Genomic Tools for the Identification of Loci Associated with Facial Eczema in New Zealand Sheep

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Abstract: Facial eczema (FE) is a significant metabolic disease that affects New Zealand ruminants. Ingestion of the mycotoxin sporidesmin leads to liver and bile duct damage, which can result in photosensitisation, reduced productivity and death. Strategies used to manage the incidence and severity of the disease include breeding. In sheep, there is considerable genetic variation in the response to FE. A commercial testing program is available for ram breeders who aim to increase tolerance, determined by the concentration of the serum enzyme, γ-glutamyltransferase 21 days after a measured sporidesmin challenge (GCT21). Genome-wide association studies were carried out to determine regions of the genome associated with GCT21. Two regions on chromosomes 15 and 24 are reported, which explain 5% and 1% of the phenotypic variance in the response to FE, respectively. The region on chromosome 15 contains the β-globin locus. Of the significant SNPs in the region, one is a missense variant within the haemoglobin subunit β (HBB) gene. Mass spectrometry of haemoglobin from animals with differing genotypes at this locus indicated that genotypes are associated with different forms of adult β-globin. Haemoglobin haplotypes have previously been associated with variation in several health-related traits in sheep and warrant further investigation regarding their role in tolerance to FE in sheep. We show a strategic approach to the identification of regions of importance for commercial breeding programs with a combination of discovery, statistical and biological validation. This study highlights the power of using increased density genotyping for the identification of influential genomic regions, combined with subsequent inclusion on lower density genotyping platforms.

Keywords: sheep; disease; facial eczema; haemoglobin; GWAS

1. Introduction

Facial eczema (FE) is a metabolic disease of major welfare concern and economic loss to farmers in New Zealand. The disease is caused by ingestion of the mycotoxin sporidesmin, which is found in the spores of the fungus Pseudopithomyces chartarum (formerly Pithomyces chartarum) [1,2]. P. chartarum grows on the dead leaf matter at the base of pastures, and ingestion of the toxic fungal spores by ruminants leads to liver inflammation and bile duct blockage [3]. While in many cases, animals might show no clinical signs of disease [4], production losses arise from a decrease in production [5] and lowered reproduction [6–8] in both clinically
and sub-clinically affected animals. In an outbreak in 1981, the production cost to New Zealand was estimated to be NZD 266 million (inflation-adjusted to 2019 [9,10]). Clinical FE is characterised by photosensitivity in exposed areas of the animal, particularly in lightly pigmented areas such as the face, giving the disease its name [11]. Although FE outbreaks have been the most severe in the North Island of New Zealand [1], outbreaks have also been recorded in Australia [12], South Africa [13], North and South America [14,15], Europe [16] and China [17]. Outbreaks are associated with increases in dead leaf matter, temperature, and humidity in the summer and autumn months [18–20], and long-term climate change projections indicate that there will be an increase in the geographical spread of *Pse. chartarum* in New Zealand, resulting in outbreaks of FE in currently unaffected regions [21].

Current strategies for preventing the severity of FE include protecting animals through dosing with zinc compounds [22], spraying pasture with fungicide [23], pasture management or alternative feeds [24], and breeding animals for tolerance [25]. Initial research in the 1970s involved challenging animals with a measured dose of sporidesmin to assess the physiological impact. Post-challenge, scoring liver damage showed between-animal variation in susceptibility to FE, with a moderate to high heritability (h²) of 0.42 ± 0.09 [26]. Selection lines using Romney sheep were subsequently established in 1975 and were initially selected for tolerance or susceptibility based on liver damage [26]. A live-animal phenotype was reported in 1978 by Towers and Stratton [27], who showed that serum γ-glutamyltransferase (GGT) collected 2 to 3 weeks after sporidesmin challenge could be used as a measure of liver damage. Using data from the above selection lines, the heritability of log-transformed serum GGT levels at 21 days after a measured sporidesmin challenge (GGT21) was 0.45 ± 0.03 [28]. A commercial testing programme, Ramguard™, was then developed in the mid-1980s to provide New Zealand sheep breeders with sporidesmin and an ethical dosing strategy for ascertaining FE tolerance [29]. The resulting GGT21 values are used to generate breeding value estimates for tested and related animals, with 800 to 1,100 rams tested each year. Recent re-estimation of genetic parameters for FE tolerance using a Ramguard™ dataset found a comparable heritability with that estimated from the selection lines (0.44 ± 0.03), with no significant genetic correlations between GGT21 and any production traits [30].

Several approaches have been used to search for causal genes or loci underlying FE tolerance, including a candidate gene approach [31,32], genome-wide scans for quantitative trait loci (QTL) [33,34] and scanning for selective sweep signatures [35]. There was limited concordance between the results of the studies, with only two of the selective sweep regions, on chromosomes OAR1 and OAR13, found within suggestive (F-statistic value<10.2) QTL regions. The development of single nucleotide polymorphism (SNP) arrays (www.sheepmap.org), facilitated significant increases in the number of genotyped animals, driving the development of genomic selection [36]. As a result, the number of animals with SNP genotypes and FE phenotypic measurements (GGT21) has increased.

