supplementary Material online). We used the UCSC RepeatMasker track to check for presence of repetitive and transposable elements at the deletion breakpoints in the chicken reference genome and identified four long interspersed nuclear elements (supplementary Additional file 2, table S6, Supplementary Material online). Even though we observed two long interspersed nuclear elements very close to each other in a genome that is not particularly enriched for them, their location (1 kb upstream and 4 kb downstream the deletion breakpoint) suggests that is unlikely that the deletion is caused by transposable elements.

Foot feathering is associated with two genomic regions. (a) Genome-wide manhattan plot. The \(-\log_{10}(P)\) for each variant is shown in the y-axis. Two clear signals can be observed on chromosomes 13 and 15, respectively. (b) Manhattan plot of chromosome 13 (16.0–16.2 Mb). (c) Manhattan plot of chromosome 15 (12.5–12.6 Mb). Significant variants associated with protein-coding genes and lncRNAs are highlighted in green. Intergenic variants are highlighted in light blue. The significant \( P \) value threshold \((P \text{ value} < 1.0e-25)\) is identified by the red dotted line.

12,600,000), which had an average \( Z_{H \rho} \) score of 3.41 (supplementary Additional file 3, fig. S4c, Supplementary Material online). The \( Z_{H \rho} \) score for the same region was above our threshold \((Z_{H \rho} > 3.0)\) in scaled samples.

Foot Feathering Has a Single Haplotype Origin

We also performed a CNV analysis on chromosome 15 but did not observe any CNV potentially associated with the phenotype. We therefore decided to reconstruct haplotypes for each individual by taking 2 kb upstream and 2 kb downstream the intron noncoding variant with the lowest \( P \) value \((15:12,573,054)\) (supplementary Additional file 1, table S6, Supplementary Material online). The total 4-kb region \((15:12,571,054–12,575,054)\) included 44 biallelic variants, 3 of which were significantly associated with foot feathering.
All 44 biallelic variants are located in the lncRNA ENSGALG00000052717. The 4-kb haplotype also overlapped two 100-kb bins (15:12,388,192–12,633,066) estimated to have an average recombination rate of 7 cM/Mb (supplementary Additional file 3, fig. S7b, Supplementary Material online).

The phylogenetic analysis of the haplotypes clearly separated scaled from feathered samples, indicating an identical origin of haplotypes for the foot-feathered samples (fig. 4). The clear separation between scaled and foot-feathered samples was further confirmed by the fixation index ($F_{st}$) analysis performed on the 44 biallelic variants found in the same 4-kb region. We reported the highest $F_{st}$ value for the noncoding variant used to reconstruct the haplotypes ($F_{st} = 0.97$) (supplementary Additional file 2, table S7, Supplementary Material online).

**Variants Associated with Foot Feathering Are in Highly Conserved Regions**

To understand the evolution of foot feathering, we looked for presence of conserved elements (CEs) in the 23 sauropsids multiple sequence alignment, which includes 15 birds, 3 crocodilians, 4 turtles, and anole lizard (supplementary Additional file 3, fig. S8, Supplementary Material online). On chromosome 13, 115 CEs were found within the GWAS peak. Of the 36 significant variants, two (one upstream gene and one intron variant) overlapped a CE (table 1) and both were associated with the $H2AFY$ gene. In the peak region of chromosome 15, we identified 275 CEs, although only the lncRNA intronic variant with the lowest $P$ value was found in a CE of considerable size (table 1).

**TBX5 Is Upregulated in the Hindlimb of Foot-Feathered Birds**

We generated high quality RNA-seq data from 21 chicken embryos sacrificed at Hamburger–Hamilton (HH) stages 35 and 39 to test whether our candidate genes and lncRNA are significantly differentially expressed in foot-feathered samples (supplementary Additional file 2, table S8, Supplementary Material online). Overall, more than 90% of the reads were uniquely mapped with an average size of $\sim$300 bp (supplementary Additional file 2, table S9, Supplementary Material online). As expected, clustering of samples based on read counts followed the embryonic HH stage (supplementary Additional file 3, fig. S10, Supplementary Material online).

