Association of inbreeding and regional equine leucocyte antigen homozygosity with the prevalence of insect bite hypersensitivity in Old Kladruber horse

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Summary

Inbreeding depression is the reduction of performance caused by mating of close relatives. In livestock populations, inbreeding depression has been traditionally estimated by regression of phenotypes on pedigree inbreeding coefficients. This estimation can be improved by utilising genomic inbreeding coefficients. Here we estimate inbreeding depression for insect bite hypersensitivity (IBH) prevalence, the most common allergic horse disease worldwide, in Old Kladruber horse. In a deep pedigree with 3214 horses (187 genotyped), we used a generalised linear mixed model with IBH phenotype from 558 horses examined between 1996 and 2009 (1368 records). In addition to the classical pedigree information, we used the single-step approach that enabled joint use of pedigree and genomic information to estimate inbreeding depression overall genome and equine leucocyte antigen (ELA) class II region. Significant inbreeding depression was observed in all models fitting overall inbreeding coefficients (odds ratio between 1.018 and 1.074, \( P < 0.05 \)) with the exception of Kalinowski’s new inbreeding \( (P = 0.0516) \). The increase of ELA class II inbreeding was significantly associated with increased prevalence of IBH \( (P = 0.0516) \). The increase of ELA class II inbreeding was significantly associated with increased prevalence of IBH \( (P = 0.027) \). However, when fitted jointly with the overall inbreeding coefficient, the effect of ELA class II inbreeding was not significant \( (P = 0.062) \). Overall, the higher ELA class II and/or overall inbreeding (pedigree or genomic) was associated with increased prevalence of IBH in Old Kladruber horses. The single-step approach provides an efficient use of all the available pedigree, genomic, and phenotype information for estimation of overall and regional inbreeding effects.

Keywords genomics, horse, inbreeding, inbreeding depression, insect bite hypersensitivity, pedigree

Mating of relatives is unavoidable in closed and finite populations and can lead to inbreeding depression – higher incidence of defects caused by recessive alleles and redistribution of genetic variance within and between populations. Inbreeding depression is a well-documented phenomenon, known as the reduction of performance in quantitative traits under constant selection pressure such as fitness and health traits (Crnokrak & Roff 1999; Bittles & Black 2010; Leroy 2014).

In livestock populations, inbreeding depression has been traditionally estimated by regression of phenotypes on pedigree inbreeding coefficients (Baker et al. 1945; Leroy 2014). Alternatively, some other pedigree-derived coefficients, for example Kalinowski’s new inbreeding coefficient, have been successfully used (McParland et al. 2009; Doekes et al. 2019; Curik et al. 2020). Development of large scale high-throughput genotyping has enabled quantification of individual genomic inbreeding in human (McQuillan et al. 2008), wild (Santure et al. 2010), and livestock populations.
The substitution of the pedigree with genomic inbreeding coefficient, improves the estimation of inbreeding depression from several perspectives (Kardos et al. 2016; Curik et al. 2017). In livestock populations, there are often many individuals in the pedigree that have phenotype records but are not genotyped. Legarra et al. (2020) showed how to estimate inbreeding coefficients by combining pedigree and genomic relationship matrices, which is commonly referred to as the single-step approach (Legarra et al. 2009). Recently, this approach was used in the estimation of inbreeding depression for semen traits in the Basco-Béarnaise dairy sheep breed (Antonios et al. 2021).

Classical horse breeds are often bred as closed populations and are known to reach high inbreeding often accompanied by inbreeding depression. Consequently, inbreeding depression has been observed for racing performance (Klemetsdal 1998; Todd et al. 2018), reproductive traits (Van Eldik et al. 2006; Müller-Unterberg et al. 2017), morphological traits (Gómez et al. 2009; Vostřy et al. 2011), and health traits (Sánchez-Guerrero et al. 2019).

