Ten years of the horse reference genome: insights into equine biology, domestication and population dynamics in the postgenome era

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Summary

The horse reference genome from the Thoroughbred mare *Twilight* has been available for a decade and, together with advances in genomics technologies, has led to unparalleled developments in equine genomics. At the core of this progress is the continuing improvement of the quality, contiguity and completeness of the reference genome, and its functional annotation. Recent achievements include the release of the next version of the reference genome (EquCab3.0) and generation of a reference sequence for the Y chromosome. Horse satellite-free centromeres provide unique models for mammalian centromere research. Despite extremely low genetic diversity of the Y chromosome, it has been possible to trace patrilines of breeds and pedigrees and show that Y variation was lost in the past approximately 2300 years owing to selective breeding. The high-quality reference genome has led to the development of three different SNP arrays and WGSs of almost 2000 modern individual horses. The collection of WGS of hundreds of ancient horses is unique and not available for any other domestic species. These tools and resources have led to global population studies dissecting the natural history of the species and genetic makeup and ancestry of modern breeds. Most importantly, the available tools and resources, together with the discovery of functional elements, are dissecting molecular causes of a growing number of Mendelian and complex traits. The improved understanding of molecular underpinnings of various traits continues to benefit the health and performance of the horse whereas also serving as a model for complex disease across species.

Keywords ancient genomes, centromeres, complex traits, domestication, Mendelian traits, modern breeds, signatures of selection, Y chromosome

Introduction

The horse (*Equus caballus*, ECA) occupies a special place amongst farm animals. Since domestication about 5500 years ago (Outram *et al.* 2009; Librado *et al.* 2016; Gaunitz *et al.* 2018), horses have served humans in agriculture, warfare and transportation, and as valued companions. In modern times, they continue to interact with humans in many different ways and are an important

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part of the leisure industry. Humans have selectively bred horses for performance traits (speed, endurance, strength, gait), appearance (size, color, conformation) and temperament, resulting in 400-500 different breeds (Hendricks 2007; Petersen et al. 2013a). The derivation of breeds from selective breeding and the inclusion of only individuals with breed-defining characteristics have resulted in genomic features that vary among populations (Petersen et al. 2013b). Furthermore, over 130 equine hereditary traits (e.g. muscle disorders, allergies, asthma) can serve as valuable models for the study of similar human conditions (Wade et al. 2009; OMIA: https://omia.org/home/). Interest in economic, biomedical, evolutionary and basic science aspects of the horse combined with intense passion for advancing knowledge on equids have promoted organized studies of the horse genome, initiated at the First

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International Equine Gene Mapping Workshop in 1995 (reviewed by Chowdhary & Bailey 2003; Finno & Bannasch 2014). Whereas multiple important achievements mark the 25 years of horse genomics (reviewed by (Chowdhary & Bailey 2003; Chowdhary & Raudsepp 2006; Chowdhary et al. 2008; Chowdhary & Raudsepp 2008; Finno et al. 2009; Brosnahan et al. 2010; Bailey & Brooks 2013; Chowdhary 2013; Finno & Bannasch 2014; Librado et al. 2016), without doubt the most important milestone was the generation of the reference genome assembly (Wade et al. 2009), made possible through collaborative efforts of the research community. The completion of a reference genome has shaped horse genome studies during the past 10 years and, together with the ongoing revolution in genomics technologies, particularly next-generation sequencing (reviewed by van Dijk et al. 2014, has taken equine genomics to a new level.

This review will focus on these technology-driven achievements in the post-genome era of horse genomics: the development of new genomics tools and resources, the improvement of the reference sequence and functional annotation of the horse genome. We will discuss how the cutting-edge genomics technologies have improved understanding of the evolutionary history of the horse, the genetic makeup of individual horse breeds, and the study of Mendelian and complex traits.

The horse genome

EquCab2.0

In 2019, we celebrate the tenth anniversary of the publication of the genome sequence of the Thoroughbred mare, Twilight (Wade et al. 2009). The sequence of Twilight represented the first equine and Perissodactyl genome and established a reference sequence for the domestic horse (Wade et al. 2009). Sequencing was performed using the Sanger method with an average of 6.8-fold genomic coverage. The contiguity of the assembly was increased by the inclusion of end sequences of approximately 315 000 BAC clones from the CHORI-241 BAC library, which represents sequences from Twilight's half-brother Bravo (https://bacpacresources.org/; Leeb et al. 2006). Radiation hybrid and cytogenetic maps (Raudsepp et al. 2008) assisted with chromosomal anchoring and orienting the scaffolds. The result was a high-quality, 2.5 Gb draft assembly, denoted as EquCab2.0, which incorporated the 31 horse autosomes, the X chromosome and the mitochondrial genome. Sequence annotation by the ENSEMBL pipeline predicted 20 322 protein-coding genes, comparable with human, mouse and other mammals. An important part of the horse genome project was the identification of over a million SNPs, which directed the development of genomic tools for mapping in the horse. The SNPs were generated from the diploid genome of Twilight and by partial

sequencing of seven additional horses of diverse breeds. The reference assembly and the SNP map marked a turning point in horse genomics by providing resources for driving subsequent molecular, clinical and evolutionary studies in the horse (reviewed by Wade 2013; Finno & Bannasch 2014; Ghosh *et al.* 2018).

EquCab3.0

Despite its high quality, EquCab2.0 is a draft assembly and, as a product of the available technology of the time, has limitations (Kalbfleisch et al. 2018). The assembly contains numerous gaps, mainly in structurally complex genomic regions, which include segmental duplications and CNV sites (Doan et al. 2012; Ghosh et al. 2014). Approximately 0.2 Gb of sequence reads in EquCab2.0, mainly highly repetitive regions, remain unassembled and unassigned to chromosomes (Wade et al. 2009; Wade 2013). Furthermore, recent re-sequencing of the whole genome (Rebolledo-Mendez et al. 2015) of selected complex regions such as the MHC (Viluma et al. 2017) and part of the PAR (Rafati et al. 2016), together with transcriptome sequencing (Coleman et al. 2010, 2013b) and gene annotations (Hestand et al. 2015; Balmer et al. 2017; Mansour et al. 2017), have revealed several inconsistencies in the assembly. Therefore, taking advantage of new genomic technologies, the genome of Twilight was recently re-sequenced and assembled, resulting in EquCab3.0 (Kalbfleisch et al. 2018). The new assembly was built upon the solid foundation of EquCab2.0 (Wade et al. 2009), physical maps (Raudsepp et al. 2008) and BAC end sequences (Leeb et al. 2006). These were augmented with 45-fold short-read data that improved the accuracy of unique regions of the genome. Chromosome length scaffolding was achieved by including Chicago[®] and Hi-C proximity ligation data and 16-fold long-read Pacific Biosciences (PacBio) data.

The new assembly improved both the contiguity and composition of the horse reference genome. For example, the number of gaps was reduced 10-fold, from 55 Mb (2.2% of the genome) in EquCab2.0 to 9 Mb (0.34% of the genome) in EquCab3.0, and the number of assembled bases in the incorporated chromosomes improved from 2.33 to 2.41 Gb (3% increase). Contiguity improved nearly 40-fold and it is noteworthy that only ECA6 comprises two scaffolds; all other chromosomes comprise a single scaffold. In addition, the use of the Chromium 10X platform allowed for true haplotype phasing, so the final assembly has the most common and likely ancestral allele at each heterozygous site. Comparison of EquCab2.0 and EquCab3.0 mapping statistics for 13 previously published ancient DNA samples (Schubert et al. 2014; Librado et al. 2015, 2017; Gaunitz et al. 2018) showed that significantly more reads mapped to the new assembly, demonstrating its improved utility for the mapping of highly fragmented and damaged DNA samples.

Table 1 demonstrates that the size and gene content of the reference genome have marginally but consistently increased for all chromosomes. The exception is ECA5, which is smaller and has fewer genes in EquCab3.0 than in EquCab2.0, most likely owing to inconsistencies in the previous assembly. Two chromosomes, ECA12 and ECAX, increased in length by almost 4 Mb owing to incorporating previously unplaced contigs, and the number of annotated, non-coding genes has essentially increased for all chromosomes. This is probably due to improved sequence composition and better annotation pipelines, although detailed annotation of the horse genome will be the task of the ongoing Functional Annotation of ANimal Genomes (FAANG) project (described in detail below). The necessary prerequisite for FAANG is a high-quality and contiguity reference genome (Andersson et al. 2015), and EquCab3.0 serves this purpose. Nevertheless, EquCab3.0 is also a tribute to EquCab2.0 and testimony that the first Sanger assembly of the Twilight genome 10 years ago was an outstanding achievement, providing the foundation for answering genetic questions related to the horse.

Unique features of horse centromeres

Centromeres are typically not part of reference genomes because they are composed of arrays of nearly identical tandem repeats, known as satellite DNA, and are extremely difficult to assemble even with long-read sequencing technologies (Miga et al. 2014; Jain et al. 2018). An outstanding exception is horse ECA11, which has no satellite DNA and is the first sequenced example of a natural satellite-free and evolutionary 'immature' centromere (Wade et al. 2009; Wade 2013). This unusual and most interesting discovery led to in-depth studies of centromeres in horses and equids. Three centromere satellite families, 37cen, 2PI and EC137, have been isolated, characterized and localized in horse and equid chromosomes, revealing that in donkeys and zebras a large number of centromeres lack satellite DNA (Piras et al. 2010; Nergadze et al. 2014). With the help of chromatin immunoprecipitation sequencing (ChIP-seq) methodology, it was shown that 37cen satellite binds to the centromeric histone H3 variant CENPA protein, is transcriptionally active and probably required for centromere function (Cerutti et al. 2016). However, further studies of satellitefree horse ECA11 revealed that centromere is defined epigenetically by binding of the CENPA protein and not by the presence of satellite repeats (Purgato et al. 2015). Interestingly, the size and exact location of the approximately 100 kb (kilo-base) CENPA binding region differ between individuals, giving rise to epialleles, a phenomenon known as centromere sliding (Purgato et al. 2015; Giulotto et al. 2017). These observations were further strengthened by a study of donkey centromeres (Nergadze et al. 2018). The donkey genome has 16 chromosomes with satellite-free centromeres, which is perfectly compatible with genome stability and species survival. Analysis of the transmission of CENPA-binding epialleles in mules and hinnies shows that centromeric domains are inherited as Mendelian traits, although centromere sliding can happen in one generation. Overall, the discovery of satellite-free centromeres in horses and equids has provided a unique model for the study of the evolution, dynamics and molecular regulation of mammalian centromeres.

