# In-silico identification and characterisation of 17 polymorphic anonymous non-coding sequence markers (ANMs) for red grouse (Lagopus lagopus scotica) 

Marius A. Wenzel ${ }^{*}$ and Stuart B. Piertney

December 16, 2014

Institute of Biological and Environmental Sciences, University of Aberdeen, Zoology Building, Tillydrone Avenue, Aberdeen AB24 2TZ, UK

* corresponding author. email address: marius.a.wenzel.08@aberdeen.ac.uk. Phone number: +44 1224272395

Word counts: 133 (abstract), 779 (main text), 86 (legends)
Keywords: anonymous nuclear markers; non-coding DNA; neutral evolution; red grouse; Lagopus


#### Abstract

Anonymous non-coding sequence markers (ANMs) are powerful neutral genetic markers with great utility in phylogeography, population genetics and population genomics. Developing ANMs has previously relied on sequencing random fragments of genomic DNA in the target species and then querying bioinformatics databases to identify unannotated, putatively neutral fragments. Here, we describe an alternative in silico approach that is based on identifying large unannotated genomic regions in model species to provide a priori neutral targets for candidate ANMs that are remote from exonic regions. We illustrate this approach by developing a set of 17 polymorphic ANMs for red grouse (Lagopus lagopus scotica) from c. 1 Mbp noncoding chromosome regions of chicken, turkey and zebrafinch genomes. This pipeline represents a powerful and efficient approach when appropriate model genomes are available for the target species of interest.


The ability to isolate and characterise nuclear DNA sequence polymorphisms remains a major priority for studies resolving population history, estimating demographic parameters and examining the genetic basis of divergence, adaptation and speciation (Thomson et al, 2010). In non-model species, one classic approach is to use exonprimed intron-crossing markers (EPICs) or comparative anchor-tagged sequences (CATS), which target nuclear intronic sequences by anchoring primers in conserved flanking exonic regions (Backström et al, 2008). These markers are considered useful for phylogenetics, gene mapping and population genetics because of high variability, cross-species utility and presumed neutrality (Brito and Edwards, 2009; Slate et al, 2009). However, they are unlikely to be truly neutral because purifying selection on flanking exons may affect intronic polymorphism through hitchhiking (Thomson et al, 2010). In contrast, nuclear anonymous non-coding markers (ANMs) that are located in regions remote from exonic domains are unlikely to be under selection and are substantially more polymorphic than EPICs or CATS (Thomson et al, 2010). Additionally, ANMs are more abundant and easier to type than microsatellites, making them ideal tools for population genetics and phylogeography (Rosenblum et al, 2007; Lee and Edwards, 2008; Thomson et al, 2010).

Isolating ANMs is usually based on sequencing random fragments of genomic DNA following shearing (Rosenblum et al, 2007; Lee and Edwards, 2008) or enzymatic digestion (Barlow et al, 2012; Ren et al, 2013), or via whole-genome massive parallel sequencing (Bertozzi et al, 2012; Lewis et al, 2014). Non-coding sequences can then be identified from absence of annotations following BLAST (Altschul et al, 1997) queries against bioinformatics databases, and primers are designed accordingly (Bertozzi et al, 2012; Lewis et al, 2014). One issue
with this strategy is that primer design on library clone sequences may be compromised because unidentified polymorphism in binding sites may cause null-alleles, PCR failure and poor cross-species utility (Thomson et al, 2010). Most crucially, however, neutrality cannot be established from mere absence of BLAST results. Confirming remoteness from exonic domains as a criterion for neutrality requires examining the genomic context of the sequences in model genomes, but direct sequence mapping may be difficult if no taxonomically close model genome is available.

Here, we describe an alternative strategy to identifying ANMs that is purely based on available bioinformatics resources and provides a priori candidate targets for designing primers in non-coding regions that are remote from exonic regions and hence likely to be truly neutral. We illustrate this strategy by developing ANMs from avian model genomes for red grouse (Lagopus lagopus scotica), an economically important game bird endemic to upland heather moors in Scotland and northern England (Martínez-Padilla et al, 2014).