The Old Kladruber horse, an old Czech breed established in 1579 that belongs to the Spanish horse group, has been maintained in a closed population for a long time with recent average inbreeding of 0.13 (Vostrá-Vydrůvá et al. 2016). In this population, inbreeding depression has been already observed for melanoma and vitiligo grading (Hofmanová et al. 2016) as well as for reproductive traits (Šichtař et al. 2017). The breed is affected also by insect bite hypersensitivity (IBH), the most common allergic skin disease caused by the bites of insects of the genus Culicoides. There is a common consensus that IBH is genetically controlled, and the starting impulse is a bite by mites of the Culicoides spp. and less frequently Simulium spp. The subsequent allergic reaction is a result of hypersensitivity to the allergens contained in their saliva (van der Rijt et al. 2008). The main symptoms are pruritus (itchiness), scaling, and hair loss. Clinical manifestation of IBH is due to an allergic reaction sharing some similarities with human atopic dermatitis. IBH is appearing in various horse breeds all over the world as a complex disease affected by environmental and genetic factors (Unkel et al. 1987; Eriksson et al. 2008; Schurink et al. 2011).

The impact of inbreeding on the IBH was analysed in Dutch Friesian horse and found non-significant (Schurink et al. 2011). In contrast, Andersson et al. (2012) have shown that increased homozygosity across the entire equine leucocyte antigen (ELA) class II region is associated with a higher risk of IBH. Unfortunately, Andersson et al. (2012) did not analyse if overall or regional inbreeding increases the risk of higher IBH.

The aim of this study was to estimate the effects of inbreeding on the prevalence of IBH in the Old Kladruber horse by logistic linear mixed regression from data with a heterogeneous combination of pedigree, genomic, and phenotypic information. Inbreeding was based on the combined pedigree-genomic single-step approach of Legarra et al. (2009, 2020) and compared with the classic pedigree approach. We also estimated the effect of ELA class II regional inbreeding to test the results from Andersson et al. (2012). With respect to the prevalent heterogeneous combination of pedigree, genomic, and phenotypic information in many livestock populations, the application of single-step approach is empowering.

**Materials and methods**

**Phenotype, pedigree, and genomic information**

The analysis was based on 1368 IBH inspections from 558 horses. We assumed that all horses have experienced the same probability of mosquito bites as all phenotyped horses have been kept at the same stud and were exposed to the same environmental factors, including biting insects. The National Stud Kladrub is situated in a temperate climate zone, at an altitude of about 200 m above the sea level in the eastern part of Bohemia, Czech Republic. While conditions are favourable for horses, they are also favourable for biting insects such as Culicoides, for example Culicoides nubeculosus and Culicoides obsoletus (van der Rijt et al. 2008). The data on the Old Kladruber horses were provided by the stud book. The inspections were performed visually twice a year (May and October) between 1996 and 2009 by the same person on a binary scale with respect to the prevalence of the typical skin lesions (0 when clinical symptoms were absent and 1 when clinical symptoms were present). Similar IBH scoring system was also used by Grevenhof et al. (2007), Peeters et al. (2015) and Schurink et al. (2011). During the measurement period, 558 horses were inspected several times and 29% of horses (163) showed clinical symptoms of IBH. The pattern of the insect bite hypersensitivity prevalence with respect to the sex and horse age at the inspection is presented in Fig. 1.

This prevalence was much higher compared to 8.1% observed in Swedish-born Icelandic horses (Eriksson et al. 2008), 8.8% in Shetland ponies (Schurink et al. 2009), 18.2% in Dutch Friesian broodmares (Schurink et al. 2011), and 10% in Belgian Warmblood horses (Peeters et al. 2015).

The total pedigree was up to 52 generations deep and comprised of 3214 individuals, with mean complete generation equivalent (EQC, Maignel et al. 1996) equal to 15.58 for the inspected horses. Overall, 187 horses were genotyped with the Equine SNP70 BeadChip (Illumina), consisting of 65 102 evenly distributed SNPs over the equine genome. This genotyping has been performed as a part of the routine diversity management, recently introduced into the breeding programme.

SNP data were edited and analysed using PLINK 1.9 (Chung et al. 2015). Only autosomal SNPs with known chromosomal position were analysed with individual call
rate >0.9 and SNP call rate >0.9. The minor allele frequency (MAF) was filtered according to two different approaches. Filtering for MAF >0.01 was performed in the calculation of the G matrix and FG, while filtering for MAF was not performed in the calculation of runs of homozygosity (ROH-based genomic inbreeding coefficients as recommended by Meyermans et al. (2020). To define the parameters of an ROH, we set the minimum density to 0.05 Mb/SNP and the largest gap to 1 Mb. The minimal number of SNPs in each ROH was set to 20. Similar parameters were used by Todd et al. (2018). The maximum number of heterozygous SNPs allowed within ROH was 0 and maximum number of missing SNPs allowed within ROH was 1. Unfortunately, only 47 inspected horses were genotyped (seven were affected and 40 were unaffected by IBH) while the remainder of 137 horses were not phenotyped (inspected for IBH). However, all included horses were highly related and were kept in the same environment, on the National Stud in Kladruby nad Labem.