The Y chromosome

The female-based horse reference genome is incomplete because it does not include the Y chromosome. It is therefore noteworthy that, concurrently with the release of Equ-Cab3.0 (Kalbfleisch et al. 2018), the first comprehensive assembly and annotation of the male-specific region of the horse Y (MSY) was published (Janečka et al. 2018). The 9.5 Mb assembly represents the Y chromosome of a Thoroughbred stallion Bravo, a half-brother of Twilight, thus completing the Thoroughbred-based horse reference genome. The assembly provides information about the horse Y chromosome organization, sequence classes and gene content (52 genes and 174 transcripts). Notably, the study identified a novel testis-expressed XY ampliconic sequence class ETSTY7, which is shared with the parasite Parascaris genome, providing evidence for eukaryotic horizontal gene transfer. Alignment of MSY assembly with horse, donkey and mule testis transcriptome data suggests candidate genes for stallion fertility. The MSY assembly provides a needed reference toward improved understanding for the role of the Y chromosome in equine male development and fertility.

Another keen interest in the male-specific and nonrecombining Y chromosome is its inheritance exclusively through male lineages. This makes Y chromosome sequence polymorphisms excellent markers for tracing the patriline history of ancient horses and modern breeds. However, until recently, the main problem with the horse Y chromosome was the lack of sequence polymorphism (Lindgren et al. 2004; Wallner et al. 2004; Brandariz-Fontes et al. 2013), suggesting a limited number of patrilines in horse domestication and omitting the use of the Y to trace those patrilines. Even though one polymorphic microsatellite, YA16, with two alleles was detected in a few individuals of indigenous Chinese horses (Ling et al. 2010), no variants were found in other modern breeds. In contrast, sequencing just a 4 kb Y chromosome fragment from eight ancient horses revealed 28 segregating sites and eight haplotypes (Lippold et al. 2011), demonstrating considerable diversity in the ancestral horse Y and the loss of this diversity during domestication. Nevertheless, persistent search, combined with the use of high-throughput next-generation sequencing technologies, led to step-wise discovery of a limited number of variable sites also in the modern horse Y chromosome. These included two SNPs that defined six Y haplotypes in modern breeds (Wallner et al. 2013), and two

	Size, Mb		Protein co	ding genes	Non-codin	g genes
Horse chromosome	EquCab2	EquCab3	EquCab2	EquCab3	EquCab2	EquCab
ECA1	185.8	188.3	1656	1683	166	705
ECA2	120.9	121.4	1062	1077	103	450
ECA3	119.5	121.4	838	883	79	391
ECA4	108.6	109.5	750	735	90	354
ECA5	99.7	96.8	1004	999	99	322
ECA6	84.7	87.2	962	988	63	326
ECA7	98.5	100.8	1236	1367	86	386
ECA8	94.1	97.6	714	773	69	397
ECA9	83.6	85.8	451	447	39	296
ECA10	84.0	85.2	1032	1146	61	346
ECA11	61.3	61.7	1086	1141	109	282
ECA12	33.1	37.0	639	738	34	177
ECA13	42.6	43.8	657	711	44	175
ECA14	94.0	94.6	665	693	63	368
ECA15	91.6	92.9	659	664	54	376
ECA16	87.4	89.0	683	713	73	299
ECA17	80.8	80.7	352	338	44	288
ECA18	82.5	82.6	422	416	62	259
ECA19	60.0	62.7	407	418	47	208
ECA20	64.2	65.3	709	738	50	262
ECA21	57.7	59.0	376	388	37	199
ECA22	49.9	50.9	533	560	53	244
ECA23	55.7	55.6	296	294	53	251
ECA24	46.7	48.3	381	453	141	298
ECA25	39.5	40.3	523	554	42	160
ECA26	41.9	43.1	221	232	19	129
ECA27	40.0	40.3	215	232	20	110
ECA28	46.2	47.3	383	388	46	189
ECA29	33.7	34.8	181	170	25	156
ECA30	30.1	31.4	171	185	20	104
ECA31	25.0	26.0	140	154	13	86
ECAX	124.1	128.2	853	821	151	371
ECAY ¹	n/a	9.5	n/a	52		n/a
Total	2440.8	2419	15 428	21 151	2055	8964

Table 1Chromosome-wise comparison ofEquCab2.0andEquCab3.0.

Ensembl: http://www.ensembl.org/index.html for assembly size and annotated gene content. ¹Y chromosome data are from Janečka *et al.* (2018).

microsatellites-YP9 in Hucul and Mongolian horses and YN04 in a Shetland pony (Kreutzmann et al. 2014). This small but significant progress led to more systematic discoveries of Y chromosome variants based on WGS and MSY assembly. Fifty-three variants (50 SNPs and three indels) and 24 haplotypes were identified from a 1.46 Mb MSY sequence of 52 males of 21 different breeds (Wallner et al. 2017), followed by the description of another 211 variants and 58 haplotypes by screening 5.8 Mb of MSY in 130 horses of rural breeds and nine Przewalski's horses (Felkel et al. 2019). This is an awaited breakthrough in horse Y chromosome research and has already launched a number of studies to determine the time and cause(s) of the loss of Y variation (Wutke et al. 2018), as well as enabling the tracing of patrilines of modern breeds and pedigrees (Wallner et al. 2017; Felkel et al. 2018, 2019; Kakoi et al. 2018; Khaudov et al. 2018; Han et al. 2019).

Recent studies of ancient samples provide clues about when and why the horse Y chromosome lost its genetic

diversity. A study of four MSY polymorphic markers in 96 ancient stallions from early domestication indicates that the reduction of Y diversity over time was not due to genetic drift or founder effect, but the result of artificial selection that started during the Iron Age and continued during the Roman period (Wutke *et al.* 2018). This is supported and further elaborated by a more extensive study involving 105 ancient stallions and over 1500 polymorphic MSY sites showing that Y chromosome nucleotide diversity decreased steadily during the last approximately 2000 years but dropped to present levels only after 850–1350 AD (Fages *et al.* 2019)

Genomics tools and resources

SNP genotyping array

The most impactful tool and resource for horse genomics has certainly been the reference genome. As noted, EquCab2.0 has been the critical template for the discovery of millions of sequence polymorphisms from diverse horse breeds, leading to the development of three generations of SNP chips which have been utilized to map traits and understand breed diversity and signatures of selection. First- and second-generation DNA genotyping arrays, containing 54 602 and 74 500 SNP markers, respectively, became available in 2011 (reviewed in Finno & Bannasch 2014). A number of phenotypic traits of interest and disease traits were identified using these arrays, including Lavender Foal Syndrome (Brooks et al. 2010), alternate gait (Andersson et al. 2012), iris color variation (Mack et al. 2017) and ocular squamous cell carcinoma (Bellone et al. 2017; Tables 2 and 3). Additionally, these resources were used to identify breed specific signatures of selection (Petersen et al. 2013b) that aid in our understanding of the biology behind performance and other selected traits in the horse (Andersson et al. 2012; Petersen et al. 2014b).

In 2017, a third-generation SNP array was developed containing 670 805 SNP markers (MNEc670k array, Affymetrix; Schaefer *et al.* 2017). This array was designed using WGS from 156 horses representing 24 distinct breeds. Mean inter-SNP distance was estimated at 3756 bp with SNP selection aimed at tagging approximately 2 million SNPs. To date, two published studies have successfully mapped traits with the 670K array, followed by fine-mapping with WGS. The first identified a nonsense variant in *ST14* associated with Naked Foal Syndrome in the Akhal-Teke (Bauer *et al.* 2017) and the second identified genetic variants in two genes, *KRT25* and *SP6*, responsible for a curly coat in horses (Thomer *et al.* 2018).

With the recently updated reference assembly of the equine genome (Kalbfleisch *et al.* 2018), the SNP array coordinates were remapped to EquCab3.0 (Beeson *et al.* 2019). The raw reports with EquCab3.0 SNP coordinates for the MNEc670k array are hosted at https://www.anima lgenome.org/repository/pub/UMN2018.1003/. Furthermore, coordinates between the two assemblies can be easily converted now at NCBI: https://www.ncbi.nlm.nih.gov/ge nome/tools/remap. The high-density SNP array resource is undoubtedly aiding in the mapping of other important traits and we anticipate a continued increase in the number of discoveries made possible.

WGSs of individual horses

As next-generation sequencing continues to become more affordable, WGSs of horses are being generated worldwide. At the time of this writing, 1936 public WGSs are available for the horse through the NCBI Sequence Read Archive (https://www.ncbi.nlm.nih.gov/sra). This tremendous resource provides investigators with a database of horse genomes to screen potential variants, and with the continuing addition of phenotypic metadata, this resource will facilitate future studies for many years to come.

WGSs are often used to identify putative genetic variants within regions identified by GWAS. Two recent examples include the discovery of the splice site mutation in B4GALT7 associated with dwarfism and joint laxity in Friesian horses (Leegwater et al. 2016). Using the first-generation SNP array (~50K markers), a region on ECA14 was initially identified by GWAS. Four dwarf cases and three unrelated controls then underwent WGS. Pooling data from the four cases and variant calling identified the putative variant in B4GALT7, later confirmed via Sanger sequencing and demonstrated to affect splicing through analysis of cDNA from affected horses. Notably, this is the second gene with a role in protein glycosylation in which a pathogenic mutation has been identified in Friesian horses. A nonsense mutation in B3GALNT2 involved in muscular dystrophy with hydrocephalus in stillborn foals was discovered previously by the same group using similar techniques (Ducro et al. 2015). This approach was also utilized to unravel the genetic mutation responsible for both immune-mediated myositis (Finno et al. 2018) and non-exertional rhabdomyolysis (Valberg et al. 2018) in the Ouarter Horse as well as congenital hepatic fibrosis in the Swiss Franches-Montagnes (Drogemuller et al. 2014).

Many recently discovered, potential causal genetic variants in the horse have been identified through WGS alone. Public WGS was screened for putative deleterious variants associated with stallion infertility and further evaluated in a group of 337 fertile stallions across 19 breeds (Schrimpf et al. 2016). A variant in NOTCH1 (g.37455302G>A) was identified as a significant stallion fertility locus in Hanoverian stallions. Additionally, nine candidate fertility loci with missing homozygous mutant genotypes were validated. WGS has recently identified a genetic cause of occipitoatlantoaxial malformation in an Arabian horse (Bordbari et al. 2017) and dwarfism in a Miniature Shetland pony (Metzger et al. 2017). WGS data have also been used to identify novel pigmentation variants (Henkel et al. 2019). Finally, WGS has been used to investigate the characteristics of highly selected breeds (Metzger et al. 2014).