The objective of our study was to determine genomic variants associated with FE in New Zealand sheep, through a combination of discovery, statistical and biological validation using cost-effective lower density genotyping platforms.

2. Materials and Methods

2.1. Animals and Genotyping

Sheep were selected for inclusion based on GGT21 records and genotype availability. All animals were from either industry or progeny test flocks and had been tested through the Ramguard™ programme. All data were obtained from the Sheep Improvement Limited (SIL, www.sil.co.nz) database. The Ramguard™ programme [29] uses *Pse. chartarum* that is cultured in a laboratory to produce the mycotoxin sporidesmin. Animals are dosed with precise amounts of sporidesmin (made up of toxic forms of sporidesmin, specifically sporidesmin A, B and E, with a predominance (>90%) of sporidesmin A) by intra-ruminal intubation at a volume that is dependent on the animal’s live weight (mg per kg live
SNPs that reached a genome-wide significance threshold ($p < 1.04 \times 10^{-7}$) from the initial ‘discovery’ analyses (Table S1) were subsequently included in the design of the Illumina OvineLD BeadChip (15,000 markers; “15K”). A subsequent ‘validation’ dataset consisted of 2,363 animals genotyped with GGT21 records from 42 flocks, born between 2010 and 2016. As above, animals were either one of the main dual-purpose sheep breeds used in New Zealand or a composite breed. All validation animals were genotyped with the 15K chip according to the manufacturer’s protocol.

Genome coordinates for all SNP were based on the OAR_v3.1 ovine genome assembly (GenBank assembly accession GCA_000298735.1 [37]). Quality control (QC) checks excluded markers that appeared non-autosomal (including pseudo-autosomal), had a call rate below 95%, and/or had a minor allele frequency (MAF) ≤ 0.05. Individuals were excluded from the analysis if there was more than 5% genotyping failure. After marker and sample QC, 2669 phenotyped animals were available for the discovery analysis, with 481,400 of the 606,006 markers utilised. For the validation dataset, 2,342 phenotyped animals were available, with 14,747 of the 15,000 markers utilised for analysis.

2.2. Imputation

For the 2,532 animals genotyped using the 50K chip, imputation from 50K to HD genotypes for the discovery analyses was performed using Fimpute (v2.0) [38]. The imputation dataset included animals that did not have GGT21 records and consisted of 12,945 animals genotyped with the 50K chip, and 2408 animals genotyped with the HD chip. Quality control was carried out before imputation, with markers excluded that appeared non-autosomal (including pseudo-autosomal), had a call rate below 95%, had >2% reproducibility between platforms, and/or had >2% Mendelian errors. Animals included in the imputation dataset were born between 1965 and 2012, and represented 236 flocks, with the largest flock contributing 124 animals. Purebred Romney (n=616), Perendale (n=207) and Primera (n=149) genotypes were used to calculate breed-specific imputation accuracies. Imputation accuracy was calculated using sets of 100 animals from each breed against the larger reference population, with HD genotypes masked, and the remaining 50k genotypes imputed to full density. Accuracy was then calculated for each dataset. Additionally, for the Romney animals within and across breed accuracy was also calculated, where 100 animals were imputed against a random sample of 500 of the remaining 516 purebred Romney, or 500 other animals comprised of the Coopworth, Corriedale, Perendale, Texel and Wiltshire breeds. To account for biases in imputation accuracy for SNP with low MAF, the correction proposed by Hayes et al. [39] was applied.

2.3. Genome-Wide Association Studies

Both the discovery and validation GWAS was performed within SNP & Variation Suite v8.4.0 (Golden Helix, Inc., Bozeman, MT, USA, www.goldenhelix.com) using two methods. Firstly, an Efficient Mixed-Model Association eXpedited (EMMAX) analysis using a kinship matrix was performed on log-transformed serum GGT levels at 21 days after a measured sporo-sidesmin challenge (GGT21). Contemporary group (incorporating flock, year of birth, sex and mob [30]) and the first two principal components, calculated from genotypic data (to account
for population stratification), were fitted as additional covariates. Secondly, genotype association tests were performed on the residuals obtained from ASReml (v4.1) [40] after fitting recorded pedigree and the fixed effect of the contemporary group. Population stratification was again corrected by fitting the first two principal components, calculated from genotypic data, in the analysis. After Bonferroni correction, the threshold for genome-wide significance ($p < 0.05$) was $1.04 \times 10^{-7}$ and $3.39 \times 10^{-8}$ for discovery and validation analyses, respectively.

### 2.4. Validation of Markers of Interest

To calculate the proportion of variance explained by each marker, SNP effects were estimated in ASReml by fitting the SNP as a fixed effect along with the contemporary group. The predicted trait values were converted to genotypic effects as follows $a = (AA-BB)/2$ and $d = AB - ((AA+BB)/2))$, where AA, BB, and AB are the predicted trait values for each genotype class. The proportion of additive genetic variance due to each SNP was estimated as $(2pq(a+d(q-p)))V_A$, where $p$ and $q$ are the allelic frequencies at the SNP locus, and $V_A$ is the total additive genetic variance of the trait obtained when no SNP fixed effects are included in the model.