Estimation of pedigree and genomic inbreeding coefficients

We estimated three different pedigree inbreeding coefficients: classical ($F_X$); classical restricted to five generations ($F_{X5}$); and Kalinowski’s new ($F_{NEW}$) inbreeding coefficient. When pedigree depth varies considerably, estimation of inbreeding depression can be biased as individual inbreeding coefficient estimates refer to different base populations. To avoid this potential bias, we used the $F_{X5}$ as all phenotyped horses had complete pedigrees for at least five generations. Note that this inbreeding coefficient refers to the recent inbreeding. $F_{NEW}$ is defined as the probability that any allele in an individual is autozygous for the first time and is assumed to better capture inbreeding depression caused by alleles with large detrimental effects or detrimental alleles that have not been purged (Doekes et al. 2019; Curik et al. 2020). $F_X$ and $F_{X5}$ were calculated by PEDIG software (Boichard 2002), while $F_{NEW}$ (Kalinowski et al. 2000) was calculated by gene dropping with 1 000 000 repetitions as suggested by (Baumung et al. 2015) and with corrections implemented in the GRAIN 2.2 software (Doekes et al. 2020).

Following the idea presented in Legarra et al. (2020) we estimated genomic inbreeding coefficients for all horses, by combining pedigree and genomic information with the single-step approach. This approach projects genomic relationships (G matrix; VanRaden 2008) to all ungenotyped individuals in a pedigree (A matrix; Legarra et al. 2009; Christensen & Lund 2010). The resulting projected matrix is often denoted as the I matrix. The projection requires careful consideration to account for different base populations of pedigree and genomic information. To address this, we used one ‘meta-founder’ for pedigree and expected allele frequency equal to 0.5 (Legarra et al. 2015). With this approach, we estimated three genomic inbreeding coefficients as the diagonal of I matrix minus one: (a) $F_H$ representing whole genome; (b) $F_{ELA-H}$ representing the ELA class II region with 87 SNPs between 32 and 34 Mb on ECA20 (Brinkmeyer-Langford et al. 2013); and (c) $F_{noELA-H}$ representing whole genome without the ELA class II region.

In addition, for the genotyped animals we also calculated overall and regional ROH inbreeding coefficient (McQuillan et al. 2008; Curik et al. 2014), to enable comparison of the inbreeding level in other horse populations. Calculated ROH inbreeding coefficient ($F_{ROH-SNPS}$) was defined as the proportion of the genome in ROH of a certain minimal length
relative to the whole autosomal genome \((F_{\text{ROH}} = I_{\text{ROH}}/I_{\text{AUTOSOMAL}})\). ELA class II ROH inbreeding coefficient \((F_{\text{ELA-H}})\) was defined in the same way but only for the ELA class II region. We estimated all genomic inbreeding coefficients with the \(r\) software, except for ROH inbreeding coefficients that were estimated with PLINK 1.9 (Chang et al. 2015) following settings of Todd et al. (2018).

Note that genomic inbreeding coefficients derived from \(H\) are defined relative to a population with allele frequency of 0.5 for all loci, due to the meta-founder single-step approach (Legarra et al. 2015). At this allele frequency, we expect 50% heterozygous loci. Therefore, these inbreeding coefficients express deviations from this expectation and an individual with less (more) than 50% heterozygous loci will have positive (negative) inbreeding coefficient. In contrast, pedigree and ROH based inbreeding coefficients are defined relative to a past base population (founders in the pedigree or conceptual base population defined by the ROH length). Therefore, these inbreeding coefficients are expressed as probabilities and strictly between 0 and 1.

