

1 **Complex and long-range linkage disequilibrium and its relationship**
2 **with QTL for Marek's Disease resistance in chicken populations**

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13 **ABSTRACT**

14 Chicken long-range linkage disequilibrium (LRLD) and LD blocks, and their
15 relationship with previously described Marek's Disease (MD) quantitative trait loci
16 regions (QTLRs), were studied in an F₆ population from a full-sib advanced intercross
17 line (FSAIL), and in eight commercial pure layer lines. Genome wide LRLD was studied
18 in the F₆ population by random samples of non-syntenic and syntenic marker pairs
19 genotyped by Affymetrix HD 600K SNP array. To illustrate the relationship with
20 QTLRs, LRLD and LD blocks in and between the MD QTLRs were studied by all
21 possible marker pairs of all array markers in the QTLRs, using the same F₆ QTLR
22 genotypes and genotypes of the QTLR elements' markers in the eight lines used in the
23 MD mapping study. LRLD was defined as $r^2 \geq 0.7$ over a distance ≥ 1 Mb, and 1.5% of
24 all syntenic marker pairs were classified as LRLD. Complex fragmented and
25 interdigitated LD blocks were found, over distances ranging from a few hundred to a few
26 million bases. Vast high, long-range, and complex LD was found between two of the MD
27 QTLRs. Cross QTLRs STRING networks and gene interactions suggested possible
28 origins of this exceptional QTLRs' LD. Thus, causative mutations can be located at a
29 much larger distance from a significant marker than previously appreciated. LRLD range
30 and LD block complexity may be used to identify mapping errors, and should be
31 accounted for while interpreting genetic mapping studies. All sites with high LD with a
32 significant marker should be considered as candidate for the causative mutation.

33

34 **Keywords:** Chicken; long-range linkage disequilibrium; QTL; F₆; LD blocks

35

36 INTRODUCTION

37 Linkage disequilibrium (LD) refers to correlations among alleles of different genomic
38 sites. It quantifies the informativity between different sites [Reich et al., 2001; Ardlie et
39 al., 2002; Aerts et al., 2007]. Useful LD indicate non-random association of alleles at
40 different loci [Aerts et al., 2007]. Appreciable LD is commonly found between pairs of
41 loci close to one another, and LD decreases rapidly as distance between the loci increases
42 [e.g., Aerts et al., 2007; García-Gómez et al., 2012]. This provides the basis for genome
43 wide association studies (GWAS) to map quantitative trait loci (QTL).

44 Nevertheless, during any one-time snapshot of a population, long-range LD (LRLD)
45 can also be found among loci that are well-separated from one another, over millions of
46 bp [Aerts et al., 2007; Corbin et al., 2010; García-Gómez et al., 2012; Koch et al., 2013;
47 Skelly et al., 2015; Park 2019; Peters et al., 2021]. Rarely, LD above background level
48 can be found between non-syntenic markers on different chromosomes (chr) [García-
49 Gómez et al., 2012].

50 LRLD could just be a matter of sampling variation, especially in small populations
51 [Skelly et al., 2015]. Alternatively, LRLD could be a result of genome assembly errors
52 where SNP locations are misidentified, and thus LRLD may help identify such assembly
53 errors [Utsunomiya et al., 2016]. This phenomenon may, however, also have genuine
54 biological origins, such as co-evolution of genomic sites (coding and non-coding genes,
55 long- and short-range regulatory sites), gene conversion, copy number variation, or
56 demographic factors such as selection, population bottlenecks, nonrandom mating, and
57 epistasis.

58 LD blocks are runs of genomic sites all having appreciable LD with one another.
59 However, high LD in general and LRLD in particular are not always continuous. Rather,
60 they are often separated by genomic sites with which they have practically no LD [Allabi
61 et al., 2005; Aerts et al., 2007; Lipkin et al., 2013; O'Brien et al., 2014]. Thus, not only
62 can LD be found among distant loci, but also its pattern may be complex, comprised of
63 fragmented blocks.

64 LRLD and LD complexity present concerns for GWAS mapping, as a significant
65 association may be found between a causative locus and markers far removed from it, thus

66 falsely placing the putative causative locus at a site far away from its actual location
67 [Skelly et al, 2015]. On the other hand, LRLD may point to interactions between
68 unlinked regions in the genome (e.g., a receptor and its ligand or a gene and its regulator).
69 Furthermore, LRLD can identify co-evolution of different genomic regions affected by
70 the same selection, natural or artificial.

71 The objectives of this study were to characterize LRLD and LD blocks in multiple Hy-
72 Line chicken lines previously used to map quantitative trait loci regions (QTLRs) for MD
73 resistance [Smith et al., 2020]. This will give a genomic view of the LD complexity, and
74 illustrate its importance to mapping results by assessing the relationship between LRLD
75 and LD blocks and MD QTLRs.

76

77 **MATERIALS AND METHODS**

78 **Populations**

79 All procedures carried out on the birds involved in this study were conducted in
80 compliance with Hy-Line International Institutional Animal Care and Use Committee
81 guidelines.

82 Nine populations described by Smith et al. (2020) were again used in the present
83 study. These comprised five families of F₆ birds from a Full Sib Advanced Intercross
84 Line (FSAIL) used by Smith et al. (2020) to map QTLRs affecting MD resistance, and
85 eight pure lines used in the same study to test these QTLRs.

86 A priori, it is expected that LRLD in F₆ will be at higher frequency than in pure lines,
87 because the families start with only 4 chromosomes each, and there are only a few
88 generation of intercrossing to break up the haplotype blocks. However, this study did not
89 compare populations, families or lines. Rather, it aim to present the phenomena of LD
90 long range and complexity.

91 **Trait**

92 The trait for which these QTLR are associated is resistance to the avian oncogenic
93 alpha herpes virus, Marek's Disease (MD virus [Smith et al., 2020]). This trait association

94 data set was used, as the phenotype and genotype information was available. It is used as
95 an illustration for the QTLR and LD associations that were identified.

96 **Genotypes**

97 Only the autosomal genotypes as used by Smith et al. (2020) to map and test QTLRs
98 were used in the present study. Genotypes on the Z chromosome will be analyzed in
99 detail in a different manuscript. Genotypes were obtained from the HD 600K Affymetrix
100 SNP chicken array [Kranis et al., 2013] in the F₆ population, and marker genotypes used
101 to test the QTLRs by the eight pure lines [Smith et al., 2020]. The only difference was
102 that instead of a minimum $MAF \geq 0.01$ used for the association tests by Smith et al.
103 (2020), a threshold of 0.10 was used here for the LD analysis, to avoid spurious high LD
104 due to rare alleles [Skelly et al., 2015]. However, to test LD with exactly the same
105 markers used for association tests in the eight pure lines, the threshold of $MAF \geq 0.01$
106 was also used for LD for these lines.

107 **Genome assemblies and remapping QTLRs**

108 As described in Smith et al. (2020), the initial analysis in this study was based on the
109 Galgal4 genome build. The [Lift Genome Annotations](#) tool [Haeussler et al., 2019] within
110 the [UCSC](#) Browser was used to remap markers from Galgal4 to Galgal6 (GRCg6a; Acc.
111 No.: GCA_000002315.5). The new coordinates were then used to remap the F₆ QTLRs as
112 was done by Smith et al. (2020). The results were very similar on both assemblies, and
113 hence only the results from Galgal6 will be presented here. Nevertheless, there was one
114 change worth noting, detailed in the Appendix.

115 **Linkage disequilibrium (LD)**

116 **LD Measure.** LD r^2 within each F₆ family and each pure line were obtained using JMP
117 Genomics software (JMP Genomics, Version 9, SAS Institute Inc., Cary, NC, USA,
118 1989–2019).

119 **Non-syntenic LD.** Background LD over all autosomes was estimated utilizing
120 Affymetrix 600K genotypes of two random combined samples with return of non-
121 syntenic marker pairs from each of the five F₆ families.

122 **Long-Range LD (LRLD).** Koch et al. (2013) used all pairs of SNPs on each human
123 chromosome and defined LRLD between haplotype-blocks rather than between SNP
124 pairs. However, due to computer limitations, we used samples of random SNP pairs on
125 the Affymetrix 600K SNP array to assess LRLD in the autosomes of the five F₆ families.
126 Koch et al. (2013) defined LRLD in human as high LD (their low p_D) over a distance \geq
127 0.25 cM (\approx 0.25 Mb). Vallejo et al. (2018) used a very relaxed LD threshold of $r^2 > 0.25$
128 in rainbow trout studies, but a larger minimum distance of 1.0 Mb. Conservatively, and
129 based on the present results, in this study we defined LRLD as a marker pair with $r^2 \geq 0.7$
130 over a distance ≥ 1.0 Mb. As noted in the Introduction, LRLD between elements in a
131 QTLR and across QTLRs can indicate a relationship between the element and between
132 the QTLRs. Hence, to illustrate its importance, LRLD in and between MD QTLRs [Smith
133 et al., 2020] was studied by all F₆ Affymetrix genotypes in the QTLRs.

134 **F₆ random LRLD and QTLRs.** F₆ LRLD of random marker pairs were aligned by
135 chromosomal location with the F₆ MD QTLRs [Smith et al., 2020], and all overlaps
136 between the LRLDs and the QTLRs were counted.

137 **LD blocks.** High LD blocks were defined as a group of markers located on the same
138 chromosome having $r^2 \geq 0.7$ with each other. The definition was applied even if markers
139 with low LD appeared between the LD markers. This definition allowed a "look over the
140 horizon" and identification of fragmented and interdigitated blocks.

141 **Gene network analysis**

142 To investigate the genes underlying QTLRs 4 and 5, the BioMart tool within the
143 Ensembl database (<https://www.ensembl.org/info/data/biomart/index.html>) was used to
144 identify genes in these regions. These identified genes were then subject to network
145 analysis using the STRING database (v11). [Jensen et al., 2009] which provides an
146 overview of known protein interactions.

147

148 **RESULTS**

149 **Remapping QTLRs from Galgal4 to Galgal6.** The new coordinates of the markers on
150 Galgal6 and association results obtained by Smith et al. (2020) were used to remap the

151 MD QTLRs identified with Galgal4 by Smith et al. (2020). The same 38 QTLRs were
152 found on each genome build (Table 1). Most changes were negligible, except one
153 movement of a fragment on Chr 1 over about 70 Mb from QTLR 1 to QTLR 4, including
154 a QTLR lncRNA tested by Smith et al. (2020). This change is detailed in the Appendix.
155 The new QTLR coordinates on Galgal6 were used for the LD analyses in this study.

156

157 **Table 1.** *Remapping Galgal4 QTLRs onto Galgal6.*

158

159 **Linkage disequilibrium in the F₆ population based on 600K genotyping SNP array**

160 **Non-syntenic random LD.** A total of 923,183 random non-syntenic pairs of markers
161 from different autosomes were used to assess the background level of LD, with an
162 average of 184,636.6 pairs in a family (Table 2). LD averaged 0.011 ± 0.016 , comparable
163 to previous reports in chicken [Lipkin et al., 2013; Seo et al., 2018], but about ten times
164 higher than values reported in mammals; horse [Corbin et al., 2010], sheep [García-
165 Gámez et al., 2012], and cattle [Khatkar et al., 2008; Sargolzaei et al., 2008] populations.
166 These differences may represent experimental design, or population sample, size, history
167 and structure or biological differences between birds and mammals.

168 Mean family LDs and standard deviations (SDs) had significant high negative
169 correlation with the size of the population sample (i.e., Ind/pair), $r = -0.895$ ($P = 0.040$)
170 and $r = -0.884$ ($P = 0.046$). These correlations are also in accord with our previous report
171 in chicken [Lipkin et al., 2013], and the expectation of Sved (1971).