The UCSC Table Browser (Karolchik et al, 2004) provides tabulated annotations from published genomes. RefSeq annotations were downloaded for the chicken genome (Gallus gallus galGal4 assembly) and analysed using custom scripts in R 3.0.3 (R Core Team, 2014). The table fields txStart and txEnd were used to calculate genomic distances (bp) between consecutive transcription blocks across each autosome. The maximum region size per autosome ranged from 0.1 Mbp to 5.1 Mbp (median 1 Mbp ) and a total of 113,19 and 7 regions of at least $1 \mathrm{Mbp}, 2 \mathrm{Mbp}$ and 3 Mbp , respectively, were available across all autosomes (Figure 1). Nine c. 1 Mbp regions in nine autosomes were arbitrarily selected as candidate target regions (Figure 1). The central 10 kbp portion of these regions was extracted from GENBANK chromosome sequences, and homologous sequences in turkey (Meleagris gallopavo melGal1 assembly) and zebrafinch (Taeniopygia guttata taeGut1 assembly) genomes were identified using the BLast-like alignment tool Blat (Kent, 2002). Alignments of all three species and also chicken and turkey alone were generated in GEnEIous v5.6.3 (Drummond et al, 2012). Non-degenerate primers (200-800 bp amplicon size, 18-27 bp primer length, $20-80 \%$ GC content, $50-64{ }^{\circ} \mathrm{C}$ melting temperature) were then designed opportunistically on small conserved regions using PRIMER3 (Rozen and Skaletsky, 2000) as implemented in GENEIOUS. Primer specificity was tested using UCSC in-SILICO PCR amplicon prediction (Hinrichs et al, 2006) on the chicken, turkey and zebrafinch genomes.

Sequence polymorphism was ascertained in three red grouse individuals from locations that maximise geographic variation across a network of grouse moors in north-east Scotland (Glenlivet $57.29{ }^{\circ} \mathrm{N} 3.18{ }^{\circ} \mathrm{W}$, Mar Lodge $56.95^{\circ} \mathrm{N} 3.66^{\circ} \mathrm{W}$ and Invermark $\left.56.89^{\circ} \mathrm{N} 2.88^{\circ} \mathrm{W}\right)$. PCR conditions followed Wenzel et al (2014), with annealing temperatures as detailed in Table 1. Amplicons were Sanger sequenced in both directions, sequences were aligned in Geneious and heterozygote sites were coded as IUPAC degenerate bases. Absence of exonic annotations was re-confirmed using BLASTN against the GENBANK NT database (Altschul et al, 1997). Polymorphic sites, numbers of haplotypes, nucleotide diversity, haplotype diversity and Tajima's $D$ were then computed on reconstructed haplotypes derived from the PHASE algorithm in DNASP v5 (Librado and Rozas, 2009).

Twenty-two out of thirty primer pairs ( $73 \%$ ) amplified in red grouse, demonstrating a high success rate of


Figure 1: Numbers of unannotated genomic regions of particular minimum sizes in chicken autosomes, based on distances between consecutive transcription blocks. Black dots represent candidate regions selected for ANM design (Table 1).
our development strategy. Polymorphic sequence alignments were obtained for seventeen loci (57\%), containing 1-18 SNPs that define 2-6 haplotypes with no evidence of deviation from neutral sequence evolution (Table 1). Insertions/deletions of $1-10 \mathrm{bp}$ were present in five loci. These polymorphic ANMs provide a valuable resource for a range of population genetics or genomics applications in red grouse. The zebrafinch genome impeded primer design in many cases due to its taxonomic distance (Table 1), but considering the taxonomic distance between red grouse, chicken and turkey, these markers should be conserved and hence useful across a range of closely related galliform species.

## Acknowledgements

This study was funded by a BBSRC studentship (MA Wenzel) and NERC grants NE/H00775X/1 and NE/D000602/1 (SB Piertney). We are grateful to Mario Röder, Keliya Bai and Marianne James for help with fieldwork, and all grouse estate factors, owners and keepers, most particularly Alistair Mitchell, Shaila Rao, Christopher Murphy, Richard Cooke and Fred Taylor, for providing access to estate game larders.