Modelling of inbreeding depression

Depending on the information source used in the estimation of inbreeding coefficients, three different modelling approaches were used. Models from the first approach were based on the pedigree inbreeding coefficients \((F_X, F_{XX} \text{ and } F_{NEW})\). In the second approach, three models were based on inbreeding coefficients derived from the single-step \((F_{H}, F_{AX}, \text{ and } F_{ELA-H})\) with \(F_{noELA-H}\).

Calculations were performed with the GLIMIX procedure, which fitted logistic linear mixed model for repeated measurements (SAS Inst. Inc.). Here, the dependent variable \((y_i)\) has value of 1 (occurrence of IBH clinical symptoms) with probability \(\pi_i\) or 0 (absence of IBH clinical symptoms) with probability \(1 - \pi_i\) for observation \(i\). We modelled this probability as described in Citek et al. (2017) with the addition of inbreeding coefficients as covariates:

\[
\log \left( \frac{\pi_i}{1 - \pi_i} \right) = \text{Year}_i + \text{Age}_j + \text{Sex}_k + bF_i + p_i + a_i,
\]

where \(\text{Year}_i\) was a fixed effect of the \(i\)th year of inspection \((i = 1996, \ldots, 2009)\), \(\text{Age}_j\) is a fixed effect of the \(j\)th year of age of the horse \((j = 1, \ldots, 16)\), \(\text{Sex}_k\) is a fixed effect of the \(k\)th sex \((k = \text{stallion or mare}; \text{geldings were not included})\), \(p_i\) is a random permanent environment of the \(i\)th animal with \(p \sim N(0,\sigma_p^2)\), \(I_p\) is the identity matrix of the appropriate dimensions and \(\sigma_p^2\) is the permanent environmental variance, \(a_i\) is a random additive genetic effect of the \(i\)th animal with \(a \sim N(0,\sigma_a^2)\), \(Q\) is one of the following: pedigree relationship matrix \(A\) (Henderson 1976) when pedigree inbreeding coefficients were fitted; or single-step relationship \(H\) matrix (Legarra et al. 2009, 2015) when single-step inbreeding coefficients were fitted, while \(b\) presents regression coefficient for the various inbreeding coefficients. Separate models were fitted for each inbreeding coefficient \((F_X, F_{XX}, F_{NEW}, F_{H}, \text{ and } F_{ELA-H})\). We also modelled joint effects of the ELA class II regional inbreeding and overall inbreeding \((F_{H} \text{ with } F_{noELA-H})\). The suitability of fixed and random effects included in the model was discussed in Citek et al. (2017).

The goodness of fit of each of the analysed models was tested by a second-order bias correction of Akaike information criterion \((\text{AICc})\), which is corrected based on samples size (Burnham & Anderson 2002):

\[
\text{AICc} = -2\log\text{likelihood} + 2(k + 1)/\left(n - k - 1\right),
\]

where \(k\) is the number of estimated parameters and \(n\) is the number of observations.

Results and discussion

Inbreeding level

Descriptive statistics of various inbreeding coefficients, equivalent complete generations, and inspected horses for IBH are presented in Table 1. The observed pedigree inbreeding level in the Old Kladruber horse \((0.139)\), with increase of inbreeding equal to 0.009 per EQC, was higher than inbreeding level, ranging from 0.103 to 0.125, estimated in other horse breeds with very deep and complete pedigrees such as Thoroughbred (Mahon & Cunningham 1982), Lipizzan (Curik et al. 2003), Andalusian (Valera et al. 2005), and Lusitano (Vicente et al. 2012). Somewhat lower inbreeding, but with higher increase in inbreeding level per EQC \((0.0134)\), was observed for the recent \((F_{XX})\) as well as for the new inbreeding coefficient \((F_{NEW}; \text{Table 1})\). Results reveal that the genetic diversity was affected by bottleneck that occurred in the first half of the 20th Century. At that time, there was a substantial reduction in the size of Old Kladruber population. After this period, from 1973, the regeneration process of the Old Kladruber breed began (Votrá-Vydrová et al. 2016).