172

173 **Table 2.** *Summary statistics of non-syntenic random LD between 600K markers in the F₆*
174 *families.*

175

176 With a mean LD of 0.011 ± 0.016 , any $r^2 > 0.043$ is above the background LD. Indeed,
177 combining all five F₆ families together, only 3.5% of the r^2 values were above 0.05
178 (Supplemental Table 1). A single high LD of $r^2 = 0.991$ was found in Family 2. Without

179 any replication, this was treated as a sampling effect. Based on these results, a
180 conservative critical LD value of $r^2 \geq 0.15$ was chosen for defining significant LD.

181

182 *Supplemental Table 1. Distribution of non-syntenic random LD values among the F_6*
183 *families.*

184

185 **Syntenic random LD.** A total of 1,008,823 random syntenic marker pairs were used to
186 assess the level of random LDs on the same chromosome (Table 3). Distance between
187 markers in the random pairs varied from 11 to 197,038,449 bp, with an average of
188 28,976,195.6 bp. Means of r^2 were all in close range around 0.11, averaging 0.114, ten
189 times the means obtained for the non-syntenic LD. Though obtained by random marker
190 pairs, some of which are at a long distance from one another, these means suggest the
191 presence of large number of LDs above the background LD of 0.011 (Table 2). The
192 expected negative correlation between distance and LD was again obtained in all five
193 families (Table 3).

194

195 *Table 3. Summary statistics of syntenic random LD between 600K markers in the F_6*
196 *families.*

197

198 In all families, about two-thirds of the LD values were up to 0.05, dropping rapidly
199 thereafter (Table 4). Interesting, for all families, there was an increase in the range of $r^2 >$
200 0.85, suggesting existence of large high-LD blocks. Pooled over all families, the
201 proportion of $r^2 \geq 0.15$, set conservatively as a threshold of significance by the non-
202 syntenic LD, was almost 0.2 (Table 4), while less than 5% of the LD values were above
203 0.7. Hence, the range of $0.15 \leq r^2 < 0.7$ was set as low to moderate LD and used to define
204 moderate LD blocks, and $r^2 \geq 0.7$ was set as high LD and used to define LRLD and high
205 LD blocks.

206

207 **Table 4.** *Distribution of syntenic random LD values among the F₆ families.*

208

209 **Long-range LD (LRLD)**

210 **Estimating LRLD by random samples of syntenic marker pairs.** Pooled over all
211 families, 418,075 pairs had a distance above 20 Mb; as expected, no high LD of $r^2 \geq 0.7$
212 was found beyond 20 Mb (Supplemental Table 2).

213

214 **Supplemental Table 2.** *Distribution of distances among random marker pairs in the F₆*
215 *families.*

216

217 Detailed inspection of the distances up to 20 Mb showed that all high LDs were in fact
218 within 10 Mb (Supplemental Table 3).

219

220 **Supplemental Table 3.** *Distribution of distances among random syntenic pairs with $r^2 \geq$*
221 *0.7 separated by up to 20 Mb in the F₆ families.*

222

223 Pooled over all families together, a total of 50,100 random marker pairs qualified
224 within the LRLD definition, namely $r^2 \geq 0.7$ over a distance ≥ 1 Mb. These LRLDs
225 constitute 30.9% of all pairs within 20 Mb, and 1.5% of the total number of syntenic pairs
226 tested (Supplemental Table 2).

227 Among the syntenic pairs, 0.016 had $r^2 > 0.95$, almost 15-times the proportion of the
228 single LD value in this range (0.000001) found among the non-syntenic pairs
229 (Supplemental Table 1). Thus, the proportion of syntenic high LD was not negligible.
230 LRLDs were distributed over all autosomes in all five F₆ families (Supplemental Table 4,
231 Figure 1; LD matrices in Genetics figshare portal). No LRLD was found on
232 Chromosomes 22 in any of the families.

233

234 **Supplemental Table 4.** *Distribution of random LRLDs over autosomal chromosomes in*
235 *the F₆ families.*

236

237 Though these LRLDs were obtained by random sampling of marker pairs, repeated
238 similar locations of marker pairs suggest the existence of many LRLD blocks. This was
239 indeed found by the LD analysis of the MD QTLRs (see below).

240

241 **F₆ MD QTLRs and random LRLD.** To check for a possible relationship between the
242 LRLDs found here and the F₆ MD QTLRs mapped in the same population (Table 1),
243 LRLDs and QTLRs were aligned together (e.g., Figure 1 and LD matrices in Genetics
244 figshare portal), and overlaps were counted (Supplemental Table 5).

245

246 **Figure 1.** *Distribution of random LRLDs over Chr 1 and overlaps with QTLRs (Table 1)*
247 *in F₆ Family 1.*

248

249 **Supplemental Table 5.** *Number of random F₆ LRLDs overlapping F₆ MD QTLRs.*

250

251 As noted above, with all markers in an interval less than 1 Mb, no LRLD could be
252 found on Chr 16 (Supplemental Table 4); hence, QTLR 32 was not included in any
253 further analyses. Of the remaining 37 QTLRs, overlaps between 28 QTLRs and LRLDs
254 were found in all families (the non-zeros under ‘Families’ in Supplemental Table 5). It
255 seems remarkable that, even though only 1.5% of the random LD values were LDLR, no
256 less than 75.7% of the mapped MD QTLRs overlapped LRLDs. Then again, in Galgal6,
257 QTLRs averaged 1.4 Mb (Table 1), and random LRLDs averaged 2.2 Mb, from 1 to
258 above 12 Mb (Supplemental Table 6). Thus, such overlap may not be so surprising, but a
259 result of the abundance and size of the QTLRs and LRLDs.

260

261 **Supplemental Table 6.** *The distribution of random F₆ LRLDs length.*

262

263 Zooming in on QTLRs clearly showed the overlap between the LRLDs and QTLRs
264 (Figure 2). Not only was LRLD found within QTLRs, but LRLD was found between
265 QTLRs 4 and 5 in all 5 families. The similar locations seen in Figure 2 suggest the
266 presence of LD blocks shared by both QTLRs.

267

268 **Figure 2.** *Overlaps between QTLRs 4-5 and LRLDs in all F₆ families.*

269

270 **LD in the QTLRs in the F₆ families**

271 The overlaps found in the F₆ families between random LRLDs and the MD QTLRs,
272 led us to examine in more detail the LRLD and LD blocks in these QTLRs, with all
273 informative markers of the five F₆ families (note that this part used *all* pairs of
274 informative markers in the QTLRs, and not only a *sample* of random pairs as in the first
275 LD analysis).

276 Chromosomes 1, 2, 4, 5, 6 and 14, harbored more than one QTLR (Table 1), thus
277 enabling examination of LD in and between QTLRs. In each F₆ family, Affymetrix SNP
278 array genotypes were used to calculate LD between all possible pairs of all markers in the
279 21 QTLRs on those chromosomes.

280 Hundreds of thousands of LRLDs were found in and between the tested QTLRs
281 (Supplemental Table 7). Total number of marker pairs ranged from below 8 to above 10
282 million in a family, to a total of more than 43 million pairs. Of these, pooled over all
283 families, 830,182 were LRLDs (62,103 - 227,015 LRLDs in a family). These constitute
284 0.7 - 2.6% of all pairs in a family, a total of 1.9%, higher than the 1.5% found among the
285 random pairs over all autosomes (Supplemental Table 3).

286 A total of 161,832 LRLDs were found between QTLRs (Supplemental Table 7),
287 19.5% of all LRLDs found (0.6 - 24.9 % among the families).

288 Family 5 is an outlier in Supplemental Table 7, with a much lower number and
289 proportion of total LRLDs and LRLDs across QTLRs compared to the other four

290 families. Further inspection did not identify any source of this difference. Hence, we have
291 no explanation other than sampling variation.

292

293 *Supplemental Table 7. Number of pairs and sum of QTLR LRLDs.*

294

295 Pooling all families together, LRLDs were found in all 6 chromosomes examined
296 (Table 5). No LRLD could be found *in* the QTLRs on Chromosomes 5 or 6 (Table 5), as
297 no QTLR there was larger than 1 Mb (Table 1). However, LRLDs *between* QTLRs were
298 also found in those two chromosomes.

299

300 *Table 5. Chromosomes and LRLDs in and between QTLRs.*

301

302 In all families, LRLDs were found between most pairs of QTLRs (Supplemental Table
303 8 a-f). Exceptional among all pairs of QTLRs, an extremely large number of LRLDs
304 (159,413) was found between QTLRs 4 and 5 on Chr 1 in all families, confirming the
305 results of the random samples (Figure 2). The tight LD between these two QTLRs was
306 further confirmed by the LD blocks (below).

307

308 *Supplemental Table 8. Number of arrays F₆ LRLDs in and between QTLRs in all*
309 *families.*

310

311 Thus, repeating in all F₆ families, LRLDs were found to be frequent, distributing
312 within and between QTLRs in all chromosomes tested.

313

314 **QTLR LD blocks**

315 **LD Blocks in the F₆ QTLR**

316 As shown by the data a complicated LD pattern was found in the F₆ QTLR. Large,

317 fragmented, and interdigitated LD blocks were found in all five families over all six
318 chromosomes examined (LD matrices in Genetics figshare portal). The range and
319 complexity would have been even larger if moderate LD blocks were included, with 0.15
320 $\leq r^2 < 0.7$.

321 An example of fragmented interdigitated blocks is presented in Figure 3a. Close
322 examination of the LD found in Family 1 in this region shows the presence of 3 high LD
323 blocks, all fragmented and all interdigitated with one another: Block 1 includes markers
324 with ID numbers 134-141, 143, 145-149, and 151; Block 2 includes markers 142, 144
325 and 152; Block 3 includes markers 150 and 337. The fact that, despite their apparent
326 fragmentation, these are indeed genuine blocks is shown in Figures 3 c-d. If the markers
327 in Blocks 2 and 3 were not included in the analysis, (e.g., because they were not on the
328 SNP array or were filtered out by the quality control or were not polymorphic in this
329 family), then three clear unambiguous blocks would have been identified.

330 Note the distance between the markers in block 3, is above 0.5 Mb. Should the
331 criterion of 0.25 Mb [Koch et al., 2013] been used, this block would be defined as LRLD.
332

333 **Figure 3.** *Fragmented interdigitated blocks in QTLR2 on chr 1 found in F₆ Family 1.*

334

335 **Blocks shared by QTLRs 4 and 5 in the F₆ families.** In accordance with the random
336 sampling of marker pairs and LRLDs in and between QTLRs in the F₆, large and long-
337 range LD blocks were shared by QTLRs 4 and 5 in all five families, as exemplified in
338 Supplemental Figure 1 and detailed in Supplemental Tables 8 a-f. In Supplemental Figure
339 1, the high LD block distributed from the first marker of QTLR 4 to close to the end of
340 QTLR 5, over 5.7 Mb, with 412 markers included. Considering moderate LD of $0.15 \leq r^2$
341 < 0.7 , would stretch the block all the way to the end of QTLR 5, over more than 7.1 Mb.
342 Thus, the exceptional LD between QTLRs 4 and 5 indicated by the random sample of
343 pairs was confirmed in all F₆ families by both LRLDs and LD blocks between QTLRs.

344

345 **Supplemental Figure 1.** *LD blocks shared by QTLRs 4 and 5 in Family 2.*

346

347 **LD among QTLR elements in the eight pure lines**

348 LD of elements within and between the F₆ QTLRs was further examined within eight
349 Hy-Line elite pure lines. Complex LD blocks between elements within and across
350 QTLRs were found, similar to that found in the F₆ families, over distances from a few bp
351 to a few Mb (Figure 4-6; all LD matrices are in Genetics figshare portal).