With the continued reduction in the cost of WGS, we can expect to soon see many more publicly available genomes for the horse. The beauty of this resource is that, once available, the data can be used for diverse projects worldwide.

Other tools

In the past 10 years, several other array platforms have been generated to study the horse genome. On the basis of EquCab2.0, an exon array (Doan *et al.* 2012) and two whole genome tiling arrays (Ghosh *et al.* 2014; Wang *et al.* 2014) were constructed for the discovery of CNVs. However, in a short time, these platforms have given way to more comprehensive WGSs. Likewise, cDNA and oligonucleotide microarrays, developed for gene expression studies

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Table 2 Genetic variants identified for	or traits influencing	pigmentation	_				
Phenotype	Gene	Allele	Type of variant	Chromosome	Breed	Year published	PubMed ID
Chestnut	MC1R	e	Missense	m	Many	1996	8995760
Frame overo (Lethal White Foal	EDNRB	0	Missense	17	American Paint Horse, Miniature Horse,	1998	9530628
Syndrome, LWFS)					Pinto Horse, Quarter Horse Thoroughbred, Appaloosa		
Chestnut	MC1R	e ^a	Missense	ſ	Black Forest	2000	11086549
Recessive black	ASIP	а	11 bp deletion		Many	2001	11353392
Cream dilution	SLC45A2	C _C	Missense	21	Many	2003	12605854
Sabino 1	KIT	SB1	Splicing	ſ	Appaloosa, Haflinger, Lipizzan, Noriker,	2005	16284805
					Quarter Horse		
Silver (Multiple Congenital Ocular	PMEL	Ζ	Missense		American Miniature Horse, Icelandic Rocky	2006	17029645
Anomalies, MCOA)					Mountain, Kentucky Mountain Horse		
Tobiano	KIT (proposed)	То	~43 Mb inversion	m	Many	2007	18253033
Dominant white	KIT	W1	Nonsense (stop-gain)	m	Franches-Montagnes	2007	17997609
Dominant white	KIT	W2	Missense	c	Thoroughbred	2007	17997609
Dominant white	KIT	W4	Missense	c	Camarillo White Horse	2007	17997609
Dominant white	KIT	W3	Nonsense (stop-gain)	c	Arabian	2007	17997609
Grey (melanoma susceptibility)	STX17	ט	4.6 kb intronic duplication	25	Many	2008	18641652
Champagne dilution	SLC36A1	Ch	Missense	14	Spanish Mustang, Tennessee Walking Horse,	2008	18802473
)					Quarter Horse, and pony breeds		
Dominant white	KIT	W11	Splicing	c	South German Draft	2009	19456317
Dominant white	KIT	W8	Splicing	c	Icelandic	2009	19456317
Dominant white	KIT	W7	Splicing	c	Thoroughbred	2009	19456317
Dominant white	KIT	W5	1 bp deletion, frameshift	£	Thoroughbred	2009	19456317
Dominant white	KIT	6M	Missense	£	Holstein	2009	19456317
Dominant white	KIT	W10	4 bp deletion, frameshift	£	Quarter Horse	2009	19456317
Dominant white	KIT	W6	Missense	£	Thoroughbred	2009	19456317
Lavender foal syndrome	MY05A	LFS	1 bp deletion, frameshift		Arabian	2010	20419149
Dominant white	KIT	W12	5 bp deletion	œ	Thoroughbred	2010	https://doi.org/
							10.11111/j.1365-
							2052.2010.
							02135.x
Dominant white	KIT	W13	Splicing	ε	American Miniature Horse, Quarter Horse	2011	21554354
Dominant white	KIT	W16	Missense	ε	Oldenburg	2011	21554354
Dominant white	KIT	W14	54 bp deletion	£	Thoroughbred	2011	21554354
Dominant white	KIT	W17b	Missense	с	Japanese Draft	2011	21554354
Dominant white	KIT	w17a	Missense	c	Japanese Draft	2011	21554354
Dominant white	KIT	W15	Missense	c	Arabian	2011	21554354
Macchiato	MITF	macchiato	Missense	16	Franches-Montagnes	2012	22511888
Splashed white	MITF	SW3	5 bp deletion, frameshift	16	Quarter Horse	2012	22511888
Splashed white	MITF	SW1	Insertion 11 bp, regulatory	16	American Miniature Horse, American Paint	2012	22511888
					Horse, Appaloosa, Icelandic, Morgan, Old-		
					Tori, Quarter Horse, Shetland Pony,		
					Trakehner		

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Table 2 (Continued)							
Phenotype	Gene	Allele	Type of variant	Chromosome	Breed	Year published	PubMed ID
Splashed white	PAX3	SW2	Missense	6	Lipizzan, Noriker, Quarter Horse	2012	22511888
Dominant white	KIT	W18	Splicing	S	Swiss Warmblood	2013	23659293
Dominant white	KIT	W20	Missense	m	American Paint Horse, Appaloosa, German Riding Pony, Gipsy, Noriker, Old-Tori, Oldenburg, Quarter Horse, Thoroughbred, Warmhlood, Welsh Pony	2013	23659293
Dominant white	KIT	W19	Missense	ſ	Arabian	2013	23659293
Splashed white	PAX3	SW4	Missense	9	Appaloosa	2013	23659293
Leopard Complex Spotting (Congenital Stationary Night	TRPM1	Γb	Insertion 1378 bp	~-	American Miniature Horse, Appaloosa, Australian Snotted Pony Britich Snotted	2013	24167615
Blindness, CSNB)					Pony, Knabstrupper, Noriker, Pony of the Americas.		
Brindle (<i>Incontinentia pigmenti</i>)	IKBKG		Nonsense (stop-gain)	×	Quarter Horse	2013	24324710
Dominant white	KIT	W21	1 bp deletion, frameshift	£	Icelandic	2015	26059442
Non-dun with primitive markings	TBX3	nd1	Regulatory	80	Many	2016	26691985
Non-dun	TBX3	nd2	1609-bp and 8-bp deletion, regulatory	œ	Many	2016	26691985
LP nattern modifier	RFWD3	PATN1	Regulatory	cr.	American Miniature Horse. Annaloosa	2016	26568529
-					Australian Spotted Pony, British Spotted Pony, Knabstrupper, Noriker, Pony of the Amaricae		
Brindle 1	AARTPC	BLI	Sulicing		American Ouarter Horse	2016	77449517
		ANTE244Glu	Alicence	16	Amorican Standardhrod	2102	11000470
wille White lee markings	MITE		Regulatory	16	Menorca Purchred	2017	78084638
Dominant white	KIT	8 <i>CV</i> M	Splicing) - m	Arahian	2017	28378977
Dominant white	KIT	CC/M	Deletion 1898 hn	, rr	Thornightred	2017	28444912
Tiger eve	SLC24A5	TE2	Deletion 626 bp) -	Paso Fino	2017	28655738
Tiger eye	SLC24A5	TE1	Missense	,	Paso Fino	2017	28655738
Dominant white	KIT	W24	Splicing	£	Italian Trotter	2017	28856698
Dominant white	KIT	W27	Missense	c	Thoroughbred	2018	29333746
Dominant white	KIT	W26	1 bp deletion, frameshift	c	Thoroughbred	2018	29333746
Dominant white	KIT	W25	Missense	c	Thoroughbred	2018	29333746
Curly coat	KRT25	Crd	Missense	11	Bashkir Curly Horse	2018	29686323 29141579
Splashed white, blue eyes and	MITF	SW5	63 kb deletion	16	American Paint Horse	2019	30644113
deafness							
Pearl dilution	SLC45A2	C ^{pri}	Missense	21	American Paint Horse, Lusitano, Purebred Snanish horse Ouarter Horse	2019	31006892, 30968968
Sunshine dilution	SLC45A2	Csun	Missense	21	Standardbred × Tennessee Walking Horse cross	2019	31006892

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Variants influencing pigmentation with known pleiotropic effects are in bold; details for genomic, coding and protein coordinates are in Table S1.

(reviewed by Coleman *et al.* 2013a), have been completely replaced by RNA-seq. Nevertheless, a few genomics resources from the past, such as the genomic BAC library CHORI-241 (https://bacpacresources.org/), remain in use in the post-genome era. End sequences of CHORI-241 BAC clones (Leeb *et al.* 2006) helped to validate the EquCab3.0 assembly (Kalbfleisch *et al.* 2018); BAC tiling paths are used to re-sequence complex genomic regions such as the terminal end of the pseudoautosomal region in ECAX (Rafati *et al.* 2016) and the male-specific region of ECAY (Janečka *et al.* 2018), and cytogenetic mapping of BAC clones is still a reliable method to validate copy number changes (Ghosh *et al.* 2014; Staiger *et al.* 2016a) and other large-scale structural rearrangements.

Ancient genomics and natural history of the horse

The assembly of the horse reference genome EquCab2.0 (Wade *et al.* 2009), along with success in extracting and sequencing ancient genomic DNA (Orlando *et al.* 2011), has essentially expanded our knowledge about the natural history of equids (Orlando *et al.* 2013; Jonsson *et al.* 2014), horse domestication and the dynamics of the horse genome over time, from the pre-domestication era to the present (Schubert *et al.* 2014; Librado *et al.* 2015, 2017; Gaunitz *et al.* 2018; Janečka *et al.* 2018; Wutke *et al.* 2018; Fages *et al.* 2019). The findings of ancient DNA studies were summarized and discussed in an excellent review by Librado *et al.* (2016). Therefore, here we highlight only the most notable discoveries and discuss the studies published since that review.

It is noteworthy that, with regards to ancient DNA, horses/ equids have made history three times-by being the first and the oldest, and most recently, by spanning the largest timescale of ancient genome data among non-human organisms (Fages et al. 2019). The first successful ancient DNA extraction and analysis ever was reported in 1984 from 150-yearold tissue from the extinct quagga (Higuchi et al. 1984), preceding by a year the first human ancient DNA study from a 2400-year-old Egyptian Mummy (Pääbo 1985). Likewise, to date the oldest genomic DNA sample which has been successfully sequenced is that of a 700 000-year-old Pleistocene horse (Orlando et al. 2011, 2013), exceeding the age of the oldest hominin genomic DNA extracted from 430 000year-old bones from Sima de los Huesos (Meyer et al. 2016). The current peak of ancient genomics of non-human organisms, but perhaps not the limit, is a study tracking 5000 years of horse domestication based on genome-scale data from 278 ancient animals (Fages et al. 2019).