## References

Altschul S, Madden T, Schäffer A, Zhang J, Zhang Z, Miller W, Lipman D (1997) Gapped BLAST and PSIBLAST: A new generation of protein database search programs. Nucleic Acids Res 25(17):3389-3402

Backström N, Fagerberg S, Ellegren H (2008) Genomics of natural bird populations: a gene-based set of reference markers evenly spread across the avian genome. Mol Ecol 17(4):964-980

Barlow A, Grail W, de Bruyn M, Wüster W (2012) Anonymous nuclear markers for the African adders (Serpentes: Viperidae: Bitis). Conserv Gen Res 4(4):967-969

Bertozzi T, Sanders KL, Sistrom MJ, Gardner MG (2012) Anonymous nuclear loci in non-model organisms: making the most of high-throughput genome surveys. Method Biochem Anal 28(14):1807-1810

Brito PH, Edwards SV (2009) Multilocus phylogeography and phylogenetics using sequence-based markers. Genetica 135(3):439-455

Drummond A, Ashton B, Buxton S, Cheung M, Cooper A, Duran C, Field M, Heled J, Kearse M, Markowitz S, Moir R, Stones-Havas S, S S, Thierer T, Wilson A (2012) Geneious v5.6.3. Available from http://www.geneious.com

Hinrichs AS, Karolchik D, Baertsch R, Barber GP, Bejerano G, Clawson H, Diekhans M, Furey TS, Harte RA, Hsu F, et al (2006) The UCSC genome browser database: update 2006. Nucleic Acids Res 34(suppl 1):D590-D598

Karolchik D, Hinrichs AS, Furey TS, Roskin KM, Sugnet CW, Haussler D, Kent WJ (2004) The UCSC Table Browser data retrieval tool. Nucleic Acids Res 32(suppl 1):D493-D496

Kent WJ (2002) BLAT-The BLAST-Like Alignment Tool. Genome Res 12:656-664

Lee JY, Edwards S (2008) Divergence Across Australia's Carpentarian Barrier: Statistical Phylogeography of the Red-Backed Fairy Wren (Malurus melanocephalus). Mem New York Botan G 62(12):3117-3134

Lewis CJ, Maddock ST, Day JJ, Nussbaum RA, Morel C, Wilkinson M, Foster PG, Gower DJ (2014) Development of anonymous nuclear markers from Illumina paired-end data for Seychelles caecilian amphibians (Gymnophiona: Indotyphlidae). Conserv Gen Res 6(2):289-291

Librado P, Rozas J (2009) DnaSP v5: a software for comprehensive analysis of DNA polymorphism data. Method Biochem Anal 25(11):1451-1452
Table 1: Characterisation of 17 anonymous non-coding sequence markers (ANMs) for red grouse. Primer GC content, melting temperature $\mathrm{T}_{\mathrm{m}}$ and annealing temperature $\mathrm{T}_{\mathrm{a}}\left({ }^{\mathrm{TD}}=\right.$ TouchDown) are presented alongside genomic locations in three bird genomes and sequence diversity statistics derived from three red grouse individuals (segregating sites S, nucleotide diversity $\pi$, haplotypes H , haplotype diversity $\mathrm{H}_{\mathrm{d}}$, Tajima's $D$ ).