352 **LD within one QTLR gene.** Figure 4 present an example of LD blocks within the QTLR
353 gene *TRANK1* in Line WL1. Despite the short distances (390 bp to 14.5 Kb), a complex
354 pattern was found, with 2 LD blocks, one of which is fragmented around the other. There
355 was high to complete LD between markers 5, 8 and 13-36. These markers had practically
356 no LD with markers 11 and 12, which were in complete LD with one another. Thus, in
357 the gene *TRANK1* in Line WL1, Block 1 starts before, but ends after Block 2. The
358 association test P values [Smith et al., 2020] completely matched the LD blocks, with the
359 same or close P values in each block. This match was found in all other combinations of
360 QTLR - line (Figures 5 - 7).

361

362 *Figure 4. LD within one QTLR gene.*

363

364 **LD between QTLR elements.** An example of a more complex LD pattern with
365 interdigitated blocks is shown in Figure 5, this time across QTLR elements (3 lncRNAs).

366

367 *Figure 5. LD blocks across QTLR elements.*

368

369 Careful inspection of Figure 5 shows 2 interdigitated blocks: Block 1 includes
370 Markers 6-8, 15, and 19-20; Block 2 comprise of Markers 11-14, 16-17 and 30-33. Thus,
371 the high LD Block extend over the 3 QTLR lncRNAs. The middle lncRNA05 is split
372 among the 2 blocks. Some of the markers are in LD with upstream lncRNA02, while
373 other markers of the same lncRNA05 form a block with the downstream lncRNA02. The

374 2 groups of lncRNA05 are interdigitated. That is, Markers 6-8 of lncRNA02 are in the
375 same block with 2 separate regions in the next lncRNA05 - Markers 15 and then 19-20
376 but not with the other markers in the same lncRNA; Markers 12-14 and 19-20 of
377 lncRNA05 are in LD with all 4 markers of lncRNA04. It would be interesting to find out
378 what are the sources of such complex LD patterns.

379 LD was found between other types of QTLR elements as well. Figure 6 present such
380 LD between the QTLR genes *TLR4* and *BRINP1* in QTLR 33 on Chr 17. The first marker
381 of *TLR4* has high LD to the first 2 markers of *BRINP1*, and the 3 markers are not linked
382 to other markers of their own gene. The other 6 markers of *TLR4* form a tight LD block.
383 Complexing it even further, the last marker of *BRINP1* (Marker 15) had low to moderate
384 LD with all markers in QTLR 33, both genes included.

385

386 **Figure 6.** LD between QTLR genes.

387

388 **LD between QTLRs 4 and 5.** Markers on both QTLRs 4 and 5 were informative only in
389 Lines WL3, WPR1, WPR2, and RIR1, up to only 4 markers in a line in QTLR 4 (Lines'
390 LD matrices in Genetics figshare portal). Thus, information on the LD between the
391 QTLRs was limited in this dataset. Nevertheless, in accord with the random LDs in the F₆
392 families (Figures 1 and 2) and cross QTLRs LRLD in these families (Supplemental
393 Tables 8 b-f), moderate LD blocks among elements in these QTLRs crossed their
394 boundaries in Lines WPR1, WPR2 and RIR1. In Line WRP1, 2 clear high LD blocks
395 were found, one in each QTLR (Figure 7). However, the 2 QTLR blocks had moderate
396 LDs of $r^2 = 0.478$ among them, thus forming one moderate LD block. Note that the
397 distances between the cross QTLR pairs, varied from 5.135 to 5.138 Mb.

398

399 **Figure 7.** LD between QTLRs 4 and 5 in a pure line.

400

401 Looking for a source of such vast, high, and complex long-range LD between QTLRs
402 4 and 5, a bioinformatics search found 10 and 68 genes in QTLRs 4 and 5, respectively

403 (Figure 8 and Table 6). STRING network analysis revealed five networks of 2 to 28
404 genes. Two of the networks ('Net' 2 and 3 in Table 6), are comprised of genes from both
405 QTLRs (Figure 8 and Table 6). Of 'Net' 2, the 2 genes in QTLR 4 and 17 of the 26 genes
406 in QTLR 5 are located in the LD blocks extending over the two QTLRs found in F₆ ('+' in
407 the column 'B4-5' in Table 6). Both genes in 'Net' 3 are in those blocks. Finally, the two
408 networks with genes from both QTLRs included 6 genes interacting with a gene from
409 another QTLR (Figure 8), all of which located in the cross QTLRs LD blocks. The gene
410 networks and interactions shared by both QTLRs could be the origin of the LD between
411 QTLRs 4 and 5. In fact, the phenomenon of genes whose products work together tending
412 to be on the same chromosomal region is quite common. For example, the Major
413 Histocompatibility Complex (MHC) on chicken chromosome 16 and the Regulators of
414 Complement Activation cluster (RCA) on chromosome 26 [Oshiumi et al, 2005;
415 Hosomichi et al, 2008; Michilak, 2008]. In fact, the networks presented in Figure 8 is a
416 good examples for this collocation of genes working together.

417

418 **Table 6.** *Genes and protein networks in QTLRs 4 and 5.*

419

420 **DISCUSSION**

421 Chicken LD over a range of distances, and patterns of LD blocks, were studied in five
422 F₆ families from a FSAIL, and eight commercial pure layer lines, thus allowing to study
423 the repetition of the results. LRLD was studied in the F₆ population by random non-
424 syntenic and syntenic samples of marker pairs genotyped by the Affymetrix HD SNP
425 array. In face of the LRLD results, and to illustrate the importance of LRLD to QTL
426 mapping results, LRLD and LD blocks were studied with all possible marker pairs of all
427 markers in previously described MD QTLRs [Smith et al., 2020].

428 This study started with SNP location information from the previous chicken genome
429 build Galgal4, and was subsequently updated to the Galgal6 assembly. This change
430 necessitated remapping of the QTLRs described in Smith et al. (2020), resulting in
431 negligible changes of most QTLR coordinates. Nevertheless, the change of genome

432 versions moved a segment of Galgal4 QTLR 1 (chr 1) to Galgal6 QTLR 4 (chr 1)
433 (Appendix). The moved segment included one QTLR lncRNA described in Smith et al.
434 (2020), thus emphasizing the importance of basing genomic analyses on the most updated
435 genome version. These results also present the power of LD to identify mapping errors,
436 as already noted by Utsunomiya et al. (2016).

437 Long-range LD (LRLD) was defined as $r^2 \geq 0.7$ over a distance ≥ 1 Mb. These criteria
438 are more restricted than previously used [Koch et al., 2013; Vallejo et al., 2018].
439 Nevertheless, repeated appreciable numbers of LRLDs within chromosomes were found
440 repeated in all five F₆ families by the random sampling of syntenic marker pairs, far
441 above the numbers of high LD found between non-syntenic markers from different
442 chromosomes. The LRLDs were further found in all five F₆ families by all QTLR array
443 markers on chromosomes with more than one QTLR. These results could be an
444 underestimate, as the F₆ population was designed to fragment the genome for high-
445 resolution QTL mapping [Heifetz et al. 2007, 2009].

446 High LD blocks were defined as a group of markers located on the same chromosome,
447 having $r^2 \geq 0.7$ with each other, even if markers with low LD appeared between them.
448 This definition allowed "a look over the horizon" and identification of complex blocks.
449 The phenomenon of fragmented and interdigitated LD blocks were repeatedly found in
450 all five F₆ families and in all eight pure lines over a vast range of distances, from
451 hundreds of bp to mega bases. The FSAIL population was composed of five families, and
452 they showed similar results. Strength of this analysis was that it repeated five times. Then
453 the same phenomenon was seen in the eight elite lines, further adding strength to the
454 validity of these results, which also agree with previous studies from us and others [e.g.,
455 Aerts et al., 2007; Allabi et al., 2005; Lipkin et al., 2013; O'Brien et al., 2014].

456 A strong linkage was found between QTLRs 4 and 5. LRLD between them was found
457 while analyzing the F₆ random samples of SNPs within all autosomes. High LD blocks
458 were found in all five F₆ families, comprised of markers from both QTLRs. Moderate LD
459 blocks between QTLRs 4 and 5 were also found in 3 of the 8 pure lines by QTLR
460 elements' markers. These results raise the question of what elements on both QTLRs are
461 in high LD over such large distances. Of course, being MD QTLRs, the LD between the

462 QTLRs could be a result of a co-selection for MD resistance. But what make these two
463 QTLRs so different from all other pairs of QTLRs? Why is their LD so exceptional?

464 To answer this, we looked at the gene content of QTLRs 4 and 5. Ten and 68 genes
465 were found in QTLRs 4 and 5, respectively. STRING protein network analysis revealed
466 five gene networks. Two of the networks include genes from both QTLRs, most of which
467 are located within the LD blocks extending over the two QTLRs found in the F₆. All 6
468 genes interacting with a gene from another QTLR are in the cross QTLRs LD blocks.
469 Obviously, the shared gene networks and interactions could be the origin of the LD
470 between QTLRs 4 and 5. However, assessing the uniqueness of the LD between the
471 QTLRs necessitates further study on the distribution of cross QTLR networks and
472 interactions among other pairs of QTLRs with less LD among them. Furthermore,
473 assessing the real effect of the gene networks and interactions needs more molecular,
474 quantitative and population studies. All of these are beyond the scope of the present
475 study.

476 In general, the complex LD found in this study could stem from technical reasons such
477 as sampling variation. It can also be a result of mapping errors, as was indeed found in
478 this study for regions on chromosome 1 with build 4 (Appendix). However, it could have
479 genuine biological meaning, through processes such as co-evolution of genomic sites (as
480 a result natural or artificial selection), gene conversion, copy number variation,
481 population bottlenecks, non-random mating, and epistasis. The repeatability in different
482 analyses, different datasets and different populations, and the agreement with previous
483 reports strengthen the case for the present results as being a genuine biological
484 phenomenon.

485 The observation of both LRLD and fragmented interdigitated blocks imply that the
486 causative element is not necessarily the closest, or even close at all to the significant
487 marker, and maybe not even to the significant block of markers. Thus, mapping results
488 and searches for causative elements should be taken with caution, by considering the
489 complexity of the LD. All sites with high LD with a significant marker are in effect
490 candidates to be the causative element.

491

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494

495 **Availability of Data:** Data have been submitted to the European Nucleotide Archive
496 (ENA) at EMBL-EBI under study accession numbers PRJEB39142 (WGS) and
497 PRJEB39361 (RNAseq). LD matrices are available at figshare.