Horse ancestry and domestication

Genomics-based searches into the wild ancestry of the domestic horse and the origins of horse domestication have

been ongoing for decades. Yet the answers have only recently started to emerge, largely thanks to the contribution of WGS from ancient equine samples (Schubert et al. 2014; Librado et al. 2015, 2016; Gaunitz et al. 2018; Fages et al. 2019). These studies reveal that, in addition to the two extant horses, the domestic (Equus caballus) and the Przewalski's horse (E. przewalskii) (Der Sarkissian et al. 2015), there existed other, now extinct, horse lineages at the time of early domestication. One lineage, that was initially identified from approximately 43 000- to 5000year-old bones from the Holarctic region (Schubert et al. 2014), extended to Southern Siberia (Fages et al. 2019). This lineage shares morphological similarities with an extinct horse described as Equus lenensis (Boeskorov et al. 2018; Fages et al. 2019). Recent mtDNA and Y haplotype analyses of an approximately 24 000-year-old specimen from the Tuva Republic suggest that there may have been another genetically divergent lineage of horses in Siberia and the New Siberian Islands, although the genetic contact between E. lenensis and this 'ghost' lineage remains unknown (Fages et al. 2019). Finally, Iberian samples from the third and early second millennia BCE, cluster separately from E. caballus, E. przewalskii and E. lenensis, have extremely divergent Y and mtDNA, and therefore suggest that there was a different, now extinct, horse lineage in Iberia during the early phase of horse domestication (Fages et al. 2019). Earlier it was reported that that the Holarctic horse (E. lenensis) contributed 12.9% to the genetic makeup of domestic horses (Schubert et al. 2014). However, the most recent study of 278 ancient equids and modern horses shows that none of the above extinct horse lineages contributed significantly to modern horse diversity (Fages et al. 2019).

Another question of interest is the genetic relationship between the two surviving horses-the domestic horse and the Przewalski's horse. Until recently, the overall consensus, based on modern and ancient WGS, was that the two are separate species, diverged approximately 45 000 years ago (Goto et al. 2011; Schubert et al. 2014; Der Sarkissian et al. 2015), with extensive bi-directional gene flow (Der Sarkissian et al. 2015; Librado et al. 2016). All studies agree that the domestic horse is not a direct descendant of the Przewalski's horse (Librado et al. 2016). These views were recently shaken by a study of over 40 ancient horse genomes from Eurasia, providing striking evidence that the Przewalski's horse is not truly wild, but rather a feral horse descended from the horses domesticated by Botai culture some 5500 years ago (de Barros Damgaard et al. 2018; Gaunitz et al. 2018). At the same time, all studied domestic horses dated from 4000 years ago to present show only 2.7% Botai ancestry, suggesting they descended from a different lineage of wild horses that subsequently went extinct (Fages et al. 2019).

Ancient DNA studies have also attempted to decipher, but have not completely resolved, the timeline and geography of

horse domestication. Current archeological and DNA evidence suggests multiple sites of horse domestication of which the earliest (~5000-5500 years ago) were in the Western Eurasian steppes: Botai culture in Northern Kazakhstan and the Pontic-Caspian steppe (Outram et al. 2009; Warmuth et al. 2012; Librado et al. 2016; Gaunitz et al. 2018; Fages et al. 2019), followed by additional candidate sites in Iberia, Eastern Anatolia, Western Iran, Levant and Eastern Europe (Hungary; Gaunitz et al. 2018). It is noteworthy that native breeds from the main domestication sites such as the Pontic-Caspian steppes still represent hotspots of genetic diversity for horses (Warmuth et al. 2011; Librado et al. 2016). However, even though the main candidate sites for domestication have been identified, the geographic origins of the modern domestic horse remain unknown. Given that the ancient Botai and Iberian lineages did not contribute substantially to modern domesticates and the temporal origins of the modern horse are modeled within the third and fourth millennium BCE, future studies of this timeline in other candidate regions of early domestication are needed (Fages et al. 2019).

Genetic cost of domestication

Domestication reduces overall fitness, known as 'the genetic cost of domestication' (Lu *et al.* 2006; Moyers *et al.* 2018). However, because of the extinction of wild horses, it has been difficult to evaluate the extent of this cost for the horse. Once again, the major breakthrough came with ancient DNA sequencing, providing a comparison with horses predating domestication (Schubert *et al.* 2014) and from early stages of domestication (Librado *et al.* 2017; Fages *et al.* 2019). Genetic changes associated with horse domestication can be summarized as follows:

- 1 Extreme mitochondrial DNA (mtDNA) diversity. Modern horses have high mtDNA diversity and lack phylogeographic mitochondrial structure, resulting in limited correspondence between mtDNA haplotypes, breeds and geography (Vila et al. 2001; Achilli et al. 2012; Librado et al. 2016). Sequencing horse genomes from early domestication sites show that similar mtDNA diversity was present in Scythian horses some 2300 years ago (Librado et al. 2017) and perhaps even earlier. Mitochondrial Bayesian Skylines reconstructed from 211 mitochondrial genomes suggest horse demographic expansion about 4500 years ago (Gaunitz et al. 2018; Fages et al. 2019). Reasons for the lack of mitochondrial structure are thought to be many, including sex-biased restocking from the wild and human management (Librado et al. 2016).
- 2 *Extreme lack of Y chromosome diversity*. Contrasting the diversity of mtDNA, the Y chromosome of domestic horses has become almost homogeneous with just a few haplotypes segregating in modern populations (Wallner

et al. 2017; Wutke *et al.* 2018). At the same time, Y sequences of ancient Scythian horses indicate that a large diversity of domestic male founders contributed to early domestication (Librado *et al.* 2017). Thus, the observed decline in Y chromosome variation happened only in the past few thousand years, probably because of the human-mediated reduction in the stallion population size and selective breeding (Librado *et al.* 2017; Wutke *et al.* 2018; Fages *et al.* 2019).

- 3 *Genetic load.* As a side-effect of human-driven selective breeding, the genome of the domestic horse has an increased level of deleterious mutations compared with pre-domestication genomes (Schubert *et al.* 2014). However, as mutation load is also lower in early domesticates from ancient Sintashta and Scythia, the excess of deleterious mutations in present day horses is probably a consequence of the past 2300 years of selective breeding (Librado *et al.* 2017), whereas the most striking changes have occurred during the past 200 years (Fages *et al.* 2019).
- 4 Domestication-associated genetic changes. One of the first comparisons between ancient pre-domestic horse genomes and those of modern breeds revealed 125 candidate genes that underwent episodes of positive selection during domestication (Schubert et al. 2014). These include genes involved in the cardiac and circulatory system, bone, limb and face morphogenesis, brain development and behavior, and coat color (Schubert et al. 2014; Librado et al. 2016). Genetic changes during horse domestication agree with the neural crest hypothesis and involve developmental changes affecting tissues and cell types of neural crest origin (Librado et al. 2017). However, the genomic cost of domestication and modern breeding is best illustrated by a striking discovery of a recent comparative study of ancient and modern horse genomes, showing that early breeders managed to maintain genetic diversity for millennia after domestication, and that the genetic diversity of the modern horse has dropped by 16% during just the past 200 years (Fages et al. 2019). It is not a coincidence that the last 200 years also cover the time of the development of horse breeds, the establishment of studbooks and the implementation of extensive selective breeding.

Genetic makeup of modern horse breeds

Since domestication, the genetic diversity present in ancient horse populations has been exploited for selective breeding for a wide range of phenotypes. However, the creation of the about 500 specialized horse populations or breeds by intense artificial selection and the establishment of (closed) studbooks happened only during the past 100–200 years (Hendricks 2007). Owing to the wealth of available genome-wide tools and resources (see above), these breeds

can now be studied in detail for genetic makeup, signatures of selection and relatedness to other breeds. The number of publications in the field is growing almost exponentially and these include a few seminal studies encompassing a global collection of breeds (McCue et al. 2012; Petersen et al. 2013a,b; Jagannathan et al. 2019), and many studies specializing in a single breed or a group of related breeds, for example, the American Quarter Horse (Petersen et al. 2014a; Avila et al. 2018; Marchiori et al. 2019; Pereira et al. 2019), the Hanoverian and other German Warmblood breeds (Nanaei et al. 2019; Nolte et al. 2019), the Arabian and related Middle Eastern breeds (Almarzook et al. 2017; Sadeghi et al. 2019), and some native/primitive breeds such as Hucul and Konik (Gurgul et al. 2019), Chinese native horses (Zhang et al. 2018), Japanese native breeds (Tozaki et al. 2019), the Yakutian Horse (Librado et al. 2015) and Korean horses (Seong et al. 2019). This is by no means an exhaustive list and it is even longer when including breeds that have recently been studied based on a genome-wide collection of microsatellite markers, such as Estonian horse breeds (Sild et al. 2019) or Konik (Szwaczkowski et al. 2016).

Despite the large number of studies, the core findings are rather similar. Collectively, modern horse breeds are characterized by high inter-breed and low intra-breed genetic diversity (McCue et al. 2012; Petersen et al. 2013a). Genomes of modern horses show multiple regions with signatures of selective pressures on performance traits and phenotypes. Among these, the most prominent are the MSTN gene in ECA18 for muscle fibers in racing breeds (Petersen et al. 2013b; Avila et al. 2018), the DMRT3 gene in ECA23 to perform alternative gaits in many breeds (Petersen et al. 2013b), and a region in ECA11 for body size in draught breeds and Miniature Horses (Petersen et al. 2013b). Clear signatures of selection are also found at known coat color loci. For example, the recessive chestnut coat color locus at MC1R is defined by a conserved approximately 750 kb haplotype across all breeds studied (McCue et al. 2012) and robust signatures of selection at the TBX3 locus in ECA8 are found in Konik horses, known to be selected for the dun color (Gurgul et al. 2019). Finally, selective breeding for the desired traits in modern breeds has unwillingly introduced accumulation of deleterious mutations (Librado et al. 2016, 2017; Jagannathan et al. 2019). For example, recent WGSs of 88 horses across 25 breeds identified heterozygotes for two potentially deleterious recessive alleles: a nonsense variant in the PALB2 gene, which is essential for mesoderm development, and a nonsense variant in the PLEKHM1 gene, necessary for osteoclast functions (Jagannathan et al. 2019).

