COMPUTATIONAL ANALYSIS OF SIGNATURES OF
SELECTION IN GENES FOR EGG PRODUCTION AND
GROWTH IN POULTRY

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Computational Analysis of Signatures of Selection in Genes for Egg Production and Growth in Poultry

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A thesis submitted in partial fulfillment for the Degree of Master of Science in Bioinformatics and Molecular Biology in the Jomo Kenyatta University of Agriculture and Technology

2017
DECLARATION

This thesis is my original work and has not been presented for degree in any other institution.

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This thesis has been submitted for examination with our approval as university supervisors.

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DEDICATION

Special dedication to my parents Dorcas Wainaina and Stephen Wainaina. To my sister Joan, brothers Charles, Philip, James, Joram, and to my niece Julie.
ACKNOWLEDGEMENTS

I would like to take this opportunity to thank God for giving me the strength during the entire difficult period of this study. I am also extremely grateful to my first supervisor Dr. Sheila Ommeh from the Institute for Biotechnology Research, JKUAT for her guidance, patience and her immense sacrifice during the project. My other supervisors, Professor Kariuki and Dr. Lichoti, I am also grateful for your guidance and positive criticism.

I am also thankful to the Indigenous Chicken Consortium for giving me an opportunity to be part of their research and also the RPE for the financial assistance which enabled the smooth running of the project.

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# TABLE OF CONTENTS

DECLARATION...................................................................................................................... ii
DEDICATION........................................................................................................................ iii
ACKNOWLEDGEMENTS......................................................................................................... iv
TABLE OF CONTENTS.......................................................................................................... v
LIST OF TABLES .................................................................................................................... x
LIST OF FIGURES ................................................................................................................ xi
LIST OF APPENDICES ......................................................................................................... xiii
LIST OF ABBREVIATIONS AND ACRONYMS ................................................................... xiv
ABSTRACT ............................................................................................................................ xvi
CHAPTER ONE ..................................................................................................................... 1

## INTRODUCTION

1.1 Background of the study ................................................................................................. 1
1.2 Chickens for egg and meat production .......................................................................... 2
1.3 Statement of the problem ............................................................................................... 7
1.4 Justification ..................................................................................................................... 8
1.5 Research hypothesis ...................................................................................................... 9
1.6 Objectives ..................................................................................................................... 9
   1.6.1 General objective .................................................................................................. 9
   1.6.2 Specific objectives ............................................................................................... 9

CHAPTER TWO .................................................................................................................. 10

## LITERATURE REVIEW

2.1 Candidate genes and their functions ............................................................................ 10
2.1.1 Prolactin ........................................................................................................10
2.1.2 Vasoactive intestinal peptide 1 and vasoactive intestinal peptide receptor 1 ....12
2.1.3 Growth hormone and growth hormone receptor ..............................................13
2.1.4 Insulin-like growth factor 1 and insulin-like growth factor 1 receptor ..........15

2.2 Gene signatures of selection ..............................................................................18
2.2.1 Long haplotype ................................................................................................20
2.2.2 Population differentiation ..............................................................................20
2.2.3 Reduction in genetic diversity ........................................................................21
2.2.4 High frequency derived alleles .....................................................................21
2.2.5 High proportion of function altering mutation ..............................................22

2.3 Lineage specific positive selection .................................................................23

2.4 Site specific positive selection .........................................................................24

2.5 Protein 3D structure prediction ........................................................................25

2.6 Protein structure alignment ................................................................................27

CHAPTER THREE ........................................................................................................29

MATERIALS AND METHODS ..................................................................................29

3.1 Retrieval of candidate gene sequences ..............................................................29
3.2 Search for homologs and retrieval .....................................................................29
3.3 Multiple sequence alignment ............................................................................30
3.4 Phylogeny construction ....................................................................................30
3.5 Substitution model selection and rate heterogeneity testing ................................31
3.6 Input preparation in PAML ................................................................................31
3.7 Test for molecular signatures of selection .......................................................32
3.8 Protein 3D structure prediction .................................................................32
3.9 Comparative 3D structure analysis ...............................................................33

CHAPTER FOUR ..................................................................................................34

RESULTS .........................................................................................................34

4.1 Homology search ..........................................................................................34
4.2 Analysis of signatures of selection at candidate genes for egg production in poultry
.................................................................................................................................34
  4.2.1 Multiple sequence alignment of prolactin................................................34
  4.2.2 Multiple sequence alignment of vasoactive intestinal peptide 1 (VIP1)........35
  4.2.3 Multiple sequence alignment of vasoactive intestinal peptide receptor 1 (VIPR1)
.................................................................................................................................37
  4.2.4 Lineage selection in vasoactive intestinal peptide receptor 1 (VIPR1) ........38
  4.2.5 Lineage selection in prolactin gene ..........................................................40
4.3 Analysis of signatures of selection at candidate genes for growth in poultry ......41
  4.3.1 Multiple sequence alignment of growth hormone (GH) .............................41
  4.3.2 Multiple sequence alignment of growth hormone receptor (GHR) ..........43
  4.3.3 Multiple sequence alignment of insulin-like growth factor 1 (IGF1) gene ....45
  4.3.4 Multiple sequence alignment of insulin-like growth factor 1 receptor (IGF1R)
gene .................................................................................................................................46
  4.3.5 Lineage selection in growth hormone (GH) gene .....................................49
  4.3.6 Lineage selection in insulin-like growth factor 1 receptor (IGF1R) gene ......50
  4.3.7 Lineage selection in growth hormone receptor (GHR) gene ....................52
  4.3.8 Amino acid site selection ...........................................................................54
4.4 Protein 3D structure prediction for genes implicated in egg production ...............57
  4.4.1 Predicted 3D structures of prolactin protein .............................................57
  4.4.2 Predicted 3D structures vasoactive intestinal peptide 1 protein ..................58
  4.4.3 Predicted 3D structures of vasoactive intestinal peptide receptor 1 protein ......58
4.5 Protein 3D structure prediction for genes implicated in growth .......................59
  4.5.1 Predicted 3D structures of growth hormone protein ....................................59
  4.5.2 Predicted 3D structures of growth hormone receptor protein .........................60
  4.5.3 Predicted 3D structures of insulin-like growth factor 1 protein .....................61
  4.5.4 Predicted 3D structures of insulin-like growth factor 1 receptor protein ..........62
4.6 Predicted 3D structure comparison of proteins implicated in egg production .......64
  4.6.1 Comparison of predicted 3D prolactin structures ........................................64
  4.6.2 Comparison of predicted 3D vasoactive intestinal peptide 1 structures ...........65
  4.6.3 Comparison of predicted 3D vasoactive intestinal peptide receptor 1 structures
  .............................................................................................................................66
4.7 Predicted 3D structure comparison of proteins implicated in growth ..................67
  4.7.1 Comparison of predicted 3D growth hormone structures .................................67
  4.7.2 Comparison of predicted 3D growth hormone receptor structures ..................68
  4.7.3 Comparison of predicted 3D insulin-like growth factor 1 structures ................69
  4.7.4 Comparison of predicted 3D insulin-like growth factor 1 receptor structures ....70

CHAPTER FIVE .................................................................................................................72

DISCUSSION ...................................................................................................................72

5.1 Analysis of signatures of selection in genes for egg production .......................72
  5.1.1 Prolactin ............................................................................................................72
5.1.2 Vasoactive intestinal peptide receptor 1 ........................................73
5.2 Analysis of signatures of selection in genes for growth ..............................74
  5.2.1 Growth hormone ...........................................................................74
  5.2.2 Growth hormone receptor .................................................................76
  5.2.3 Insulin-like growth factor 1 receptor ................................................77
5.3 Prediction of secondary and tertiary structures for egg production proteins ......79
  5.3.1 Prolactin ......................................................................................79
  5.3.2 Vasoactive intestinal peptide 1 and vasoactive intestinal peptide receptor 1 ....81
5.4 Prediction of secondary and tertiary structures for growth proteins .................82
  5.4.1 Growth hormone and growth hormone receptor ...................................82
  5.4.2 Insulin-like growth factor 1 and Insulin-like growth factor 1 receptor ........83
5.5 Predicted 3D structure comparison of prolactin, vasoactive intestinal peptide
  receptor 1, growth hormone, growth hormone receptor, and insulin-like growth factor 1
  proteins .................................................................................................85
5.6 Predicted 3D structure comparison of vasoactive intestinal peptide 1 and insulin-like
  growth factor 1 receptor proteins ..............................................................88

CHAPTER SIX ..........................................................................................90
CONCLUSION AND RECOMMENDATIONS .............................................90
  6.1 Conclusion .......................................................................................90
  6.2 Recommendations ...........................................................................93
REFERENCES .........................................................................................94
APPENDICES .........................................................................................110
LIST OF TABLES

**Table 1.1:** Index ranking of traits perceived by farmers, marketers and consumers to be of economic importance..........................................................5

**Table 4.1:** Likelihood ratio tests (LRT) to detect selection in lineages and amino acid sites.................................................................................................48

**Table 4.2:** Predicted 3D structures of prolactin protein. ........................................................................57

**Table 4.3:** Predicted protein 3D structures of vasoactive intestinal peptide 1. ...............58

**Table 4.4:** Predicted protein 3D structures of vasoactive intestinal peptide receptor 1. 59

**Table 4.5:** Predicted protein 3D structures of growth hormone.............................................60

**Table 4.6:** Predicted protein 3D structure of growth hormone receptor. .......................61

**Table 4.7:** Predicted protein 3D structure of insulin-like growth factor 1. ....................62

**Table 4.8:** Predicted protein 3D structure of insulin-like growth factor 1 receptor......63
LIST OF FIGURES

Figure 4.1: Multiple sequence alignment of prolactin protein. .............................................35

Figure 4.2: Multiple sequence alignment of vasoactive intestinal peptide 1 (VIP1) protein. .........................................................36

Figure 4.3: Multiple sequence alignment of vasoactive intestinal peptide receptor 1 (VIPR1) protein. ..............................................................38

Figure 4.4: A phylogram showing evolution of vasoactive intestinal peptide receptor 1 (VIPR1) gene. .................................................................39

Figure 4.5: Phylogeny showing evolution of prolactin (PRL) gene. ...........................................41

Figure 4.6: Multiple sequence alignment of growth hormone (GH) protein. ............................43

Figure 4.7: Multiple sequence alignment of growth hormone receptor (GHR) protein. ...............44

Figure 4.8: Multiple sequence alignment of insulin-like growth factor 1 (IGF1) protein. .................................................................46

Figure 4.9: Multiple sequence alignment of insulin-like growth factor 1 receptor (IGF1R) protein. ............................................................................47

Figure 4.10: Phylogenetic tree showing evolution of growth hormone (GH) gene.....................50

Figure 4.11: Phylogenetic tree of the insulin-like growth factor 1 receptor (IGF1R) gene. ..........52

Figure 4.12: Phylogenetic tree showing evolution of growth hormone receptor (GHR) gene. .................................54

Figure 4.13: Multiple sequence alignment of insulin-like growth factor 1 receptor (IGF1R) with site under positive selection. .................................................55
Figure 4.14: The 3D structure of insulin-like growth factor 1 receptor showing the different domains. .................................................................56

Figure 4.15: Predicted 3D structure comparison of peafowl, common pheasant, common quail, helmeted guinea fowl, Japanese quail, knob-billed, ostrich, wild chicken, and wild duck prolactin...............................................................65

Figure 4.16: Predicted 3D structure comparison of wild chicken 1 and wild chicken 2 vasoactive intestinal peptide 1.................................................................66

Figure 4.17: Predicted 3D structure comparison of Japanese quail and wild chicken vasoactive intestinal peptide receptor 1.................................................................67

Figure 4.18: Predicted 3D structure comparison of common quail, greylag goose, wild chicken, and wild duck growth hormone..................................................................68

Figure 4.19: Predicted 3D structure comparison of greylag goose, Southern ostrich, wild chicken, and wild duck growth hormone receptor........................................69

Figure 4.20: Predicted 3D structure comparison of Japanese quail, southern ostrich, wild chicken, and wild duck insulin-like growth factor 1. ..........................70

Figure 4.21: Predicted 3D structure comparison of Japanese quail, southern ostrich, wild chicken, and wild duck insulin-like growth factor 1 receptor. ...........71
LIST OF APPENDICES

Appendix 1: Homologues for prolactin gene...........................................110

Appendix 2: Homologues for vasoactive intestinal peptide 1 (VIP1) gene........112

Appendix 3: Homologues for vasoactive intestinal peptide receptor 1 (VIPR1) gene.................................................................113

Appendix 4: Homologues for growth hormone (GH) gene..........................115

Appendix 5: Homologues for growth hormone receptor (GHR) gene..........116

Appendix 6: Homologues for insulin-like growth factor 1 (IGF1) gene.........118

Appendix 7: Homologues for insulin-like growth factor 1 receptor (IGF1R) gene..................................................................................121

Appendix 8: Predicted protein 3D structures for prolactin............................123

Appendix 9: Predicted protein 3D structures for vasoactive intestinal peptide 1 (VIP1)...........................................................................124

Appendix 10: Predicted protein 3D structures for vasoactive intestinal peptide receptor 1 (VIPR1)..............................................................125

Appendix 11: Predicted protein 3D structures for growth hormone (GH)........126

Appendix 12: Predicted protein 3D structures for Homologues for growth hormone receptor (GHR)..........................................................127

Appendix 13: Predicted protein 3D structures for insulin-like growth factor 1 (IGF1).................................................................................128

Appendix 14: Predicted protein 3D structures for insulin-like growth factor 1 receptor (IGF1R).................................................................129
LIST OF ABBREVIATIONS AND ACRONYMS

3D  Three Dimensions
BLAST  Basic Local Alignment Search Tool
BLOSUM62  Blocks Substitution Matrix 62
bp  base pair
cAMP  Cyclic Adenosine Mono Phosphate
CRFs  Conditional Random Fields
Cu-Zn SoD  Copper-Zinc superoxide dismutase
DAG  Di-acyl glycerol
dN/dS  Ratio of rate of non-synonymous mutations to synonymous mutations
DNA  Deoxyribonucleic Acid
E Value  Expectation Value
Fe SoD  Iron Superoxide dismutase
GDP  Gross Domestic Product
GH  Growth hormone
GHR  Growth Hormone Receptor
IGF1  Insulin-like growth factor 1
IGF1R  Insulin-like growth factor 1 receptor
IRS1  Insulin Receptor Substrate 1
kDa  Kilo daltons
LD  Linkage disequilibrium
LRH  Long Range Haplotypes
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
</tr>
</thead>
<tbody>
<tr>
<td>LRT</td>
<td>Likelihood Ratio Test</td>
</tr>
<tr>
<td>MAFFT</td>
<td>Multiple Alignment using Fast Fourier Transform</td>
</tr>
<tr>
<td>MAPK</td>
<td>Mitogen-Activated Protein Kinase</td>
</tr>
<tr>
<td>MnSoD</td>
<td>Manganese Superoxide dismutase</td>
</tr>
<tr>
<td>mRNA</td>
<td>messenger Ribonucleic acid</td>
</tr>
<tr>
<td>mRNA</td>
<td>messenger Ribonucleic Acid</td>
</tr>
<tr>
<td>MSA</td>
<td>Multiple Sequence Alignment</td>
</tr>
<tr>
<td>MTT</td>
<td>Multiple Template Threading</td>
</tr>
<tr>
<td>MUSCLE</td>
<td>Multiple Sequence Comparison by Log Expectation</td>
</tr>
<tr>
<td>nr</td>
<td>non redundant</td>
</tr>
<tr>
<td>PAML</td>
<td>Phylogenetic Analysis using Maximum Likelihood</td>
</tr>
<tr>
<td>PDB</td>
<td>Protein Data Bank</td>
</tr>
<tr>
<td>PKC</td>
<td>Protein Kinase C</td>
</tr>
<tr>
<td>PRANK</td>
<td>Probabilistic Alignment Kit</td>
</tr>
<tr>
<td>PRL</td>
<td>Prolactin</td>
</tr>
<tr>
<td>SNP</td>
<td>Single Nucleotide Polymorphism</td>
</tr>
<tr>
<td>STAT</td>
<td>Signal Transducers and Activators of Transcription</td>
</tr>
<tr>
<td>VIP1</td>
<td>Vasoactive Intestinal Peptide 1</td>
</tr>
<tr>
<td>VIPR1</td>
<td>Vasoactive Intestinal Peptide Receptor 1</td>
</tr>
</tbody>
</table>
ABSTRACT