$PITX1$ was significantly downregulated in the hindlimb of foot-feathered birds at HH35 ($q$-value: 1.79e-03), but not at HH39 ($q$-value: 0.38) (supplementary Additional file 3, fig. S11, Supplementary Material online). We observed a similar pattern in expression for $H2AFY$ at HH35 ($q$-value: 0.016) compared with HH39 ($q$-value: 0.88) (supplementary Additional file 3, fig. S12, Supplementary Material online). On the contrary, $TBX5$ was always significantly upregulated in foot-feathered birds at both embryonic stages (HH35 $q$-value: 2.49e-14; HH39 $q$-value: 6.87e-03) (fig. 5). At HH35, among the first ten most significant differentially expressed genes, we also identified $ZIC1$ ($q$-value: 2.42e-21), a transcription factor acting as scale-feather converter (Wu, Yan, et al. 2018). The FEELnc program classified our candidate lncRNA ENSGALG00000052717 as belonging to the set of mRNAs, since its coding potential was above the cutoff estimated by a 10-fold cross validation procedure that maximizes
both sensitivity and specificity (Wucher et al. 2017). We think that the low lncRNA read counts observed in our samples may have affected the FEELnc classification.

Discussion
Foot Feathering Has a Parallel Genetic Origin
Researchers have repeatedly questioned the genetic basis of parallel evolution. Despite this long-lasting interest, for many traits that evolved in parallel little is known whether these are mirrored in underlying genes or mutations. Foot feathering is an interesting case since, although it is a very recognizable trait that can be very easily selected to fixation in breeds, it is in fact not a monogenic trait.

The molecular basis of foot feathering has so far only been studied in detail in domestic pigeon (Domyan et al. 2016; Boer et al. 2019). Chicken and pigeon diverged more than 89 Ma (Jarvis et al. 2014) and are currently classified as belonging to two separate subgroups within the Neognathae clade, the Galloanserae and Neoaves, respectively (Jarvis et al. 2014; Brusatte et al. 2015). Since domestication, both species have experienced selection for a variety of traits that are remarkably different or absent in the wild ancestor (Tixier-Boichard et al. 2011; Domyan and Shapiro 2017). Foot feathering has been under artificial selection since ancient times and
nowadays extensive variation can be observed among breeds (i.e., groused feet, slippered legs, muffed legs, and vulture hocks). As for many other traits, ptilopody has been artificially selected to fixation and has become a breed characteristic in both species (Bartels 2003). For parallel evolution to occur, loci associated with similar regulatory pathways and likely to generate variation should be targeted by selection. Our findings and those of Domyan et al. (2016) corroborate this hypothesis, as the independent evolution of foot feathering in chicken and pigeon involves a similar genetic basis and set of genes that not only generate outstanding variation, but this variation is also targeted for recurring artificial selection. In this study, we showed that artificial selection has left clear signatures of positive selection in the genome of foot-feathered birds, as indicated by the negative \( Z_{	ext{H}} \) scores.

These results also illustrate how combining genome-wide association studies and signature of selection analyses forward our understanding of the genomic basis of traits, providing support for the role of, in this case, chromosomes 13 and 15 in foot feather development. Similar signatures of selection were also identified in pigeon by Domyan et al. (2016). However, compared with chicken, pigeons homozygous for the deletion on scaffold 79 showed elevated levels of haplotype homozygosity relative to scaled birds, whereas positive selection was only observed among heavily feathered birds (i.e., muff phenotype) on scaffold 70 (Domyan et al. 2016).

Interestingly, foot feathering is also observed in avian wild species, including snowy owl, golden eagle, and rock ptarmigan (Bartels 2003; Boer et al. 2017). Even though in raptor and boreal species ptilopody has entirely evolved by natural selection, the occurrence of the phenotype suggests that the same underlying genes and mutations can evolve in different species under different types of selection and selection pressure. However, studies on both wild and domesticated avian species are required to further validate this hypothesis.