To quantify the predictive ability of markers of interest, 5-fold cross-validation was performed with random sampling ($n = 50$ iterations) within SNP & Variation Suite v8.4.0 for genomic best linear unbiased prediction (GBLUP). Only animals that had been genotyped on either the Ovine Infinium® HD SNP BeadChip or the Illumina OvineLD BeadChip ($n = 2999$) were used for the analysis, and fixed effect coefficients included contemporary group. K-fold cross-validation was run including all markers ($n=12,500$), and after removing markers in the regions of interest on chromosomes 15 and 24 ($n=12,477$). Accuracies were calculated by dividing the Pearson product-moment correlation coefficient by the square root of the heritability.

To test whether SNP markers were predictive of haemoglobin phenotype (Hb-A, Hb-AB or Hb-B), blood was collected from 12 animals with different genotypes in the β-globin locus, using SNPs rs405755938 and rs402069107 (Table 1). Proteomic analyses are described in Supplementary Methods (Methods S1).

#### Table 1. SNPs that reached genome-wide significance levels for Facial Ecema tolerance. EMMAX p-values, predicted log-transformed GGT21 values, allele substitution effects ($\alpha$), dominance effects ($d$) and proportion of total additive ($V_A$) and phenotypic ($V_F$) variance explained by each marker.

| RSID      | Chr | Position ¹ | −log_{10} p-value ² | MAF (allele) ³ | Predicted GGT21 Values ⁴ | $\alpha$ | $d$ | Prop $V_A$ | Prop $V_F$ |
|-----------|-----|------------|----------------------|----------------|--------------------------|---------|-----|-----------|-----------|
| rs398930318 | 15  | 47,489,709 | 6.09                 | 0.45 (A)       | AA (±SE) 4.39 (0.06) AB (±SE) 4.63 (0.07) | -0.21   | 0.07 | 0.04      | 0.02      |
| rs423653664 | 15  | 47,504,887 | 11.93                | 0.48 (A)       | AA (±SE) 4.92 (0.06) AB (±SE) 5.04 (0.07) | -0.28   | 0.10 | 0.07      | 0.04      |
| rs398614689 | 15  | 47,505,272 | 12.00                | 0.48 (A)       | AA (±SE) 4.92 (0.06) AB (±SE) 5.04 (0.07) | -0.28   | 0.11 | 0.07      | 0.04      |
| rs429709432 | 15  | 47,513,135 | 11.09                | 0.46 (A)       | AA (±SE) 4.51 (0.07) AB (±SE) 4.92 (0.06) | -0.28   | 0.16 | 0.08      | 0.04      |
| rs400809788 | 15  | 47,519,231 | 10.07                | 0.49 (B)       | AA (±SE) 4.49 (0.06) AB (±SE) 4.92 (0.06) | 0.30    | 0.14 | 0.08      | 0.04      |
| rs405755938 | 15  | 47,519,931 | 13.07                | 0.50 (A)       | AA (±SE) 4.49 (0.06) AB (±SE) 4.92 (0.06) | -0.30   | 0.13 | 0.08      | 0.05      |
| rs427105378 | 15  | 47,564,204 | 13.69                | 0.50 (B)       | AA (±SE) 4.49 (0.06) AB (±SE) 4.92 (0.06) | 0.30    | 0.13 | 0.08      | 0.05      |
| rs402069107 | 15  | 47,567,105 | 13.69                | 0.50 (A)       | AA (±SE) 4.49 (0.06) AB (±SE) 4.92 (0.06) | -0.30   | 0.13 | 0.08      | 0.05      |
| rs411410654 | 15  | 47,569,499 | 13.69                | 0.50 (B)       | AA (±SE) 4.49 (0.06) AB (±SE) 4.92 (0.06) | 0.30    | 0.13 | 0.08      | 0.05      |
| rs430842113 | 15  | 47,570,178 | 12.51                | 0.50 (A)       | AA (±SE) 4.49 (0.06) AB (±SE) 4.92 (0.06) | 0.30    | 0.13 | 0.08      | 0.05      |
| rs425052505 | 15  | 47,609,978 | 12.70                | 0.50 (B)       | AA (±SE) 4.49 (0.06) AB (±SE) 4.92 (0.06) | 0.30    | 0.13 | 0.08      | 0.05      |
| rs425270036 | 24  | 1,155,234  | 5.70                 | 0.33 (B)       | AA (±SE) 4.49 (0.06) AB (±SE) 4.92 (0.06) | 0.19    | 0.08 | 0.04      | 0.01      |
| rs409675199 | 24  | 1,156,263  | 5.76                 | 0.33 (A)       | AA (±SE) 4.49 (0.06) AB (±SE) 4.92 (0.06) | -0.19   | 0.08 | 0.04      | 0.01      |