| Primer name | Primer sequence ( $5^{\prime} \rightarrow 3^{\prime}$ ) | GC <br> (\%) | $\begin{aligned} & \mathrm{T}_{\mathrm{m}} \\ & \left({ }^{\circ} \mathrm{C}\right) \end{aligned}$ | $\mathrm{Ta}_{\mathrm{a}}\left({ }^{\circ} \mathrm{C}\right)$ | in silico amplification (genomic location and predicted amplicon size) |  |  | in vitro amplification |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  | Chicken | Turkey | Zebrafinch | Size | s | $\pi$ | H | ${ }^{\mathrm{H}} \mathrm{d}$ | D | GENBANK accession |
| Lls_ANM_6_1F | ACCCTTGTGAGCtGAGAGCTT | 52 | 59.7 | $60-50^{\text {TD }}$ | chr6:5360100+5360504 | chr8:2879375+2879779 | - | 406 | 9 | 0.011 | 4 | 0.800 | 0.793 | KM379116 |
| Lls_ANM_6_1R | tcacacctatggcanacaiancac | 41 | 56.7 |  | 405 bp | 405 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_8_1F | TGGCCAGGGTtatctgangtac | 59 | 63.3 | 68 | chr8:9422697+9423060 | chr10:1058050-1058413 | - | $287^{\text {ab }}$ | 3 | 0.005 | 4 | 0.867 | 0.600 | KM379117 |
| Lls_ANM_8_1R | tGCCCTCTGAAGAAGCCATtTGA | 47 | 59.8 |  | 364 bp | 364 bp | - |  |  |  |  |  |  |  |
| Lls_ANM - $^{\text {- }}$ 2F | TCTGTCACTGTtctcacattte | 36 | 52.1 | ${ }_{60-50}{ }^{\text {TD }}$ | chr8:9424802+9425186 | chr10:1055959-1056338 | - | $388{ }^{\text {b }}$ | 2 | 0.002 | 2 | 0.333 | ${ }^{-1.132}$ | KM379118 |
| Lls_ANM_8_2R | Cactcantttgatttectcagtancc | 34 | 52.5 |  | 385 bp | 380 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_9_1F | AGTCTGAGACATTtTCCCCATCC | 47 | 57.6 | 65 | chr9:20994525+20994916 | chr 11:21699903+21700290 | - | $390{ }^{\text {b }}$ | 5 | 0.007 | 4 | 0.867 | 0.708 | KM379119 |
| Lls_ANM_9_1R | AGAACTCATtCtGCtttgcagc | 45 | 56.8 |  | 392 bp | 388 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_-9_2F | tGAaAtGtacttcctancacatge | ${ }^{37}$ | 53.4 | $60-50^{\text {TD }}$ | chr9:20992310+20992689 | chr 11:21697593+21697976 | - | 385 | 3 | ${ }^{0.004}$ | 4 | 0.867 | 1.386 | KM379120 |
| Lls_ANM_9_2R | tGtttttctittctanttatgtgal | 26 | 50.8 |  | 380 bp | 384 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_9_3F | CtCCagGatactcangccaca | 52 | 57.5 | 65 | chr9:20999542+20999958 | chr 11:21704861+21705255 | - | 407 | 2 | 0.002 | 3 | 0.733 | -0.050 | KM379121 |
| Lls_ANM_9_3R | tCCttGcagtttttagacttgan | 39 | 54.6 |  | 417 bp | 395 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_10_1F | CACTCAGCCCtCtgtatangtge | 56 | 61.4 | $65-55^{\text {TD }}$ | chr 10:15610965-15611270 | chr12:16100168-16100473 | chr 10:16580442-16580734 | 305 | 1 | 0.001 | 2 | 0.333 | -0.933 | KM379122 |
| Lls_ANM_10_1R | TGAGTtGTtagaccacacgcha | 50 | 59.6 |  | 306 bp | 306 bp | 292 bp |  |  |  |  |  |  |  |
| Lls_ANM_10_2F | ACtCGCGTGTGGtctancancte | 52 | 60.4 | $65-55^{\text {TD }}$ | chr 10:15610580-15610988 | chr12:16099786-16100191 | chr 10:16580040-16580466 | ${ }^{393}$ | 2 | 0.002 | 3 | 0.733 | 0.311 | KM379123 |
| Lls_ANM_10_2R | ACtGCatgGtghagantgcca | 57 | 63.7 |  | 409 bp | 406 bp | 426 bp |  |  |  |  |  |  |  |
| Lls_ANM_10_3F | TTGCTGCCTGCCACTGCTTA | 55 | 61.3 | ${ }_{65-55^{\text {TD }}}$ | chr 10:15611228+15611646 | chr12:16100431+16100850 | - | $278{ }^{\text {ab }}$ | 18 | 0.034 | 6 | 1.000 | 0.723 | KM379124 |
| Lls_ANM_10_3R | agccacacctcccocattca | 60 | 63.1 |  | 419 bp | 420 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_11_1F | AGTtGACATCAAAGTGGAGACA | 40 | 54.3 | ${ }_{65-55^{\text {TD }}}$ | chr 11:5087733+5088037 | chr 13:5398213+5398522 | chr 11:9467370+9467669 | 304 | 4 | 0.004 | 2 | ${ }^{0.333}$ | -1.295 | KM379125 |
| L1s_ANM_11_1R | gtgtctggtttcacatctgge | 52 | 57.5 |  | 305 bp | 310 bp | 299 bp |  |  |  |  |  |  |  |
| Lls_ANM_13_1F | gGacatttagcancangatcaca | 43 | 56.1 | $65-55^{\text {TD }}$ | chr 13:5734379+5734751 | chr 15:5954643+5955002 | - | ${ }^{373}$ | 3 | 0.004 | 3 | 0.800 | 1.124 | KM379126 |
| Lls_ANM_13_1R | Ggatgattaggetgtgtanaicci | 45 | 56.6 |  | 373 bp | 360 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_ ${ }^{13}$ _3F | tgtggatgtatactacctggca | 45 | 55.8 | ${ }_{60-50}{ }^{\text {TD }}$ | chr 13:5741550+5741896 | chr 15:5961829+5962181 | - | $244^{\text {ab }}$ | 3 | 0.006 | 3 | 0.733 | 0.338 | KM379127 |
| Lls_ANM_13_3R | GCtGatacctttatanactttgatge | 34 | 53.3 |  | 347 bp | 353 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_18_1F | TGGAAGCCATGAGGAAGGGGA | 57 | 62.2 | ${ }_{6} 7$ | chr 18:8776322+8776692 | chr20:6558242-6558622 | - | 379 | 7 | 0.009 | 4 | 0.867 | 0.508 | KM379128 |
| L1s_ANM_18_1R | AGGAAGGAAGAATGCAAGGCA | ${ }^{47}$ | 57.8 |  | 371 bp | 381 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_18_2F | tCAGCCAATATtGCttcanagg | 40 | 54 | ${ }_{60-50}{ }^{\text {TD }}$ | chr 18:8779401+8779815 | chr20:6555203-6555600 | - | $322^{\text {a }}$ | 7 | 0.008 | 4 | 0.867 | -0.631 | KM379129 |
| L1s_ANM_18_2R | tcCantganatgangctgtatge | 39 | 53.9 |  | 415 bp | 398 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_20_2F | ATtCCtCgCtGgttgetgac | 60 | 62.6 | 68 | chr 20:4665027+4665427 | chr $22: 4275127+4275522$ | - | 403 | 9 | 0.008 | 3 | 0.600 | -0.818 | KM379130 |
| Lls_ANM_20_2R | CTGCACtTGTGGGCAGACCC | 65 | 63.5 |  | 401 bp | 396 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_22_2F | GCGATGCTACACCCTCCAAG | 60 | 59.9 | ${ }^{60-50}{ }^{\text {TD }}$ | chr 22:3864116+3864470 | chr $24: 3873443+3873798$ | - | 357 | ${ }^{2}$ | 0.002 | ${ }^{2}$ | ${ }^{0.333}$ | $-1.132$ | KM379131 |
| L1s_ANM_22_2R | ACAAAATGCTACTGACAAATCTGA | ${ }^{33}$ | 52.6 |  | 355 bp | 356 bp | - |  |  |  |  |  |  |  |
| Lls_ANM_22_3F | GCtttccctcetctatttccttc | 47 | 56 | ${ }^{66}$ | chr22:3862528+3862926 | chr 24:3871853+3872244 | - | 397 | ${ }^{6}$ | 0.008 | ${ }^{3}$ | 0.733 | 1.392 | KM379132 |
| L1s_ANM_22_3R | AGAATCCCAAAGCCTtTCCCT | 47 | 57.4 |  | 399 bp | 392 bp | - |  |  |  |  |  |  |  |