Poultry are highly variable phenotypically as a result of natural selection and domestication. This leaves behind signatures of selection which may be used for genetic improvement of poultry through selective breeding. The aim of this study was to perform analysis of signatures of selection at candidate genes for egg production and growth. Genes selected for egg production were prolactin, vasoactive intestinal peptide 1 and vasoactive intestinal peptide receptor 1. Genes selected for growth were growth hormone, growth hormone receptor, insulin-like growth factor 1, and insulin-like growth factor 1 receptor. A reciprocal BLASTp using BLOSUM62 substitution matrix was performed to identify the homologs. Orthologs with an expectation value greater than 1e-10 were selected for further analysis. Thereafter, multiple sequence alignment was performed using MUSCLE which is based on an iterative algorithm. Phylogeny construction was then done using Distance-based FastME followed by analysis using codon-based models in PAML. Likelihood ratio test was used to detect positive selection followed by posterior probability using Bayes Empirical Bayes Analysis to identify the sites under selection. Web-based servers: Raptor X and DeepAlign were used to predict 3D structures and compare the structures, respectively. This led to identification of purifying selection in all lineages in vasoactive intestinal peptide receptor 1, while in prolactin, there was positive selection in poultry. Additionally, purifying selection was detected in poultry lineages for Growth Hormone and Growth Hormone Receptor genes. Insulin-like growth factor 1 receptor (IGF1R) had positive selection on amino acid Isoleucine at position 460 on Receptor L domain.
The positive selection on IGF1R may be used as a molecular marker in improving growth of poultry through molecular breeding. The computational approach is fast and accurate and may be used as an additional tool in genetic improvement of poultry.
CHAPTER ONE
INTRODUCTION

1.1 Background of the study

Chickens, quails and guinea fowls are poultry which are domesticated by humans for production of eggs, meat, or feathers. Poultry is classified in the Kingdom Animalia, Phylum Chordata, Subphylum Vertebrata, Class Aves, Order Galliformes (Al-Nasser et al., 2007). The chickens belong to the Family Phasianidae, subfamily Phasianinae, Genus Gallus. There are four species of the genus Gallus which include Gallus gallus, Gallus varius, Gallus lafayettei and Gallus sonnerati and three sub species of Gallus gallus which are Gallus gallus gallus, Gallus gallus spadiceus and Gallus gallus bankira (Al-Nasser et al., 2007).

Though the introduction of chicken domestication in Africa has not been documented well, there are postulations that chickens were introduced to Africa first via Egypt then they dispersed southwards through River Nile (Mwacharo et al., 2013).

In East Africa, there is archaeological evidence of chickens in the mid seventeenth century BC in Sudan (Houlihan & Goodman, 1986), 800 AD in coastal Kenya (Marshall, 2000) and in Akameru and Cyinkomane in Rwanda (MacDonald, 1992). Subsequent dispersion to other parts of Africa however remains unclear (Mwacharo et al., 2013).
On the other hand, the quail has its origin in Northern and southern Asia and its dispersal to other parts of the world is through its ability to fly in short distances (Hosomichi et al., 2006). There are two avian Coturnix species: *Coturnix japonica* (Japanese quail) and *coturnix coturnix* (common quail). Of the two quail species, the Japanese quail is the domesticated one, while the common quail remained a wild bird (Hosomichi et al., 2006). The Japanese quail first migrated to North America, Europe and Middle East in the 20th century (Hosomichi et al., 2006).

The guinea fowl is a native of Africa. There are footprints and a clutch of eggs that have a close comparison to those of the guinea fowl which were found in Laetoli in Tanzania (Leakey and Hay, 1979). Therefore, the guinea fowl is believed to have originated from Tanzania.

1.2 Chickens for egg and meat production

Chickens account for the largest livestock species reared by man probably because they have little to less cultural or religious restrictions (Tadelle et al., 2003). There are currently 32 million poultry in Kenya (Agricultural sector development strategy, 2010-2020). Indigenous chickens form 76%, 22% is comprised of layers, broilers, and the breeding stock. The rest of the domestic birds like quails, guinea fowls, ducks, geese, turkeys, and pigeons make up 2% of the poultry population (Olwande et al., 2010).

In Kenya 70% of the population resides in the rural areas with 90% engaging in poultry production (FAO, 2014). Majority of the poultry farmers are women and the youth (Kingori et al., 2010; FAO, 2014).
According to FAO (2010), indigenous chickens lay an average of 40-60 eggs per year. These eggs are laid in 3-4 clutches with each clutch consisting of 12-20 eggs. Each egg weighs 25-49 g with the exception of the egg laid by the naked neck which is 52g (Olwande et al., 2010). Indigenous chickens are also slow in maturing consequently influencing meat production (FAO, 2010).

The low production of eggs and meat in indigenous chickens has been attributed to poor housing, diseases, poor nutrition, and lack of appropriate breeds (Okeno et al., 2012).

Kenya produces about 1.3 billion poultry eggs and 20 tonnes of meat in a year (Agricultural sector development strategy, 2010). Despite the indigenous chicken having a low production, they contribute 47% and 55% to the Kenya’s egg and meat production respectively (King’ori, 2010).

There has been an increase in the demand for poultry products owing to the increase in human population, higher incomes, and higher standards of living (FAO, 2002).

Understanding the various production systems is important in identifying the needs of farmers, marketers, and consumers. This is crucial in formulating a holistic breeding program that aims at improving indigenous poultry (Okeno et al., 2012). The production is however not meeting the demand by the population (USAID, 2010).

However, indigenous poultry are highly variable phenotypically in size, earlobes, beaks, plumage color, feather morphology and pattern, skin color, comb type, live weights, and
egg production, among other traits (Kingori et al., 2010). This variation provides an opportunity for improvement through selective breeding.

The distinct ecotypes of indigenous chickens have been named according to their phenotypic descriptions and they include, naked neck, frizzled feathered, barred feathered, feathered shanks, bearded, dwarf sized, kuchi, bantam, and rumpleless (Moraa et al., 2015).

Most indigenous poultry are reared under small extensive production system (Okeno et al., 2012). In an attempt to improve production of eggs and meat in this system, local farmers use various observable traits like selection of egg size while others use selection of chicken for their preferred traits based on the performance history to influence successive generations. In some cases, cross breeding of indigenous poultry with exotic breeds is done which has not been successful because although production is generally improved, there is loss of broodiness making the poultry incapable of reproducing naturally (Okeno et al., 2012) (FAO, 2014). This crossbreeding is also accompanied by dilution of the appearance of the indigenous poultry which leads to rejection of the products by consumers (FAO, 2014). The costs of production also go up as the resultant crossbreeds are not adapted to scavenging conditions and there are also higher mortalities. These challenges are attributed to inappropriate technologies resulting from lack of understanding of the characteristics of the different production systems and the lack of information on consumer preferences and market dynamics (Okeno et al., 2012). Indiscriminate cross breeding is also threatening the existence of indigenous poultry which may lead to their extinction (Besbes, 2009).
The table below shows the preferred chicken traits by farmers, marketers and consumers that may help breeders in improving indigenous chickens to meet the demands (Okeno et al., 2012).

Table 1.1: Index ranking of traits perceived by farmers, marketers and consumers to be of economic importance.

<table>
<thead>
<tr>
<th>Traits</th>
<th>Index ranking</th>
<th>Index ranking</th>
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</tr>
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<tbody>
<tr>
<td></td>
<td>Farmers</td>
<td>Marketers</td>
<td>Consumers</td>
</tr>
<tr>
<td></td>
<td>1  2  3  Sum</td>
<td>1  2  3  Sum</td>
<td>1  2  3  Sum</td>
</tr>
<tr>
<td>Growth rate</td>
<td>12.64 7.03 6.86 26.53 0.512</td>
<td>- - - - - - - - - -</td>
<td>- - - - - - - - - -</td>
</tr>
<tr>
<td>Body size</td>
<td>5.51 9.06 11.52 26.10 0.503</td>
<td>8.33 15.68 27.02 51.03 0.152</td>
<td>10.38 18.24 20.44 49.06 0.154</td>
</tr>
<tr>
<td>Egg number</td>
<td>7.63 7.96 10.70 26.29 0.507</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Fertility</td>
<td>15.92 6.10 3.68 25.70 0.500</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Prolificacy</td>
<td>5.73 13.02 - 19.08 0.368</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Disease resistance</td>
<td>14.87 6.973 4.25 26.08 0.503</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Heat tolerance</td>
<td>3.17 5.17 10.62 18.97 0.366</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Drought tolerance</td>
<td>1.17 4.13 4.90 10.20 0.200</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Broodiness</td>
<td>3.40 9.18 6.21 18.79 0.362</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Mothering ability</td>
<td>4.51 6.10 7.92 18.54 0.357</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Temperament</td>
<td>- 1.80 2.45 5.14 0.099</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Plumage colour</td>
<td>3.06 1.31 7.19 11.76 0.227</td>
<td>1.83 1.74 3.11 6.68 0.021</td>
<td>- 1.37 4.48 5.85 0.018</td>
</tr>
<tr>
<td>Chicken shape</td>
<td>1.17 3.49 12.17 16.83 0.525</td>
<td>- - - - - - - -</td>
<td>- - - - - - - -</td>
</tr>
<tr>
<td>Meat quality</td>
<td>4.34 9.94 2.89 16.57 0.319</td>
<td>11.50 9.76 13.35 34.61 0.094</td>
<td>8.63 6.69 10.49 33.80 0.099</td>
</tr>
<tr>
<td>Egg size</td>
<td>12.58 5.06 3.10 20.74 0.400</td>
<td>34.33 13.59 1.24 49.16 0.192</td>
<td>37.45 12.46 - 49.91 0.205</td>
</tr>
<tr>
<td>Egg shell colour</td>
<td>- - - 1.26 0.024</td>
<td>11.83 20.49 3.73 46.05 0.172</td>
<td>12.26 31.91 4.20 48.38 0.180</td>
</tr>
<tr>
<td>Egg yolk colour</td>
<td>- - - 1.38 0.027</td>
<td>- - - 1.17 0.010</td>
<td>1.12 - 1.36 2.48 0.010</td>
</tr>
</tbody>
</table>

Source: Okeno et al., 2012.
The aim of this study is to identify signatures of selection in production genes for eggs and growth in poultry. The molecular approach is an accurate method of selecting preferred traits compared to traditional selection methods (Dekkers, 2002). This is because the genetic architecture of a phenotype is known compared to traditional selection method that relies on observation of the phenotype which may be influenced by external factors such as the environment. (Naqvi, 2007). This molecular approach reduces the generation interval as DNA is available at any stage of the animal unlike in traditional selection where the phenotype is recorded at a certain stage. Using the molecular approach in addition to traditional selection methods may lead to success in improving poultry production (Dekkers, 2002).

Agriculture accounts for 25-26% of the Gross Domestic Product (GDP) in Kenya with poultry farming contributing 30% of agricultural contribution to the country’s GDP (Nyaga, 2007).

Indigenous poultry can thus alleviate poverty and the Millenium Development Goals (MDG) of halving the number of poor people and empowering women will be met, if high producing genotypes are also identified (Moreki et al., 2010; FAO, 2014).
1.3 Statement of the problem

Production traits have not been successfully improved using the traditional selection methods. These methods are based on the use of phenotypic selection of traits that are observable and measurable (Naqvi, 2007). Animals with superior performance are used as the breeding stock (Naqvi, 2007). The traditional methods are difficult to improve traits for egg and meat production which have low heritability.

Improving these traits with low heritability is also costly as the phenotype is measurable only later in life (Dekkers, 2002). In this case, a large number of animals have to be raised so that a few are selected for breeding.

It is also difficult to improve traits that are not easily measurable like those which are recorded in one sex only or those that can be recorded after the death of an animal using the traditional selection methods (Dekkers, 2002; Naqvi, 2007). Crossbreeding of indigenous poultry with exotic genotypes is also used which leads to genetic erosion of the indigenous genotypes (FAO, 2014). The purpose of this study was to use the in silico approach to identify signatures of selection in production genes for egg and meat. This may be used as an additional tool to phenotypic selection by the government and other stakeholders to enhance selection by identifying superior breeds for the genetic improvement of economic traits in poultry (Dekkers, 2002).
1.4 Justification

As the computation approach uses genetic information, there is higher heritability because genetic information is not influenced by the environment (Naqvi, 2007). The use of genetic information allows for early selection and the generation intervals are reduced since the molecular information is available as early as at the embryo stage (Dekkers, 2002). Genetic information is available on all selection candidates: sex-limited traits, traits which are difficult to measure, or traits which require the animal to be slaughtered (Dekkers, 2002). As the use of this approach enhances selection intensity and accuracy with reduction in generation intervals, the rate of genetic change is high (Dekkers, 2002).

Using the molecular approach in addition to traditional selection methods may lead to success in improving poultry production (http://www.ccsi.ca/Meetings/canmore/jack_dekkers). This will benefit the resource-limited farmers who form the majority of poultry farmers.

Poultry farming has the potential to alleviate poverty (Moreki et al., 2010). When poultry production is improved, the economy of the country will also improve as poultry contributes to 30% of agricultural contribution to the country’s GDP (Nyaga, 2007).
1.5 Research hypothesis

**Null hypothesis:** There are no signatures of selection in genes for egg production and growth in poultry.

**Alternative hypothesis:** There are signatures of selection in candidate genes for egg production and growth in poultry.

1.6 Objectives

1.6.1 General objective

To perform computational analysis of molecular signatures of selection in candidate genes involved in egg production and growth in poultry.

1.6.2 Specific objectives

1. To analyze and identify molecular signatures of selection in Prolactin, Vasoactive intestinal peptide 1, and vasoactive intestinal peptide receptor 1 genes implicated in egg production in poultry.

2. To analyze and identify molecular signatures of selection in Growth hormone, Growth hormone receptor, Insulin-like growth factor 1, and Insulin-like growth factor 1 receptor genes implicated in growth in poultry.

3. To predict secondary and tertiary structures of proteins encoded by the selected candidate genes for growth and egg production in poultry.

4. To compare the putative 3D structures of the proteins encoded by the candidate genes between the homologs.
CHAPTER TWO
LITERATURE REVIEW

2.1 Candidate genes and their functions

2.1.1 Prolactin

Avian species have a single prolactin-encoding gene which is located on chromosome 2 (Alipanah et al., 2011). The prolactin gene has been cloned in different avian species, examples including turkey, quail, duck, pigeon, and domestic chicken. The avian prolactin gene consists of five exons and four introns (Yousefi et al., 2012). The gene has two regulatory regions, the proximal and distal enhancers. The distal segment is responsible for most of the transcription activity. Expression of the avian prolactin gene occurs in the ovary, oviduct, and hypothalamus. The highest expression is found in the pituitary gland (Li et al., 2009).

The prolactin hormone is the expressed product of the prolactin gene. It is composed of 199 amino acids (Wilkanowska et al., 2013). The hormone is a polypeptide which is secreted by the anterior pituitary gland and by other numerous tissues and cells (Freeman et al., 2000). It has more than 300 functions in various vertebrates (Bole-Feyesot et al., 1998). The functions of prolactin can be divided into the following categories: 1) Water and electrolyte balance 2) Growth and development 3) Endocrinology and metabolism 4) Brain and behavior 5) Reproduction 6) Immunoregulation and protection.
In avian species, the induction of broodiness and incubation is one of the most important functions of prolactin (Jiang et al., 2005). Riddle et al., (1935) demonstrated that injection of prolactin induces incubation behavior in birds. However, changes in the promoter binding site of prolactin due to polymorphism influences mRNA expression conversely influencing incubation behavior and egg production. For example, an insertion of a 24 bp at the promoter region of prolactin is associated with egg production (Cui et al., 2006 ; Usman et al., 2014). To perform any of its functions, prolactin interacts with its receptor in various target cells leading to a cascading activation of intracellular events (Freeman et al., 2000).