Investigating Mendelian traits

The investigation of Mendelian traits in the horse began in the early 1900s. Because of the relative ease of tracking

simply inherited phenotypes across generations, some of the first Mendelian traits to be studied involved variations in pigmentation. In fact, Alfred Sturtevant, an undergraduate student working on mapping traits in Drosophila under Thomas Hunt Morgan, was among the first to publish on inheritance of coat color in harness horses (Sturtevant 1920). However, it was almost 100 years later when the genetic mechanism proposed by Sturtevant for the chestnut coat color was identified at the molecular level (Table 2). This and other variants identified in the early 2000s influencing pigmentation were discovered using candidate gene approaches. To date, 58 variants affecting pigmentation have been described, including 27 in the KIT gene that contribute to the dominant white phenotype. The identification of the majority of these pigmentation variants was made possible by the high quality of the horse reference genome sequence and available SNP array tools (Tables 2 and S1 and described above). For example, the initial horse linkage map developed by the horse genomics community (Penedo et al. 2005) allowed for the mapping of several pigmentation traits with pleiotropic effects, such as leopard complex spotting (Terry et al. 2004) and gray depigmentation (Pielberg et al. 2005). However, the discovery of causal variants was possible only through the availability of a reference genome that enabled both whole genome and RNA-seq data analyses to uncover large structural variants (Rosengren Pielberg et al. 2008; Bellone et al. 2013).

The first genetic disorder to be identified at the molecular level in the horse was hyperkalemic periodic paralysis, reported in 1992. Much like the early work investigating pigmentation in the horse, this was discovered by a candidate gene approach. At the time the first draft of the horse genome sequence was complete, nearly 15 years later, only nine disease causing mutations had been discovered (Tables 3 and S2). In the last several years, there has been acceleration in the discovery of causal or highly associated variants for Mendelian traits, with 14 variants reported in the last four years. This acceleration in discovery is due to advances in genomic tools and resources. DNA testing for these Mendelian traits has enabled markerassisted selection to be utilized by horse breeders and in some cases is helping to assist in the clinical management of disease.

Genetics of complex traits

Traits considered complex are those which are not determined by one or a few genomic variants but rather by small contributions of many and perhaps hundreds of variants across the genome. These traits are generally lowly heritable, as the phenotypic outcome is determined not only by the underlying genetic variation but also by a significant impact of the environment (e.g. nutrition, training). The role of the environment complicates both the identification and understanding of the genetic components driving these

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Disease	Gene	Type of variant	Mode of Inheritance	Chromosome	Breed	Year published	PubMed ID
Hyperkalemic periodic paralysis	SCN4A	Missense	Dominant	11	American Quarter Horse and related	1992	1338908
Ovotesticular disorder of sexual development (DSD)	SRY	Large deletion of the DNA- binding domain of the SRY	Y-linked	~	breeds Standardbred	1995	7558880
Severe combined immunodeficiency disease	PRKDC	gene Deletion 5 bp	Recessive	6	Arabian	1997	9103416
טטטט) (אפון) איז	LAMC2	Insertion 1 bp	Recessive	5	Belgian and Italian draft horse	2002	12230513
Malignant hyperthermia (MH)	RYR1	Missense	Dominant	10	American Quarter Horse	2004	15318347
Glycogen branching enzyme deficiency (GBED)	GBE1	Nonsense (stop-gain)	Recessive	26	American Quarter Horse and related breeds	2004	15366377
Thrombasthenia	ITGA2B	Missense	Recessive	11	American Quarter Horse & Thoroughbred	2006	16407493
Thrombasthenia	ITGA2B	Deletion 10 bp	Recessive	11	American Quarter Horse	2007	17338169
Hereditary equine regional dermal asthenia	PPIB	Missense	Recessive	~	American Quarter Horse	2007	17498917
Polysaccharide storage myopathy (PSSM1)	GYS1	Missense	Incompletely dominant	10	American Quarter Horse, American Paint Horse, Appaloosa, Draft, Pony of the America. and Warmblood	2008	18358695
Junctional epidermolysis bullosa (JEB2)	LAMA3	Deletion 6589 bp	Recessive	5	American Saddlebred	2009	19016681
Racing distance	MSTN	Insertion 227 bp, regulatory		18	Thoroughbred	2010	20098749 25160752 20278862
Cerebellar abiotrophy (CA)	МИТҮН	Regulatory	Recessive	7	Arabian, Bashkir Curly Horse, Danish Sport Horse, Trakehner, and Welsh Ponv	2011	29103988
Foal immunodeficiency syndrome in the Fell and Dales nonv (FIS)	SLC5A3	Missense	Recessive	26	Dales Pony and Fell Pony	2011	21750681
Androgen insensitivity syndrome (AIS)	AR	Regulatory	X linked	×	American Quarter Horse	2012	22095250
Myotonia	CLCN1	Missense	Recessive	4	New Forest Pony	2012	22197188
Permissive to gait	DMRT3	Nonsense (stop-gain)	Dominant	23	Numerous breeds	2012	22932389
Warmblood fragile foal syndrome (WFFS) or Ehlers-Danlos syndrome. type VI	PLOD1	Missense	Recessive	2	Warmblood	2015	25637337
Hoof wall separation syndrome	SERPINB11	Insertion 1 bp	Recessive	00	Connemara	2015	25875171
Hydrocephalus	B3GALNT2	Nonsense (stop-gain)	Recessive	-	Friesian	2015	26452345
Androgen insensitivity syndrome (AIS)	AR	Missense	X linked	×	Thoroughbred	2016	27073903
Skeletal atavism	SHOX	2 over lapping deletions 160 = 180 kb and 60–80 kb	Recessive	X and Y PAR	Shetland	2016	27207956
Dwarfism, Friesian	B4GALT7	Missense	Recessive	14	Friesian	2016	27793082
Dwarfism, ACAN-related D3*	ACAN	Missense	Recessive	1	Miniature Shetland	2017	27942904
Occipitoatlantoaxial malformation (OAAM)	НОХДЗ	Deletion 2.7 kb	Recessive	18	Arabian	2017	28111759
Androgen insensitivity syndrome (AIS)	AR	Deletion 25 bp	X linked	×	Warmblood	2017	28192783

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Disease	Gene	Type of variant	Mode of Inheritance	Chromosome	Breed	Year published	PubMed ID
Naked foal syndrome Ocular squamous cell carcinoma (ocular SCC)	5T14 DDB2	Nonsense (stop-gain) Missense	Recessive Recessive	7 12	Akhal-Teke Belgian, Haflinger, Percheron, Rocky Mountain Horse	2017 2017	28235824 28425625
Immune-mediated myositis (IMM/MYH1)	NYH1	Missense	Recessive	11	American Quarter Horse	2018	29510741
Curly coat with hypotrichosis Crd	KRT25	Missense	Dominant	11	Bashkir Curly Horse	2018	29686323 29141579
Curly coat without hypotrichosis	SP6	Missense	Dominant	11	American Bashkir Curly Horse and Missouri Foxtrotter	2018	29686323
Dwarfism, ACAN-related D4	ACAN	Deletion 42 bp	Recessive	~	Miniature	2018	30058072
Dwarfism, ACAN-related D2	ACAN	Missense	Recessive	~	Miniature	2018	30058072
Dwarfism, ACAN-related D1	ACAN	Deletion 1 bp	Recessive	-	Miniature	2018	30058072
Genomic coding and protein sequence coordinat	tes are in Table	5 C)					

phenotypes. That said, complex traits are of significance to the industry, both economically and for animal wellbeing and include measures of athletic performance, growth and body size, and disorders such as metabolic syndrome, laminitis, equine asthma or recurrent airway obstruction (RAO) and osteochondrosis (OC), among others. Given the availability of genome-wide genotyping technologies, researchers have been working to piece together the role of genetics in these complex traits. What is known regarding the genetic basis of several of the most highly studied complex traits in horses is outlined below.

Size

Unlike many complex traits, the size of the horse is highly heritable with estimates for wither height ranging from 0.52 to 0.78 (Molina et al. 1999; Zechner et al. 2001; Suontama et al. 2009; Signer-Hasler et al. 2012). Not surprisingly, positive genetic and phenotypic correlations are reported between wither height and other growth phenotypes such as body length, heart girth and cannon bone circumference (Molina et al. 1999; Sadek et al. 2006). Providing insight into the biology underlying size, Makvandi-Nejad et al. (2012) employed the 50K SNP array to identify four loci that explained a majority of variation in size across-breeds. The loci identified in this work include LCORL/NCAPG, HMGA2, ZFAT and LASP1, most of which have also been shown to play a role in size and growth phenotypes in other species (Snelling et al. 2010; Weikard et al. 2010; Lindholm-Perry et al. 2013; Saatchi et al. 2014). Additional work has supported the association of LCORL/NCAPG (Signer-Hasler et al. 2012; Tetens et al. 2013; Staiger et al. 2016a; Tozaki et al. 2017), ZFAT (Signer-Hasler et al. 2012; Tozaki et al. 2017), and HMGA2 (Frischknecht et al. 2015) with height in various other populations. Whereas the means by which these loci act to alter growth traits is not fully understood, a missense mutation in HMGA2 was reported to alter DNA binding, which was attributed to reduced size in ponies (Frischknecht et al. 2015) and was also recently correlated with metabolic syndrome in Welsh ponies (Norton et al. 2019a). Functionally, variation of LCORL was shown to alter its expression, which may explain its role in determining an individual's size (Metzger et al. 2013). Interestingly, LCORL has also been associated with other complex disorders, including recurrent laryngeal neuropathy (Boyko et al. 2014) and OC (Corbin et al. 2012), serving as another example of the pleiotropic effects of loci involved in complex traits. As height has been associated with both OC and recurrent laryngeal neuropathy (Brakenhoff et al. 2006; McGivney et al. 2019), these associations are biologically relevant. The power of high-density genome-wide genotyping arrays has also enabled the identification of loci with more minor effects than those found by Makvandi-Nejad et al. (2012) or which may be breed specific (Meira et al.

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Table 3 (Continued)

2014b; Frischknecht *et al.* 2016; Metzger *et al.* 2018). It is important that height is treated distinctly from dwarfism, which is simply inherited in several populations including the Miniature Horse (Metzger *et al.* 2017; Eberth *et al.* 2018) and Friesian (Leegwater *et al.* 2016; Table 3).