498

499 REFERENCES

- 500 1. Aerts J, Megens HJ, Veenendaal T, Ovcharenko I, Crooijmans R, Gordon L, Stubbs
501 L, Groenen M. Extent of linkage disequilibrium in chicken. *Cytogenet Genome Res.*
502 2007;117(1-4):338-45. doi: 10.1159/000103196. PMID: 17675876.
- 503 2. Allabi AC, Horsmans Y, Issaoui B, Gala JL. Single nucleotide polymorphisms of
504 ABCB1 (MDR1) gene and distinct haplotype profile in a West Black African
505 population. *Eur J Clin Pharmacol.* 2005 Apr;61(2):97-102.
- 506 3. Ardlie, K., Kruglyak, L. & Seielstad, M. Patterns of linkage disequilibrium in the
507 human genome. *Nat Rev Genet* **3**, 299–309 (2002). <https://doi.org/10.1038/nrg777>.
- 508 4. Corbin LJ, Blott SC, Swinburne JE, Vaudin M, Bishop SC, Woolliams JA. Linkage
509 disequilibrium and historical effective population size in the Thoroughbred horse.
510 *Anim Genet.* 2010 Dec;41 Suppl 2:8-15. doi: 10.1111/j.1365-2052.2010.02092.x.
511 PMID: 21070270.
- 512 5. García-Gámez, E., Sahana, G., Gutiérrez-Gil, B., & Arranz, J. J. (2012). Linkage
513 disequilibrium and inbreeding estimation in Spanish Churra sheep. *BMC genetics*, *13*,
514 43. <https://doi.org/10.1186/1471-2156-13-43>.
- 515 6. Haeussler M, Zweig AS, Tyner C, Speir ML, Rosenbloom KR, Raney BJ, Lee CM,
516 Lee BT, Hinrichs AS, Gonzalez JN, Gibson D, Diekhans M, Clawson H, Casper J,
517 Barber GP, Haussler D, Kuhn RM, Kent WJ. [The UCSC Genome Browser](#)
518 [database: 2019 update](#). *Nucleic Acids Res.* 2019 Jan 8;47(D1):D853-D858.
519 PMID: [30407534](#); PMC: [PMC6323953](#)
- 520 7. Heifetz, E.M.; Fulton, J.E.; O’Sullivan, N.P.; Arthur, J.A.; Wang, J.; Dekkers, J.C.;

- 521 Soller, M. Mapping quantitative trait loci affecting susceptibility to Marek's disease
522 virus in a backcross population of layer chickens. *Genetics* **2007**, *177*, 2417–2431.
523 [[CrossRef](#)] [[PubMed](#)]
- 524 8. Heifetz, E.M.; Fulton, J.E.; O'Sullivan, N.P.; Arthur, J.; Cheng, H.; Wang, J.; Soller,
525 M.; Dekkers, J.C.M. Mapping QTL affecting resistance to Marek's disease in an F6
526 advanced intercross population of commercial layer chickens. *BMC Genom.* **2009**,
527 *10*, 20. [[CrossRef](#)]
- 528 9. Hosomichi K, Miller MM, Goto RM, Wang Y, Suzuki S, Kulski JK, Nishibori M,
529 Inoko H, Hanzawa K, Shiina T. Contribution of mutation, recombination, and gene
530 conversion to chicken MHC-B haplotype diversity. *J Immunol.* 2008 Sep
531 1;181(5):3393-9. doi: 10.4049/jimmunol.181.5.3393. Erratum in: *J Immunol.* 2010
532 May 1;184(9):5415. PMID: 18714011; PMCID: PMC2657362.
- 533 10. Jensen LJ, Kuhn M, Stark M, Chaffron S, Creevey C, Muller J, Doerks T, Julien P,
534 Roth A, Simonovic M, Bork P, von Mering C. STRING 8--a global view on proteins
535 and their functional interactions in 630 organisms. *Nucleic Acids Res.* 2009
536 Jan;37(Database issue):D412-6. doi: 10.1093/nar/gkn760. Epub 2008 Oct 21. PMID:
537 18940858; PMCID: PMC2686466.
- 538 11. Khatkar, M.S., Nicholas, F.W., Collins, A.R. *et al.* Extent of genome-wide linkage
539 disequilibrium in Australian Holstein-Friesian cattle based on a high-density SNP
540 panel. *BMC Genomics* **9**, 187 (2008). <https://doi.org/10.1186/1471-2164-9-187>.
- 541 12. Koch E, Ristroph M, Kirkpatrick M (2013) Long Range Linkage Disequilibrium
542 across the Human Genome. *PLoS ONE* 8(12): e80754.
543 <https://doi.org/10.1371/journal.pone.0080754>.
- 544 13. Kranis, A.; Gheyas, A.A.; Boschiero, C.; Turner, F.; Yu, L.; Smith, S.; Talbot, R.;
545 Pirani, A.; Brew, F.; Kaiser, P.; et al. Development of a high density 600K SNP
546 genotyping array for chicken. *BMC Genom.* **2013**, *14*, 59. [[CrossRef](#)]
- 547 14. Ehud Lipkin, Marlies Dolezal, Alessandro Bagnato, Janet Fulton, Petek
548 Settar, Jesus Arango, Neil O'Sullivan, Attilio Rossoni, Theo Meuwissen,
549 Kellie Watson, Morris Soller. 2013. Linkage Disequilibrium and Haplotype
550 Blocks in Cattle and Chicken Populations. *Proceedings of the Plant &*

- 551 Animal Genomes XXI Conference, San Diego USA, P0680: Genome
552 Mapping, Tagging & Characterization: General Comparative.
- 553 15. Michalak P. Coexpression, coregulation, and cofunctionality of
554 neighboring genes in eukaryotic genomes. *Genomics*. 2008 Mar;91(3):243-
555 8. doi: 10.1016/j.ygeno.2007.11.002. Epub 2007 Dec 21. PMID: 18082363.
- 556 16. Pérez O'Brien, A.M., Utsunomiya, Y.T., Mészáros, G. et al. Assessing
557 signatures of selection through variation in linkage disequilibrium between
558 taurine and indicine cattle. *Genet Sel Evol* 46, 19 (2014).
559 <https://doi.org/10.1186/1297-9686-46-19>.
- 560 17. Oshiumi H, Shida K, Goitsuka R, Kimura Y, Katoh J, Ohba S, Tamaki Y,
561 Hattori T, Yamada N, Inoue N, Matsumoto M, Mizuno S, Seya T.
562 Regulator of complement activation (RCA) locus in chicken: identification
563 of chicken RCA gene cluster and functional RCA proteins. *J Immunol*.
564 2005 Aug 1;175(3):1724-34. doi: 10.4049/jimmunol.175.3.1724. PMID:
565 16034113.
- 566 18. Park, L. Population-specific long-range linkage disequilibrium in the human genome
567 and its influence on identifying common disease variants. *Sci Rep* **9**, 11380 (2019).
568 <https://doi.org/10.1038/s41598-019-47832-y>.
- 569 19. Peters SO, Kızılkaya K, Ibeagha-Awemu EM, Sinecen M, Zhao X. Comparative
570 accuracies of genetic values predicted for economically important milk traits,
571 genome-wide association, and linkage disequilibrium patterns of Canadian Holstein
572 cows. *Journal of Dairy Science*. 2021 Feb;104(2):1900-1916. DOI: 10.3168/jds.2020-
573 18489.
- 574 20. Reich, D., Cargill, M., Bolk, S. *et al*. Linkage disequilibrium in the human
575 genome. *Nature* **411**, 199–204 (2001). <https://doi.org/10.1038/35075590>.
- 576 21. M. Sargolzaei, F.S. Schenkel, G.B. Jansen, L.R. Schaeffer, Extent of Linkage
577 Disequilibrium in Holstein Cattle in North America, *Journal of Dairy Science* 91,
578 Issue 5, 2008, Pages 2106-2117, ISSN 0022-0302, [https://doi.org/10.3168/jds.2007-](https://doi.org/10.3168/jds.2007-0553)
579 [0553](https://doi.org/10.3168/jds.2007-0553).

- 580 22. Seo D, Lee DH, Choi N, Sudrajad P, Lee SH, Lee JH. Estimation of linkage
581 disequilibrium and analysis of genetic diversity in Korean chicken lines. *PLoS One*.
582 2018 Feb 9;13(2):e0192063. doi: 10.1371/journal.pone.0192063. PMID: 29425208;
583 PMCID: PMC5806858.
- 584 23. Skelly, D. A., Magwene, P. M., & Stone, E. A. (2016). Sporadic, Global Linkage
585 Disequilibrium Between Unlinked Segregating Sites. *Genetics*, 202(2), 427–437.
586 <https://doi.org/10.1534/genetics.115.177816>.
- 587 24. Smith, J.; Lipkin, E.; Soller, M.; Fulton, J.E.; Burt, D.W. Mapping QTL Associated
588 with Resistance to Avian Oncogenic Marek’s Disease Virus (MDV) Reveals Major
589 Candidate Genes and Variants. *Genes* 2020, 11,
590 1019. <https://doi.org/10.3390/genes11091019>.
- 591 25. Sved J.A. (1971) Linkage disequilibrium and homozygosity of chromosome segments
592 in finite populations. *Theoretical Population Biology* 2, 125–41.
- 593 26. Utsunomiya, A.T.H., Santos, D.J.A., Boison, S.A. et al. Revealing misassembled
594 segments in the bovine reference genome by high resolution linkage disequilibrium
595 scan. *BMC Genomics* 17, 705 (2016). <https://doi.org/10.1186/s12864-016-3049-8>.
- 596 27. Roger L. Vallejo, Rafael M. O. Silva, Jason P. Evenhuis, Guangtu Gao, Sixin Liu,
597 James E. Parsons, Kyle E. Martin, Gregory D. Wiens, Daniela A. L. Lourenco,
598 Timothy D. Leeds, Yniv Palti. Accurate genomic predictions for BCWD resistance in
599 rainbow trout are achieved using low-density SNP panels: Evidence that long-range
600 LD is a major contributing factor. *J Anim Breed Genet*. 2018; 135(4):263–74.
601 <https://doi.org/10.1111/jbg.12335>.

602

603

APPENDIX - Genome assemblies

604

As stated in the Methods, this analysis started based on the Galgal4 genomic build.

605

Then [Lift Genome Annotations](#) tool [Haeussler et al., 2019] of [UCSC](#) Browser was used

606

to move from Galgal4 to Galgal6 (GRCg6a). The new coordinates were then used to

607

redefine the F₆ QTLRs as was done by Smith et al. (2020). The same LD analyses were

608

carried on both assemblies, results were similar, and hence only the results on Galgal6 are

609

presented in the body of the paper. Nevertheless, there was one change worthy of note,

610

and here we show the effect of updating the genome build (moving from Galgal4 to

611

Galgal6), and thus the ability to locate assembly errors by LD.

612

Using *Galgal4*, almost all high syntenic LDs of $r^2 \geq 0.7$ were below 20 Mb, as

613

expected (Appendix Table 1).

614

Appendix Table 1. Galgal4. Distance distribution of high LD syntenic marker pairs

615

randomized within chromosomes.

Distance (Mb)	Fam1	Fam2	Fam3	Fam4	Fam5	All
≥ 20	10,710	9,792	9,740	9,986	10,193	50,421
>20 - ≥ 30	0	0	0	0	0	0
>30 - ≥ 40	0	0	0	0	0	0
>40 - ≥ 50	0	0	0	0	0	0
>50 - ≥ 60	0	0	0	0	0	0
>60 - ≥ 70	1	3	4	3	2	13
>70 - ≥ 80	0	0	0	0	0	0
>80 - ≥ 90	0	0	0	0	0	0
>90 - ≥ 100	0	0	0	0	0	0
>100 - ≥ 110	0	0	0	0	0	0
>110 - ≥ 120	0	0	0	0	0	0
>120 - ≥ 130	0	0	0	0	0	0
>130 - ≥ 140	0	0	0	0	0	0
>140 - ≥ 150	0	0	0	0	0	0
>150 - ≥ 160	0	0	0	0	0	0
>160 - ≥ 170	0	0	0	0	0	0
>170 - ≥ 180	0	0	0	0	0	0
>180 - ≥ 190	0	0	0	0	0	0
>190 - ≥ 200	0	0	0	0	0	0
Sum	10,711	9,795	9,744	9,989	10,195	50,434

616

617

Unexpectedly, in all 5 families few high LDs were found with distances from above

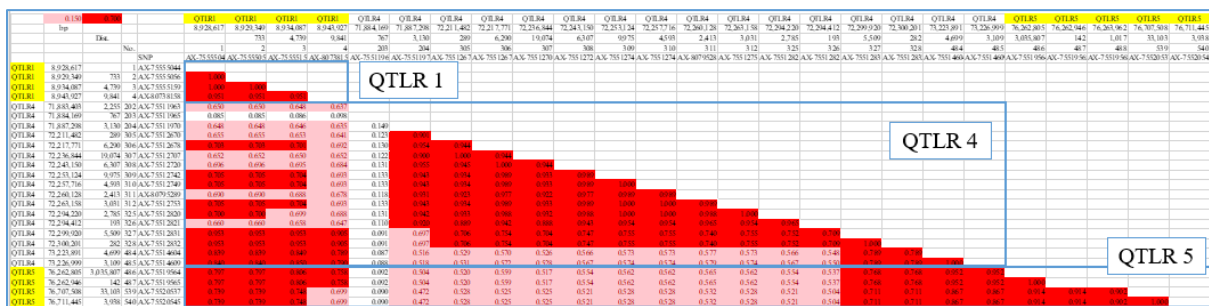
618

60 to 70 Mb (bolded in Appendix Table 1). This was the only interval with high LD

619 above 20 Mb. Detail examination of those 13 extreme long range LDs (LRLDs), showed
 620 that all markers in these very distant pairs involved the same 3 regions on chromosome 1.
 621 Intriguing, all markers were within or very close to 3 of Smith et al. (2020) F₆ MD QTL
 622 regions (QTLRs). In each pair, one marker was from QTLR 1, the other from QTLRs 4
 623 or 5. Although these 3 QTLRs were significant only in Family 3, the extreme LRLDs
 624 were found in all 5 families.