¹Position is based on OARv3.1. ²Combined EMMAX analysis using animals genotyped on either the Ovine Infinium® HD SNP BeadChip or the Illumina OvineLD BeadChip ($n = 2999$; 13,212 shared markers). After Bonferroni correction, the threshold for genome-wide significance ($p < 0.05$) was $3.78 \times 10^{-8}$ (log-transformed $= 5.42$). ³Minor allele frequencies (MAF), with minor allele shown in brackets. Alleles are shown in AB format. ⁴Predicted log-transformed serum GGT levels at 21 days after a measured sporidesmin challenge (GGT21) values estimated using ASReml.
3. Results

3.1. Imputation

Imputation accuracy, corrected for MAF (IAMAF), was consistent among the datasets tested. Romney, which was the largest of the purebred groups (n=616) had a marginally higher accuracy of 0.987, compared to the smaller group of purebred Perendale (n=207; IAMAF=0.983). Primera (n=149), which can be considered a composite breed and therefore more closely reflects the entire reference set, had the highest imputation accuracy of 0.989. When the 100 Romney animals were imputed against a reference of 500 of the remaining Romney, or 500 purebred animals (excluding Romney and composite breeds), IAMAF reduced to 0.962 and 0.932, respectively.

3.2. Genome-Wide Association Studies

To investigate regions of the genome that are associated with FE, imputed HD genotype data were combined with log-transformed serum GGT levels at 21 days after a measured sporidesmin challenge (GGT21) for additive GWAS analysis (Figure 1A,B and Figure S1). In both EMMAX and association analyses a significant quantitative trait locus (QTL) was observed on chromosome 15, with a smaller peak apparent on chromosome 24. Significant SNPs from both QTL (17 SNPs on chromosome 15 and 5 SNPs on chromosome 24) were subsequently included in the design of the Illumina OvineLD BeadChip (Table S1). Figure 2 shows the EMMAX results from the discovery analysis, using genotypes that had been imputed from 50K to HD (600K). When these results are reduced to only the ~41k SNPs that are present on both BeadChips, no significant QTL were observed.

Low-density genotyping (Illumina OvineLD BeadChip) of additional animals with FE phenotypes confirmed the QTL on chromosome 15, using both EMMAX and association analyses (Figure 1C,D). The SNPs in this region (Figure 2) are not present on the Illumina OvineSNP50 BeadChip, therefore only animals that had been genotyped on either the Ovine Infinium® HD SNP BeadChip or the Illumina OvineLD BeadChip (n = 2999; 13,212 shared markers) were used to repeat the EMMAX analysis (Figure 3A), calculate linkage disequilibrium (R²) between markers (Figure 2) and allele frequency by breed (Figure 2).
Figure 1. Manhattan plots of genome-wide association analysis for Facial Eczema in New Zealand sheep. Efficient Mixed-Model Association eXpedited (EMMAX) analyses using a kinship matrix (A,C) were conducted using log-transformed serum GGT levels at 21 days after a measured sporidesmin challenge (GGT21), with the contemporary group and the first two principal components fitted as covariates. Genotype association tests (B,D), fitting two principal components, were performed using residuals obtained from ASReml after fitting pedigree and the fixed effect of the contemporary group. Animals had either high-density (genotyped using either the Ovine Infinium® HD SNP BeadChip or the Illumina OvineSNP50 BeadChip, and imputed to HD) (A,B) or low-density (Illumina OvineLD BeadChip) (C,D) genotypes. After Bonferroni correction, the threshold for genome-wide significance ($P < 0.05$) was $1.04 \times 10^{-7}$ (log-transformed = 6.98) and $3.39 \times 10^{-8}$ (log-transformed = 5.47) for high-density and low-density analyses, respectively.
Figure 2. Comparison of imputed HD and 50K genotype data. Efficient Mixed-Model Association eXpedited (EMMAX) analysis of log-transformed serum GGT levels at 21 days after a measured sporidesmin challenge (GGT21), fitting contemporary group and the first two principal components fitted as covariates. Blue diamonds (♦) show p-values for the ~41k SNPs from the Illumina OvineSNP50 BeadChip. Black diamonds (♦) show p-values for the additional Ovine Infinium® HD SNP BeadChip SNPs.

Figure 3. Detailed examination of the region underlying the GWAS peak on chromosome 15 (47,490 Kb – 47,650 Kb). (A) Efficient Mixed-Model Association eXpedited (EMMAX) analysis using animals genotyped with either the Ovine Infinium® HD SNP BeadChip or Illumina OvineLD BeadChip. (B) The previously reported [41] ~37-kb gap in the v3.1 sheep genome assembly is indicated by the red line. The ~40-kb region of low sequence identity between haplotype A and B (reference genome) is shown by the two black lines. Linkage disequilibrium (R²), from 1 (red) to 0 (white). (C) Allele frequency (based on overall minor allele) by breed: Composite (blue); Coopworth (green); Highlander (orange); Perendale (grey); Romney (red); Wiltshire (purple), based on overall minor allele. (D) Schematic diagram of the β-globin locus (duplication of an ancestral four-gene set, ε-ε-ψ-β), showing the arrangement of haplotype A and B. While both haplotypes A and B contain the adult β-globin gene, HBB, the two are considered allelic variants (βA and βB).
3.3. Regions of interest

A total of 11 SNPs, all on chromosome 15, reached genome-wide significance in all analyses (Figure 1 and Table 1). In addition, two SNPs on chromosome 24 reached genome-wide significance in both the high-density genotype analyses (Figure 1A,B) and the combined analysis (Table 1). Minor allele frequencies ranged from 0.33 to 0.5. Dominance effects were significant for all SNPs aside from rs398930318. The proportion of the phenotypic variance explained by individual SNPs ranged from 1% to 5%.