The prolactin-receptor binding happens in a two-step process: 1) one site of prolactin binds to one receptor molecule 2) a second receptor molecule binds on a second site of prolactin (Bole-Feysot et al., 1998, Freeman et al., 2000). This forms a homodimer of prolactin and the two receptor molecules. The prolactin associates with JKA2 which is a cytoplasmic tyrosine kinase (Freeman et al., 2000). On dimerization, tyrosine phosphorylation is induced followed by activation of the JAK kinase and phosphorylation of the receptor. The phosphorylation of cytoplasmic stat proteins, which dimerize and translocate to the nucleus then bind specific promoter elements on PRL-responsive genes is a major pathway of signaling. The PRL also activates the Ras/Raf/MAP kinase pathway which may be involved in the proliferation effects of the hormone. IRS-1, P1-3 kinase, SHP-2, PLCγ, PKC and intracellular Ca^{2+} are also potential mediators (Bole-Feysot et al., 1998).
2.1.2 Vasoactive intestinal peptide 1 and vasoactive intestinal peptide receptor 1

The vasoactive intestinal peptide 1 (VIP1) gene is 1,069 bp long. The avian VIP1 gene is found on chromosome 3 (https://www.ncbi.nlm.nih.gov/nuccore/NM_205366.2).

The expressed protein is vasoactive intestinal peptide 1 (VIP1) which is a 28 amino acid, regulatory peptide whose actions are through its interaction with specific receptors which are coupled to adenyl cyclase and the production of cAMP (Igarashi et al., 2011). It has regulatory effects on the circulatory, reproductive, immune and gastrointestinal systems. Chicken VIP1 is widely distributed in the peripheral organs and the brain. It is found notably in the endocrine glands where it exerts pleiotropic physiological functions.

In avian species, it stimulates prolactin release although it is not the main regulator and also stimulates prolactin gene expression (Chaiseha et al., 2004). In the avian reproductive cycle, VIP1 controls prolactin (PRL) secretion at the hypothalamus while vasoactive intestinal peptide receptor 1 (VIPR1) partly controls it in the pituitary (Chaiseha et al., 2004).

It also occurs in high concentrations in the hypothalamus. A study done by Tong et al., (1998) on turkey hens showed that VIP1 regulates the expression of PRL gene by increasing its concentration by acting both at the transcriptional level and at the level of PRL mRNA stability. On the other hand, El-Halawani et al., (2000) demonstrated the action of VIP1 on incubation behavior through the active immunization of turkey hens against VIP1 which resulted in inhibition of incubation behavior and subsequent
increase in egg production. The role of VIP1 is regulated through the VIPR1 gene expression in the pituitary (Zhou et al., 2008).

The chicken vasoactive intestinal peptide receptor 1 (VIPR1) gene is 67,906 bp long (Zhou et al., 2008). It is located on p3.2 of chromosome 2 (Kansaku et al., 2001).

The chicken VIPR1 gene is composed of 13 exons. The exons range in size from 45 to 1,031 bp. The VIPR1 gene is expressed in the pituitary and the hypothalamus. The differential mRNA expression of the VIPR1 gene is associated with changes in reproduction (Chaiseha et al., 2004).

The VIPR1 is a glycoprotein that belongs to the class II subfamily of the 7-transmembrane G-protein-coupled receptors superfamily (Zhou et al., 2008). It has a large hydrophilic extracellular N-terminus followed by 7 highly conserved hydrophobic transmembrane helices and a cytoplasmic C-terminus. In avian species, VIPR1 is found in the hypothalamus, on the surface membranes of the anterior pituitary, the small intestines and the granulosa cells (Zhou et al., 2008). DNA variation of VIPR1 is associated with broodiness in avian species (Zhou et al., 2008).

2.1.3 Growth hormone and growth hormone receptor

The chicken growth hormone (GH) gene is about 4 kbp (Tanaka et al., 1992). It is located on the long arm of chromosome 1q4. The chicken GH gene consists of five exons and four introns (Tanaka et al., 1992). Although the chicken GH gene is analogous to mammalian GH genes, the size is significantly larger. The intron size of the
chicken GH gene makes it bigger by 3.5 kbp. Primer extension analysis determined the
start point of transcription of the GH gene to be 56 bp upstream from the start codon
(Tanaka et al., 1992).

Growth hormone (GH) is a 190 amino acid protein that is synthesized and secreted in the
anterior pituitary gland (Buggiotti and Primmer, 2006). It performs many physiological
actions among which is controlling growth by regulating insulin-like growth factor 1
(IGF1) concentrations, stimulating lipolysis, providing FFAs, and glycerol as substrates
for energy metabolism and also inhibiting insulin-induced suppression of hepatic
gluconeogenesis (Clemmons, 2004).

The growth hormone also maintains carbohydrate and lipid homeostasis under normal
conditions. Many of these functions are exerted by regulation of the transcription of
other genes including IGF1 transcription factors and metabolic enzymes (Clemmons, 2004).

On the other hand, the chicken growth hormone receptor (GHR) gene is located on
exons.

Growth hormone receptor (GHR) belongs to the vertebrate somatotropic axis which
regulates postnatal growth and metabolism (Varvio et al., 2008). It is also a member of
the large cytokine receptor superfamily and it has 3 domains which are an extracellular
ligand-binding domain, a transmembrane domain and a cytoplasmic domain. Growth
hormone binds to the extracellular domains of the two GHR molecules leading to GHR
dimerization, activation of the GHR-associated JAK2 (Janus-Family Tyrosine Kinase-2), and tyrosyl phosphorylation of both JAK2 and GHR (Frank, 2001). This leads to activation of a variety of signaling molecules including MAPKs (Mitogen-Activated Protein Kinases), IRS1 (Insulin Receptor Substrate-1), PI3K (Phosphatidylinositol-3-Phosphate-Kinase), DAG (Diacylglycerol), PKC (Protein Kinase-C), Ca$^{2+}$ (intracellular calcium), and STATs (Signal Transducers and Activators of Transcription).

The signaling molecules contribute to the changes in enzyme activity, transport function and gene expression all of which are GH-induced. This ultimately leads to changes in growth and metabolism (Varvio et al., 2008).

2.1.4 Insulin-like growth factor 1 and insulin-like growth factor 1 receptor

The insulin-like growth factor 1 (IGF1) gene is 50 kbp long (Kajimoto and Ratowein, 1991). Genetic linkage analysis has determined the chicken IGF1 gene to be located on the short arm chromosome 1 near the centromere (Klein et al., 1996). The expression of IGF1 gene is ubiquitous including in the kidney, testes, liver, muscle, heart, ovary, brain, intestines, among other tissues (Tanaka et al., 1996).

IGF1 plays a role in various physiological and metabolic processes for example stimulating myogenesis, inhibiting apoptosis, increasing the absorption of glucose, activation of cell cycle genes and it also intervenes in the synthesis of DNA, RNA, protein and cell proliferation (Etherton, 2004).
The IGF1 gene is hence used as a candidate gene in the prediction of growth and meat quality traits in genetic improvement programs (Andrade et al., 2008). The fundamental role of IGF1 in prenatal and postnatal development and other physiological roles are through the binding to IGF1 receptor (Delafontaine et al., 2004).

On the other hand, the insulin-like growth factor 1 receptor (IGF1R) gene is 10,818 bp long. It is located on chromosome 10. Chicken IGF1R has 21 exons (https://www.ncbi.nlm.nih.gov/gene/395889).

The insulin-like growth factor 1 receptor (IGF1R) protein belongs to the receptor tyrosine kinase family (Delafontaine et al., 2004). It is phosphorylated following the binding to IGF1 (Delafontaine et al., 2004). IGF1R is a large transmembrane glycoprotein dimer which consists of several structural domains. The N terminal of the ectodomain contains two leucine-rich repeat domains separated by a cysteine-rich region while the C-terminal of the ectodomain consists of three fibronectin type III domains (Adams et al., 2000). The IGF1R is widely expressed in many cells in the fetal and postnatal tissue where it stimulates growth. IGF1R binds IGF1 and also IGF2 although with a lower affinity. IGF1, IGF2 and insulin ligands share a common 3D architecture and can bind IGF1R in a competitive manner. On activation, the IGF1R functions in cell growth and survival control.

When the IGF1R binds with its ligand, the receptor kinase is activated leading to receptor autophosphorylation, and tyrosine phosphorylation of multiple substrates which function as signaling adapter proteins including, the insulin-receptor substrates (Insulin
receptor substrate (IRS)1/2, Shc and 14-3-3 proteins (Delafontaine et al., 2004). The phosphorylation of IRSs proteins leads to the activation of two main signaling pathways: the PI3K-AKT/PKB pathway and the Ras-MAPK pathway. Activation of the MAPK pathway leads to increased cellular proliferation, whereas activating the PI3K pathway inhibits apoptosis and stimulates protein synthesis. Phosphorylated IRS1 can activate the 85 kDa regulatory subunit of PI3K (PIK3R1), leading to activation of several downstream substrates, including protein AKT/PKB. AKT phosphorylation, in turn, enhances protein synthesis through mTOR activation and triggers the anti-apoptotic effects of IGFIR through phosphorylation and inactivation of BAD.

IPI3K-driven signaling leads to recruitment of Grb2/SOS by phosphorylated IRS1 or Shc leads to recruitment of Ras and activation of the ras-MAPK pathway. IGF1R also signals through the Janus kinase/signal transducer and activator of transcription pathway (JAK/STAT). Phosphorylation of JAK proteins can lead to phosphorylation/activation of signal transducers and activators of transcription (STAT) proteins. Particularly, STAT3 activation may be essential for the transforming activity of IGF1R. The JAK/STAT pathway activates gene transcription and may be responsible for the transforming activity. JNK kinases can also be activated by the IGF1R. The activation can be inhibited by IGF1 through phosphorylation and inhibition of MAP3K5/ASK1, which is able to associate with the IGF1R directly. When present in a hybrid receptor with INSR, it binds IGF1 (Delafontaine et al., 2004).
2.2 Gene signatures of selection

Variation of genes in organisms both within and between species comes about by several evolutionary forces among which are natural selection, single nucleotide polymorphism, recombination, genetic drift and migration among others. This genetic polymorphism may lead to phenotypic polymorphism.

Natural selection

Natural Selection is the process by which some phenotypes adapt to their environment, survive and reproduce (Aquadro et al., 2001).

There are three forms of natural selection (Nielsen, 2005) i) positive selection which favors alleles that are advantageous ii) neutral/balancing selection which favors both alleles in a population hence maintaining polymorphism iii) negative/purifying selection which eliminates mutations which are unfavorable.

The different modes of selection are a response to the external pressure with each leading to change in allele frequency (Oleksyk et al., 2010). Positive selection on favoring advantageous alleles leads to a decrease in genetic variation. On the other hand, purifying selection on eliminating deleterious mutations maintains the integrity of functional sequences while balancing selection acts to maintain polymorphism (Oleksyk et al., 2010).
There are several postulations on the nature of variation among living organisms (Nielsen, 2005). According to the neutral theory, a large proportion of variation both within and between species is selectively neutral.

Meaning that new mutations that arise do not confer a fitness advantage as the loci are not affected by selection (Kimura, 1983). New mutations (due to random factors) which may not confer any fitness advantage may occur and increase in frequency in the population. The process by which the frequency of alleles change in a population due to random factors is known as genetic drift (Nielsen, 2005). Another postulation maintains that most of the variation observed does affect the fitness of the organism and is subject to positive selection (Gillespie, 1991). There is however an unsettled debate on the true postulation (Nielsen, 2005).

The regions with DNA sequences which are involved in variation of phenotype due to natural or artificial selection are called selection signatures (Qanbari et al., 2012). Each mode of selection leaves a specific signature on the genome architecture (Oleksyk et al., 2010).

The search for molecular signatures of selection is motivated by: 1. Interest in species’ evolutionary past and the mechanism of the evolution 2. The association of the signatures with traits of interest has adaptive or functional importance (Nielsen et al., 2007).

The studies are possible due to development of various tools and methods (Lopez et al., 2015). Negative and purifying selection may be of great interest because it may help
detect regions or residues of functional importance, but positive selection generates more interest because it is associated with formation of new functions and adaptation (Nielsen, 2005). When alleles that have undergone positive selection increase in a population, they become fixed in the population bringing about genetic variation and subsequent phenotypic variation.

The signatures of positive selection are divided into five main groups:

2.2.1 Long haplotype

A positively selected allele can become prevalent rapidly such that recombination does not substantially break the association with alleles at nearby loci on the ancestral chromosome (Sabeti et al., 2006). This collection of alleles in a chromosomal region that occur together is called a haplotype. Selective sweeps may produce an allele with high prevalence and long range associations with other alleles. The long range associations are a long haplotype which have not been broken down by recombination (Sabeti et al., 2006). This signature detects positive selection that occurred <30,000 years ago (Sabeti et al., 2006). The method used to detect this is the LD and (Long-range haplotype) LRH (Toomajian et al., 2003).

2.2.2 Population differentiation

This signature detects positive selection that happened between <50,000-75,000 years ago (Sabeti et al., 2006).
Population differentiation can only arise when populations living in separate geographical regions are subject to different environmental or cultural pressure, positive selection may change the frequency of an allele in one of the populations (Sabeti et al., 2006). Positive selection may lead to genetic variation in one population and not the other. Indices used to detect this include the breeding coefficient $F_{ST}$ (Akey et al., 2002).

2.2.3 Reduction in genetic diversity

When an allele increases in population frequency, variants at nearby locations on the same chromosome also rise in frequency. This so called “hitchhiking” leads to a selective sweep that alters the genetic variation pattern of the region (Sabeti et al., 2006). When a complete sweep occurs, the allele that has been selected together with the closely linked variants becomes fixed eliminating diversity. This diversity is restored when new slow mutations occur. These selective sweeps are detected in genetic variation within species. SNP is the most commonly used variant. This signature is used to detect positive selection that happened <250,000 years ago (Sabeti et al., 2006). Statistical tests used for this detection include deviations from HWE, Tajima’s D, HLA test and Fu and Li’s D and D*, Fu and Li’s F and F*, heterozygosity and Ewens Watterson homozygosity (F) (Hudson et al., 1987).

2.2.4 High frequency derived alleles

Derived or non-ancestral alleles arise as a result of new mutation and usually have lower allele frequencies compared to ancestral alleles (Watterson and Guess, 1977). Derived alleles that are linked to a beneficial allele can hitchhike to high frequency in a selective
sweep (Sabeti et al., 2006). Many of the derived alleles will not reach complete fixation thus a signature of a region that contains many high-frequency derived alleles is created by positive selection.

This signature detects positive selection that occurred <80,000 years ago (Sabeti et al., 2006). The test commonly used to detect this is the Fay and Wu’s H (Fay and Wu, 2000).

2.2.5 High proportion of function altering mutation

Molecular variations that alter the function of a protein are deleterious and are less likely to become fixed compared to mutations that do not lead to a change in protein function (Sabeti et al., 2006). Fixation of function altering mutations that are beneficial can be increased when positive selection occurs over a prolonged period (Li et al., 1985).

The changes can be measured by comparing DNA sequences between species. Detection of the increase may be done by comparing the rate of non-synonymous to the rate of synonymous mutations (Li et al., 1985). This signature can be determined using tests such as ka/ks ratio, McDonald and Kreitman tests (MK tests), \( d_{NS}/d_S \) ratios and codon substitution models implemented in the PAML package (McDonald and Kreitman 1991). This signature is studied the most and is used to detect adaptive evolution.