625 In the face of these results, we LD together all three regions involved (regions of the
 626 markers in the 13 LDLRs, not the entire QTLR). As presented in the body of the paper
 627 for Galgal6, a very complex LD pattern was obtain in Family 1. A representative example
 628 is presented in Appendix Figure 1. The entire matrix of 540 SNPs is too large to present,
 629 hence only a sample of markers is presented the give general impression of the LD
 630 pattern obtained.

631 **Appendix Figure 1. Galgal4.** Representative matrix in Family 1 of the 3 regions with
 632 markers from LRLDs > 60 Mb. These regions are parts of QTLRs 1, 4 and 5. Yellowed
 633 on the flanks, QTLRs 1 and 5, with the white QTLR 4 between them; blue frame, a
 634 QTLR; Red, $r^2 \geq 0.7$; purple, $0.15 < r^2 < 0.7$.



635
 636 On Galgal4 there were over 62.2 Mb between QTLRs 1 and 4, and over 3.0 Mb
 637 between 4 and 5. Nevertheless, very high to complete LD values were obtained. As
 638 presented in the body of the paper, fragmented interdigitated LD blocks were found,
 639 shown by the purple and white rows and columns interfering the red regions.

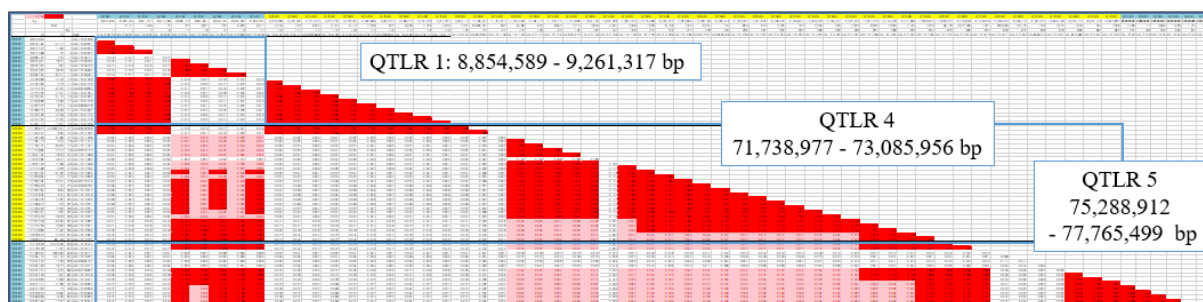
640 In light of these results we, LD together the entire 3 QTLRs in each 5 F₆ family.
 641 Appendix Figure 2 present an examples of the analysis of the whole QTLRs, again in
 642 Family 1. Similar results with some differences were found in all families.

643 Close examination of Appendix Figure 1 show 2 high LD blocks cover together all

644 QTLR 1, completely unlinked with one another (shown by the white rectangle below and
 645 left to the upper left and the lower right red triangles). One of the QTLR 1 blocks is
 646 fragmented, covering the start and the end of this QTLR.

647 The fragmented block of QTLR1 is completely linked with the start of QTLR 4, 62
 648 Mb downstream. The second un-fragmented block of QTLR 1 has moderate to high LD
 649 of $0.15 \geq r^2 < 0.7$ with the fragmented interdigitated blocks of QTLRs 4 and 5.

650 **Appendix Figure 2. Galgal4.** Representative matrix of Family 1 of the entire QTLRs 1,
 651 4 and 5 together. The entire matrix of 602 SNPs is too large to present, hence only a
 652 sample of markers is presented to give a panoramic view of the LD pattern obtained.
 653 Blued on the flanks, QTLRs 1 and 5 with yellowed QTLR 4 between them; blue frame,
 654 QTLR; Text boxes present the boundaries of the QTLRs; Red, LD $r^2 \geq 0.7$; purple, $0.15 <$
 655 $r^2 < 0.7$.



656

657 Moving from Galgal4 to Galgal6, more than 99% of the markers were mapped
 658 successfully in Galgal6. Few hundreds markers were deleted in the new assembly, 0.3%
 659 of the markers submitted to Lift. Very few markers even changed chromosomes.

660 After QTLR mapping as in Smith et al. (2020), the same 38 QTLRs reported by Smith
 661 et al. (2020) were found. Nevertheless, lnrRNA01 that was located in QTLR 1 on
 662 Galgal4 was located in QTLR 4 on Galgal6. Furthermore, Galgal6 completely eliminated
 663 the LRLD between QTLR 1 and the other 2 QTLRs, and zeroed that 60 - 70 Mb high
 664 LDs (Appendix Table 2).

665 **Table 4. Galgal6.** Distance distribution of high LD syntenic marker pairs randomized
 666 within chromosomes.

Distance (Mb)	Fam1	Fam2	Fam3	Fam4	Fam5	All
≥ 20	10,643	9,736	9,687	9,927	10,115	50,108

Distance (Mb)	Fam1	Fam2	Fam3	Fam4	Fam5	All
>20 - ≥30	0	0	0	0	0	0
>30 - ≥40	0	0	0	0	0	0
>40 - ≥50	0	0	0	0	0	0
>50 - ≥60	0	0	0	0	0	0
>60 - ≥70	0	0	0	0	0	0
>70 - ≥80	0	0	0	0	0	0
>80 - ≥90	0	0	0	0	0	0
>90 - ≥100	0	0	0	0	0	0
>100 - ≥110	0	0	0	0	0	0
>110 - ≥120	0	0	0	0	0	0
>120 - ≥130	0	0	0	0	0	0
>130 - ≥140	0	0	0	0	0	0
>140 - ≥150	0	0	0	0	0	0
>150 - ≥160	0	0	0	0	0	0
>160 - ≥170	0	0	0	0	0	0
>170 - ≥180	0	0	0	0	0	0
>180 - ≥190	0	0	0	0	0	0
>190 - ≥200	0	0	0	0	0	0
Sum	10,643	9,736	9,687	9,927	10,115	50,108

667

668

669

These results show the limitations of any genome build, and present the ability of LD analysis to find assembly errors.

1 **Complex and long-range linkage disequilibrium in chicken populations**

2 **Tables**

3 **Table 1.** Remapping Galgal4 QTLRs on Galgal6. QTLR, ordinal number of the QTLR
 4 [Smith et al., 2020]; Chr, chromosome; Start/End, QTLR coordinates of the first and last
 5 SNP in the QTLR; Length, size of the QTLR (bp).

QTLR	Chr	Galgal4			Galgal6		
		Start	End	Length	Start	End	Length
1	1	8,854,589	9,261,317	406,729	9,510,148	9,902,036	391,889
2	1	13,268,810	14,294,877	1,026,068	13,994,599	14,950,768	956,170
3	1	52,012,381	52,517,953	505,573	52,166,588	52,643,244	476,657
4	1	71,738,977	73,085,956	1,346,980	71,892,917	73,277,481	1,384,565
5	1	75,288,912	77,765,499	2,476,588	75,513,671	79,029,197	3,515,527
6	1	91,482,501	91,803,475	320,975	93,533,567	93,853,587	320,021
7	1	101,761,945	104,477,713	2,715,769	103,738,415	106,416,920	2,678,506
8	1	109,395,871	110,913,451	1,517,581	111,372,866	112,400,685	1,027,820
9	1	169,656,958	172,228,091	2,571,134	171,680,812	174,306,953	2,626,142
10	1	174,394,951	175,634,386	1,239,436	176,474,702	177,748,402	1,273,701
11	1	194,193,118	194,788,548	595,431	196,152,404	196,750,875	598,472
12	2	15,960	883,257	867,298	48,621	959,053	910,433
13	2	46,423,161	46,868,789	445,629	45,786,534	46,247,754	461,221
14	2	105,493,833	108,988,920	3,495,088	105,791,822	109,334,178	3,542,357
15	2	125,168,753	127,089,304	1,920,552	125,532,963	127,219,187	1,686,225
16	2	138,767,830	139,701,774	933,945	139,198,404	140,160,087	961,684
17	3	108,220,206	109,260,649	1,040,444	108,593,746	109,643,999	1,050,254
18	4	8,308,498	11,268,107	2,959,610	8,328,709	11,309,259	2,980,551
19	4	84,393,173	88,579,400	4,186,228	84,829,085	89,057,374	4,228,290
20	5	7,568,851	8,147,837	578,987	8,388,371	8,967,466	579,096
21	5	18,806,924	19,673,354	866,431	19,753,005	20,610,009	857,005
22	6	2,077,640	2,709,412	631,773	3,323,132	3,946,659	623,528
23	6	29,536,109	29,817,337	281,229	30,954,349	31,233,344	278,996
24	6	31,006,769	31,448,342	441,574	32,440,880	32,888,648	447,769
25	7	13,062,871	16,436,053	3,373,183	13,563,779	16,986,311	3,422,533
26	10	22,643	1,713,384	1,690,742	1,025,523	2,668,959	1,643,437
27	11	7,397,790	8,440,259	1,042,470	7,912,510	8,959,749	1,047,240
28	12	8,996,686	9,432,693	436,008	9,414,714	9,845,036	430,323
29	13	10,363,430	12,176,727	1,813,298	11,756,937	13,566,822	1,809,886
30	14	8,085,563	9,335,685	1,250,123	8,499,374	9,745,708	1,246,335
31	14	13,138,194	15,087,518	1,949,325	13,542,085	15,384,231	1,842,147
32	16	1,630	490,907	489,278	1,852,095	2,669,032	816,938
33	17	3,442,598	5,634,042	2,191,445	3,808,082	5,932,858	2,124,777
34	18	3,196,488	4,093,129	896,642	3,221,049	4,118,252	897,204

QTLR	Chr	Gagal4			Galgal6		
		Start	End	Length	Start	End	Length
35	24	4,489,675	5,514,833	1,025,159	4,160,414	5,498,172	1,337,759
36	26	4,378,168	5,036,699	658,532	4,438,584	5,002,302	563,719
37	27	1,540,112	2,270,461	730,350	3,930,559	4,689,821	759,263
38	28	1,282,726	1,571,011	288,286	1,447,725	1,687,264	239,540

6

7 **Table 2.** Summary statistics of non-syntenic random LD between 600K markers in the F₆
8 families. Pairs, number of pairs r² values obtained; Ind/pairs, average number of
9 individuals used to calculate a markers pair LD in a family; r²: Avg, average; SD, standard
10 deviation; Min, minimum; Max, maximum; All: Pairs, average number of pairs in all
11 families combined; Ind/pairs r², means weighted by the number of Pairs.