The significant region on chromosome 15 (15:47504887-47609978; Figure 3) has previously been identified as the location of the β-globin locus [41]. Of the significant SNPs in the region (Table 1), one (rs402069107) is a missense variant within the HBB gene, with an A>G change at position 359 of the coding sequence resulting in a change from Aspartagine (N) to serine (S) at amino acid 120. In sheep, the repetitive β-globin locus has two different haplotypes, the longer haplotype A and the shorter haplotype B. While both haplotypes A and B contain the adult β-globin gene, HBB, the two are considered allelic variants (βA and βB) [42,43] differing structurally by at least seven amino acids [44–46]. Version 3.1 of the ovine genome (OAR_v3.1), which all of the SNPs were designed from and mapped to, contains the shorter haplotype B [41]. There were, therefore, no SNPs on any of the genotyping platforms used in this study that were only present in the longer haplotype A.

The two SNPs immediately on either side of the β-globin region that uniquely mapped to multiple public genome assemblies (Table 2), rs405755938 and rs402069107, were used to estimate the β-globin haplotype. All haplotype A animals were called CC (the B allele in Table 1) at rs405755938, and GG (the B allele in Table 1) at rs402069107, with the haplotype B animals homozygous for the alternate allele (T and A, respectively). The rs402069107 alleles are consistent with HBBA (UniProt ID Q1KY77) and HBB (UniProt ID P02075) protein sequences; the G allele results in an S at amino acid 120 of the HBB protein, as seen in HBBA (βB). At both positions, the predicted GGT21 values from ASReml were lower for the haplotype A animals (Table 1).

Table 2. Genomic position of markers used to predict β-globin haplotype.

| Genome Assembly | Genbank Accession | Chromosome | rs405755938 | rs402069107 | Region Distance |
|-----------------|------------------|------------|-------------|-------------|-----------------|
| OAR_v3.1        | GCA_000298735.2   | 15         | 47,519,931  | 47,567,105  | 65,291          |
| Oar_v4.0        | GCA_000298735.2   | 15         | 47,414,827  | 47,462,139  | 65,429          |
| Oar_rambouillet_v1.0 | GCA_002742125.1 | 15         | 51,898,128  | 51,984,579  | 104,993         |
| ARS-UI_Ramb_v2.0 | GCA_016772045.1  | 15         | 47,951,894  | 48,038,330  | 104,963         |
| White Dorper    | GCA_000298735.2   | 15         | 47,750,476  | 47,797,960  | 65,855          |
| Romanov         | GCA_000298735.2   | 15         | 48,107,973  | 48,194,400  | 104,969         |

1 Ben Rosen, pers. comm. [47].

The significant genome region on chromosome 24 is approximately 230 Kb long. This region reached significance in both high-density genotype analyses, however, only one SNP reached significance in the EMMAX analysis of the low-density genotypes, and this SNP was almost 300Kb upstream of the region identified by the high-density genotypes. Of the significant SNPs on chromosome 24 from the combined EMMAX analysis (Table 1), one (rs425270036) is a missense variant within the PTX4 gene.

Including markers in the regions of interest from chromosomes 15 and 24 increased GBLUP accuracy from 0.491 to 0.505, although this was not significant (Table 3).
Table 3. Summary statistics from K-fold (K = 5) cross-validation (n = 50 iterations) with random sampling (n = 50 iterations).

| Markers                                         | Fixed Effects       | PPMCC (± SD) | h²   | GBLUP Accuracy |
|-------------------------------------------------|---------------------|--------------|------|---------------|
| All SNPs (n = 12,500)                           | Contemporary group  | 0.335 ± 0.007| 0.44 | 0.505         |
| Removing markers in regions of interest (n = 12,477) | Contemporary group  | 0.326 ± 0.007| 0.44 | 0.491         |

1 Contemporary group (incorporating flock, year of birth, sex and mob) were fitted as covariates in all analyses. 2 Product-Moment Correlation Coefficient (PPMCC). 3 GBLUP accuracy calculated as PPMCC/h².