Foot Feathering Is Associated with a Single, Identical Haplotype in Chicken

As we showed, foot feathering has evolved independently in chicken and pigeon as a result of human-driven selection and this selection pressure has resulted in similar causal mutations. In chicken, foot-feathered birds were also found to share an identical 4-kb haplotype on chromosome 15 independently on whether the individual has the 17-kb deletion on chromosome 13. Sharing of an identical haplotype was also reported in pigeon (i.e., scaffold 70), though a clear

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**Table 1. Variants Associated with Foot Feathering Are Highly Conserved.**

| Chr | CE Start | CE End | Size | Strand | Variant | Maj/Min Allele | \( P \) Value |
|-----|----------|--------|------|--------|---------|----------------|-------------|
| 13  | 16,112,206 | 16,112,215 | 9    | +      | 16,112,207 | C/T           | 1.822e-29   |
| 13  | 16,128,762 | 16,128,774 | 12   | +      | 16,128,768 | C/A           | 5.395e-30   |
| 15  | 12,572,741 | 12,573,218 | 477  | +      | 12,573,054 | C/T           | 1.144e-43   |

**Note:** The allele associated with the phenotype is underlined. Chr, chromosome; CE, conserved element; Maj, major allele; Min, minor allele. The \( P \) value is that of the genome-wide association analysis.
clustering was only observed among heavily feathered birds (Domyan et al. 2016). The presence of a single, identical underlying haplotype suggests that foot feathering is caused by mutations that occurred only once in the domestication history of chicken. These causal mutations have then been selected in multiple breeds, in many cases by deliberately crossing foot-feathered birds with scaled birds of a different breed. Because of repeated crossing, the causal mutations underlying pitlopody have been recycled many times since domestication and because the genetic basis is strikingly the same among breeds, the underlying genes can easily be detected by an across-breed GWAS. The mutations found in the 4-kb haplotype, which are clearly related to domesticated populations, are likely to have first appeared in Asia and have later been introgressed into Europe through human migration.

Haplotype length can provide important information on the age of the haplotype. This means that longer haplotypes are of more recent origin (“younger”), as recombination events did not break them down into smaller tracks over time. A negative correlation is, therefore, expected between haplotype length and recombination. The relative small size of the haplotype reported on chromosome 15 supports our conclusions on the single, and likely old, occurrence of the mutations underlying pitlopody on chromosome 15. In fact, it is likely that repeated crossing has not only allowed the spreading of the causal mutation but also contributed to breaking down the original haplotype at each generation, which, by means of artificial selection, is now fixed in all foot-feathered birds considered in this study. The intact haplotype length is also explained by the local recombination rate reported in bins of 100 kb on chromosome 15, if we consider that recombination rate in microchromosomes (50–100 kb/cM) is nearly three times higher than that of macrochromosomes (~300 kb/cM) (Megens et al. 2009). The limited number of SNPs found in the 4-kb haplotype makes, however, the estimation of the substitution model and mutation rate required to infer the haplotype age a challenging task. A possible solution would be to analyze the same candidate region in ancient samples to better estimate the age of the haplotype.

The long-noncoding variant (15:12,573,054) used for the haplotype analysis is found upstream TBX5, a gene encoding a key transcriptional regulator of forelimb identity and development (Logan et al. 1998; Logan and Tabin 1999; Rodriguez-Esteban et al. 1999; Takeuchi et al. 1999). Interestingly, the same mutation was associated with foot feathering in a parallel study in chicken by Li et al. (2020). In chicken embryos, TBX5 is normally expressed in the forelimb, but its misexpression in the hindlimb at early embryonic stages can induce a partial wing-like transformation, including the formation and development of feathers on the feet (Takeuchi et al. 1999). Similarly to pigeon, the absence of fixed nonsynonymous coding mutations in TBX5 confirms the role of expression changes in the determination of feathered versus scaled feet in chicken as well. We could observe misexpression in the hindlimb of feathered embryos at both HH35 and HH39, meaning that misexpression starts at a very early embryonic stage (in the study of Domyan, similar expression changes were observed at HH25) and is maintained almost up to the end of the embryonic development. Among our most significantly upregulated genes in the hindlimb at HH35, we also found the novel scale-feather converter ZIC1, a transcription factor whose overexpression in feather forming regions (i.e., wings and tail feathers) is sufficient to initiate the invagination step required to form the follicle, but not to form mature follicles (Wu, Yan, et al. 2018).

The 17-kb Deletion on Chromosome 13 Likely Acts as Qualitative Molecular Driver

Even though ectopic expression of TBX5 is associated with foot feathering in chicken, extensive variation in feather type and distribution is often observed. In our study, the 17-kb deletion on chromosome 13 was homozygous in 16 birds, whereas absent in three birds (supplementary Additional file 1, table S4, Supplementary Material online). The absence of the deletion suggests that this locus may affect the qualitative variation in epidermal appendages. This means that the deletion is important for an individual to display variation in the type and extent of feathers, such as enlarged feathers on the feet or wing-like feathers on the feet and toes, but is not essential to determine the localized development of feathers on the feet, which seems the function of TBX5. Therefore, the qualitative role of the 17-kb deletion reasonably explains the discrepancies observed in the Marans, Sundheimer, and German Faverolles breed.