Martínez-Padilla J, Redpath SM, Zeineddine M, Mougeot F (2014) Insights into population ecology from longterm studies of red grouse Lagopus lagopus scoticus. J Anim Ecol 83(1):85-98

R Core Team (2014) R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing, Vienna, Austria, URL http://www.R-project.org

Ren QP, Fan Z, Zhou XM, Jiang GF, Wang YT, Liu YX (2013) Identification and characterization of anonymous nuclear markers for the double-striped cockroach, Blattella bisignata. B Entomol Res 103(01):29-35

Rosenblum E, Belfiore N, Moritz C (2007) Anonymous nuclear markers for the eastern fence lizard, Sceloporus undulatus. Mol Ecol Notes 7(1):113-116

Rozen S, Skaletsky HJ (2000) Primer3 on the WWW for general users and for biologist programmers. Meth Mol Biol 132:365-386

Slate J, Gratten J, Beraldi D, Stapley J, Hale M, Pemberton JM (2009) Gene mapping in the wild with SNPs: guidelines and future directions. Genetica 136(1):97-107

Thomson RC, Wang IJ, Johnson JR (2010) Genome-enabled development of DNA markers for ecology, evolution and conservation. Mol Ecol 19(11):2184-2195

Wenzel MA, Webster LMI, Paterson S, Piertney SB (2014) Identification and characterisation of 17 polymorphic candidate genes for response to parasitic nematode (Trichostrongylus tenuis) infection in red grouse (Lagopus lagopus scotica). Conserv Gen Res