3.4. Proteomic analysis

The A and B variants of sheep β-globin (Figure S2) were characterised by the following amino acid changes from A (UniProt ID: Q1KYZ7) → B (UniProt ID: P02075), respectively: S99 → N, A307 → P, VQ73 → MK, S128 → N, E128 → D and R143 → K. These amino acid changes are found in four tryptic peptides from haemoglobin β. In the A variant, these peptides are defined by the following residues: 40-58, 65-81, 116-131 and 132-143. In the B variant, the change of Q75 → K results in a shorter peptide 65-75 but the other identifier tryptic peptides are of the same length. Following the LC-MS run, genotypes (rs405755938 and rs402069107) at the β-globin locus predicted β-globin peptides (βA, βAB or βB) for 11 of the 12 animals, with one animal, predicted to be Hb-B, 5745, also showing polymorphism of the seven variable peptides (Table 4).

Table 4. Unique β-globin peptides detected by mass spectrometry from sheep blood samples from genotyped animals.

| Predicted β-Globin Type | βA | βB |
|-------------------------|----|----|
| βA (n = 3)              |    |    |
| 001 MLTAEEKAAV TGFVGKVKVDD EVGAELGR1 LVVYPWTQRF | 001 MLTAEEKAAV TGFVGKVKVDD EVGAELGR1 LVVYPWTQRF |
|                        | 041 FEHFDLSSA DAVMNNAKVK AHGKKVLDSF SNGVQLDDL | 041 FEHFDLSSA DAVMNNAKVK AHGKKVLDSF SNGVQLDDL |
|                        | 081 KGTFAQSL HCDKLHVDP EFRLLGNVL VVLARHHGSE | 081 KGTFAQSL HCDKLHVDP EFRLLGNVL VVLARHHGSE |
|                        | 121 FTPVQAEFQ KVVAGVANA ALHRYH | 121 FTPVQAEFQ KVVAGVANA ALHRYH |
| βAB (n = 3)             |    |    |
| 001 MLTAEEKAAV TGFVGKVKVDD EVGAELGR1 LVVYPWTQRF | 001 MLTAEEKAAV TGFVGKVKVDD EVGAELGR1 LVVYPWTQRF |
|                        | 041 FEHFDLSSA DAVMNNAKVK AHGKKVLDSF SNGVQLDDL | 041 FEHFDLSSA DAVMNNAKVK AHGKKVLDSF SNGVQLDDL |
|                        | 081 KGTFAQSL HCDKLHVDP EFRLLGNVL VVLARHHGSE | 081 KGTFAQSL HCDKLHVDP EFRLLGNVL VVLARHHGSE |
|                        | 121 FTPVQAEFQ KVVAGVANA ALHRYH | 121 FTPVQAEFQ KVVAGVANA ALHRYH |
| βB (n = 1)              |    |    |
| 001 MLTAEEKAAV TGFVGKVKVDD EVGAELGR1 LVVYPWTQRF | 001 MLTAEEKAAV TGFVGKVKVDD EVGAELGR1 LVVYPWTQRF |
|                        | 041 FEHFDLSSA DAVMNNAKVK | 041 FEHFDLSSA DAVMNNAKVK |


4. Discussion

In this study, we used large phenotypic datasets in conjunction with high- and low-density genotype data to interrogate the sheep genome for regions associated with variability in tolerance to Facial Eczema (FE). Two QTL on chromosomes 15 and 24, were discovered using imputed high-density genotypes. The significant SNPs were subsequently included on a bespoke low-density ovine SNP chip for validation using industry-phenotyped animals. This study highlights the power of using increased density genotyping for the identification of influential genomic regions. SNPs from the Illumina OvineSNP50 BeadChip were not associated with the QTL on chromosome 15, likely due to the high levels of diversity in the breeds represented in this study, and the resulting reduced linkage disequilibrium between markers [48]. When the variants of importance are subsequently included on more widely used, lower density genotyping platforms, this enables validation in a larger number of animals at a lower cost, through alignment with industry-collected genotypes and phenotypes.

The 13 SNPs examined had largely additive effects, although there was some evidence for dominance. If these genotypes are also used in genomic selection, they may increase breeding value accuracy, although this is largely captured by GBLUP. Given the subclinical manifestation of the disease, further work is needed to understand the impact of these effects on the phenotype.

The significant region (P < 1.04x10\(^{-7}\)) on chromosome 24 from the analysis using the high-density genotypes is approximately 230 Kb. Of the SNPs that reached significance in the high-density SNP analysis, two are missense variants within the PTX4 gene. PTX4 is reported to be widely expressed at low levels, with the highest levels in the liver in mice, and the small intestine and testes in humans [49], however, expression in sheep has been detected primarily in the cerebellum, at low levels [37,50]. PTX4 belongs to the pentraxin (PTX) superfamily of multifunctional conserved proteins; some PTXs are part of the humoral arm of innate immunity and behave as functional ancestors of antibodies by medi-

| \(\beta^A (n = 2)\) | 041 FEHFGDLSNA DAVMNPKVK AHGKKVLDSE SNGMKHLDLDL  
|----------------|-----------------------------------------------|
|                | 081 KGTFAQLSHEL HCDKLHVDPE NFRLGNVVLV VVLARHHGNE  
|                | 121 FTPVLQADFQ KVAGVANAL AHKYH  

\(^1\) Predicted using genotypes from rs405755938 and rs402069107 (Table 1). \(\beta\)-globin sequence taken from UniProt (A: Q1KY2Z; B: P02075).
ating agglutination and complement activation, however, PTX4 mRNA expression is distinct from other PTX family members [49], and there is no current evidence supporting the function of PTX4 in innate immunity [51].