Family	Pairs	Ind/ pair	r ²			
			Avg	SD	Min	Max
1	190,232	176.6	0.014	0.019	0.000	0.368
2	182,366	231.7	0.012	0.016	0.000	0.991
3	184,399	354.0	0.008	0.011	0.000	0.372
4	187,406	219.5	0.010	0.014	0.000	0.246
5	178,780	200.7	0.012	0.016	0.000	0.243
All	184,636.6	236.3	0.011	0.016	0.000	0.443

12

13

14 **Table 3.** Summary statistics of syntenic random LD between 600K markers in the F₆
 15 families. Pairs: No., number of pairs r² values obtained; Ind, average number of
 16 individuals used to calculate a markers pair LD in a family; bp: average, minimum and
 17 maximum bp between markers in a pair; r²: Avg, average; SD, standard deviation; Min,
 18 minimum; Max, maximum; r(bp-r²), correlation between the distance and r² of a marker
 19 pair; All: Pairs, total number of pairs in all families combines; Ind, bp and r²: means of
 20 weighted by the number of pairs.

Family:		1	2	3	4	5	All
Pairs	No.	207,770	200,411	200,866	204,453	195,323	1,008,823
	Ind	176.6	231.7	354.0	219.5	200.7	236.2
bp	Avg	28,745,980.9	30,164,000.4	28,092,640.5	29,039,387.5	28,844,817.9	28,976,195.6
	Min	11	11	40	64	165	57.3
	Max	196,356,451	197,038,449	196,167,202	196,407,037	196,676,774	196,526,525.6
r ²	Avg	0.119	0.112	0.109	0.110	0.119	0.114
	SD	0.221	0.218	0.216	0.218	0.224	0.220
	Min	0.000	0.000	0.000	0.000	0.000	0.000
	Max	1.000	1.000	1.000	1.000	1.000	1.000
r(bp-r ²)		-0.346	-0.346	-0.347	-0.336	-0.351	-0.345

21

22

23 **Table 4.** Distribution of syntenic random LD values among the F₆ families. All, all
 24 families combine; $r^2 \geq$ (last 2 rows), frequencies of r^2 above the indicated value.

r^2	Fam1	Fam2	Fam3	Fam4	Fam5	All
≤ 0.05	0.6429	0.6778	0.6920	0.6920	0.6920	0.6707
$>0.05 - \leq 0.10$	0.1036	0.0898	0.0757	0.0757	0.0757	0.0879
$>0.10 - \leq 0.15$	0.0495	0.0451	0.0423	0.0423	0.0423	0.0457
$>0.15 - \leq 0.20$	0.0351	0.0277	0.0292	0.0292	0.0292	0.0315
$>0.20 - \leq 0.25$	0.0242	0.0192	0.0221	0.0221	0.0221	0.0219
$>0.25 - \leq 0.30$	0.0180	0.0165	0.0171	0.0171	0.0171	0.0174
$>0.30 - \leq 0.35$	0.0165	0.0138	0.0152	0.0152	0.0152	0.0152
$>0.35 - \leq 0.40$	0.0120	0.0110	0.0120	0.0120	0.0120	0.0122
$>0.40 - \leq 0.45$	0.0101	0.0101	0.0103	0.0103	0.0103	0.0102
$>0.45 - \leq 0.50$	0.0094	0.0092	0.0081	0.0081	0.0081	0.0091
$>0.50 - \leq 0.55$	0.0076	0.0087	0.0079	0.0079	0.0079	0.0080
$>0.55 - \leq 0.60$	0.0064	0.0079	0.0064	0.0064	0.0064	0.0070
$>0.60 - \leq 0.65$	0.0066	0.0076	0.0072	0.0072	0.0072	0.0069
$>0.65 - \leq 0.70$	0.0067	0.0069	0.0062	0.0062	0.0062	0.0066
$>0.70 - \leq 0.75$	0.0056	0.0070	0.0066	0.0066	0.0066	0.0062
$>0.75 - \leq 0.80$	0.0061	0.0062	0.0063	0.0063	0.0063	0.0060
$>0.80 - \leq 0.85$	0.0060	0.0063	0.0067	0.0067	0.0067	0.0063
$>0.85 - \leq 0.90$	0.0074	0.0068	0.0065	0.0065	0.0065	0.0068
$>0.90 - \leq 0.95$	0.0080	0.0073	0.0082	0.0082	0.0082	0.0081
$>0.95 - \leq 1.00$	0.0180	0.0151	0.0139	0.0139	0.0139	0.0162
Sum	1.0000	1.0000	1.0000	1.0000	1.0000	1.0000
$r^2 > 0.15$	0.2040	0.1872	0.1900	0.1886	0.2089	0.1957
$r^2 > 0.70$	0.0512	0.0486	0.0482	0.0486	0.0518	0.0497

25

26

27 **Table 5.** Chromosomes and LRLDs in and between QTLRs. Number of F₆ array LRLDs
 28 in and between QTLRs on the six chromosomes with more than one QTLR. Chr,
 29 chromosome; Fam, family; All, All families combined; In QTLRs, LRLDs within the MD
 30 QTLRs; Across QTLRs, LRLDs between QTLRs; Total, total number of LRLDs.

LRLDs	Chr	Fam1	Fam2	Fam3	Fam4	Fam5	All
In QTLRs	1	83,455	52,985	55,025	39,612	0	231,077
	2	15,854	5,195	0	0	0	21,049
	4	87,859	106,654	70,245	88,091	61,605	414,454
	5	0	0	0	0	0	0
	6	0	0	0	0	0	0
	14	503	511	489	116	151	1,770
	Sum	187,671	165,345	125,759	127,819	61,756	668,350
Between QTLRs	1	39,244	54,674	33,243	33,937	321	161,419
	2	52	122	1	1	0	176
	4	26	35	24	33	22	140
	5	1	1	0	0	0	2
	6	2	2	2	2	0	8
	14	19	32	8	24	4	87
	Sum	39,344	54,866	33,278	33,997	347	161,832
Total	1	122,699	107,659	88,268	73,549	321	392,496
	2	15,906	5,317	1	1	0	21,225
	4	87,885	106,689	70,269	88,124	61,627	414,594
	5	1	1	0	0	0	2
	6	2	2	2	2	0	8
	14	522	543	497	140	155	1,857
	Sum	227,015	220,211	159,037	161,816	62,103	830,182

31

32 **Table 6.** Genes and protein networks in QTLRs 4 and 5, ordered by location. Q, QTLR;
 33 Start, End, genes' coordinated on Galgal6; Net, arbitrary number of a network seen in Figure
 34 8, given by order of location of the first gene (not by order of appearance in Figure 8); Net
 35 bolded, net comprise of genes from both QTLRs; B4-5, location in high LD blocks
 36 extending over the two QTLRs in the five F₆ families.

Q	Gene	Start	End	Net	B4-5	Q	Gene	Start	End	Net	B4-5
4	DUSP16	71,872,763	71,904,322	1	+	5	ING4	77,734,990	77,749,769		+
4	CREBL2	71,960,834	71,970,851	1	+	5	ZNF384	77,755,770	77,781,779		+
4	GPR19	71,980,272	71,988,795		+	5	PIANP	77,803,621	77,808,412		+
4	CDKN1B	72,102,062	72,105,400	1	+	5	COPS7A	77,817,350	77,820,482		+
4	MRPS35	72,622,696	72,644,857	2	+	5	MLF2	77,831,588	77,841,435	5	+
4	MANSC4	72,649,269	72,664,039		+	5	PTMS	77,844,882	77,848,395		+
4	KLHL42	72,671,873	72,682,871	3	+	5	LAG3	77,862,795	77,868,711	2	+
4	PTHLH	72,752,038	72,764,874	2	+	5	CD4	77,873,930	77,885,897	2	+
4	CCDC91	72,867,675	73,073,035		+	5	GPR162	77,894,615	77,899,113		+
4	FAR2	73,191,857	73,320,608			5	P3H3	77,900,316	77,910,271		+
5	SLC2A14	75,548,453	75,558,943			5	GNB3	77,917,296	77,922,102		+
5	NANOG	75,593,243	75,596,024	2		5	CDC43	77,922,769	77,924,912	5	+
5	AICDA	75,632,084	75,637,754	2		5	USP5	77,924,975	77,939,984	5	+
5	MFAP5	75,647,640	75,660,701	2		5	TPI1	77,940,005	77,943,711	2	+
5	RIMKLB	75,676,381	75,723,185	2		5	LRRC23	77,945,286	77,949,479		+
5	PHC1	75,865,514	75,884,654			5	ENO2	77,952,924	77,962,832	2	+
5	M6PR	75,883,096	75,891,432	2		5	C1H12ORF57	77,991,895	77,993,033		+
5	OVST	76,362,509	76,397,638			5	PTPN6	77,994,636	78,014,698	2	+
5	MAN1A2	77,147,561	77,281,468		+	5	PHB2	78,015,335	78,020,461	2	+
5	CD86	77,307,980	77,318,936	2	+	5	EMG1	78,020,566	78,022,842	2	+
5	CASR	77,388,128	77,429,662	2	+	5	LPCAT3	78,023,023	78,038,708	5	+
5	CSTB	77,433,979	77,438,548	2	+	5	C1S	78,045,961	78,055,022	4	
5	CSTA	77,439,994	77,445,026		+	5	C1R	78,058,485	78,065,761	4	
5	CCDC58	77,454,326	77,464,115	2	+	5	RBP5	78,070,071	78,071,702		
5	FAM162A	77,463,747	77,472,850		+	5	CLSTN3	78,071,723	78,086,627	5	
5	KPNA1	77,477,877	77,514,773	2	+	5	PEX5	78,099,886	78,110,916	5	
5	FBXO40	77,526,063	77,541,005	3	+	5	EPHA1	78,162,527	78,195,976		
5	TAPBPL	77,546,387	77,554,991		+	5	ZYX	78,201,045	78,212,460	5	
5	gga-mir-6553	77,567,295	77,567,395		+	5	FAM131B	78,257,533	78,259,472		
5	SCNN1A	77,568,659	77,576,825		+	5	CLCN1	78,276,084	78,333,916		
5	VAMP1	77,579,970	77,584,568		+	5	CASP2	78,336,971	78,362,958		
5	MRPL51	77,585,845	77,587,396	2	+	5	TMEM139	78,364,767	78,369,487		
5	NCAPD2	77,587,505	77,611,310	5	+	5	RAP1GAP1	78,380,088	78,418,997		
5	SCARNA10	77,588,152	77,588,472		+	5	GSTK1	78,425,057	78,436,557	5	
5	CNP1	77,615,840	77,617,557		+	5	TAS2R40	78,481,425	78,482,360		
5	GAPDH	77,619,214	77,623,350	2	+	5	TRPV6	78,604,630	78,633,785	2	
5	IFFO1	77,633,594	77,642,717		+	5	EPHB6	78,678,054	78,732,359		
5	NOP2	77,649,727	77,654,873	2	+	5	PRSS2	78,802,240	78,805,418	2	
5	LPAR5	77,694,435	77,702,350	2	+	5	PRSS3	78,879,681	78,923,402	2	

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Complex and long-range linkage disequilibrium in chicken populations

Figures

Figure 1. Distribution of random LRLDs over Chr 1 and overlaps with QTLRs (Table 1) in F₆ Family 1. Location of marker pairs plotted against r^2 . Start, End, locations of the markers in a pair (for each Start dot there is a matched End dot; see Figure 2 for clarity); numbers, QTLR numbers (Table 1).