The \( d_{NS}/d_S \) ratio is the most popular test to model positive natural selection in the gene sequences in terms of positive adaptive selection (Ngandu et al., 2008). This is the relative rate of non- synonymous to synonymous substitutions in the evolutionary history of a sample of sequences.
The $d_N/d_S$ ratio can detect positive selection between and within species. The codon substitution models of Nielsen and Yang (1998), as implemented in the PAML package promoted $d_N/d_S$ to the status of a parameter which they called omega ($\omega$). Values of omega $<1$ indicate negative or purifying selection while values of omega $>1$ indicate positive or diversifying selection (Yang, 2007). When the value of omega is equal to 1, this is an indication of neutral selection (Yang, 2007).

### 2.3 Lineage specific positive selection

An approach for detecting positive selection is by use of comparative data that detects positive selection within and between species. For lineage specific positive selection, two codon based models are commonly used, Model 0 which is the one-ratio model and the free-ratio model (Model 1). The one-ratio model assumes an equal $\omega$ ratio for all the branches in the phylogeny while the free-ratio model assumes an independent $\omega$ ratio for each branch (Yang, 1998).

The two models are then compared by the likelihood ratio test (LRT) using the following formula: $\text{LRT} = 2 (\ln LM_0 - \ln M_1)$ (Yang, 1998). The result is then compared to a $\chi^2$ distribution with N-1 degrees of freedom where N is the number of branches. A significant difference in the LRT value is an indication that the $\omega$ results obtained are reliable (Yang, 1998).
2.4 Site specific positive selection

For site specific positive selection, again two codon based models are commonly used, the model 7 (beta) which uses a beta distribution hence allows sites with $\omega$ values of 0 to 1 and Model 8 (beta and $\omega$) adds an extra class of sites to model 7 with the proportion and the $\omega$ ratio estimated from the data hence allowing for sites with $\omega > 1$ (Yang et al., 2000). The two models are also compared by the LRT i.e. LRT = 2 (lnLM$_7$ - lnM$_8$). For the sites specific models, posterior probabilities are also calculated for sites under positive selection and values >0.95 indicate that the results for the amino acid sites under selection are reliable (Nielsen & Yang, 1998; Yang et al., 2005).

Few studies have been done to detect positive selection at candidate genes that influence production of eggs and meat in livestock. When positive selection is detected on the genes that influence the general production of eggs and meat, the phenotype can be confirmed by carrying out in vitro and in vivo studies.

Zhu et al., (2010) used the codon substitution models to identify positive selection in the mx gene of the galliform and anseriform avian. The mx proteins confer different antiviral activity in many different species (Zhu et al., 2010). In the study, the positively selected sites were distributed in the N-terminal end of the mx gene and GTPase domain which are important for the antiviral activity of mx protein. These may be used as molecular markers for breeding poultry with improved antiviral activity (Zhu et al., 2010).
2.5 Protein 3D structure prediction

The structure of a protein is valuable in determining the functions of a protein like enzymatic activity or the interaction with other proteins (Bairoch, 2000). The solving of protein structures experimentally such as those in Protein Data Bank (PDB) is time consuming and costly and can be sometimes technically difficult (Kallberg et al., 2012). There are different computational protocols that have been developed to solve the problem of the many protein sequences that have been deposited without solved structures. There are three categories of computational structure prediction methods: Comparative/template modeling, threading and template free/ab initio modeling (Kallberg et al., 2012).

In comparative modeling, the structure is constructed by aligning the sequence or sequence profile of the target protein to an evolutionary related protein which has a solved structure (template) to find equivalent residues (Marti-Renom et al., 2000). In threading, the target sequence is matched to a solved 3D structure template which has no evolutionary relationship (Jones et al., 1992). Threading is based on the principle that evolutionary distant or protein sequences that are unrelated often share common structural elements. Structure predictions based on existing template structures like comparative modeling and threading produce high resolution models. However, these models are only useful if close templates are available (Kallberg et al., 2012). When there are no solved structures for homologs, ab initio methods construct models without using such information. However, this method can successfully create models from sequences which are <120 residues (Wu et al., 2007).
Accuracy of the predicted model is important for it to be biologically useful. The accuracy is largely dependent on the availability of close structural templates and the prediction method.

The RaptorX is a software that can create models in which there are no close templates (Peng & Xu, 2011). It optimizes the modeling strategy by using the profile entropy scoring method that considers the number of non-redundant homologs which are available for the target and the template structure to assess the information in sequence profiles (Peng & Xu, 2010). The software also integrates biological signals in a nonlinear score function using conditional random fields (CRFs) (Peng & Xu, 2009).

In addition, RaptorX uses a Multiple-Template Threading (MTT) procedure which uses multiple templates to model a target sequence (Multiple-Template Threading) procedure (Peng & Xu, 2011). It is possible to have many good templates for a given target which may improve the modeled structure. Using the MTT, the alignment between individual pairwise sequence-template alignments is improved through the use of a probabilistic consistency transformation (Källberg et al., 2012).

Thiel et al. (2015) used Raptor X to predict the structure of CD45 in primates. CD45 has a crucial role in the immune system where it is central to the development of T cells and also determines T and B lymphocytes activation (Thiel et al., 2015). Viruses interfere with the immune system through the interaction of viral proteins with the CD45. This interaction is the driving force for the evolution of CD45 (Thiel et al., 2015). In this
study, they modeled the ectodomain of the CD45 which consists of three fibronectin III-like repeats and a cysteine-rich domain.

Sites under positive selection were distributed in the different domains. The sites were found at the surface of the ectodomain where pathogens could easily interact with the CD45 through the viral proteins. This adaptive selection is important to the host in that it prevents the virus from evading the immune system (Thiel et al., 2015).

2.6 Protein structure alignment

A protein structure alignment method consists of two major components: a scoring function measuring protein similarity and a search algorithm optimizing the scoring function (Wang et al., 2013).

There are different programs for automatic structure alignment whose accuracy is low compared to manually-curated alignments which consider the geometric similarity, evolutionary and functional relationship (Mayr et al., 2007). Most of these programs produce alignments that are not biologically meaningful. DeepAlign is one of the programs that produces biologically meaningful alignments as its scoring function considers the 3D geometric similarity, evolutionary information at the sequence and local substructure levels as well as hydrogen-bonding similarity (Wang et al., 2013). Proteins in a family share an overall shape but their structures are highly flexible due to evolution at the local substructure and the sequence level. DeepAlign uses amino acid and local substructure substitution matrices to quantify the local conformational change resulting from evolutionary events (Wang et al., 2013). Block substitution matrix
(BLOSUM) is the commonly used substitution matrix. It is not sensitive for remote homologs as it is derived from close homologs.

DeepAlign hence uses conformation letter substitution matrix (CLESUM) which is a local substitution matrix to measure the evolutionary distance of two proteins at the local substructure level (Zheng, 2008; Wang et al., 2013). In addition, to make alignments that are more biologically meaningful especially for beta-strands, DeepAlign uses the hydrogen-bonding similarity. Conformation letter substitution matrix (CLESUM) favors the alignment of two loop regions that are evolutionary related but disfavors that of unrelated helices.

To reduce the running time used to find an optimal structure alignment, DeepAlign identifies a set of initial alignments. It uses both amino acid and local substructure mutation scores to identify evolutionary related structural fragment pairs then builds initial alignments from these pairs (Wang et al., 2013).

In order to generate biologically meaningful structural alignments, all four factors are necessary (Wang et al., 2013). However the 3D geometric similarity which includes the TMscore, RMSD and uGDT is the major factor for determining similarity of proteins. TMscore is better than RMSD as it is independent of the length of the protein and is not biased by only a few residues that are badly aligned (Wang et al., 2013). Structure alignments with a TM value of > 0.5 are mostly in the same fold while those with TM<0.5 are not in the same fold (Xu & Zhang, 2010).
CHAPTER THREE
MATERIALS AND METHODS

3.1 Retrieval of candidate gene sequences

The full coding sequence mRNA sequences of chicken prolactin, vasoactive intestinal peptide 1, and vasoactive intestinal peptide receptor 1 implicated for egg production and growth hormone, growth hormone receptor, insulin-like growth factor 1, insulin-like growth factor 1 receptor implicated for growth were retrieved from GenBank (https://www.ncbi.nlm.nih.gov/genbank/) using ENTREZ searches across the databases. The reference genome used was the chicken genome available at ENSEMBL (Hubbard et al., 2002).

3.2 Search for homologs and retrieval

A reciprocal BLAST was done using the algorithm BLASTp which is available at NCBI to infer homology. The amino acid sequence of each candidate gene in the fasta format was used as the query in performing pairwise sequence alignments in non-redundant (nr) databases. The substitution matrix BLOSUM62 was used. For egg production, homologues selected were birds and amphibians with an E value greater than 1e-10. For meat production, the homologues selected were animals that are commonly eaten by man with an E value of greater than 1e-10. Amino acid sequences of the homologues were converted to their corresponding coding sequences and the stop codons manually removed to prevent interference with later analysis.
3.3 Multiple sequence alignment

ClustalX2 (Larkin et al., 2007) which is a Multiple sequence alignment software that uses a progressive algorithm which is heuristic in nature was used to align the different homologues for each candidate gene to confirm homology and as a preceding step in phylogenetic analysis. PRANK that also uses the progressive algorithm and MAFFT (Katoh & Standley, 2013) were used. This was followed by multiple sequence alignment using MUSCLE version1.3.8.31-1 (Edgar, 2004). Jalview version 2.8 (Waterhouse et al., 2009) and Seaview were used to view and edit the alignments. On comparing of the alignments from the three softwares, alignments from MUSCLE were found to be better and were used for subsequent analysis.

3.4 Phylogeny construction

MEGA6 was used to model the amino acid substitution and the rate heterogeneity within the various homologues (Tamura et al., 2013). The model was selected using the Lowest Bayesian Information Criterion. Phylogenetic trees were then inferred using Nearest Neighbour Interchange with subtree pruning and regrafting of FastME2 (Lefort et al., 2015). 1000 bootstraps were used to test for confidence of the inferred relationships. The inferred trees were saved in Newick format. Figtree version 1.4.2 (Rambaut, 2014) was used for graphical visualization of the trees.
3.5 Substitution model selection and rate heterogeneity testing

Codon-based substitution models which were developed by Nielsen and Yang (1998) and Yang (2007), which are implemented in CODEML program of PAML were used to detect selection in lineage and in sites.

For lineage, 2 codon-based models: Model 0 or one ratio Model and Model 1 also known as the free ratio Model were used. The one ratio model assumes an equal $\omega$ ratio for all branches in the phylogeny. The free ratio model assumes an independent $\omega$ ratio for each branch. For the site specific, 2 codon-based models were used: Model 7(beta) and Model 8(beta and $\omega$). Model 7 allows for sites with values $\omega$ of 0 to 1 while Model 8 allows for sites with $\omega > 1$.

3.6 Input preparation in PAML

The Perl script copygaps_single.pl was then used to align the aligned protein sequence with the corresponding codon sequences. The output file for the analysis was saved as infile.nuc. At the UNIX prompt, three directories were created: Model0, Model1 and Sites. The codeml.ctl file was edited according to each directory corresponding to the analyses that were to be performed.

For Model0 directory, Model 0 was selected, for Model 1 directory, Model 1 was selected while for the Sites directory, Model 7 and 8 were selected. The phylogeny tree that was saved in Newick format was then moved into the intree.trees file. The
codeml.ctl which is the control file, infile.nuc which is the output from the Perl script and intree.trees which contained the tree file were each copied into the three directories.

3.7 Test for molecular signatures of selection

The PAML5 package (Yang, 2007) was then used for phylogenetic analysis using maximum likelihood. CODEML was run on the command prompt. This was done in two parts, first was the branch or lineage analysis and the second was the amino acid or site analysis. After the analysis, the log likelihood ratio test (LRT) (Yang, 1998) was used to compare two nested models: a null model that does not allow for any codons with ω ratio of greater than 1 against a more general model that does. Twice the log likelihood difference between the two models was compared against a chi squared ($\chi^2$) distribution to test the significance of the result obtained. For Lineage analysis, LRT was also used to compare Model7 (beta) with ω ratios of between 0 and 1 and Model8 (beta & ω) that adds a class of sites with dN/dS of >1 were used. Bayes Empirical Bayes analysis was done to infer the category an amino acid site belongs to (Yang et al., 2005).

Amino acid sites with a predetermined posterior probability of >0.95 belong to a site class that has a dN/dS ratio of > 1 and thus believed to be under positive selection (Yang et al., 2005).

3.8 Protein 3D structure prediction

The secondary and tertiary structures of the proteins encoded by the candidate genes for growth and egg production for poultry were modeled by sending the sequences to
RaptorX (http://raptorx.uchicago.edu/) which is a webserver that models protein structures using the homology algorithm. Pymol version 1.7.6 (DeLano, 2006) was used to view the 3D structures of the proteins.

3.9 Comparative 3D structure analysis

The sequences for the genes for egg production and growth for poultry were submitted to DeepAlign (http://raptorx.uchicago.edu/DeepAlign) which is a server for comparative structure analysis.
CHAPTER FOUR

RESULTS

4.1 Homology search

From the reciprocal BLASTp performed in the non-redundant databases, the orthologs (shown in Appendix 1-8) were selected. These had an E value lesser than 1e-10 for the different genes.

4.2 Analysis of signatures of selection at candidate genes for egg production in poultry

4.2.1 Multiple sequence alignment of prolactin

On using Muscle software to perform the Multiple sequence alignment of prolactin homologous, the alignment in Figure 4.1 was produced. The domain for prolactin is highly conserved across the poultry (greylag goose, wild duck, wild chicken, grey jungle fowl, Japanese quail, common quail, chicken breed lohmann, chicken breed yangshan, chicken breed 15 Hubbard, chicken cob 500, chicken breed avian 48) with a few substitutions. However, there were notable amino acids substitutions in common quail, Japanese quail, helmeted guinea fowl. In position 24 of common quail and Japanese quail, serine was substituted by glycine. In position 26 of common quail, arginine was substituted by threonine. In addition, in position 124 of common quail, Japanese quail, and common quail, glycine was substituted by alanine, in position 171 of common pea fowl, serine was substituted by alanine. For helmeted guinea fowl, threonine substituted
arginine in position 213. In position 253 of Japanese quail and common quail, glycine substituted alanine.

Figure 4.1: Multiple sequence alignment of prolactin protein. The alignment shows the degree of identity between the species. The sequences of different poultry species are highly conserved with few substitutions.

4.2.2 Multiple sequence alignment of vasoactive intestinal peptide 1 (VIP1)

On doing a Multiple sequence alignment of the VIP1 homologs using the Muscle software, the alignment below was produced (Figure 4.2). The domain for VIP1 is also conserved in different poultry species (ostrich, wild turkey, southern ostrich, swan...
goose, wild chicken) as shown in Figure 4.2. However, this sequence has many indels that are widely spread across the sequence even in the poultry species. Interestingly, the ostrich sequence is highly variable compared to the other poultry sequences.

In position 159 of the ostrich sequence, aspartate substituted asparagine, alanine substituted threonine in position 186, valine substituted proline in position 188, aspartate replaced asparagine in position 190, glycine substituted serine in position 253, glycine replaced arginine in position 241 and glutamate was replaced by glycine in position 293.

**Figure 4.2: Multiple sequence alignment of vasoactive intestinal peptide 1 (VIP1) protein.** The alignment shows that there have been many insertions/deletions that have occurred. The sequence of the ostrich is highly variable from other poultry sequences.
4.2.3 Multiple sequence alignment of vasoactive intestinal peptide receptor 1 (VIPR1)

From the Multiple Sequence Alignment of VIPR1 homologous sequences done using Muscle software, the alignment (shown in Figure 4.3) was produced.

VIPR1 gene is conserved with minimal variation within the poultry (wild turkey, wild duck, wild chicken, Japanese quail and Southern ostrich). The Southern ostrich and the wild duck are the most varied of the poultry species.