Genomic ROH inbreeding level \((F_{\text{ROH-SMB}})\) was lower than the estimated pedigree inbreeding level \((\text{Table 1})\) but with comparable increase of inbreeding per generations \((0.010)\) assuming that 5 Mb long ROH estimates are expected to descend from ancestors that are 10 generations back in the pedigree. The genomic ROH inbreeding was higher than observed by in Grilz-Seger et al. (2019) in Lipizzan \((0.035)\), Austrian Haflinger \((0.021)\), Italian Haflinger \((0.051)\), and Noriker \((0.029)\) horse breeds but similar to the Bosnian Mountain horse \((0.101)\) observed by Drum et al. (2018).

A much higher inbreeding estimate, 0.399 \((F_H)\), was obtained from the relationship matrix \(H\) due to a distant (ancient) base population. There was no notable difference between overall estimates with or without 87 SNPs from the
ELA class II region as the only small proportion of the genome (results not shown). All estimates of the ELA class II regional inbreeding were lower than overall inbreeding estimates \( F_{\text{ELA-H}} = 0.343 \) vs. \( F_h = 0.399 \), and \( F_{\text{ELA-ROH-SMB}} = 0.051 \) vs. \( F_{\text{ROH-SMB}} = 0.099 \), which might be a consequence of the heterozygote advantage (overdominance) in the ELA class II region.

Relationship among inbreeding coefficients

Correlations between the various inbreeding coefficients are presented in Fig. 2. High correlations, ranging from 0.505 to 0.955, were observed among all overall inbreeding coefficients \( (F_X, F_{XS}, F_{NEW}, F_H \text{ and } F_{ROH-SMB}). \text{ Low correlations were observed between all overall and regional ELA class II inbreeding coefficients, while relatively high correlation was observed between two regional estimates } F_{\text{ELA}} \text{ and } F_{\text{ELA-ROH-SMB}} (0.651). \)

Inbreeding depression

The results of the inbreeding depression analyses for the prevalence of IBH in Old Kladruber Horse are presented in Table 2. Significant overall inbreeding depression, that is the increase of overall inbreeding increased the prevalence of IBH, was estimated in models with \( F_X \), \( F_{XS} \), and \( F_H \), while the estimate with \( F_{NEW} \) was in the same direction and close to significant \( (P = 0.0516) \).

When fitted alone in the model, ELA class II inbreeding was significantly associated with increasing the prevalence of IBH, corroborating the negative impact of inbreeding on IBH. Interestingly, according to AIC\textsubscript{c} values, the model with \( F_{\text{ELA-H}} \) as a single covariate was the best model in this study. All models based on the single-step inbreeding fitted the data better than models based on pedigree inbreeding (Table 2) pointing to the advantage of utilising genomic information in addition to pedigree information.

The presence of a polygenic effect on IBH has been observed in several studies – narrow sense heritability \( (h^2) \) on an IBH status (four categories) was estimated at 0.33 in Swedish-born Icelandic horses (Eriksson et al. 2008) and at 0.16 on a liability scale in Dutch Friesian Broodmares (Schurink et al. 2011), between 0.08 and 0.24 in the Dutch Shetland breeding mares (Schurink et al. 2009) and between 0.36 and 0.63 in the Old Kladruber horse (Citek et al. 2017). The wide range in the last two studies was due to different model definitions. The polygenic nature of IBH was also manifested in several GWAS studies, while several candidate genes located on \( ECA7 \), \( 9 \), \( 11 \) and \( 20 \) were also reported (Schurink et al. 2012, 2013; Shrestha et al. 2015; Velle et al. 2016). The most consistent association with IBH across various breeds was observed for the major histocompatibility complex located on \( ECA20 \), or more precisely for the ELA class I and II regions (Martí et al. 1992; Andersson et al. 2012; Schurink et al. 2012).

In contrast to other genetic analyses of IBH, this is the first study where significant overall inbreeding depression is reported for IBH. Up to now, the effect of inbreeding on IBH has been estimated only in Dutch Friesian horse (Schurink et al. 2011), but no significant estimates were observed. Our results support the positive association of the ELA class II region homozygosity with IBH previously reported by Andersson et al. (2012). In the models where the overall and regional ELA class II inbreeding were analysed simultaneously, regional inbreeding was not significant while overall inbreeding was significant. Still, we think that in the Old Kladruber horse, both overall and ELA class II inbreeding are associated with increased prevalence of IBH. We have five arguments this. First, in almost all models overall inbreeding depression was significant, while regional inbreeding depression was significant when fitted independently and gave the best model fit according to AIC\textsubscript{c}. Second, estimates obtained with the single inbreeding covariate models were similar to estimates obtained with two inbreeding covariate models. Third, correlations between overall and regional ELA class II inbreeding coefficients were very small, which should exclude potential confounding. Fourth, the regional inbreeding depression was close to significant \( (P = 0.062) \) in the two inbreeding covariate model. Fifth, significant regional inbreeding depression was observed in other studies (Andersson et al. 2012).