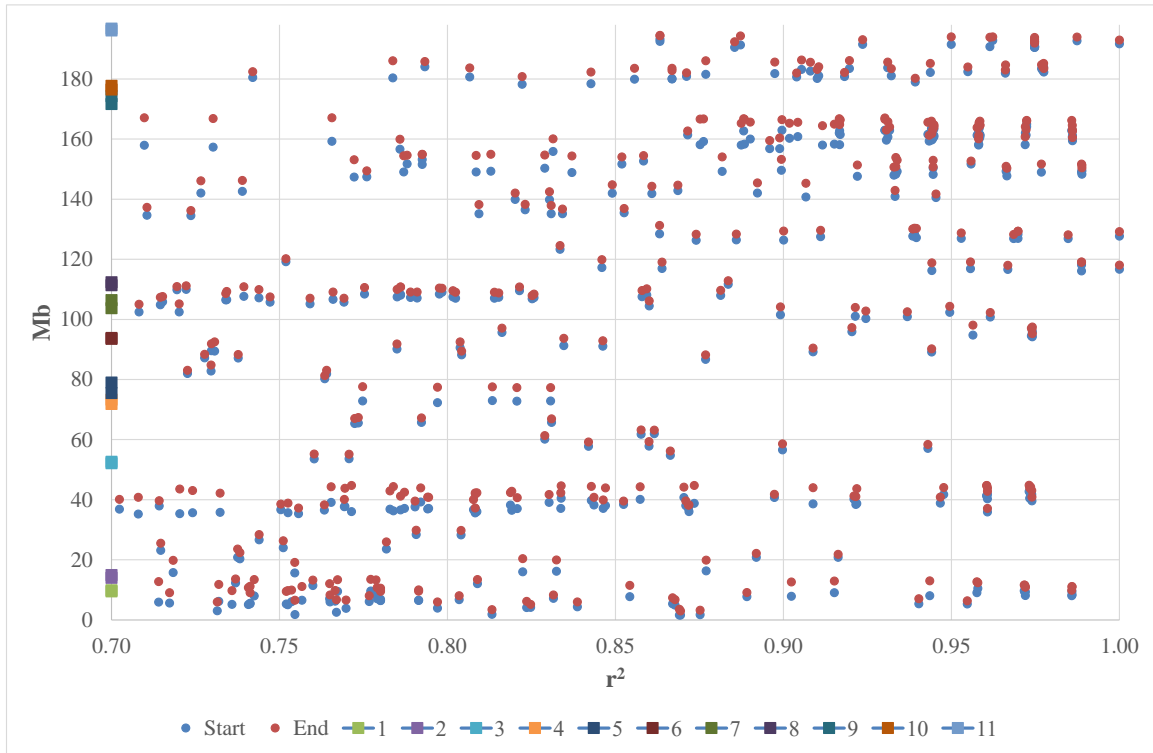
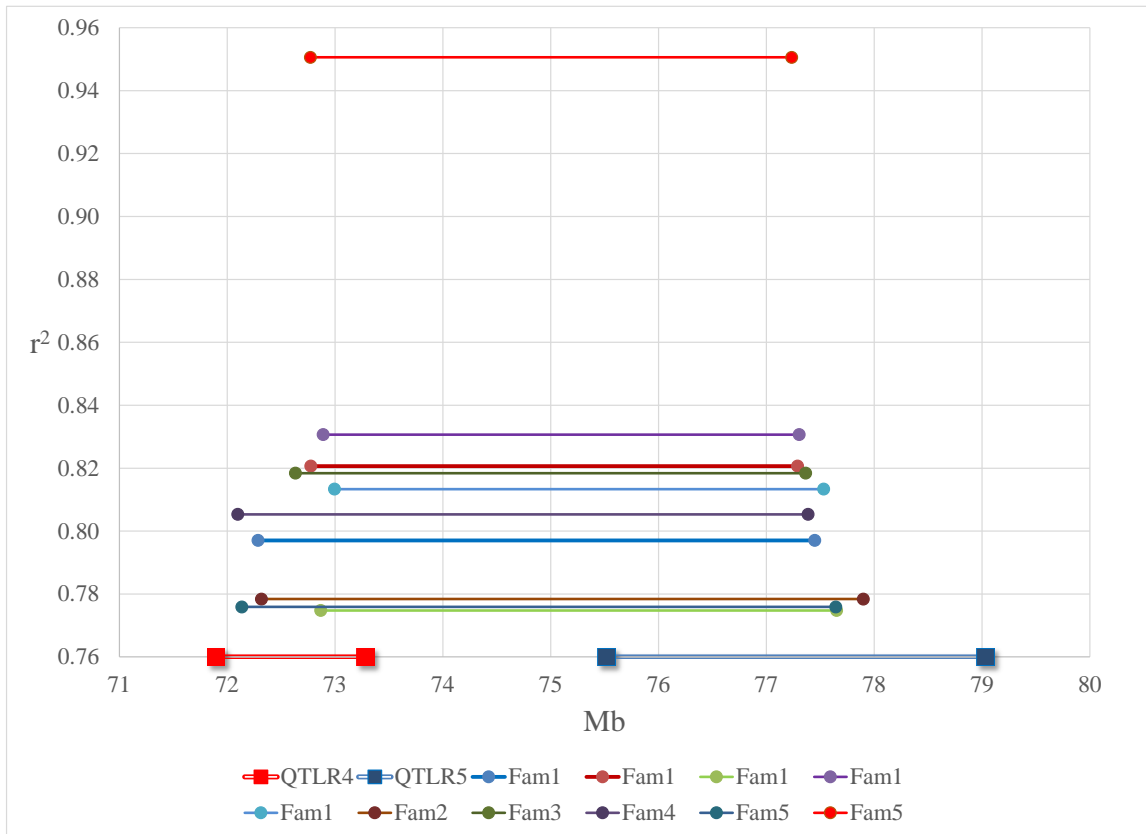


Figure 2. Overlaps between QTLs 4-5 and random LRLD in all F₆ families. Fam, family. There were 5 LRLDs in Family 1, 1 LRLD in Families 2 - 4, and 2 LRLDs in Family 5. Each series of an LRLD is composed of two circles connected by a single line; the circles represent the locations on the x-axis of 2 random markers constitute a pair, and the LD r^2 of this pair is presented on the y-axis. There could be more than one LRLD in a family. Similarly, the series of QTLs 4 and 5 are presents by squares connected by a double line; the squares represent the locations of the QTLR boundaries. QTLRs do not have an r^2 of course; they are simply presented on the x-axis, aligned with the LRLD.



- 1 **Figure 3.** Fragmented interdigitated blocks in QTLR 2 on Chr 1 found in F₆ Family 1. B, block serial number ordered by the location
- 2 of the first marker, colored by block; Mb, location on Galgal6 in Mb; Dist., distance in Mb; ..., unrepresented intermediate markers;
- 3 No., serial number of the marker; red, LD $r^2 \geq 0.7$.
- 4 a. all blocks.

B			1	1	1	1	1	1	1	1	2	1	2	1	1	1	1	3	1	2...	3		
	Mb		14.0830	14.0859	14.0934	14.0947	14.1026	14.1041	14.1077	14.1134	14.1403	14.1441	14.1460	14.1461	14.1471	14.1507	14.1536	14.1544	14.1563	14.1613	14.1739	...	14.7193
		Dist.		0.00292	0.00745	0.00133	0.00784	0.00150	0.00368	0.00568	0.02691	0.00379	0.00191	0.00004	0.00107	0.00361	0.00287	0.00081	0.00191	0.00496	0.01255	...	0.54541
		No.	134	135	136	137	138	139	140	141	142	143	144	145	146	147	148	149	150	151	152	...	337
1	14.0830		134																				
1	14.0859	0.00292	135	1.000																			
1	14.0934	0.00745	136	1.000	1.000																		
1	14.0947	0.00133	137	0.973	0.986	0.973																	
1	14.1026	0.00784	138	0.779	0.778	0.779	0.752																
1	14.1041	0.00150	139	0.778	0.778	0.778	0.751	1.000															
1	14.1077	0.00368	140	1.000	1.000	1.000	0.973	0.778	0.778														
1	14.1134	0.00568	141	1.000	1.000	1.000	0.973	0.779	0.778	1.000													
2	14.1403	0.02691	142	0.022	0.023	0.022	0.023	0.111	0.111	0.023	0.022												
1	14.1441	0.00379	143	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779	0.111											
2	14.1460	0.00191	144	0.024	0.024	0.024	0.024	0.094	0.093	0.024	0.024	0.947	0.094										
1	14.1461	0.00004	145	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779	0.111	1.000	0.094									
1	14.1471	0.00107	146	1.000	1.000	1.000	0.986	0.778	0.778	1.000	1.000	0.023	0.778	0.024	0.778								
1	14.1507	0.00361	147	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779	0.111	1.000	0.094	1.000	0.778							
1	14.1536	0.00287	148	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779	0.111	1.000	0.094	1.000	0.778	1.000						
1	14.1544	0.00081	149	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779	0.111	1.000	0.094	1.000	0.778	1.000	1.000					
3	14.1563	0.00191	150	0.003	0.003	0.003	0.001	0.005	0.005	0.003	0.003	0.002	0.005	0.001	0.005	0.003	0.005	0.005	0.005				
1	14.1613	0.00496	151	0.778	0.778	0.778	0.765	1.000	1.000	0.778	0.778	0.111	1.000	0.093	1.000	0.778	1.000	1.000	1.000	0.005			
2	14.1739	0.01255	152	0.022	0.023	0.022	0.023	0.111	0.111	0.023	0.022	1.000	0.111	0.947	0.111	0.023	0.111	0.111	0.111	0.002	0.111		
...
3	14.7193	0.54541	337	0.000	0.000	0.000	0.002	0.000	0.000	0.000	0.000	0.001	0.000	0.001	0.000	0.000	0.000	0.000	0.000	1.000	0.000	0.001	...

5

6

7 b. Block 1. Blocks 2 and 3 omitted.

B			1	1	1	1	1	1	1	1	1	1	1	1	1	1	
	Mb		14.0830	14.0859	14.0934	14.0947	14.1026	14.1041	14.1077	14.1134	14.1441	14.1461	14.1471	14.1507	14.1536	14.1544	14.1613
		Dist.		0.00292	0.00745	0.00133	0.00784	0.00150	0.00368	0.00568	0.03070	0.00195	0.00107	0.00361	0.00287	0.00081	0.00687
		No.	134	135	136	137	138	139	140	141	143	145	146	147	148	149	151
1	14.0830		134														
1	14.0859	0.00292	135	1.000													
1	14.0934	0.00745	136	1.000	1.000												
1	14.0947	0.00133	137	0.973	0.986	0.973											
1	14.1026	0.00784	138	0.779	0.778	0.779	0.752										
1	14.1041	0.00150	139	0.778	0.778	0.778	0.751	1.000									
1	14.1077	0.00368	140	1.000	1.000	1.000	0.973	0.778	0.778								
1	14.1134	0.00568	141	1.000	1.000	1.000	0.973	0.779	0.778	1.000							
1	14.1441	0.03070	143	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779						
1	14.1461	0.00195	145	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779	1.000					
1	14.1471	0.00107	146	1.000	1.000	1.000	0.986	0.778	0.778	1.000	1.000	0.778	0.778				
1	14.1507	0.00361	147	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779	1.000	1.000	0.778			
1	14.1536	0.00287	148	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779	1.000	1.000	0.778	1.000		
1	14.1544	0.00081	149	0.779	0.778	0.779	0.752	1.000	1.000	0.778	0.779	1.000	1.000	0.778	1.000	1.000	
1	14.1613	0.00687	151	0.778	0.778	0.778	0.765	1.000	1.000	0.778	0.778	1.000	1.000	0.778	1.000	1.000	1.000

8

9 c. Block 2. Blocks 1 and 3 omitted.