In the Southern ostrich, there are 6 sites with substitutions: at position 131, glutamic acid substituted glutamine, at position 134, alanine was substituted by glutamic acid, at position 151, tyrosine substituted phenylalanine, at position 256, serine substituted proline and at position 512, glycine substituted serine. The wild duck had several substitutions at position 183 where phenylalanine substituted tyrosine, at position 196, serine substituted alanine, at position 361, threonine was substituted by alanine and at position 428 threonine replaced alanine.
Figure 4.3: Multiple sequence alignment of vasoactive intestinal peptide receptor 1 (VIPR1) protein. The alignment shows the variations between the sequences.

4.2.4 Lineage selection in vasoactive intestinal peptide receptor 1 (VIPR1)

The phylogenetic tree for VIPR1 (shown in Figure 4.4) was constructed using FastME which is distance-based with 1000 bootstraps replicates to test the confidence of the topology. To test whether there was variable selective pressures acting on the lineages, the coding sequences for the homologs were fitted to the one ratio model (Model0) and the free ratio model (Model1) (Yang, 1998). For VIPR1, model 1 fitted significantly better than model 0 with P<0.05 when compared to $X^2$ distribution as seen in Table 4.1. This was an indication of heterogeneity in the selective pressures along the lineages.
Purifying selection was found to be acting on all lineages in VIPR1. All the lineages had a dN/dS<1.

**Figure 4.4:** A phylogram showing evolution of vasoactive intestinal peptide receptor 1 (VIPR1) gene. The branch lengths were estimated by maximum likelihood under the free ratio model (Model 1) which assumes an independent ω for each branch. The branch lengths are drawn in proportion to the expected numbers of nucleotide substitutions per codon.
4.2.5 Lineage selection in prolactin gene

The phylogram for prolactin shown in Figure 4.5 was constructed using FastME which is distance-based with 1000 bootstraps replicates to test the confidence of the topology.

To test whether there was variable selective pressures acting on the lineages, the coding sequences for the homologs were fitted to the one ratio model (Model0) and the free ratio model (Model1) (Yang, 1998). For prolactin, model1 fitted significantly better than model0 with P<0.05 when compared to X^2 distribution as seen in Table 4.1. This is an indication of heterogeneity in selective pressures along the lineages of the phylogeny below. The two branches leading to poultry were under positive selection. The branch leading to the ancestor of the wild chicken, quails and common peafowl had a high dN/dS ratio of 3.64. The rest of the lineages are under purifying selection.
Figure 4.5: Phylogeny showing evolution of prolactin (PRL) gene. The branch lengths were estimated by maximum likelihood under the free ratio model (Model 1) which assumes an independent $\omega$ for each branch. The branch lengths are drawn in proportion to the expected numbers of nucleotide substitutions per codon. The colored branches have $dN/dS>1$ hence indicates positive selection.

4.3 Analysis of signatures of selection at candidate genes for growth in poultry

4.3.1 Multiple sequence alignment of growth hormone (GH)

On performing Multiple sequence alignment of the growth hormone homologues using Muscle software, the alignment in Figure 4.6 below was produced.
Among the poultry species used in this alignment: wild chicken, common quail, wild duck, greylag goose, and Southern ostrich, the Southern ostrich has the most substitutions. At amino acid position 13, alanine was substituted by threonine in common quail, wild duck and greylag goose. At position 20, proline was substituted by glutamine in the greylag goose, wild duck and Southern ostrich. At position 69, tyrosine was substituted by histidine in greylag goose, wild duck and southern ostrich. At position 70, alanine substituted threonine in Southern ostrich, at position 86 valine substituted alanine, at position 87, phenylalanine substituted proline, at position 136, tyrosine substituted phenylalanine.
**Figure 4.6: Multiple sequence alignment of growth hormone (GH) protein.** The alignment shows the high level of identity between the poultry sequences. The Southern ostrich has more substitutions compared to the other poultry sequences.

**4.3.2 Multiple sequence alignment of growth hormone receptor (GHR)**

The alignment below (Figure 4.7) was produced by the Multiple Sequence Alignment software Muscle.

The sequences of greylag goose, knob-billed duck, and wild duck are highly similar with very few varying sites. At position 224, serine substituted alanine in these three species, at position 230, arginine substituted glutamine. At position 308, threonine substituted isoleucine in knob-billed duck, at position 337, glycine substituted glutamic acid in wild...
duck. At position 595, the three species have cysteine substituting tyrosine, at position 633, aspartate substituted histidine, at position 639, the greylag goose had valine being substituted by histidine while in knob-billed duck and wild duck, aspartate substituted valine, at position 641, serine substituted alanine in the three species, at position 650, alanine substituted threonine in the three species. At position 671 of the greylag goose, alanine substituted proline while leucine substituted proline in the knob-billed at the same position, at position 676 of the knob-billed duck, arginine substituted threonine while in the wild duck, serine substituted threonine.

Figure 4.7: Multiple sequence alignment of growth hormone receptor (GHR) protein. The alignment shows that there are very few sites with variations between the poultry sequences.
4.3.3 Multiple sequence alignment of insulin-like growth factor 1 (IGF1) gene

The alignment below (Figure 4.8) was produced on aligning the Insulin-like growth factor 1 homologous sequences using Muscle software.

The IGF1 gene is conserved with a few substitutions. At position 3 of the wild duck, lysine was substituted by glutamate, at position 58 of the wild turkey, alanine substituted glycine, at position 66 of Yunnan Daweishan chicken breed, serine was substituted by isoleucine, at position 117 of chicken haplotype h-12 and chicken haplotype h-5, proline substituted glutamine. In addition, at position 138 of chicken haplotype h-12 and chicken haplotype h-5, asparagine substituted lysine, at position 144 of chicken haplotype h-12 and chicken haplotype h-5, leucine substituted arginine, at position 145 of wild turkey and Southern ostrich, glycine was substituted by alanine.
4.3.4 Multiple sequence alignment of insulin-like growth factor 1 receptor (IGF1R) gene

On using Muscle software to perform a Multiple sequence alignment of IGF1R, the alignment in Figure 4.9 was obtained.

The sequences of wild duck and Southern ostrich are similar and had more substitutions compared to other poultry species: wild turkey, wild chicken and Japanese quail. At position 213 of wild turkey and the Southern ostrich, valine substituted isoleucine, at position 471 of the Southern ostrich, isoleucine substituted threonine, at position 745 of...
the Southern ostrich and wild duck asparagine substituted aspartate. At position 944 of wild duck and the Southern ostrich, threonine substituted alanine, at position 1117 of the Southern ostrich and wild duck, glycine substituted aspartate. At position 1291 of wild duck, threonine substituted alanine, at position 1331 of wild turkey, lysine substituted glutamic acid, at position 1332 of wild turkey, methionine substituted aspartate. At position 1474 of Japanese quail, alanine substituted proline.

Figure 4.9: Multiple sequence alignment of insulin-like growth factor 1 receptor (IGF1R) protein. The alignment shows that the sequences of the Southern ostrich and wild duck are highly similar but differ from the other sequences.
Table 4.1: Likelihood ratio tests (LRT) to detect selection in lineages and amino acid sites. Likelihood ratio tests were done to determine fitness of a model. Probability values of <0.05 shows that a model is significant.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Model Analysis</th>
<th>$2\Delta \text{Lineage}=\frac{2(\text{InlM0-InlM1})}{\text{Site}= \frac{2(\text{InlM7-M8})}}$</th>
<th>$X^2$ (Chi squared) value</th>
<th>Degrees of freedom</th>
<th>Probability Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Prolactin</td>
<td>Lineage Analysis</td>
<td>2(-5519.413936+5449.921133) = 138.9866</td>
<td>138.9866</td>
<td>139-71=68</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td></td>
<td>Site Analysis</td>
<td>2(-5469.7794-5466.9746) = 5.6096</td>
<td>5.6096</td>
<td>74-72=2</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>VIP1</td>
<td>Lineage Analysis</td>
<td>2(-529.499090+514.296366) = 30.41</td>
<td>30.41</td>
<td>115-59=56</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td></td>
<td>Site Analysis</td>
<td>2(-517.381707+517.979078) = 1.19</td>
<td>1.19</td>
<td>62-60=2</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>VIPR1</td>
<td>Lineage Analysis</td>
<td>2(-9226.699685+9164.329783) = 124.7</td>
<td>124.7</td>
<td>115-59=56</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td></td>
<td>Site Analysis</td>
<td>2(-9013.544+9005.761) = 15.566</td>
<td>15.566</td>
<td>62-60=2</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td>GH</td>
<td>Lineage Analysis</td>
<td>2(-4704.661507+4661.203348) = 86.9</td>
<td>86.9</td>
<td>111-57=54</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td></td>
<td>Site Analysis</td>
<td>2(-4634.798616+4634.799225) = 0.001218</td>
<td>0.001218</td>
<td>60-58=2</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>GHR</td>
<td>Lineage Analysis</td>
<td>2(-11526.92548+11486.242124) = 81.568540</td>
<td>81.568540</td>
<td>127-65=62</td>
<td>&lt;0.05</td>
</tr>
<tr>
<td></td>
<td>Site Analysis</td>
<td>2(-11295.23339+11293.456950) = 3.552886</td>
<td>3.552886</td>
<td>68-66=2</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>IGF1</td>
<td>Lineage Analysis</td>
<td>2(-2076.081066+2032.048315) = 88.065502</td>
<td>88.065502</td>
<td>151-77=74</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td></td>
<td>Site Analysis</td>
<td>2(-2017.692136+2015.465467) = 4.853338</td>
<td>4.853338</td>
<td>80-78=2</td>
<td>&gt;0.05</td>
</tr>
<tr>
<td>IGF1R</td>
<td>Lineage Analysis</td>
<td>2(-12701.06843+12641.506653) = 119.1</td>
<td>119.1</td>
<td>115-57=56</td>
<td>&lt;0.05</td>
</tr>
</tbody>
</table>
4.3.5 Lineage selection in growth hormone (GH) gene

The phylogram of GH shown in Figure 4.10 below was constructed using distance-based FastME with 1000 bootstrap replicates to test the confidence of the topology. To test whether there was variable selective pressures acting on the lineages, the coding sequences for the homologs were fitted to the one ratio model (model0) and the free ratio model (model1) (Yang, 1998). For GH, model1 fitted significantly better than model0 with P<0.05 when compared to X^2 distribution as seen in Table 1. This is an indication of heterogeneity in selective pressures along the lineages as shown in Figure 4.10.

In GH, positive selection was detected in the branch leading to cattle, sheep, goat and camel (shown in red). The dN/dS was 4.98 which is greater than 1. In the rest of the lineages, GH was under purifying selection as indicated by dN/dS of <1.
Figure 4.10: Phylogenetic tree showing evolution of growth hormone (GH) gene. The branch lengths are in proportion to the number of nucleotide substitutions per codon. There is heterogeneity in the dN/dS values in the branches. dN/dS>1 indicate positive selection (shown in red).

4.3.6 Lineage selection in insulin-like growth factor 1 receptor (IGF1R) gene

The phylogram of IGF1R (Figure 4.11) was constructed using distance-based FastME with 1000 bootstrap replicates to test the confidence of the topology.
To test whether there was variable selective pressures acting on the lineages, the coding sequences for the homologs were fitted to the one ratio model (model0) and the free ratio model (model1) (Yang, 1998). In the Lineages, P values of < 0.05 confirm heterogeneity in selection pressures. For IGF1R, model1 fitted significantly better than model0 with P<0.05 when compared to $X^2$ distribution as seen in Table 1. This is an indication of heterogeneity in selective pressures along the lineages as shown in Figure 4.11.

Purifying selection was found to be acting on all lineages in IGF1R. All the lineages had a dN/dS<1.
Figure 4.11: Phylogenetic tree of the insulin-like growth factor 1 receptor (IGF1R) gene. The tree was constructed using FastME with 1000 bootstraps. The branch lengths were estimated using maximum likelihood under the free ratio model (M1) which assumes an independent ω for each branch. The branch lengths are drawn in proportion to the expected numbers of nucleotide substitutions per codon.

4.3.7 Lineage selection in growth hormone receptor (GHR) gene

The phylogram of GHR shown in Figure 4.12 below was constructed using distance-based FastME with 1000 bootstrap replicates to test the confidence of the topology.
To test whether there was variable selective pressures acting on the lineages, the coding sequences for the homologs were fitted to the one ratio model (model0) and the free ratio model (model1) (Yang, 1998). For GHR, model1 fitted significantly better than model0 with P<0.05 when compared to $X^2$ distribution as seen in Table 1. This is an indication of heterogeneity in selective pressures along the lineages as shown in Figure 4.12.

Purifying selection was found to be acting on all lineages in GHR. All the lineages had a $dN/dS<1$. 
Figure 4.12: Phylogenetic tree showing evolution of growth hormone receptor (GHR) gene. The branch lengths were estimated using maximum likelihood under the free ratio model (M1) which assumes an independent $\omega$ for each branch. The branch lengths are drawn in proportion to the expected numbers of nucleotide substitutions per codon.

4.3.8 Amino acid site selection

To test the hypothesis of different selective pressure on the amino acid sites, twice the log likelihood difference of Model7 and Model8 was done for all the proteins as shown in Table 4.1.
The Multiple sequence alignment of insulin-like growth factor 1 receptor (IGF1R) with Isoleucine under positive selection is shown in Figure 4.13. The Likelihood ratio test showed that in GH and IGF1R genes, there are different selective pressures acting on each amino acid site. Posterior probability analysis identified positive selection on amino acid Isoleucine at site 460 of the IGF1R to be under positive selection as it had a posterior probability of 0.985. This was greater than the predetermined cut-off value of >0.95 for inferring that a site belongs to a class which has a dN/dS of > 1.

![Image of Multiple sequence alignment]

Figure 4.13: Multiple sequence alignment of insulin-like growth factor 1 receptor (IGF1R) with site under positive selection. Posterior probability analysis identified isoleucine at position 460 to be under positive selection. The site is shown in red.

The raptor X server predicted the 3D structure of insulin-like growth factor 1 receptor (IGF1R) which is shown in Figure 4.14. The amino acid site under positive selection...
was located on the 3D structure. Pymol was used to visualize the structure with the site under selection.

Figure 4.14: The 3D structure of insulin-like growth factor 1 receptor showing the different domains. The site shown in red is predicted to be under positive selection by Model 8. The site is located in the Receptor L domain which is important for binding insulin-like growth factor 1.
4.4 Protein 3D structure prediction for genes implicated in egg production

4.4.1 Predicted 3D structures of prolactin protein

In Table 4.2 below, the secondary and tertiary structures of prolactin from different poultry species were predicted. There was a small variation in the secondary structure but the tertiary structure was similar.

Table 4.2: Predicted 3D structures of prolactin protein. The table below shows the secondary and tertiary structures of prolactin from different poultry species.

<table>
<thead>
<tr>
<th>Prolactin protein structures</th>
<th>Helices</th>
<th>Beta sheets</th>
<th>Loops</th>
<th>Domain interval</th>
<th>Solvent accessibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Common peafowl</td>
<td>62%</td>
<td>0%</td>
<td>37%</td>
<td>31-229</td>
<td>50%</td>
</tr>
<tr>
<td>Common pheasant</td>
<td>63%</td>
<td>0%</td>
<td>35%</td>
<td>31-229</td>
<td>50%</td>
</tr>
<tr>
<td>Common quail</td>
<td>60%</td>
<td>0%</td>
<td>39%</td>
<td>31-229</td>
<td>50%</td>
</tr>
<tr>
<td>Helmeted guinea fowl</td>
<td>65%</td>
<td>0%</td>
<td>34%</td>
<td>31-229</td>
<td>50%</td>
</tr>
<tr>
<td>Japanese quail</td>
<td>58%</td>
<td>0%</td>
<td>41%</td>
<td>31-229</td>
<td>50%</td>
</tr>
<tr>
<td>Knob-billed duck</td>
<td>65%</td>
<td>0%</td>
<td>34%</td>
<td>31-229</td>
<td>50%</td>
</tr>
<tr>
<td>Ostrich</td>
<td>65%</td>
<td>0%</td>
<td>34%</td>
<td>31-229</td>
<td>50%</td>
</tr>
<tr>
<td>Wild chicken</td>
<td>63%</td>
<td>0%</td>
<td>36%</td>
<td>31-229</td>
<td>50%</td>
</tr>
<tr>
<td>Wild duck</td>
<td>59%</td>
<td>0%</td>
<td>40%</td>
<td>31-229</td>
<td>50%</td>
</tr>
<tr>
<td>Wild turkey</td>
<td>62%</td>
<td>0%</td>
<td>37%</td>
<td>31-229</td>
<td>50%</td>
</tr>
</tbody>
</table>
4.4.2 Predicted 3D structures vasoactive intestinal peptide 1 protein

In Table 4.3 below, the secondary and tertiary structures of vasoactive intestinal peptide 1 from different poultry species were predicted. There was some variation in the secondary structure but the tertiary structure was similar.