Athletic performance

The athletic ability of a horse, whether it be jumping, racing, cutting or pulling, is clearly complex, depending upon efficient metabolic and musculoskeletal properties as well as intricate interactions and the influence of training and husbandry. Particularly in Thoroughbreds, the genetics of racing performance has been of long-standing interest. Prior to the availability of genotyping technologies, pedigree and performance data were used to examine the genetic components of phenotypic variance for Thoroughbred, Standardbred and sport horse performance (Hintz & Vanvleck 1978; Ojala et al. 1987; Tavernier 1990; Árnason 2001; Ricard & Chanu 2001; Langlois & Blouin 2004, 2007). Heritability estimates for racing in the Thoroughbred range greatly from nearly zero to upwards of 0.75, depending upon the specific phenotype measured (e.g. race time, race length or race winnings) and the model assumed (O'Ferrall & Cunningham 1974; Gaffney & Cunningham 1988; Williamson & Beilharz 1996; Thiruvenkadan et al. 2009a). As genotyping methodologies became available, regions implicated in racing performance have been identified using candidate gene (Gu et al. 2010; Hill et al. 2010a) and genome-wide association analyses (Binns et al. 2010; Hill et al. 2010b; Tozaki et al. 2010; Shin et al. 2015) and transcriptome analyses (Park et al. 2014), and through the identification of selective sweeps in racing populations (Moon et al. 2015). Genes implicated include COX4I2 (Gu et al. 2010) and PDK4 (Hill et al. 2010a), both involved in cellular respiration. Although these studies have uncovered evidence of genetic factors involved in the racing performance of Thoroughbreds, a single locus, MSTN, has been repeatedly associated with racing performance (Binns et al. 2010; Tozaki et al. 2010). In 2010, an intronic variant of MSTN was noted to be predictive of a horse's best racing distance (Hill et al. 2010b); the use of a genetic test for this variant has been adopted as a means to tailor training programs or choose matings. Since the first publication of this variant, several lines of evidence have supported the role of a SINE insertion in the promoter of MSTN, in high linkage disequilibrium with the intronic SNP in the Thoroughbred breed, as the functional variant (Petersen et al. 2014b; Santagostino et al. 2015; Rooney et al. 2018).

Outside of the Thoroughbred, the *MSTN* variant predictive of better suitability as a sprinter is highly frequent in the Quarter Horse and associated with a higher proportion of fast-twitch muscle fibers (Petersen *et al.* 2013b, 2014b). Additional studies in the racing Quarter Horse have identified other loci associated with racing performance; as a result of these works, genes involved in muscle contractility, skeletal development and neurologic function have been suggested to be associated with sprinting (Meira et al. 2014a,c; Beltran et al. 2015). In some cases, variants for racing speed are common across breeds (Shin et al. 2015; Pereira et al. 2016), fitting a hypothesis that these horses shared selective pressures for superior metabolic and musculoskeletal traits. Efforts to understand the genetic components of variation in harness racing horses suggest that their heritability is low to moderate (reviewed in Thiruvenkadan et al. 2009b). In the harness racing populations, similar to the role of MSTN in the Thoroughbreds, a single variant was reported to impact performance to the extent that the variant is nearly fixed in trotting breeds (Promerova et al. 2014). The gene implicated, DMRT3, alters motor coordination and stride length (Andersson et al. 2012); horses homozygous for the variant perform at a higher level than those heterozygous or wt (Jaderkvist et al. 2014; Jaderkvist Fegraeus et al. 2015). Interestingly, this variant was identified not in a study of racing performance but in a study of another complex trait in the horse-the ability to perform an alternative gait (Andersson et al. 2012), again demonstrating the interplay among physiological pathways. As an aside, whereas the DMRT3 variant is deemed necessary for 'gaitedness', the basis of variations in gait is yet to be understood (Patterson et al. 2015; Staiger et al. 2016a; Fegraeus et al. 2017; Fonseca et al. 2017).

Requiring a different type of athleticism, endurance racing studied in Arabian horses identified five QTL, including genes hypothesized to be involved in neuronal and cardiac function (Ricard *et al.* 2017). As in complex traits, how the regulation of these genes or variants within them may enhance performance is an area of study. The differentially expression of microRNAs prior to and after endurance exercise is probably one means by which gene regulation is altered to allow horses to endure and excel at this type of performance (Mach *et al.* 2016).

Sport horses require yet another type of athleticism and the heritability and genetics of show jumping and dressage have been studied in several European populations. Heritability estimates, calculated for the longevity (years) of performance, have been similar across studies ranging from 0.07 to 0.18 (Ricard & FournetHanocq 1997; Braam *et al.* 2011; Seiero *et al.* 2016; Ricard *et al.* 2017). Heritability estimates for show jumping range from 0.11 (Sole *et al.* 2017) to 0.61 (Stock & Distl 2007), and in most cases are greater than estimates for dressage in the same population (Viklund *et al.* 2010; Braam *et al.* 2011), although this is not the case in the Swedish Warmblood studied by Wallin *et al.* (2003).

Finally, the use of the Illumina Equine50 SNP array to investigate the biology underlying the success of jumpers revealed a QTL explaining 0.7% of the phenotypic variance in French Warmbloods in a region including the candidate gene *RYR2* (Brard & Ricard 2015). In the Hanoverian, a

GWAS using the same genotyping platform identified six QTL including genes predicted to function in muscle structure and metabolism (Schroder *et al.* 2012).

Osteochondrosis

Osteochondrosis is a dysregulation of endochondral ossification of cartilage at the articular/epiphyseal complex most commonly occurring at the fetlock, hock and stifle joints (Jeffcott 1996). As a result, the cartilage becomes thickened and/or is retained, interfering with the normal function of the joint (Jeffcott 1996). The prevalence of OC varies by breed with relatively low incidence $(\sim 7\%)$ in Thoroughbreds (Kane et al. 2003) and moderate frequency (~30%) in Danish Warmbloods (van Grevenhof et al. 2009), and with estimates of as many as 50% of Standardbred and German coldblooded horses (Wittwer et al. 2006; Lykkjen et al. 2012) being affected. Although it can be quite common, the heritability of OC is low to moderate (reviewed in Distl 2013; Naccache et al. 2018; McCoy et al. 2019), with a variety of environmental factors identified as having a significant role in its occurrence (Lepeule *et al.* 2009, 2013; Vander Heyden et al. 2013). Incidence has been positively correlated with size (Stock et al. 2005) and LCORL, itself associated with the size of a horse (described above), has been noted as a risk factor for OC and associated with incidence in GWAS (Teyssedre et al. 2012; Orr et al. 2013; Naccache et al. 2018). In Thoroughbreds, a QTL also on ECA3, although over 20 Mb distant from LCORL, was estimated to explain over 30% of the genetic variation (Corbin et al. 2012). Several research groups have been working to identify genetic risk factors for OC. The complexity as well as hypothesized population-specific risk factors are evident in the many loci associated using genome-wide SNP assays, many of which do not overlap between populations (Dierks et al. 2007, 2010; Wittwer et al. 2007; Lampe et al. 2009a,b; Lykkjen et al. 2010; Corbin et al. 2012; Teyssedre et al. 2012; Orr et al. 2013; McCoy et al. 2016; Lewczuk et al. 2017; Table 4).

Functional studies show differential expression of the MMP-13 gene, encoding for a matrix metallopeptidase, in the cartilage of horses with OC compared with controls (Mirams et al. 2009; Riddick et al. 2012), consistent with hypothesized dysfunction in cartilage maturation and endochondral ossification. Candidate gene expression studies also suggest that chondrocyte maturation and catabolism are altered through dysregulated Wnt signaling (Kinsley et al. 2015). Finally, Mirams et al. (2016) used subtractive hybridization of the transcriptome from cartilage of affected and unaffected foals resulting in a hypothesized etiology that involves cartilage retention in subchondral bone. In addition to protein coding transcripts, differentially expressed miRNAs have been identified and may play a role in alteration of gene expression associated with OC (Desjardin et al. 2014).

Equine metabolic syndrome and laminitis

Horses affected by equine metabolic syndrome (EMS) present with insulin resistance and obesity and/or regional adiposity; additionally, hypertriglyceridemia, elevated leptin and hypertension may occur (Frank *et al.* 2010). Horses with EMS have an increased risk of laminitis or the disruption of the attachment between the distal phalanx (coffin bone) and inner hoof wall, leading to the coffin bone being rotated downward into the sole of the hoof. The incidence of EMS and associated laminitis is greatest in ponies and obese horses (Treiber *et al.* 2006; Bailey *et al.* 2008). Not all cases of laminitis, however, are attributed to EMS as it can occur in conjunction with other wellbeing issues such as colic, abdominal or reproductive infection, or excessive concussion on hard surfaces (Hood 1999).

SNP-based heritability estimates for traits associated with EMS (e.g. circulating insulin, glucose, ACTH, leptin) have been reported to be quite high (Norton *et al.* 2019b). Genetic factors contributing to risk are also supported by varied rates of prevalence among breeds and variation in insulin responsiveness by breed (Treiber *et al.* 2006; Bailey *et al.* 2008; Bamford *et al.* 2014). In a study of pedigreed ponies, Treiber *et al.* (2006) proposed that one or a few major dominant genes contribute to predisposition to laminitis.

Changes in gene expression related to the onset of laminitis have been studied as a possible means of both understanding the progression of the condition and identifying biomarkers to detect a possible bout prior to the onset of clinical symptoms. With a hypothesized inflammatory component, Tadros et al. (2013) found that circulating cytokine expression of IL-1B, IL-8 and IL-10 was elevated several hours prior to the detectable onset of laminitis. A similar inflammatory response has been identified in the laminar tissue itself in horses induced to develop laminitis compared with healthy controls (Belknap et al. 2007; Loftus et al. 2007; Leise et al. 2010). These studies show a systemic inflammatory response associated with laminitis. However, whereas these studies help identify pathways involved in disease progression, they fail to answer questions of genetic risk.

Toward the goal of understanding genetic risk factors for disease, Lewis *et al.* (2017) identified a single candidate gene, *FAM174A*, in Arabian horses associated with laminitis and correlated with an increased insulin-to-glucose ratio. However, a more complete understanding of the genetic risk factors for EMS remains elusive. Finally, the influence of the microbiome on the development of EMS and the associated laminitis is a recent area of study. Characterization of the hindgut microbiome revealed differences in microbial composition in horses with chronic (Steelman *et al.* 2012) and induced (Milinovich *et al.* 2006) laminitis, as well as in horses with EMS (Elzinga *et al.* 2016) compared with healthy controls. The composition of the microbiome itself has been suggested to be heritable (Blekhman *et al.* 2015;
 Table 4 Complex equine diseases and traits with ongoing genetic studies.