Strikingly, in pigeon a similar deletion, 44 kb in size, in the exact same region as the 17-kb deletion in chicken, is present (Domyan et al. 2016). The 7-bp microhomology we identified at the deletion breakpoints in chicken indicates that this structural variant (SV) has emerged in both species multiple times independently. However, contrary to chicken, based on their QTL and WGS analyses Domyan et al. (2016) concluded that the deletion is sufficient for the development of small feathers (grouse phenotype), whereas the development of large feathers (muff phenotype) is mostly driven by the TBX5 locus.

In chicken, the 17-kb deletion is ~9 kb upstream H2AFY and ~200 kb upstream PITX1, a gene encoding a key transcriptional regulator of hindlimb identity and development. PITX1 is normally expressed in the hindlimb, but not in the forelimb, as an abnormal expression in the forelimb blocks feather development (Logan and Tabin 1999). Interestingly, in pigeon the peak on scaffold 79 is also ~200 kb upstream PITX1. Compared with H2AFY, which was downregulated at HH35 and upregulated at HH39, PITX1 was always downregulated. The key role of PITX1 in limb-type morphology determination has been demonstrated across multiple species (Logan and Tabin 1999; Takeuchi et al. 1999; Delaurier et al. 2006; Ouimette et al. 2010; Kragesteen et al. 2018) and in all species investigated the relationship between PITX1 and H2AFY is maintained. In pigeon, the 44-kb deletion spans an element orthologous to a known human limb enhancer, hs1473, which shows a strong limb-specific activity (Spielmann et al. 2012; Domyan et al. 2016). As limb patterning and morphogenesis are regulated by highly conserved
networks (Boer et al. 2019), it is reasonable to assume that, as in pigeon, also in chicken the deletion encompasses this enhancer, causing loss of PITX1 expression, thus resulting in a partial leg-to-wing homeotic transformation. However, further molecular analysis, including ChiP-seq data, is required to formally confirm this conclusion.

**Conclusions**

Foot feathering is an interesting example of a polygenic trait that has evolved by parallel evolution as its parallel evolution is mirrored in almost every detail at the molecular and, most likely, developmental level. In this study, we showed that, although chicken and pigeon diverged more than 89 Ma, in both avian species the exact same number of loci containing the exact same set of genes are involved. This similarity is even more striking as a similar deletion at one of the loci has the same outcome in regulating gene expression.

Even though genetic variants arose independently millions of years after the species divergence, it is remarkable to see that not only are the exact same genes involved, but they are affected in very similar ways, despite the many ways in which a similar phenotype conceivably could have arisen. Therefore, even under different types of selection and selection pressure, the same genes and causal mutations underlying major phenotypic changes can evolve in different lineages. Our findings provide support for the hypothesis that only a limited number of evolutionary trajectories at the molecular level are open to generate a specific outcome if developmental pathways are sufficiently constrained.

**Materials and Methods**

**Blood Collection and Animal Experiments**

Collections of blood samples was done in accordance with the German Animal Protection Law and was approved by the Committee of Animal Welfare at the Institute of Farm Animal Genetics (Friedrich-Loeffler-Institut) and the Lower Saxony State Office for Consumer Protection and Food Safety (No. 33.9-42502-05-10A064). Sample collection and data recording were also conducted strictly according to the Dutch law on animal protection and welfare (Gezondheids- en welzijnszorg voor dieren).

**Samples and Phenotype**

DNA of 169 samples from 87 traditional chicken breeds and one individual from each of the four living wild species of Gallus (i.e., G. gallus, G. sonneratii, G. lafayetii, and G. varius) was used for WGS on an Illumina HiSeq 3000 (supplementary Additional file 1, table S1, Supplementary Material online). WGS data of the 97 birds sampled in the Netherlands were previously deposited in the European Nucleotide Archive under accession number PRJEB34245 (Bortoluzzi et al. 2020). WGS data of the remaining 68 birds sampled in Germany have been deposited in ENA under accession number PRJEB36674. Detailed information on the sequenced reads can be found in supplementary Additional file 1, table S2, Supplementary Material online. Information on foot feathering was collected during sampling and confirmed by the breeding associations. The phenotype was observed in 11 breeds, for a total of 19 samples. Of these, nine showed short and tight feathers on the metatarsus and digits, whereas ten samples had extensive feather development, as well as long flight-like feathers on the posterior toes (fig. 1; supplementary Additional file 2, table S1, Supplementary Material online). The remaining 150 samples, which have scaled epidermis, were used as control.