Table 4.3: Predicted protein 3D structures of vasoactive intestinal peptide 1. The secondary and tertiary structures of VIP1 from different poultry species are shown in the table below.

<table>
<thead>
<tr>
<th>Vasoactive intestinal peptide 1 (VIP1) protein structures</th>
<th>Helices</th>
<th>Beta sheets</th>
<th>Loops</th>
<th>Domain interval</th>
<th>Solvent accessibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild chicken 1</td>
<td>48%</td>
<td>0%</td>
<td>51%</td>
<td>89-115, 129-155</td>
<td>50%</td>
</tr>
<tr>
<td>Wild chicken 2</td>
<td>33%</td>
<td>0%</td>
<td>66%</td>
<td>89-115, 129-155</td>
<td>50%</td>
</tr>
</tbody>
</table>

4.4.3 Predicted 3D structures of vasoactive intestinal peptide receptor 1 protein

In Table 4.4 below, the secondary and tertiary structures of vasoactive intestinal peptide receptor 1 from different poultry species were predicted. The secondary structure was similar but the tertiary structure varied between the species.
Table 4.4: Predicted protein 3D structures of vasoactive intestinal peptide receptor 1. The secondary and tertiary structures of VIPR1 from different poultry species are shown in the table below.

<table>
<thead>
<tr>
<th>Vasoactive intestinal peptide receptor 1 (VIPR1) protein structures</th>
<th>Helices</th>
<th>Beta sheets</th>
<th>Loops</th>
<th>Domain interval</th>
<th>Solvent accessibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Japanese quail</td>
<td>52%</td>
<td>9%</td>
<td>38%</td>
<td>50-114, 127-373</td>
<td>50%</td>
</tr>
<tr>
<td>Wild chicken</td>
<td>52%</td>
<td>9%</td>
<td>38%</td>
<td>22-86, 99-345</td>
<td>50%</td>
</tr>
</tbody>
</table>

4.5 Protein 3D structure prediction for genes implicated in growth

4.5.1 Predicted 3D structures of growth hormone protein

In Table 4.5 below, the secondary and tertiary structures of growth hormone from different poultry species were predicted. The secondary structure was similar with minimum variation in the tertiary structure.
Table 4.5: Predicted protein 3D structures of growth hormone. The secondary and tertiary structures of GH from different poultry species are shown in the table below.

<table>
<thead>
<tr>
<th>Growth hormone (GH) protein structures</th>
<th>Helices</th>
<th>Beta sheets</th>
<th>Loops</th>
<th>Domain interval</th>
<th>Solvent accessibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Common quail</td>
<td>56%</td>
<td>0%</td>
<td>43%</td>
<td>35-214</td>
<td>50%</td>
</tr>
<tr>
<td>Greylag goose</td>
<td>56%</td>
<td>0%</td>
<td>43%</td>
<td>36-214</td>
<td>50%</td>
</tr>
<tr>
<td>Wild chicken</td>
<td>55%</td>
<td>0%</td>
<td>43%</td>
<td>35-214</td>
<td>50%</td>
</tr>
<tr>
<td>Wild duck</td>
<td>56%</td>
<td>0%</td>
<td>43%</td>
<td>35-214</td>
<td>50%</td>
</tr>
</tbody>
</table>

4.5.2 Predicted 3D structures of growth hormone receptor protein

In Table 4.6 below, the secondary and tertiary structures of growth hormone receptor from different poultry species were predicted. There were notable variations in the secondary and tertiary structures in the different species.
Table 4.6: Predicted protein 3D structure of growth hormone receptor. The secondary and tertiary structures of GHR from different poultry species is shown in the table above.

<table>
<thead>
<tr>
<th>Growth hormone receptor (GHR) protein structures</th>
<th>Helices</th>
<th>Beta sheets</th>
<th>Loops</th>
<th>Domain interval</th>
<th>Solvent accessibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Greylag goose</td>
<td>6%</td>
<td>19%</td>
<td>73%</td>
<td>29-108, 123-223, 292-590</td>
<td>50%</td>
</tr>
<tr>
<td>Southern ostrich</td>
<td>9%</td>
<td>19%</td>
<td>70%</td>
<td>24-233</td>
<td>50%</td>
</tr>
<tr>
<td>Wild chicken 1</td>
<td>9%</td>
<td>21%</td>
<td>69%</td>
<td>24-230</td>
<td>50%</td>
</tr>
<tr>
<td>Wild chicken 2</td>
<td>10%</td>
<td>21%</td>
<td>68%</td>
<td>24-230</td>
<td>50%</td>
</tr>
<tr>
<td>Wild duck</td>
<td>9%</td>
<td>19%</td>
<td>17%</td>
<td>24-233</td>
<td>50%</td>
</tr>
</tbody>
</table>

4.5.3 Predicted 3D structures of insulin-like growth factor 1 protein

In Table 4.7 below, the secondary and tertiary structures of insulin-like growth factor 1 from different poultry species were predicted. There were variations in the secondary structure but the tertiary structure was similar in all the species.
Table 4.7: Predicted protein 3D structure of insulin-like growth factor 1. The secondary and tertiary structures of IGF1 from different poultry species are shown in the table below.

<table>
<thead>
<tr>
<th>Insulin-like growth factor 1 (IGF1) protein structures</th>
<th>Helices</th>
<th>Beta sheets</th>
<th>Loops</th>
<th>Domain interval</th>
<th>Solvent accessibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Japanese quail</td>
<td>32%</td>
<td>5%</td>
<td>62%</td>
<td>49-116</td>
<td>50%</td>
</tr>
<tr>
<td>Southern ostrich</td>
<td>37%</td>
<td>9%</td>
<td>53%</td>
<td>49-116</td>
<td>50%</td>
</tr>
<tr>
<td>Wild chicken</td>
<td>29%</td>
<td>5%</td>
<td>64%</td>
<td>49-114</td>
<td>50%</td>
</tr>
<tr>
<td>Wild duck</td>
<td>32%</td>
<td>5%</td>
<td>62%</td>
<td>49-116</td>
<td>50%</td>
</tr>
<tr>
<td>Wild turkey</td>
<td>31%</td>
<td>5%</td>
<td>62%</td>
<td>49-116</td>
<td>50%</td>
</tr>
</tbody>
</table>

4.5.4 Predicted 3D structures of insulin-like growth factor 1 receptor protein

In Table 4.8 below, the secondary and tertiary structures of insulin-like growth factor 1 receptor from different poultry species were predicted. There was minimum variation in the secondary structure but notable differences in the tertiary structure of the different species.
Table 4.8: Predicted protein 3D structure of insulin-like growth factor 1 receptor.

The secondary and tertiary structures of IGF1R from different poultry species is shown in the table below.

<table>
<thead>
<tr>
<th>Insulin-like growth factor 1 receptor (IGF1R) protein structures</th>
<th>Helices</th>
<th>Beta sheets</th>
<th>Loops</th>
<th>Domain interval</th>
<th>Solvent accessibility</th>
</tr>
</thead>
<tbody>
<tr>
<td>Japanese quail</td>
<td>16%</td>
<td>24%</td>
<td>59%</td>
<td>51-161, 75-330, 230-277, 351-464, 495-605, 610-653, 835-924, 993-1269, 1000-1267</td>
<td>50%</td>
</tr>
<tr>
<td>Southern ostrich</td>
<td>14%</td>
<td>27%</td>
<td>58%</td>
<td>86-672, 673-989, 1041-1350</td>
<td>50%</td>
</tr>
<tr>
<td>Wild chicken</td>
<td>17%</td>
<td>23%</td>
<td>58%</td>
<td>86-672, 673-989, 932-972, 1041-1350</td>
<td>50%</td>
</tr>
<tr>
<td>Wild duck</td>
<td>16%</td>
<td>24%</td>
<td>59%</td>
<td>18-606, 606-922, 932-972, 973-1283</td>
<td>50%</td>
</tr>
</tbody>
</table>
4.6 Predicted 3D structure comparison of proteins implicated in egg production

4.6.1 Comparison of predicted 3D prolactin structures

The DeepAlign server was used to obtain the 3D structure alignment of the prolactin structures (Figure 4.15) using their sequences.

The 3D structural comparison of common peafowl, common pheasant, common quail, helmeted guinea fowl, Japanese quail, knob-billed, ostrich, wild chicken, wild duck prolactin produced a template modeling (TM) score of 0.901. This showed that these structures were highly similar and they were likely to share a similar fold.
Lali: 117  RMSD: 0.79  uGDT(GDT): 116(81)  TMscore: 0.901

Figure 4.15: Predicted 3D structure comparison of peafowl, common pheasant, common quail, helmeted guinea fowl, Japanese quail, knob-billed, ostrich, wild chicken, and wild duck prolactin. The scores represent 3D similarity scores. Template modeling (TM) score values of >0.5 meant that the structures were similar.

4.6.2 Comparison of predicted 3D vasoactive intestinal peptide 1 structures

On sending the sequences of the vasoactive intestinal peptide 1 (VIP1) sequences to the DeepAlign server, a structural alignment of the structures was obtained as shown in Figure 4.16 below.

The 3D structures of wild chicken 1 and wild chicken 2 VIP1 were not similar and therefore do not share a similar fold as their structural alignment gave a template modeling (TM) score of 0.199.
Lali: 57  RMSD: 5.19  UGDT(GDT): 26(16)  TMscore: 0.199

Figure 4.16: Predicted 3D structure comparison of wild chicken 1 and wild chicken 2 vasoactive intestinal peptide 1. The 3D similarity scores were used to evaluate similarity of scores. Template modeling (TM) score of <0.5 meant that the structures were not similar.

4.6.3 Comparison of predicted 3D vasoactive intestinal peptide receptor 1 structures

The DeepAlign server produced the structural alignment of the vasoactive intestinal peptide receptor 1 (VIPR1) structures shown in Figure 4.17 below.

From the 3D structural comparison of Japanese quail and wild chicken VIPR1, it was evident that these structures were highly similar and shared a fold as they had a template modeling (TM) score of 0.65.
Lali: 288  RMSD: 1.96  uGDT(GDT): 270(66)  TMscore: 0.675

Figure 4.17: Predicted 3D structure comparison of Japanese quail and wild chicken vasoactive intestinal peptide receptor 1. The similarity scores shown were for evaluating similarity. Template modeling (TM) score of >0.5 meant that the structures were similar.

4.7 Predicted 3D structure comparison of proteins implicated in growth

4.7.1 Comparison of predicted 3D growth hormone structures

The different sequences of growth hormone (GH) were sent to DeepAlign for structural comparison. Figure 4.18 below shows the structural superimposition.

The template modeling (TM) score of the 3D structural comparison of common quail, greylag goose, wild chicken, and wild duck GH was 0.919. These structures were very likely to share a fold and they were similar.
Figure 4.18: Predicted 3D structure comparison of common quail, greylag goose, wild chicken, and wild duck growth hormone. The scores represent 3D similarity scores. The scores were used in evaluating similarity. Template modeling (TM) score of >0.5 meant that the structures were similar.

4.7.2 Comparison of predicted 3D growth hormone receptor structures

DeepAlign was used to compare the different growth hormone receptor (GHR) structures. Figure 4.19 below shows the structural alignment of the structures. On the alignment of the 3D structures of greylag goose, southern ostrich, wild chicken and wild duck GHR, the template modeling (TM) score was 0.880. This evidently showed that these structures were highly similar and they were likely to share a fold.
Figure 4.19: Predicted 3D structure comparison of greylag goose, Southern ostrich, wild chicken, and wild duck growth hormone receptor. The scores were used in evaluating 3D similarity. The structures were similar as they had a template modeling (TM) score of > 0.5.

4.7.3 Comparison of predicted 3D insulin-like growth factor 1 structures

The sequences of the different insulin-like growth factor 1 (IGF1) structures were sent to DeepAlign server to obtain the structural alignment shown below (Figure 4.20). The 3D structural comparison of Japanese quail, southern ostrich, wild chicken, and wild duck IGF1 had a template modeling (TM) score of 0.615. These structures were thus similar and they were likely to share a fold.
Lali: 61  RMSD: 3.35  uGDT(GDT): 43(44)  TMscore:0.615

Figure 4.20: Predicted 3D structure comparison of Japanese quail, southern ostrich, wild chicken, and wild duck insulin-like growth factor 1. The TMscore was used to evaluate 3D structural similarity with template modeling (TM) score of >0.5 indicating similarity of the structures.

4.7.4 Comparison of predicted 3D insulin-like growth factor 1 receptor structures

DeepAlign was used to compare the different insulin-like growth factor 1 receptor (IGF1R) structures. Figure 4.21 below shows the structural alignment of the structures. The 3D structures of Japanese quail, southern ostrich, wild chicken, and wild duck IGF1R were not similar and were therefore less likely to share a fold. This was evident from the structural alignment that gave a template modeling (TM) score of 0.481.
Lali: 597 RMSD: 2.77 uGDT(GDT): 522(42) TMscore: 0.481

Figure 4.21: Predicted 3D structure comparison of Japanese quail, southern ostrich, wild chicken, and wild duck insulin-like growth factor 1 receptor. The template modeling (TM) score was used to evaluate 3D structural similarity of the structures. The structures were not similar as they had a TM score of <0.5.
CHAPTER FIVE
DISCUSSION

This study supported the hypothesis that there is variation at the selected candidate genes involved in growth and the production of eggs in poultry as a result of Natural selection.

5.1 Analysis of signatures of selection in genes for egg production

5.1.1 Prolactin

In prolactin, the codon substitution models identified positive selection on the lineage leading to domesticated birds (Figure 4.5). This implies that domestication has played a role in shaping the prolactin gene in poultry. Domestication causes a rapid physical, physiological, and behavioral change (Cheng, 2010).

The domestication process is cumulative and happens over many generations (Cheng, 2010). During domestication, the animals which have been selected adapt to the new human environments by integrating the changes in their genetic makeup, psychology, physiology, and behavior (Cheng, 2010). Chickens were domesticated 5,000-10,000 years ago through the taming of a few jungle fowls (Underhill, 1997). Only 4-6 eggs are laid by the jungle fowl in a year. In contrast, a commercial laying hen lays eggs exceeding 300 in a year. This is because commercial layers have been subjected to artificial selection for higher production (Cheng, 2010). Prolactin has many and diverse roles but in birds, it is implicated in the onset and maintenance of broodiness (Wilkanowska et al., 2014). Broodiness causes avian egg production to be low (Usman
et al., 2014). Through domestication, the prolactin gene has been subjected to pressures to meet human demands. The other species in this study are not domesticated. This is probably the reason these species are under a different selection pressure.

The ratio of non-synonymous mutations to synonymous mutations was greater than 1 implying that on the poultry lineage there was an increase egg production.

5.1.2 Vasoactive intestinal peptide receptor 1

All lineages in vasoactive intestinal peptide receptor 1 (VIPR1) had a dN/dS<1 which was an indication that this protein is under purifying selection (Figure 4.4). This is an indication that VIPR1 has endured strong selection pressures during breeding. Zhou et al., (2008) also concluded that VIPR1 had endured strong selection pressures in chickens during breeding.