The significant region on chromosome 15 contains the β-globin locus. In ruminants, the β-globin locus was formed by the duplication of an ancestral four-gene set, ε-ε-γ-β, consisting of two embryonic-like genes (ε), a pseudogene (γ) and a β (β) globin gene [52]. Goats have three of the four-gene sets, where each set contains a different form of the β-globin gene. The three β-globin proteins are synthesised to make different forms of haemoglobin during development; in the foetus (βf), pre-adult (βc) and the adult (βa) [53]. In cattle, the β-globin locus contains two four-gene sets, containing βf and βc, with the foetal cluster containing an additional β-like pseudogene [54]. In sheep, the β-globin locus has two different haplotypes, A and B [42,43,55,56]. The arrangement of haplotype A is identical to that of the goat, containing three four-gene sets, whereas haplotype B, like cattle, lacks the four-gene set containing the pre-adult βc. While both haplotypes A and B contain the adult β-globin gene, HBB, the two are considered allelic variants (βA and βB) [42,43] differing structurally by at least seven amino acids [44-46]. Haemoglobin (Hb) from adult sheep with haplotype A (Hb-A; αβεγ) and B (Hb-B; αβεμ) is distinguishable electrophoretically [57].

The OARv3.1 (GCA_000298735.2) sheep genome contains the shorter haplotype B (Table 2), either as a result of the phenotypes of animals selected for sequencing or the challenges faced when assembling repetitive regions of the genome using short-read sequencing [58]. In comparison, the Rambouillet genome (ARS-UI_Ramb_v2.0; GCA_016772045.1), created using HiSeq X Ten and PacBio RS II reads, contains the longer haplotype A. There are several SNPs in the β-globin region on the high-density chip that appear to be copy number variants (CNVs) in animals with the longer haplotype A, mapping uniquely to OARv3.1 but mapping to two locations in both Oar_rambouillet_v1.0 (GCA_002742125.1) and ARS-UI_Ramb_v2.0 (Table S1). The Rambouillet genome should therefore also be used for future work searching for more accurate genetic markers for haemoglobin type in New Zealand sheep.

Various studies have shown variation in the frequency of haemoglobin A and B, indicating that there are advantages to both types [59]. Hb-A has a higher affinity for molecular oxygen than Hb-B and is, therefore, less efficient in transporting oxygen as it releases oxygen at a lower rate than normal [56,60-62]. As a result, Hb-B sheep tolerate anaemia better than Hb-A sheep due to their greater ability to liberate oxygen to tissue [56,59,60], whereas Hb-A sheep are more resistant to hypoxia [63]. Adult haplotype A sheep can switch from the synthesis of Hb-A (αβεγ) to the juvenile haemoglobin C (Hb-C; αβες) in response to hypoxia [63,64], anaemia [65-67] or erythropoietin injection [68]. It has been suggested that Hb-C is advantageous in compensating for relative tissue hypoxia of sheep and goats during severe anaemic states [69]. The β-globin locus is introduced into Tibetan sheep from argali (Ovis ammon), suggesting the importance of the locus for sheep living in high altitude and low oxygen environments [70].

Our results indicate that the β-globin locus haplotype, and therefore ovine haemoglobin type, can be estimated using SNPs within the β-globin locus, although one animal that was hypothesised to be homozygous for haplotype B (βB) appeared to also have a shortened form of the βa-globin peptide. It appears that haplotype A (haemoglobin A; Hb-A) animals are more tolerant to FE. The overall frequency of haplotype A in the animals genotyped in this study, however, is 0.5, indicating that the locus may be under balancing selection in these populations, and warrants further investigation. In crossbred ewes, Hb-A has been associated with beneficial effects for several health-related traits, including mastitis and parasite resistance, whereas Hb-B was associated with increased fertility [71]. In Norway, Hb-A lambs have been reported to be significantly more resistant to hepatogenous photosensitisation (alveld) as a result of ingestion of Natharicium ossifragum than Hb-B animals [72]. Hepatogenous photosensitisation, known as geeldikkop, also
occurs in small ruminants grazing *Tribulus terrestris* in South Africa. While the pathogenetic mechanisms underlying geeldikkop and FE differ, both affect the biliary tree, resulting in the accumulation of phylloerythrin [73]. In agreement with this study, and the Norwegian studies, Hb-A was shown to have a functional advantage in sheep affected by geeldikkop; both Hb-A and Hb-AB lambs were more resistant to serious haemolytic episodes, and a more severe temporary liver insufficiency developed in Hb-B animals than Hb-AB types [74]. Haemoglobin C (Hb-C) was present in Merino sheep recovering from a geeldikkop attack but was not present in the animals before the onset of the disease, with the authors suggesting that it may aid in recovery from the disease [75]. Of additional interest is that the frequency of haplotype A was higher in South African farms that had large scale outbreaks of haemolytic syndromes in their recent history [75]. Previous work looking at the distribution of haemoglobin in New Zealand sheep has found both types present in both Romney and Finnish Landrace breeds, with the A haplotype at a frequency of 0.6 and 0.84 in each breed, respectively [76]. Finnish Landrace lambs have also been reported to be less susceptible to a sporidesmin challenge than Romney lambs [77].