**Whole-Genome Sequencing**

Library preparation and sequencing were performed at the Institut national de la recherche agronomique, France, following their established protocols. Reads were mapped with the Burrows–Wheeler alignment (BWA-MEM) algorithm v0.7.17 (Li and Durbin 2009) to the chicken GRCg6a reference genome (GenBank accession: GCA_000003155.5) with default settings. Duplicate reads were removed with the markdup option in Sambamba v0.6.3 (Tarasov et al. 2015). Sites with mapping quality <30 and base quality <20 were discarded from further analyses (supplementary text, Supplementary Material online).
Phasing, Imputation, and Annotation
Genotypes were imputed and phased into haplotypes with Beagle v4.0 (Browning and Browning 2013) by considering in each of the ten independent cycles 20,000 markers in each sliding window, allowing 1,000 markers to overlap between sliding windows. Imputation accuracy was estimated by masking 10% of the known sites (supplementary text, Supplementary Material online). Imputed variants were annotated to the Ensembl’s G. gallus annotation database using the Ensembl Variant Effect Predictor (release 95) tool (McLaren et al. 2016) (supplementary text, Supplementary Material online).

Population Stratification
Genetic distances were analyzed with a PCA and a NJ tree, both based on a subset of phased variants filtered for a minor allele frequency <0.05. Filtering and PCA were performed in PLINK v1.9 (Purcell et al. 2007). The phylogenetic tree was generated in PHYLPAR v3.696 (Felsenstein 2005) from the distance relationship matrix estimated in PLINK.

Association Study and Annotation of Genes
We performed a standard case/control association analysis using the Fisher’s exact test to generate uncorrected and corrected P values, subsequently applying an adaptive Monte Carlo permutation test with 5,000 replications. Variants with a P value <1.0e-25 were considered to be significantly associated with the phenotype. Manhattan plots were generated using the qqman library in R v3.2.0 (R Core Team 2013; Turner 2014). Genes in genomic regions showing significant association with the phenotype were identified using the Ensembl Genes 95 Database in BioMart (Kinsella et al. 2011). Chicken QTLs were downloaded from the Animal QTL Database (Hu et al. 2013). Genomic coordinates were converted to the GRCh38a assembly in LiftOver (Rhead et al. 2009).

Signatures of Selection
Screening for signatures of selection was performed on the control (n = 150) and case (n = 19) group, separately, and only on the autosomes that showed a significant association with the phenotype. For each pool and identified SNP, we determined the number of reads corresponding to the most (n\textsubscript{MAJ}) and least abundant allele (n\textsubscript{MIN}). The pooled heterozygosity (H\textsubscript{p}) was calculated in sliding 40-kb windows following Rubin et al. (2010):

\[
H_p = 2 \frac{n_{\text{MAJ}} n_{\text{MIN}}}{(n_{\text{MAJ}} + n_{\text{MIN}})^2}.
\]

To resemble a normal distribution, H\textsubscript{p} values were normalized into ZH\textsubscript{p} scores as ZH\textsubscript{p} = (H\textsubscript{p}−μH\textsubscript{p})/σH\textsubscript{p}. Windows with at least 300 SNPs and a ZH\textsubscript{p} ≤ −3 were retained, as windows below this threshold represent the extreme lower end of the distribution (supplementary Additional file 3, fig. S3, Supplementary Material online).

Recombination Rate
We used the linkage map of Elferink et al. (2010) to estimate the recombination rate, expressed as the genetic length in centimorgans (cM) divided by the physical genomic distance in mega base pairs. Recombination rate was calculated in bins of ~100 kb after converting the genomic positions of all SNPs to the GRCh38a genome assembly.