		Tupo of gonotic	
Disease/trait (reference)	Breed	study	Genomic region(s) identified
Atrial fibrillation (Physick-Sheard <i>et al</i> . 2014; Kraus <i>et al</i> . 2017)	Standardbred	Heritability only	Probably polygenic; no region identified to date
Body size (e.g. Makvandi-Nejad <i>et al</i> . 2012)	Multiple	GWAS	Loci on ECA3, 6, 9 and 11
Bone fracture (Blott & Vaudin 2013; Blott <i>et al</i> . 2014)	Thoroughbred	GWAS	ECA18
Brachygnathia (Signer-Hasler et al. 2014)	Franches-Montagnes	GWAS	ECA13
Chronic progressive lymphedema (De Keyser <i>et al.</i> 2014), ¹	Draft breeds	Candidate gene approach	Continuing to pursue sequencing ELN
Common variable immunodeficiency ²	Various	Epigenetic investigation	RNA-seq and Methyl-Seq of E2A and PAX5
Corneal stromal loss (Lassaline-Utter <i>et al</i> . 2014; Alberi <i>et al</i> . 2018)	Friesian	Heritability only/ candidate gene	Likely heritable; excluded BGN variants
Cribbing (crib-biting) (Hemmann <i>et al.</i> 2014) Cryptorchidism ¹	Multiple Icelandic	Candidate gene Heritability only	Excluded subset of stereotypic genes Likely to be heritable
Degenerative joint disease (Welsh et al. 2013)	Thoroughbred	Heritability only	Small to moderate heritability identified
Guttural pouch tympany (Metzger <i>et al.</i> 2012)	Arabians and German Warmbloods	GWAS	ECA15 (Arabians) and ECA3 (German Warmbloods)
Insect bite hypersensitivity (Schurink <i>et al</i> . 2012; Velie <i>et al</i> . 2016)	Icelandic, Shetland and Exmoor	GWAS	ECA7, 9, 10 and 17ECA8 (Exmoor)
Recurrent laryngeal neuropathy (Dupuis <i>et al.</i> 2011; Boyko <i>et al.</i> 2014)	Thoroughbred, Warmblood, Trotter and Draft	GWAS	ECA3 (height locus; Thoroughbred),ECA21 and 31 (Multiple breeds)
Metabolic syndrome (Lewis <i>et al</i> . 2017; Norton <i>et al</i> . 2019b,a) ¹	Welsh pony, Arabian, Morgan	GWAS	ECA6 (Welsh pony),ECA14 (Arabian),multiple regions (Morgan)
Navicular disease (Diesterbeck <i>et al</i> . 2007; Lopes <i>et al</i> . 2009, 2010)	Warmbloods	GWAS	ECA2 and ECA10
Neuroaxonal dystrophy/equine degenerative myeloencephalopathy (Finno <i>et al.</i> 2013, 2014)	Quarter Horse	GWAS	ECA8 region exclusion, exclusion of <i>TTPA</i> candidate gene
Osteochondrosis/osteochondrosis dissecans (Dierks <i>et al.</i> 2007; Lampe <i>et al.</i> 2009a,b,c; Sevane <i>et al.</i> 2016, 2017)	Warmbloods, Trotters, Standardbreds, Spanish Purebred	GWAS	ECA2, 4, 5, 16, 18 (Warmblood),ECA10, 14, 21 (Standardbred),candidate gene analysis (Spanish Purebred)
Polysaccharide storage myopathy, type II ¹	Quarter Horses	GWAS	ECA18
Recurrent airway obstruction (Swinburne <i>et al</i> . 2009; Schnider <i>et al</i> . 2017; Mason <i>et al</i> . 2018)	Warmblood	GWAS	ECA 11, 13, 15 (Warmblood)
Recurrent exertional rhabdomyolysis (Fritz <i>et al</i> . 2012)	Thoroughbred, Standardbred	GWAS	ECA11, 16, 30 (Thoroughbred),ECA10, 11 (Standardbred)
Recurrent uveitis (Kulbrock <i>et al</i> . 2013; Fritz <i>et al</i> . 2014)	Appaloosa, German Warmblood	Candidate gene/ GWAS	ECA1, 20 (Appaloosa),ECA18, 20 (German Warmblood)
Sarcoid (Christen <i>et al.</i> 2013; Staiger <i>et al.</i> 2016b)	Franches-Montagnes, Quarter Horse, Thoroughbred	GWAS	ECA20, 22 (QH, TB)
Stallion subfertility owing to impaired acrosome reaction (Raudsepp <i>et al.</i> 2012)	Thoroughbred	Susceptibility gene	ECA13: FKBP6
Stallion fertility (Schrimpf et al. 2015)	Hanoverian	GWAS	ECA13: FKBP6
Stallion fertility (Schrimpf <i>et al</i> . 2016)	19 European breeds	GWAS	High-impact variants in CFTR (ECA4), OVGP1 (ECA5), FBXO43 (ECA9), TSSK6 (ECA21), PKD1 (ECA13), FOXP1 (ECA16), TCP11 (ECA20), SPATA31E1 (n/a), NOTCH1 (ECA25)
Swayback (lordosis) (Cook et al. 2010)	Saddlebred	GWAS	ECA20

ECA, Equus caballus chromosome; GWA, genome-wide association.

¹Abstract presented at the Dorothy Russell Havemeyer Foundation International Equine Genome Mapping Workshop.

²Abstract presented at the Plant and Animal Genome Conference.

Goodrich *et al.* 2016), associated with genes involved in metabolism and immunity; these data connect host genetics to yet another means by which risk for EMS and/or laminitis may be amplified.

Equine asthma or RAO

Equine asthma, also known as RAO or heaves, is a chronic disease of the lower airway, particularly problematic in

environments where air flow is limited and in which bedding or hay has high levels of dust or other respiratory irritants (reviewed in Woods et al. 1993; Ramseyer et al. 2007; Pirie 2014). The prevalence of RAO is estimated to be 14% in a sample of British horses (Hotchkiss et al. 2007) and 10% in Swiss Warmbloods (Ramseyer et al. 2007). Often compared with asthma in humans, affected individuals have difficulty breathing, neutrophil and mucus accumulation in the airway, cholinergic bronchospasm and coughing, and are especially sensitive to inhaled allergens (Robinson et al. 1996; Gerber et al. 2004). Whereas symptoms can be mediated by removing the horse from the problematic environment (Vandenput et al. 1998a, b), observations of increased risk in foals with affected parents, or foals of particular families (Marti et al. 1991; Ramsever et al. 2007), have suggested a heritable component to its etiology.

Before genomic tools, a hypothesis derived from the clinical accumulation of mucus in RAO-affected horses led to the investigation of mucin gene variation as a possible risk factor. As a result, the equine ortholog of *MUC5AC* was identified as being upregulated in affected horses (Gerber *et al.* 2003). A candidate gene study also found an isoform of *MYH11* to be overexpressed in affected horses (Boivin *et al.* 2014). The origin of the *MYH11* isoform identified in Boivin *et al.* (2014) has been associated with alternative regulatory mechanisms (Issouf *et al.* 2018), providing an example of how variable genome regulation contributes to important phenotypes associated with animal health and wellbeing.

In perhaps the most highly studied populations of horses, and prior to the genome assembly, the candidate gene IL4R was investigated in two half-sibling families of Warmbloods using microsatellite genotyping (Jost et al. 2007). In one family, a significant association with RAO was found with a haplotype in ECA13 near the cytokine IL4R, and a recessive mode of inheritance was suggested. However, this association was not consistent in the other family, where an association with RAO was identified in ECA15 with a predicted dominant mode of inheritance. In a GWAS of these families using microsatellites, QTL were identified on ECA13, and whereas no association was found with the positional candidate gene ITGAX, IL4R was noted to be proximal to the hit (Swinburne et al. 2009). In Shakhsi-Niaei et al. (2012) the same research group used SNP50 to fine-map the region, which resulted in the identification of a signal on ECA13 in both the family from which the original QTL was identified and a group of unrelated horses; however, the result was not statistically significant and no clear casual mutation or genes were identified. Chromosome 13 was again the strongest region of association in a GWAS repeated on these horses with the high-density Affymetrix SNP array after it became available (Schnider et al. 2017); the authors note positional candidate gene, TXNDC11, from those analyses.

Sequencing of IL4R and expression from bronchoalveolar lavage fluid identified increased expression in horses from which the association was derived but not in other horse families (Klukowska-Rotzler et al. 2012), further supporting its hypothesized role in RAO in this population. Additional follow-up studies included quantitative PCR of candidate genes (Lanz et al. 2013) and RNA-seq (Pacholewska et al. 2015) of peripheral blood mononuclear cells (PBMCs) derived from RAO-affected and control horses that were stimulated with irritants such as hay dust or lipopolysaccharide. The role of IL4R in the population in which the QTL was identified was supported whereas other data continued to support different mechanisms of disease between the two families (Lanz et al. 2013). RNA-seq of horses from the sample families found CXCL13 to be upregulated along with cell cycle regulatory transcripts such as CDC20, and genes involved in immune function and development (Pacholewska et al. 2015). Finally, Mason et al. (2018) conducted expression QTL studies of experimentally treated PBMCs from horses with prior RAO and healthy controls: whereas the studies supported prior SNP associations in these families, the identification of functional risk markers remains elusive.

Lastly, considering the possible impact of variations other than nucleotide substitutions, genomic copy number variants were analyzed using a tiling array to compare variants of RAO with control horses (Ghosh *et al.* 2014). Over 700 CNVs were identified across samples with the RAO horses found to have, on average, more CNVs than the controls. Although no significant associations were identified, *NME7*, involved in ciliary function of the lung, had a suggestive (P = 0.06) association with the RAO phenotype, warranting further investigation (Ghosh *et al.* 2016).

Reproduction

Compared with other complex traits, the genomics of equine reproduction has been given relatively less attention, even though reproductive performance is of high economic importance for purebred horses and vital for survival in feral populations (Raudsepp et al. 2013; Metzger et al. 2015). Only a few candidate loci or genomic regions have been associated with various fertility parameters and phenotypes (Raudsepp et al. 2013). Among these, the FKBP6 gene is of particular interest because of contrasting associations: in Thoroughbreds it is associated with subfertility owing to impaired acrosome reaction (Raudsepp et al. 2012), but in Hanoverians it is associated with improved conception rates (Schrimpf et al. 2015). Like for other complex traits, genome-wide SNP- or WGS-based analyses are expected to also make a difference in fertility research. A good example is a recent whole-genome screening that revealed high-impact variants in nine putative male fertility genes (Schrimpf et al. 2016; Table 4).

On a different note, breeding animals are typically selected on the basis of their pedigrees, athletic performance and appearance, but not for their reproductive potential. This suggests that there are no strong signatures of selection for reproductive performance. Surprisingly, this is not true as shown by a whole-genome analysis of runs of homozygosity in six diverse breeds, including commercial breeds such as Hanoverian and Thoroughbred, and native breeds such as Sorraia (Metzger *et al.* 2015). The findings suggest a significant artificial as well as natural positive selection on reproduction performance in all types of horse populations.