The underlying role of haemoglobin type in the response to FE is not straightforward. The mechanism of toxicity of sporidesmin, the causative mycotoxin, is believed to be through the generation of reactive oxygen species including superoxide radicals, leading to liver damage [78]. Other modes of action have also been postulated and may warrant further examination [79]. Antioxidants scavenge reactive oxygen species, and research in mice and human cell lines suggest haemoglobin may function as an antioxidant, suppressing oxidative stress and protecting cells from oxidative damage. Overexpression of haemoglobin in both murine kidney [80] and dopaminergic [81] cell lines impacted genes involved in oxidative stress and O2 homeostasis, respectively, and suppressed oxidative stress in human liver cell lines [82]. Antioxidant enzymes including superoxide dismutase (SOD), glutathione peroxidase (GPx) and catalase (CAT) have all been investigated as potential indicators of FE resistance. In the FE selection line animals, there was an overlap between enzyme activity distributions of the lines, therefore differences in enzyme activity were not large enough to be a reliable indicator of FE tolerance [83]. Analysis of allele frequencies in the FE selection line animals found significant differences between markers flanking the catalase gene (*CAT*) between the resistant and susceptible line, however, there was no evidence of linkage in outcross pedigrees, and it was concluded that the effect of catalase was probably recessive or minor [31]. Of interest is that the catalase gene is also located on chromosome 15, albeit 15.5 Mb from the β-globin locus. Additionally, a genome-scan experiment to screen for QTL affecting FE tolerance in Romney sheep identified a suggestive (F-statistic value=10.2) QTL on chromosome 15 (23.4–73.8 Mb), which, while large, overlaps the β-globin locus.

While the frequency of the haemoglobin haplotypes in the FE selection lines is unknown, in an analysis of 120 New Zealand Romney ewes of known haemoglobin type, Hb-B sheep had significantly higher levels of plasma copper, whole-blood selenium, and erythrocytic GPx than Hb-A sheep [76]. Previous studies also found Hb-B sheep had higher levels of plasma copper than type A sheep [84–86], and a strong positive relationship between copper and GGT levels post-exposure to sporidesmin has recently been reported [87]. A QTL for plasma copper concentration has been reported to be close to the haemoglobin locus in sheep [88]. The biological links between exposure to sporidesmin, haemoglobin type, antioxidant enzymes and copper do not appear to be straightforward and warrant further investigation.
5. Conclusions

Two loci for tolerance to facial eczema in sheep are reported on chromosomes 15 and 24. Markers at each locus explain 5% and 1% of the phenotypic variance in GGT21 levels after a sporidesmin challenge, respectively. The significant region on chromosome 15 overlaps the β-globin locus; mass spectrometry of haemoglobin from animals with differing genotypes in this region indicated that genotypes are associated with different forms of adult β-globin. Haplotype A animals appear to be more tolerant to FE. Given the importance of the β-globin region and observed intermediate haplotype frequencies of the animals in this study, the region may be under balancing selection. Further work is needed to explore this in a larger dataset. Haemoglobin haplotypes have previously been associated with variation in several health-related traits in sheep, and therefore warrant further investigation regarding their role in tolerance to facial eczema in sheep. This study highlights the power of using increased density genotyping for the identification of influential genomic regions, combined with subsequent inclusion on lower density genotyping platforms for rapid validation and industry uptake.

Supplementary Materials: The following are available online at www.mdpi.com/article/10.3390/genes12101560/s1, Table S1: FE discovery GWAS SNPs included in the design of the Illumina OvineLD BeadChip, mapped to publicly available ovine genome assemblies, Methods S1: Proteomic analysis of blood samples from genotyped animals, Figure S1: Quantile-quantile (QQ) plots of genome-wide association analysis for Facial Eczema in New Zealand sheep, Figure S2: Separation of haemoglobins from sheep blood by 4-20%T Tris-Tricine SDS-PAGE.

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Institutional Review Board Statement: Animals were managed following the provisions of the New Zealand Animal Welfare Act 1999, and the New Zealand Codes of Welfare developed under sections 68–79 of the Act. The collection of blood samples was approved by the AgResearch Animal Ethics Committee (Application number 14842) and complied with the institutional Codes of Ethical Conduct for the Use of Animals in Research, Testing and Teaching, as prescribed in the New Zealand Animal Welfare Act of 1999 and its amendments.

Data Availability Statement: The data used in this study are in the Sheep Improvement Limited (https://www.sil.co.nz/) database. These data were collected from commercial sheep and thus the information is accessible only with permission from the flock owners.

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