Overall, models based on the single-step relationship matrix combining genomic and pedigree information outperformed models based only on pedigree information.
The utilisation of all available information in the estimation of inbreeding depression seems intuitively logical, although computer simulations are required to better explore all properties and assumptions of the applied models. With a recent estimate of inbreeding depression for semen traits in the Basco-Béarnaise dairy sheep breed (Antonios et al. 2021), this is the first study to use the single-step approach to estimate total and regional inbreeding depression. This option might be important for analysing traits where genes with large detrimental impact are participating in the inbreeding depression, particularly as the regional inbreeding might substantially differ from the expected inbreeding estimated by the pedigree inbreeding coefficients (Kardos et al. 2016; Curik et al. 2017).

**Conclusions**

The overall pedigree or genomic and regional ELA class II inbreeding were positively associated with the prevalence of insect bite hypersensitivity in Old Kladruber horses. The utilisation of single-step inbreeding coefficients that combines all the available genomic and pedigree information is a powerful tool for analysing traits where genes with large detrimental impact are participating in the inbreeding depression.
useful approach in the estimation of detrimental inbreeding effects.

Acknowledgements
L.V. and H.V.V. acknowledge support from the Ministry of Agriculture of the Czech Republic (Project Nos. QK1910156 and MZE-RO0719). J.C. acknowledges support from the Ministry of Agriculture of the Czech Republic (Project No. QK1910320). I.C. acknowledges support from the Croatian Science Foundation (Project ANAGRAMS-IP-2018-01-8708) while G.G. acknowledges support from the BBSRC to The Roslin Institute (BBS/E/D/30002275) and The University Of Edinburgh’s Data-Driven Innovation Chancellor’s Fellowship.

Data availability statement
Phenotypic and genotypic data were obtained from National Stud in Kladruby nad Labem. These datasets were used under a material transfer agreement, and hence, are not publicly available. However, data are available upon request to LV and with the permission of National Stud in Kladruby nad Labem.

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Additional supporting information may be found online in the Supporting Information section at the end of the article.
Table S1. Estimated environmental and inbreeding effects with significant impact on insect bite hypersensitivity prevalence in Old Kladruber Horse: model is based on pedigree relationship matrix and five generations inbreeding coefficient ($F_{X5}$); model AICc value was equal to 5541.

Table S2. Estimated environmental and inbreeding effects with significant impact on insect bite hypersensitivity prevalence in Old Kladruber Horse: model is based on pedigree relationship matrix and classical inbreeding coefficient ($F_X$); model AICc value was equal to 5546.

Table S3. Estimated environmental and inbreeding effects with significant impact on insect bite hypersensitivity prevalence in Old Kladruber Horse: models is based on pedigree relationship matrix and Kalinowski’s new inbreeding coefficient ($F_{NEW}$); model AICc value was equal to 5551.

Table S4. Estimated environmental and inbreeding effects with significant impact on insect bite hypersensitivity prevalence in Old Kladruber Horse: models is based on single-step relationship matrix and inbreeding coefficient representing the ELA class II region ($F_{ELA-H}$); model AICc value was equal to 5319.

Table S5. Estimated environmental and inbreeding effects with significant impact on insect bite hypersensitivity prevalence in Old Kladruber Horse: model is based on single-step relationship matrix and inbreeding coefficients representing the ELA class II region ($F_{noELA-H}$) and the whole genome without the ELA class II region ($F_{noELA-H}$); model AICc value was equal to 5361.

Table S6. Estimated environmental and inbreeding effects with significant impact on insect bite hypersensitivity prevalence in Old Kladruber Horse: model is based on single-step relationship matrix and inbreeding coefficient representing the whole genome ($F_{ELA-H}$); model AICc value was equal to 5368.