Broodiness is a trait which is polygenic (Romanov et al., 2002). The role of prolactin is initiating and maintaining incubation in birds (Zhou et al., 2008). At the hypothalamus, the secretion of prolactin is controlled by vasoactive intestinal peptide 1 while at the pituitary level the secretion is under the control of VIP receptor 1. Thus VIPR1 may be a candidate gene for broodiness in avian species (Zhou et al., 2008). In a study done by Zhou et al., 2008, there was variation in VIPR1 in different chicken populations. These variations were associated with broodiness. Broodiness has been a challenge to eradicate even with domestication and artificial selection (Zhou et al., 2008).
Purifying selection has also been detected in several protein-coding sequences in HIV-1. Many of these sequences are involved in regulating gene expression in HIV. Some of the sequences include 3’ Long terminal repeat (LTR) which encode the Nef protein (Das et al., 1998). It also contains some regulatory elements. For example in the pol gene, there is an intragenic nuclease hypersensitive region which has a role in regulating gene expression.

Additionally, in the env gene is the rev-responsive element (RRE) which transports unspliced or RNA that partially spliced from the nucleus of the infected cell to the cytoplasm (Ngandu et al., 2008). These regions are important for viral viability hence are under purifying selection to preserve their functions (Ngandu et al., 2008).

Likewise in VIPR1 as in HIV-1, purifying selection acts to preserve the function. As a result of purifying selection, the function of VIPR1 on prolactin secretion is maintained. Although broodiness leads to a decrease in egg production, it is an important condition when one wants to use the natural process to produce chicks (FAO, 2010).

5.2 Analysis of signatures of selection in genes for growth

5.2.1 Growth hormone

In this study, the evolution of growth hormone (GH) is different in the various taxonomic groups. Positive selection was identified in the lineage leading to ruminants and artiodactyls, while purifying selection was detected in the other lineages (Figure
4.10). This is an indication that the gene has had different patterns of evolution in different species.

There are two features of interest in the molecular evolution of GH: (i) Variation in evolutionary rate (ii) Multiple duplications of the gene (Forsyth and Wallis, 2002). The sequence of GH is very conserved in mammals (Forsyth and Wallis, 2002). The differences in the sequences of mammalian species occurs at a few amino acid sites. The rate of GH evolution has been very slow in eutherians with the rate having increased 25-50 times during the evolution of primates and during the evolution of artiodactyls (Forsyth and Wallis, 2002).

The evolution of GH in mammals has been slow with most changes (~85%) having occurred during the periods of rapid change. As a result of the bursts of evolution, GH shows specie specificity. The specificity is due to single substitutions in the GH and its receptor (Forsyth and Wallis, 2002). For example GH from mammals differs from that of non-primate mammals at 35% of the residues. This is reflected in the multiple sequence alignment of GH in Figure 4.6. Further, the sequences of the different artiodactyls differ at a few amino acid sites while their sequences differ at multiple amino acid sites from other species. Hence GH from non-primates is non-functional in mammals as it binds very poorly to the receptor. The specie specificity correlates with differences in the binding of the growth hormone receptor (GHR) and the biological actions (Buggiotti and Primmer, 2006). These bursts of change occurred for a short period after which the rate fell back to the slower rate. Although there are three possible causes for the variation in the evolution rate: (a) The variability being an artifact because
of comparing non-orthologous proteins (b) The variability reflected periods of evolution of GH when the function was lost so the sequence change was caused by neutral mutations, and (c) The bursts of change were driven by positive natural selection. The third explanation was found to be the most compelling. One argument supporting the third explanation was that the episodes of rapid change occurred on the mature protein. In the mature GH protein, the rate of non-synonymous to synonymous mutation rose from 0.02 (basal) to 1.37 (artiodactyls) and 1.27 (primates) (Buggiotti and Primmer, 2006). This was evidence that the driving force was positive selection. In my study, the ratio of the non-synonymous to synonymous changes in artiodactyls was 4.98. It is evident that during this period there was accelerated rate of evolution which was driven by positive selection (Forsyth and Wallis, 2002).

The pattern of evolution in the avian growth hormone is different from mammals because of the difference in function (Buggiotti and Primmer, 2006). In avian species, the growth hormone is involved in secondary features such as egg production, aging and reproduction (Zhao et al., 2004). Buggioti and Primmer (2006) identified many amino acids in avian growth hormone to be under purifying selection and concluded that growth hormone is under purifying selection following the bursts of rapid changes.

5.2.2 Growth hormone receptor

In this study, growth hormone was found to be under purifying selection in all lineages (Figure 4.12). However, previous studies have found that variable evolution rates in growth hormone (GH) correlate with changes in evolution rate of growth hormone
receptor (GHR). This is probably because in the receptor, the pattern is different for the extracellular and intracellular domains. In the extracellular domain, the evolution was slow followed by bursts of change during the evolution of artiodactyls, primates and rodents (Zhenfang et al., 2005). The two occasions are equivalent to the evolution that occurred in GH. In the intracellular domain, the evolution rate was high but constant. Acceleration was only found in the case of rodents. Previous studies also found the lineages leading to artiodactyls, primates, and rodents to be under positive selection. (Zhenfang et al., 2005).

5.2.3 Insulin-like growth factor 1 receptor

Maximum Likelihood Models identified one amino acid site in IGF1R to be positively selected. This site is on the L2 domain. This domain is important in binding IGF1 to bring about growth and development (Ward et al., 2000).

There are only a few amino acid sites responsible for the positive selection acting on a gene (Yokoyama and Yokoyama (1996). This is because different amino acid sites are under different selective pressure because the sites have different biological functions and structural roles.

The ectodomains of insulin-like growth factor 1 receptor (IGF1R) has two homologous domains (L1 residues 1-150 and L2 residues 300-460) which are on the N-terminal (Ward et al., 2000). These are separated by a cysteine-rich region. This region covers two-thirds the IGF1R molecule, and makes contact with the L1 domain but a slight contact with the L2 domain. This creates a space in the middle which is sufficient to
accomodate the ligands (Ward et al., 2000). Insulin-like growth factors are involved in prenatal and postnatal growth and development (Delafontaine et al., 2004). In avian species, growth hormone binds growth hormone receptor indirectly on the liver membrane. This then activates insulin-like growth factor 1 (IGF1) secretion. When IGF1R binds IGF1, the physiological response to this ligand is initiated in vivo (LeRoith et al., 1995). This stimulates growth through differentiation and proliferation of the bone and muscle cells (Kuhn et al., 2002). The location of the positively selected site on this domain means that this receptor-ligand binding will be enhanced and so will the growth. Domestication probably played a role in the positive selection whereby growth was selected for.

Maximum likelihood methods have also been successfully used to identify amino acid sites subject to positive selection in mammalian alpha defensins. Defensins are antimicrobial peptides which are cationic in nature (Lynn et al., 2004). They are broad spectrum acting against fungi, enveloped viruses and both gram-positive and gram-negative bacteria. Their mode of action is by disruption of microbes’ membrane integrity (Lynn et al., 2004). Alpha defensins are variable in their action towards different pathogens. Some microbes are sensitive to some defensins and not others. This can be caused by small changes in the primary structure of the potent defensins. These sites have been found to be under positive selection. Defensins are encoded first as prepropeptides which undergo proteolytic cleaving. The prorregion is cytoprotective and is inactive. On cleaving they release the C-terminal antimicrobial peptide which is active. The sites under selection are on the mature antimicrobial peptide region.
As mammals evolved to occupy new environments, it is likely that they encountered new different pathogens. The pathogens have evolved mechanisms to escape the antimicrobial molecules. This ongoing ‘arms race’ with pathogens may be the significant force driving the evolution of alpha defensins (Lynn et al., 2004). Positive selection would be of selective advantage to the antimicrobial peptides as they would have sensitivities to new infectious agents.

As with the mammalian alpha defensins, positive selection on the IGF1R also occurred on functionally important regions.

One of the objectives of studies such as these is the expectation that there is an association between these regions under selection and phenotypes of interest (Nielsen et al., 2007).

In a study done by Ojeda et al., 2008, he identified a signature of section in the IGF2 gene in three pig breeds (Pietrain, Hampshire, and Duroc). These breeds are commonly used to improve growth and meat leanness in pigs.

5.3 Prediction of secondary and tertiary structures for egg production proteins

5.3.1 Prolactin

In this study, prolactin was predicted as an alpha protein with 4 long alpha helices arranged in an anti-parallel fashion (Table 4.2, Appendix 8). It also has more helices than loops. This is consistent with nuclear magnetic resonance experimental studies.
done before that described the secondary structure of prolactin to be made of 50% Helices and the rest being Loops (Freeman et al., 2000). Teilum et al., (2005) also found the secondary structure of human prolactin to be composed of four alpha helices in his studies. It has been proposed that the 4 helices of prolactin are packed in a similar way to the porcine growth hormone. The structural similarity is based on the fact that these two genes arose from a common ancestor through gene duplication followed by divergent evolution (Forsyth and Wallis, 2002).

In this study, I found prolactin protein to have one domain which has a predicted role in receptor binding. Prolactin carries out its biological functions through binding with the prolactin receptor in a one-to-two complex (Freeman et al., 2000). This is regulated by tertiary structures (Teilum et al., 2005).

Prolactin binds the first subunit of the receptor with high-affinity followed by association with the second subunit which has lower affinity. This leads to an intracellular cross-phosphorylation cascade. It has been hypothesized that binding to the first subunit causes a conformational change in the hormone leading to an increased affinity for the second subunit (Freeman et al., 2000). When the hormone is in the unbound state, subunit two is considered incompetent. In prolactin, helix 1, helix 4, and the second half of loop 1 contain residues important for binding site 1 (Freeman et al., 2000).

The binding site is concave with residues that are aromatic and have large hydrophobic and polar residues which are positively charged (Clackson and Wells, 1995). These
complement the tryptophan and negatively charged residues of the hot spots of the receptor (Clackson and Wells, 1995).

The second receptor binding site in human prolactin is outlined by a flat hydrophobic channel which is formed between helix 1 and helix 3 (Clackson and Wells, 1995). When site 1 binds a receptor, the affinity of binding a receptor by site 2 is significantly increased approaching that of site 1 (Sivaprasad et al., 2004).

5.3.2 Vasoactive intestinal peptide 1 and vasoactive intestinal peptide receptor 1

From this study, vasoactive intestinal peptide 1 (VIP1) structures were found to be composed mainly of alpha helices (Table 4.3, Appendix 9). Couvineau et al., (2012) carried out nuclear magnetic resonance (NMR) studies on human VIP1 that also revealed that the structure has more of the alpha helices except sequence 1-5 of the N terminal which lacks a defined structure when in solution.

The biological functions of VIP1 are triggered on interaction with the vasoactive intestinal peptide receptor 1 (VIPR1) (Zhou et al., 2008). The N terminal 1-5 plays an important role of receptor activation mainly through the adenylyl cyclase activation.

Studies have revealed that residues in position 0, 6, 22, 24, or 28 are in contact with residues glutamine\textsuperscript{135}, aspartate\textsuperscript{107}, glycine\textsuperscript{116}, cysteine\textsuperscript{122}, and lysine\textsuperscript{127} respectively in the N-ted of the VIPR1. During binding, the central and C-terminal residues of VIP1 forms contact with N-ted (Couvineau et al., 2012). The N-terminus of VIP1 is in contact with the transmembrane domains and the extracellular loop of the receptor.
Histidine$^{112}$, leucine$^{131}$, and glutamine$^{134}$ residues present in the N-ted are among the residues involved in the VIP1-receptor binding affinity.

These three play a role in the interaction of C-terminal and the central parts, and N-ted of VIP1. The binding interaction of lysine$^{143}$, threonine$^{144}$, and threonine$^{147}$ residues of VIPR1 and histidine$^1$ of VIP1 is an important step in the adenylyl cyclase activation (Couvineau et al., 1984).

5.4 Prediction of secondary and tertiary structures for growth proteins

5.4.1 Growth hormone and growth hormone receptor

In this study, I used Raptor X to predict the structure of growth hormone (GH). I found growth hormone to have a higher proportion of alpha helix compared to loops (Table 4.5, Appendix 11). It has a four-helix bundle which is consistent with physicochemical studies which found growth hormone to consist of a high proportion of alpha helix. This is consistent with studies done by Forsyth and Wallis, (2002) who determined the 3D structure of pig GH by X-ray to consist of a four-helix bundle. The structure of a protein is valuable in determining the functions of a protein like enzymatic activity or the interaction with other proteins (Bairoch, 2000).

GH performs many physiological actions among which is controlling growth by regulating IGF1 concentrations, stimulating lipolysis, providing FFAs, and glycerol as substrates for energy metabolism and also inhibiting insulin-induced suppression of hepatic gluconeogenesis (Clemmons, 2004). GH binds to GHR leading to growth and
metabolism (Varvio et al., 2008). From the computational structure prediction, GH has one domain (Somatotropin-like) which runs from residues 35-214. The function of this domain is in binding the receptor.

Helical cytokines like GH and their cognate receptors have related structures and they are likely to have a similar binding mechanism (Wells, 1996). These hormones have at least two sites for binding and oligomerizing their receptors (Wells, 1996). The GH binds and dimerizes two receptors using two different sites (Site 1 and Site 2) (Wells, 1996). GH reacts with the first receptor by using Site 1 (found at the helix IV) then binds to the second receptor using Site 2. Electrostatic and hydrophobic interactions play a significant role in the formation of the signal transduction complex (Demchuk et al., 1994). Although hydrophobic contacts are crucial in binding, and polar residues appear less important for affinity, polar residues are important for solubility and specificity in the binding interaction (Wells, 1996). Electrostatic properties of molecules are also important in the specificity of the hormone-receptor binding (Demchuk et al., 1994).

5.4.2 Insulin-like growth factor 1 and Insulin-like growth factor 1 receptor

The predicted secondary and 3D structure of IGF1R is composed of helices, beta sheets, and loops (Table 4.8, Appendix 14). The following domains were in the IGF1R structure: receptor L domain, Furin-like domain, furin-like repeats, receptor L domain, Fibronectin type 3, fibronectin type 3, fibronectin type 3, PTKc-InsR-like, and Protein tyrosine kinase.
IGF1R binds its ligand (IGF1) with high affinity consequently initiating the physiological response to the ligand (LeRoith et al., 1995). Comparative sequence analysis followed by confirmation through X-ray analysis has also identified the domain boundaries of the IGF1R. This has identified the N terminal half of IGF1R ectodomain to contain two homologous domains (L1 and L2) separated by a Cys-rich region (Cysteine 152-Cysteine 298) (Ward et al., 2000). The C terminal half of IGF1R ectodomains is made up of 3 fibronectin type III (FnIII) domains, with the second containing a large insert domain of ~120-130 residues (Ward et al., 2000). Intracellularly, the IGF1R has a tyrosine kinase catalytic domain (973-1229) which is flaunted by two regulatory regions: a juxtamembrane region (930-972), a C-tail with 108 residues, and residues 1230-1337 which contain the phosphotyrosine binding sites for signalling molecules (Ward et al., 2000).

Alanine scanning studies that were done on the IGF1R ectodomain indicated that L1 and CR domains and the carboxy-terminal peptide in the alpha subunit are components of the binding site of IGF1. Residues 131-315 in the Cys-rich plus flanking regions of L1 and L2 were found to be significant determinants of IGF1 binding. 10 residues in the L1 domain and 4 residues in the Cys-rich region which are implicated in IGF1 binding are located in two regions which are discontinuous (Ward et al., 2000).

The first site has amino acids asparagine11, tyrosine28, histidine30, leucine33, leucine56, phenylalanine58, arginine59, and phenylalanine90. These are distributed across the first four repeats of the L1 domain. The second functional epitope is made of Tryptophan79 which is situated in a bulge in the fourth turn of the L1 beta helix and
Arginine240, Phenylalanine241, Glutamate242, and Phenylalanine251 in the CR domain. The putative IGF1R binding site has residues which are acidic found in the corner where the Cys-rich region separates from L1. Other acidic residues of the receptor are inside the Cys-rich domain and the loop residues 255-263 which extend from module 6. Electrostatics play an important part in IGF1 binding. The C-region binds to the acidic patch of the Cys-rich region which is near L1 and the acidic patch which is in the other side of IGF1 directed to a small patch with basic residues (residues 307-310) on the L2 terminal end (Ward et al., 2000).