B			2	2	2
	Mb		14.1403	14.1460	14.1739
		Dist.		0.00570	0.02783
		No.	142	144	152
2	14.1403		142		
2	14.1460	0.00570	144	0.947	
2	14.1739	0.02783	152	1.000	0.947

10

11

12 d. Block 3. Blocks 1 and 2 omitted.

B				3	3
	Mb			14.1563	14.7193
		Dist.			0.56293
			No.	150	337
3	14.1563		150		
3	14.7193	0.56293	337	1.000	

13

14 **Figure 4.** LD within one QTLR gene. Line WL1, the gene TRANK1 in QTLR 13 on
 15 chromosome 2. B, block serial number ordered by the location of the first marker (same
 16 LD block have the same color); Mb, location on Galgal6 in Mb; Dist., distance in Mb;
 17 Red, $r^2 \geq 0.7$; similar P values has similar colors.

B			1	1	2	2	1	1	1	1	1	1
	Mb		45.9001	45.9006	45.9013	45.9021	45.9025	45.9151	45.9157	45.9188	45.9214	45.9359
		Dist.		0.0005	0.0007	0.0008	0.0004	0.0126	0.0007	0.0031	0.0026	0.0145
		No.	5	8	11	12	13	18	20	28	29	36
1	45.9001		5									
1	45.9006	0.0005	8	0.895								
2	45.9013	0.0007	11	0.016	0.041							
2	45.9021	0.0008	12	0.016	0.041	1.000						
1	45.9025	0.0004	13	1.000	0.895	0.016	0.016					
1	45.9151	0.0126	18	1.000	0.895	0.016	0.016	1.000				
1	45.9157	0.0007	20	1.000	0.895	0.016	0.016	1.000	1.000			
1	45.9188	0.0031	28	1.000	0.895	0.016	0.016	1.000	1.000	1.000		
1	45.9214	0.0026	29	1.000	0.895	0.016	0.016	1.000	1.000	1.000	1.000	
1	45.9359	0.0145	36	0.895	1.000	0.041	0.041	0.895	0.895	0.895	0.895	0.895
Association test P:			3E-02	5E-02	5E-01	5E-01	3E-02	3E-02	3E-02	3E-02	3E-02	5E-02

18

19 **Figure 5.** LD blocks across QTLR elements. Line WL3, QTLR 3 on chromosome 1. B,
 20 block serial number ordered by the location of the first marker (same LD block have the
 21 same color); Mb, location on Galgal6 in Mb; Dist., distance in Mb; E, QTLR element
 22 [Smith et al., 2020]; Red, $r^2 \geq 0.7$; similar P values has similar colors.

B			1	1	1	2	2	2	2	1	2	2	2	1	1	2	2	2	2	
	Mb		52.2768	52.2769	52.2769	52.3375	52.3532	52.3826	52.4001	52.4082	52.4281	52.4684	52.4745	52.6418	52.6419	52.6422	52.6424			
		Dist.		0.0001	0.0000	0.0605	0.0157	0.0294	0.0175	0.0081	0.0199	0.0403	0.0060	0.1674	0.0001	0.0003	0.0002			
		E	lncRNA02					lncRNA05					lncRNA04							
		No.	6	7	8	11	12	14	15	16	17	19	20	30	31	32	33			
1	52.2768		6																	
1	52.2769	0.0001	7	1.000																
1	52.2769	0.0000	8	1.000	1.000															
2	52.3375	0.0605	11	0.006	0.006	0.006														
2	52.3532	0.0157	12	0.006	0.006	0.006	1.000													
2	52.3826	0.0294	14	0.016	0.016	0.016	0.959	0.959												
1	52.4001	0.0175	15	1.000	1.000	1.000	0.006	0.006	0.016											
2	52.4082	0.0081	16	0.016	0.016	0.016	0.958	0.958	0.998	0.016										
2	52.4281	0.0199	17	0.016	0.016	0.016	0.945	0.945	0.985	0.016	0.986									
1	52.4684	0.0403	19	1.000	1.000	1.000	0.006	0.006	0.016	1.000	0.016	0.016								
1	52.4745	0.0060	20	1.000	1.000	1.000	0.006	0.006	0.016	1.000	0.016	0.016	1.000							
2	52.6418	0.1674	30	0.015	0.015	0.015	0.921	0.921	0.960	0.015	0.961	0.978	0.015	0.015						
2	52.6419	0.0001	31	0.015	0.015	0.015	0.921	0.921	0.960	0.015	0.961	0.978	0.015	0.015	1.000					
2	52.6422	0.0003	32	0.015	0.015	0.015	0.921	0.921	0.960	0.015	0.961	0.978	0.015	0.015	1.000	1.000				
2	52.6424	0.0002	33	0.015	0.015	0.015	0.921	0.921	0.960	0.015	0.961	0.978	0.015	0.015	1.000	1.000	1.000			
Association test P:			7E-01	7E-01	7E-01	1E-01	1E-01	2E-01	7E-01	2E-01	2E-01	7E-01	7E-01	3E-01	3E-01	3E-01	3E-01			

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24

25 **Figure 6.** LD between QTLR genes. Line WL2, QTLR 3 on chromosome 17. B, block
 26 serial number ordered by the location of the first marker (same LD block have the same
 27 color); Mb, location on Galgal6 in Mb; Dist., distance in Mb; E, QTLR element [Smith et
 28 al., 2020]; Red, $r^2 \geq 0.7$; purple, $0.15 \leq r^2 < 0.7$; same LD block have the same color;
 29 similar P values has similar colors.

B			1	2	2	2	2	2	2	2	1	1		
	Mb		3.9390	3.9418	3.9420	3.9421	3.9428	3.9435	3.9443	3.9447	4.3588	4.3642	4.3689	
		Dist.		0.0029	0.0001	0.0001	0.0007	0.0008	0.0007	0.0005	0.4141	0.0054	0.0047	
		E	TLR4									BRINP1		
		No.	2	4	5	6	7	8	9	10	11	14	15	
1	3.9390		2											
2	3.9418	0.0029	4	0.135										
2	3.9420	0.0001	5	0.135	1.000									
2	3.9421	0.0001	6	0.135	1.000	1.000								
2	3.9428	0.0007	7	0.135	1.000	1.000	1.000							
2	3.9435	0.0008	8	0.135	1.000	1.000	1.000	1.000						
2	3.9443	0.0007	9	0.135	1.000	1.000	1.000	1.000	1.000					
2	3.9447	0.0005	10	0.135	1.000	1.000	1.000	1.000	1.000	1.000				
1	4.3588	0.4141	11	0.830	0.102	0.102	0.102	0.102	0.102	0.102	0.102			
1	4.3642	0.0054	14	0.830	0.102	0.102	0.102	0.102	0.102	0.102	0.102	1.000		
	4.3689	0.0047	15	0.157	0.661	0.661	0.661	0.661	0.661	0.661	0.661	0.171	0.171	
Association test P:				3E-01	7E-01	7E-01	7E-01	7E-01	7E-01	7E-01	7E-01	2E-01	2E-01	8E-01

30



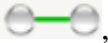




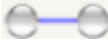
31

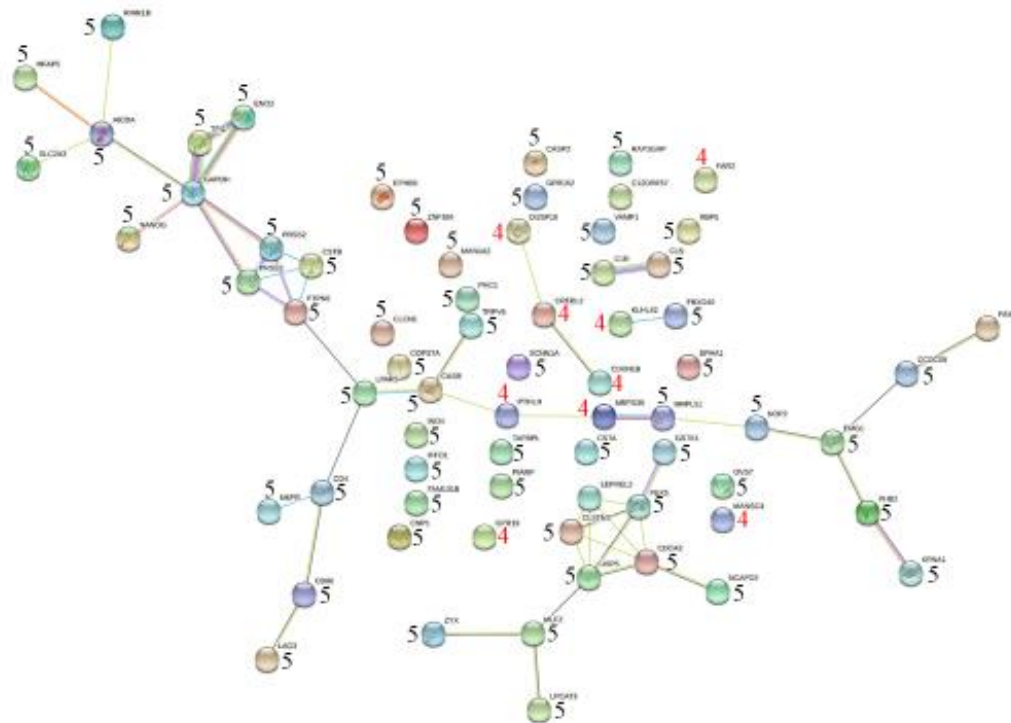
32 **Figure 7.** LD between QTLRs 4 and 5. Line WPR1, the lncRNA01 in QTLR 4 and the
 33 gene CSTA in QTLR 5. Q, QTLR serial number (Table 1); B, high LD block serial
 34 number ordered by the location of the first marker (same LD block have the same color);
 35 Mb, location on Galgal6 in Mb; Dist., distance in Mb; E, QTLR element [Smith et al.,
 36 2020]; Red, $r^2 \geq 0.7$; purple, $0.15 \leq r^2 < 0.7$; same LD block have the same color; similar P
 37 values has similar colors.

Q			4	4	4	4	5	5	5		
B			1	1	1	1	2	2	2		
	Mb			72.3073	72.3074	72.3076	72.3076	77.4421	77.4449	77.4449	
	Dist.				0.0001	0.0002	0.0000	5.1345	0.0028	0.0000	
		E		lncRNA01.03				CSTA			
				No.	34	36	37	38	42	44	45
4	1	72.30728	lncRNA01.03	34							
4	1	72.30737		36	1.000						
4	1	72.30762		37	1.000	1.000					
4	1	72.30763		38	1.000	1.000	1.000				
5	2	77.44215	CSTA	42	0.478	0.478	0.478	0.478			
5	2	77.44490		44	0.478	0.478	0.478	0.478	1.000		
5	2	77.44493		45	0.478	0.478	0.478	0.478	1.000	1.000	
Association test P:					8E-02	8E-02	8E-02	8E-02	4E-01	4E-01	4E-01

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40 **Figure 8.** STRING network of genes under QTLRs 4 and 5. Network nodes represent
41 proteins; Colored nodes, query proteins and first shell of interactors; Node content: empty,
42 protein of unknown 3D structure; filled, some 3D structure is known or predicted; Known
43 interactions: ; from curated databases; , experimentally determined;
44 Predicted interactions: , gene neighborhood; , gene fusions; ,
45 gene co-occurrence; Others: , text mining; , co-expression; ,
46 protein homology. The number of the QTLR is shown next to the gene. As can be seen,
47 some gene associations include genes across each QTLR.



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