Structural Variants
SVs calling and genotyping were performed using Smoove (https://github.com/brentp/smoove, last accessed November 11, 2019). Smoove makes use of various existing tools to call SVs and improves specificity by removing noise from spurious alignment signals. First, discordantly mapped and split reads were extracted from the alignment by Samblaster (Faust and Hall 2014). Next, Lumpy software (Layer et al. 2014) was used to call SVs, and genotyping was performed by SVtyper (Chiang et al. 2015). To further filter SV calls, Mosdepth (Pedersen and Quinlan 2018) was used to discard reads from regions where the sequence depth of split or discordant reads was >1,000 to remove regions that contribute to spurious calls. Duphold (Pedersen and Quinlan 2019) was subsequently used to annotate depth changes within and on the breakpoints of SVs.

PCR-Based Screens for Genomic Rearrangement
A set of four PCR primers was designed to amplify two bands around the deletion breakpoints and two bands over the deletion. More information on the primers and PCR protocol are reported in supplementary Additional file 4, Supplementary Material online. Gel image with the presence (control, scale-footed samples) or absence (case, feather-footed samples) of PCR product is reported in supplementary Additional file 4, Supplementary Material online.

Phylogenetic Analysis of Haplotypes
For each sample, we extracted and considered the two alternative haplotypes as separate haplotypes, so that haplotypes belonging to the same individual did not necessarily cluster together. No missing alleles were present in the phased haplotypes since missing sites were imputed with Beagle. Haplotypes were reconstructed considering only biallelic sites. We then constructed a NJ tree based on the distance matrix estimated in PLINK from all haplotypes.

DNA Sequence Conservation
CEs were predicted from the 23 sauropsids multiple whole-genome alignment generated by Green et al. (2014) (supplementary text, Supplementary Material online). CEs were predicted using PhastCons (Siepel et al. 2005) from a neutral evolutionary model estimated from 114,709 four-fold degenerate (4D) sites in Phylofix (Siepel and Haussler 2004; Green et al. 2014). After filtering for assembly gaps, a total of 1.14 million CEs covering 73 Mb of the chicken genome were retained (supplementary text, Supplementary Material online).
Tissue Collection, RNA Isolation, and RNA Sequencing
Forelimb and hindlimb buds were harvested from 21 chicken embryos sacrificed at HH stage 35 (n = 11) (Hamburger and Hamilton 1951) and HH39 (n = 10) (supplementary Additional file 3, fig. S9, Supplementary Material online). Detailed information on the breeds and samples can be found in supplementary Additional file 2, table S8, Supplementary Material online. Sequencing was performed at BGI, China, following the manufacturer’s protocol.

RNA-seq Analysis
Clean reads were mapped to the chicken GRCg6a reference genome using STAR v2.4.0 (Dobin et al. 2013) with the chicken reference genome and its annotation file as guide, both downloaded from Ensembl (release 95). Quality of mapped RNA-seq data was assessed using Deeptools v3.3.1 (Ramirez et al. 2014). RSEM was used to quantify expression of RNA transcripts and genes (Li and Dewey 2011), whereas StringTie v2.0.3 (Pertea et al. 2015) was used for gene model using the Ensembl gene annotation file as reference. Transcripts of all samples were afterward combined using the merge option in StringTie and used as input file in the FLEXible Extraction of Long noncoding RNA (FEELnc) program (Wucher et al. 2017) to predict and annotate IncRNAs (SI Text).

We used DESeq2 (Love et al. 2014) to test whether the genes/IncRNAs identified by the genome-wide association analysis were differentially expressed in foot-feathered birds (case) compared with scaled birds (control). The differential expression analysis was performed for each embryonic stage and for the forelimb (F) and hindlimb (H), separately, considering only genes with at least 20 reads. Protein-coding genes and IncRNAs were considered to be significantly differentially expressed only if their adjusted P value was <0.05 (Benjamini–Hochberg adjustment).

Supplementary Material
Supplementary data are available at Molecular Biology and Evolution online.

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Author Contributions
C.B., M.B., and H.-J.M. conceived the study. C.B. performed the analysis. M.F.L.D. performed the structural variant analysis. B.D., K.L., and S.W. performed the PCR. S.W. performed the blood collection of the German samples. C.B., B.D., K.L., and R.P.M.A.C. performed the sampling of eggs used for the extraction of limb buds. M.B., H.-J.M., M.A.M.G., and R.P.M.A.C. supervised the project. C.B., M.B., and H.-J.M. wrote the manuscript. All authors read and contributed to the manuscript. All authors agreed on the final manuscript.

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