There has not been done an extensive mapping of structure-function relationships of the IGF1 and IGF1R molecules as of insulin (Meyts and Whittaker, 2002).

Mutagenesis studies of IGF1 have shown that Alanine8, Aspartate12, Phenylalanine23, and Tyrosine24 in the B-domain, Tyrosine31, Arginine36, and Arginine37 in the C-peptide and Methionine59, Tyrosine60, and Alanine62 in the A-domain are important in the high-affinity binding to the IGF1R (Meyts and Whittaker, 2002).

5.5 Predicted 3D structure comparison of prolactin, vasoactive intestinal peptide receptor 1, growth hormone, growth hormone receptor, and insulin-like growth factor 1 proteins

On doing the structure comparison of the different prolactin structures, the template modeling score (TM) value of >0.5 showed that the predicted structures of common peafowl, common pheasant, common quail, helmeted guinea fowl, Japanese quail, knobby-billed duck, ostrich, wild chicken, and wild duck prolactin were highly similar. Structure
alignments with a TM value of > 0.5 are mostly in the same fold while those with TM<0.5 are not in the same fold (Xu and Zhang, 2010). Structural similarity implies that the prolactin protein may have similar functions in the different species (Xu and Zhang, 2010).

The structural alignment of Japanese quail and wild chicken vasoactive intestinal peptide receptor 1 (VIPR1) structures also showed high similarity as shown by TM value of 0.675.

The structural comparison of common quail, greylag goose, wild chicken, and wild duck growth hormone (GH) gave a TM>0.5 showing that these structures were similar. This is an indication that there has been minimal variation in the overall fold of the different species’ GH. Although there has not been a direct determination of the GH structure from other species, a homology model for the sheep growth hormone suggested that there is minimal variation in the overall fold despite the significant difference in the sequences of different species (Forsyth and Wallis, 2002).

A structural alignment of greylag goose, southern ostrich, wild chicken, and wild duck growth hormone receptor (GHR) showed that these structures were highly similar as shown by TM value of 0.880.

The 3D structural comparison of Japanese quail, southern ostrich, wild chicken, and wild duck insulin-like growth factor 1 (IGF1) showed that these structures were highly similar as shown by TM value of 0.615.
Comparative analysis of protein structure is important in enabling the study of evolutionary and functional relationships between proteins (Balasubramanian et al., 2012). Notably, the sequences of the different species of prolactin, VIPR1, GH, GHR, and IGF1 which had similar structures had variations in some sites of the sequences. Despite these variations, their structures were similar to each other. This is an indication that evolution conserves the structure of a protein more than the sequence (Balasubramanian et al., 2012). However, there are cases where similar or identical sequences have different 3D structures (Kinch and Grishin, 2002). The 3D structure determines the biological role of a protein (Bairoch, 2000).

In addition to the different proteins having similar functions in different species, the proteins can probably bind to each others’ receptors and carry out their biological functions. The similarity in structure may also mean that the proteins from the different species share a common ancestor (Kinch and Grishin, 2002).

This is probably so as is the conclusion from a study done by Balasubramanian et al., (2012). In their study they compared the structures of three Superoxide dismutases from Oryza sativa: manganese superoxide dismutase (Mn SoD), iron superoxide dismutase (Fe SoD), and copper-zinc superoxide dismutase (Cu-Zn SoD). Mn SoD and Fe SoD were found to have similar structures with corresponding similar functions while Cu-Zn SoD had a different tertiary structure from the two. Cu-Zn SoD was also found to have evolved independently from Mn SoD and Fe SoD which were closely related as they had a common phylogenetic origin. From this study, the two enzymes with similar structures can also bind each other’s ligands (Balasubramanian et al., 2012).
5.6 Predicted 3D structure comparison of vasoactive intestinal peptide 1 and insulin-like growth factor 1 receptor proteins

The predicted 3D structures of Japanese quail, southern ostrich, wild chicken, and wild duck insulin-like growth factor 1 receptor (IGF1R) had differing structures. For vasoactive intestinal peptide 1 (VIP1), the structural comparison of the two wild chicken sequences found these to be different. On comparison of the structures, the template modeling score (TM) values were less than 0.5. This suggests that the genes in the different species have different functions.

The structures of proteins evolve through different mechanisms, often including gene duplication then mutation and selection (Kinch and Grishin, 2002). A combination of different mechanisms can bring about mutation at the structural level. Most of these mechanisms are insertion/deletion/substitution of secondary structure, circular permutation, hairpin swap, and strand invasion (Kinch and Grishin, 2002).

The sequences of the two wild chicken VIP1 structures that were compared had many indels. The indels present in the sequences analyzed may have accumulated over time resulting to the structural differences and subsequently a difference in functions (Kinch and Grishin, 2002).

It could also be possible that for the species that were compared, there is species-specificity whereby the protein cannot bind to the ligand of another specie’s protein and vice versa. The variation of 3D leading to species-specificity is in support of a hypothesis from a study done by Govindaraj et al., (2011) on toll-like receptor 8.
(TLR8). In their study, they discovered that there was specificity of ligand recognition between rodent and non-rodent TLR8 (Govindaraj et al., 2011). They hypothesised that structural differences between rodent and non-rodent TLR8s lead to this species-specificity. TLR8 plays a vital role in innate immune response by recognizing ssRNA derived from viruses and small molecular weight ligands respectively (Govindaraj et al., 2011). The mechanism of antiviral activity occurs through the binding of TLR8 through its ligand binding cavity with the ligand (Govindaraj et al., 2011). In their study, they discovered an undefined region LRR14-15 (residues 438-442) which varied in 3D structures. Although this region is not involved in ligand interaction, it is required for TLR8 activation. Despite both non-rodent and rodent TLR8s being able to bind their ligands, rodent TLR8 have signalling initiators which are weaker compared to those of non-rodent TLR8. Non-rodent ligands can bind with rodent TLR8 although the signalling mechanism is not activated.

Therefore, they hypothesised that variation in the 3D structure caused the species-specificity in ligand recognition as the LRR14-15 is in close proximity to the binding region (Govindaraj et al., 2011).

Although structural similarity may be used in elucidating function, there are cases whereby similar structures may have a divergence in function and non-similar structures have a convergence of the functions (Kinch and Grishin, 2002).
CHAPTER SIX

CONCLUSION AND RECOMMENDATIONS

6.1 Conclusion

The null hypothesis that there are no signatures of selection in candidate genes for egg production and growth in poultry was rejected by the findings in this study.

The findings supported the alternative hypothesis that there are signatures of selection in candidate genes for egg production and growth in poultry.

The general objective has also been met as I identified molecular signatures of selection at candidate genes involved in growth and egg production in poultry through the computational approach.

- Analysis of molecular signatures of selection at production genes for eggs

Positive selection was identified in the prolactin gene in the lineage leading to domestic birds (Figure 4.5). Human beings have attempted to increase egg production through domestication activities (Cheng, 2010). This has been a major force in shaping the DNA sequence of poultry prolactin.

On the other hand, purifying selection was identified in all lineages of vasoactive intestinal peptide receptor 1 (VIPR1) gene (Figure 4.4). When synonymous mutations occur, the sequence of the amino acid is not altered. It was long thought that these mutations had no effect on the function of the protein encoded or advantageous to the organism. However, the genes with this kind of mutation are important, highly
expressed, and are involved in key pathways (Ngandu et al., 2008). Therefore, VIPR1 could be an important gene in poultry egg production (Zhou et al., 2008).

- **Analysis of molecular signatures of selection at production genes for growth**

  In growth hormone, positive selection was detected in ruminants and artiodactyls lineages (Figure 4.10). Growth is an important economic trait in Livestock production which is influenced by various genes. Artificial selection of growth characteristics has been practiced mainly in ruminants and artiodactyls. Hence, it has been the major driving force in shaping the growth characteristics of these species.

  IGF1R gene has one amino site which is positively selected. IGF1R binds IGF1 leading to pre-natal and post-natal growth and development (Delafontaine et al., 2004). As the positive selected site is on the L2 domain which is important for binding IGF1, the different species have enhanced growth. This may be an important marker in breeding livestock with improved growth.

- **Prediction of secondary and tertiary structures of production genes for eggs and growth**

  Protein structure prediction is important in understanding the effects of selection on the protein function (Bairoch, 2000). It is useful in designing in vitro studies for studying the effects of selection because the mechanism that determines the function is well known.
• **Comparison of the structures of production genes for eggs and growth**

For prolactin, vasoactive intestinal peptide receptor 1 (VIPR1), growth hormone, growth hormone receptor, and insulin-like growth factor 1 (IGF1), the structures of the species compared were similar. Therefore, for each gene that showed similarity, the function of the gene is conserved across the species.

The same breeding program formulated for one species may be applied in all the other species which have similarity in structure. This will save on resources that would have been used for formulating individual breeding programs. Unfortunately for insulin-like growth factor 1 receptor (IGF1R) which had a positively selected site, the species had different 3D structures hence breeding programs for improving growth should be formulated for individual species.
6.2 Recommendations

- Experimental studies in genes under purifying selection should be carried out to determine the sites under selection and the roles they play on the function of the protein.
- Breeders should perform *in vitro* and *in vivo* studies to ascertain the effect of the amino acid selection on Insulin-like growth factor 1 receptor (IGF1R) on the phenotype. IGF1R is a potential marker for improving growth in poultry.
- *In vivo* and *in vitro* studies should be performed to determine whether genes of the different species whose predicted 3D structures are similar share the same function. Structures that are not similar could still share a function. Therefore, studies should be carried out to find out if the vasoactive intestinal peptide 1 and insulin-like growth factor 1 receptor genes with dissimilar structures could have the same function in the different species.
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*RaptorX: a Web Portal for Protein Structure and Function Prediction.*


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APPENDICES

Appendix 1: Homologues for prolactin gene.

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Appendix 2: Homologues for vasoactive intestinal peptide 1 (VIP1) gene.

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**Appendix 3: Homologues for vasoactive intestinal peptide receptor 1 (VIPR1) gene.**

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Appendix 4: Homologues for growth hormone (GH) gene.

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<th>GH gene</th>
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<th>Accession Number</th>
<th>Expectation value</th>
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<td>Common quail</td>
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<td>Rock dove</td>
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<td>Wild canary1</td>
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15. Gold collared manakin | XP 003686878.1 | 5e-33  
16. Wild canary2 | XP 002435767.0 | 1e-25  
17. Ground tit | XP 005524208.1 | 2e-29  
18. Collared flycatcher | XP 005059628.1 | 1e-25  
19. White throated sparrow2 | XP 006576477.8 | 4e-27  
20. Zebrafinch2 | XP 002187284.1 | 3e-28  
21. European rabbit | XP 007636368.7 | 2e-28  
22. Wild Bactrian camel | XP 006177464.1 | 4e-23  
23. Zebu | XP 001122366.7 | 2e-23  
24. Chinese sturgeon | XP 006356356.0 | 2e-23  
25. Russian sturgeon | ABK74674.6 | 9e-24  
26. Sheep1 | ABK59498.1 | 2e-23  
27. Sheep2 | ABO21737.1 | 5e-23  
28. Cattle | ABK67647.0 | 5e-23  
29. Goat | ADX66303.1 | 9e-24

**Appendix 5: Homologues for growth hormone receptor (GHR) gene.**

<table>
<thead>
<tr>
<th>GHR gene</th>
<th>Species</th>
<th>Accession Number</th>
<th>Expectation value</th>
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<tbody>
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116
<p>| 2. Chicken breed Yunnan Wuding | KC242241.1 | 0 |
| 3. Commercial chicken | M74057 | 0 |
| 4. Wild chicken1 | AGG38006.1 | 0.0 |
| 5. Wild chicken2 | NP001001293.1 | 0.0 |
| 6. Dalmatian pelican | XP 0876532.0 | 0.0 |
| 7. Greylag goose | ACY38605.1 | 0.0 |
| 8. Saker falcon | XP 005433804.1 | 0.0 |
| 9. Peregrine falcon | XP 005242027.1 | 0.0 |
| 10. Wild duck | ACT 20710.1 | 0.0 |
| 11. Knob billed duck | ACT 20711.1 | 0.0 |
| 12. Rock dove | EMC76968.1 | 0.0 |
| 13. Southern ostrich | EMC9876.0 | 0.0 |
| 14. Anna’s humming bird | ACY2165.0 | 0.0 |
| 15. Golden collared manakin | ACY3476.0 | 0.0 |
| 16. Red throated loon | ACY5876.1 | 0.0 |
| 17. White throated sparrow | XP 005493766.1 | 0.0 |
| 18. Zebra finch | XP 002193695.2 | 0.0 |
| 19. Medium groundfinch | XP 005422066.1 | 0.0 |
| 20. Wild canary | XP 3454267876.0 | 0.0 |
| 21. The great cormorant | NP6473676787.0 | 0.0 |
| 22. Collared flycatcher | AA018173.1 | 0.0 |</p>
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<tr>
<th>Species</th>
<th>Accession Number</th>
<th>Expectation value</th>
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<td>25. European rabbit</td>
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<td>26. Wild Bactrian camel</td>
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<td>27. Alpaca</td>
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<td>28. Carolina anole1</td>
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<td>29. Carolina anole2</td>
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<td>31. Zebu</td>
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Appendix 6: Homologues for insulin-like growth factor 1 (IGF1) gene.

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<th>IGF1 gene</th>
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Appendix 7: Homologues for insulin-like growth factor 1 receptor (IGF1R) gene.

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<td>9. Medium ground finch2</td>
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<td>18. Atlantic canary</td>
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Appendix 8: Predicted protein 3D structures of prolactin.
KEY:

a= Predicted 3D structure of common peafowl prolactin.
b= Predicted 3D structure of common pheasant prolactin.
c= Predicted 3D structure of common quail prolactin.
d= Predicted 3D structure of helmeted guinea fowl prolactin.
e= Predicted 3D structure of wild turkey prolactin.
f= Predicted 3D structure of Japanese quail prolactin.
g= Predicted 3D structure of knob-billed duck prolactin.
h= Predicted 3D structure of ostrich prolactin.
i= Predicted 3D structure of wild duck prolactin.
j= Predicted 3D structure of wild chicken prolactin.

Appendix 9: Predicted protein 3D structures of vasoactive intestinal peptide 1 (VIP1).

KEY:

a= Predicted 3D structure of wild chicken 1 VIP1.
b= Predicted 3D structure of wild chicken 2 VIP1.
Appendix 10: Predicted protein 3D structures of vasoactive intestinal peptide receptor 1 (VIPR1).

**KEY:**

a= Predicted 3D structure of Japanese quail VIPR1.

b= Predicted 3D structure of wild chicken VIPR1.
Appendix 11: Predicted protein 3D structures of growth hormone (GH).

KEY:

a= Predicted 3D structure of common quail GH.

b= Predicted 3D structure of wild chicken GH.

c= Predicted 3D structure of greylag goose GH.

d= Predicted 3D structure of wild duck GH.
Appendix 12: Predicted 3D structure for growth hormone receptor (GHR).

<table>
<thead>
<tr>
<th>a</th>
<th>b</th>
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**KEY:**

a= Predicted 3D structure of wild chicken 1 GHR.

b= Predicted 3D structure of wild chicken 2 GHR.

c= Predicted 3D structure of wild duck GHR.
Appendix 13: Predicted protein 3D structures of insulin-like growth factor 1 (IGF1).

KEY:

a= Predicted 3D structure of Japanese quail IGF1.
b= Predicted 3D structure of southern ostrich IGF1.
c= Predicted 3D structure of wild chicken IGF1.
d= Predicted 3D structure of wild duck IGF1.
e= Predicted 3D structure of wild turkey IGF1.
Appendix 14: Predicted protein 3D structures of insulin-like growth factor 1 receptor (IGF1R).

**KEY:**

a= Predicted 3D structure of Japanese quail IGF1R.
b= Predicted 3D structure of southern ostrich IGF1R.
c= Predicted 3D structure of wild chicken IGF1R.
d= Predicted 3D structure of wild duck IGF1R.