Horse as a large animal model for humans

Unlike rats or mice, horses are large and expensive animals with long generation intervals, and therefore are not typical model species for studying human disease and physiology. However, some equine conditions, such as obesity, respiratory disease, orthopedic disease, equine recurrent uveitis and certain cancers translate into similar human conditions better than those from classical model species. For example, because of unique similarities between human and horse insulin resistance response to overfeeding, EMS has the potential to serve as a model for human obesity (Frank et al. 2010; Jacob et al. 2018). Likewise, naturally occurring equine asthma is recognized as a model for some forms of asthma in humans (Bullone & Lavoie 2017: Bond et al. 2018). As an athletic species, the horse is considered as an important large animal model for cardiovascular disease (Tsang et al. 2016) and musculoskeletal disorders, including osteoarthritis (McCoy 2015), as well as a model for articular cartilage repair and regeneration studies (Dias et al. 2018). The horse has also been proposed as a potential model for immune response for infections, such as acute synovitis and septic arthritis (Ludwig et al. 2016), and autoimmune disorders like recurrent uveitis (Witkowski et al. 2016). Furthermore, studies of melanoma in gray horses are expected to help dissect the molecular mechanisms underlying melanoma as well as vitiligo in humans (Rosengren Pielberg et al. 2008; van der Weyden et al. 2016).

Functional annotation of the equine genome

Despite the recent and rapid successes in identifying causative mutations for these mostly simple inherited traits, the genetic investigation of complex traits has not been as straightforward. Although there is strong evidence for the heritability of many complex diseases and traits (described above), despite a significant financial investment by the equine industry in phenotyping and genotyping large numbers of horses, functional genetic mutations have not been discovered for most complex traits examined (Table 4).

Finding genetic associations of chromosomal segments to a specific phenotype without identifying functional or

causative mutations within protein-coding genes is not a problem unique to the horse. The idea that much of the non-coding genome is 'junk DNA' (Ohno 1972) and uninteresting to consider further has been reconsidered after overwhelming evidence from the human and murine Encyclopedia of DNA Elements (ENCODE) projects, demonstrating that these regions are functionally important (Consortium et al. 2007; Yue et al. 2014). As many as 93% of the associated and potentially causative variants from human GWAS publications are outside of annotated protein-coding regions (Hindorff et al. 2009). It is unlikely that these associations are aberrant, but rather they are identifying genomic regions involved in processes other than protein coding, such as regulation of gene expression (Maurano et al. 2012). In addition to the recent discovery that the majority of the genome is transcribed (Consortium et al. 2007), it has also been shown that 92–94% of human protein-coding genes express multiple mRNA variants or isoforms (Wang et al. 2008). Many of the variants responsible for mapped equine diseases and traits in Table 3, in addition to others, may be regulatory, non-coding mutations.

The need for defining the tissue-specific gene expression and regulation (i.e. functional annotation) across domestic animal species was acknowledged by the establishment of the International FAANG project (Andersson et al. 2015). The ultimate goal of this initiative is to provide high-quality functional annotation of animal genomes in a coordinated effort that facilitates data sharing and analysis, while establishing standards for assay quality and continuity for metadata analysis. In an effort to establish a more direct connection between genome function and phenotype, the FAANG consortium has focused on initially assaying tissues from one to two individual animals representing a breed with minimal genetic diversity within a species (Andersson et al. 2015). The resulting data from this annotation across species will provide power to associate phenotypes with functional data, making it possible to generate and test hypotheses regarding the functional mechanisms underlying associations (Tuggle et al. 2016; Giuffra et al. 2019).

As demonstrated by the human and murine ENCODE projects, five types of assays uncover the majority of variation in tissue-specific gene expression and epigenetic modifications:

- 1 *Expression*—RNA-seq identifies expression levels of primarily protein-coding genes.
- 2 RNA expression—small RNA-seq identifies expression of miRNAs, non-coding RNAs that function primarily in RNA silencing.
- 3 *Histone modifications*—ChIP-seq identifies genome-wide patterns of histone modifications using antibodies against the modified histone proteins. These prioritized marks have been standardized across species and those focused upon for the FAANG initiative and the genomic features they identify include H3K4me1 (enhancers and

Table 5 Biobank tissues collected from two Thoroughbred mares.

Musculoskeletal system:	Nervous system:	Digestive system:
Cartilage — Fetlock	Cerebellum Vermis	Cecum
Cartilage — Stifle	Cerebellum Lateral Hemisphere	Duodenum
Coronary Band	Cornea	Epiglottis
Deep Digital Flexor	Corpus Callosum	Esophagus
Gluteal Muscle	C1 Spinal Cord	lleum
Lamina	C6 Spinal Cord	Jejunum
Long Bone Marrow	Dorsal Root Ganglia	Left Dorsal Colon
Longissimus Dorsi	Dura Mater	Left Ventral Colon
Metacarpal Bone Diaphysis	Frontal Cortex	Right Dorsal Colon
Rib Bone Marrow	Hypothalamus	Right Ventral Colon
Sacrocaudalis Muscle	L1 Spinal Cord	Small Colon
Sesamoid Bone	L6 Spinal Cord	Stomach
Superficial Digital Flexor	Occipital Cortex	Tongue
Suspensory Ligament	Parietal Cortex	Abdominal/thoracic organs:
Cardiovascular system:	Pituitary	Adrenal Cortex
Aortic Valve	Pons	Adrenal Medulla
Heart Left Atrium	Retina	Kidney Cortex
Heart Left Ventricle	Sciatic Nerve	Kidney Medulla
Heart Right Atrium	Temporal Cortex	Larynx
Heart Right Ventricle	Thalamus	Liver
Left Lung	T8 Spinal Cord	Lymph Node
Mitral Valve	Cell types:	Pancreas
Pulmonic Valve	Fibroblasts (culture)	Spleen
Trachea	Keratinocytes (culture)	Thyroid
Tricuspid Valve	PBMCs	Integumentary system:
Urogenital System:	Body fluids:	Dorsum Skin (over back)
Cervix	Plasma	Gluteal Adipose
Mammary Gland	Serum	Loin Adipose
Ovary	Cerebrospinal fluid	Neck Skin
Oviduct	Synovial fluid	
Urinary Bladder		
Uterus		
Vagina		

Updated from Burns *et al.* (2018). Prioritized tissues for study are in bold. Additional tissues from which RNA-seq data have been collected as funded by outside collaborators are shown in italics.

distal regulatory elements), H3K4me3 (active promoters and enhancers), H3K27me3 (gene silencing) and H3K27ac (active regulatory elements; Lee *et al.* 2014).

- 4 *Chromatin accessibility*—the Assay for Transposase-Accessible Chromatin using sequencing (ATAC-seq) identifies regions of open chromatin.
- 5 *DNA methylation*—reduced representation bisulfite sequencing identifies DNA methylation across the genome.

In line with the priorities of the FAANG initiative, in 2016, an equine biobank was created based on the sampling and preservation of 86 tissues, two cell lines and fluids from two Thoroughbred mares (Table 5; Burns *et al.* 2018). Tissues were flash frozen, preserved for histopathology, fixed for chromatin-immunoprecipitation, and in 16 tissues, nuclei isolation was conducted. This biobank is available for all researchers to access for assays appropriate to the goals of the FAANG initiative. Notably, extensive ante- and post-mortem evaluation by veterinarians was conducted on these two horses to provide the highest standard of a true 'reference' database for researchers to utilize. In addition, both mares represented in the biobank

had normal karyotypes. Utilizing the strict standard of phenotyping allows for both association of phenotype with genomic data and standardization of future sampling efforts, enhancing the utility of the data generated (Burns *et al.* 2018).

Eight tissues were prioritized in the initial equine annotation efforts (Table 5) owing to their cross-species application (e.g. skeletal muscle, liver, lung and ovary) as well as importance to the horse (laminae). Additionally, as an international collaborative initiative, 24 individuals representing 16 research institutions in 10 countries voluntarily participated to support RNA-seq of additional tissues by providing funding for transcriptome characterization of the tissue(s) most relevant to their own research; RNA-seq datasets from these tissues are publicly available (EMBL: http://www.ebi.ac.uk/embl/). Seven laboratories also conducted additional assays, such as karyotype analyses, centromere mapping of fibroblast cells, reduced representation bisulfite sequencing of the eight priority tissues, fibroblast functional assays, functional assays on tissues of the suspensory apparatus and further phenotyping through the sequencing of microbiome samples.

ChIP assays for four histone modification marks were recently completed on the eight prioritized tissues. At this time, ChIP assays for the major insulator-binding protein in vertebrates, CCCTC-binding factor and ATAC-seq experiments are underway. All datasets will then be fully integrated and correlated with gene expression data and made publicly available as an equine-specific tissue atlas to the entire equine research community.

Future directions

With the sheer volume of sequencing efforts currently underway in the horse, in addition to the FAANG efforts to define regulatory regions of the genome, the genetic contributions to complex traits can be discovered. Many of these more complex diseases (Table 4) probably have strong environmental contributions to the overall phenotype. Therefore, educating veterinarians and horse owners on the proper use of these genetic 'risk factors' in breeding management will be essential to advance the health of horses.

Concluding remarks and future perspectives

In this review, we demonstrate that the 10 years of postgenome era in equine genomics have been unparalleled and decorated with outstanding achievements in almost all conceivable directions. These include improved characterization of the structure and function of the horse genome, delineating the genetic makeup of breeds and populations, deciphering the evolutionary ancestry of horses and continuing search for molecular causes of Mendelian and complex traits and diseases. The central pillar of support for this success is definitely the high-quality horse reference genome combined with unprecedented advances in genomics technologies and global collaborations between researchers of diverse disciplines, clinicians, breeders and horse owners.

Whereas similar achievements characterize the postgenome era of all main domestic species, it must be emphasized that the collection of genome-level sequence data from hundreds of ancient horses from the past few thousand years is unparalleled and not available for any other domestic or non-primate species. This unique resource allows researchers to track the evolutionary past of any genomic features, particularly the sequence variants that are associated with equine traits of importance, such as performance, coat color, disease and adaptations. This also demonstrates that the history of domestic animals cannot be fully understood without ancient genomic data.

The enhanced knowledge about the horse genome, biology and populations also highlights the areas that require critical improvement. Among these, high expectations are put on the equine FAANG initiative in order to identify important functional elements in the horse genome, particularly those underlying simple and complex traits. Also, the growing number of available WGSs of individual horses of diverse breeds and phenotypes allows for a comprehensive catalog of common and rare variants in the horse genome. This in turn is the foundation for equine precision medicine, which should identify novel genetic mutations in a small number of individuals and connect the variation with function.

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Conflict of interests

The authors declare no conflict of interests.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article. **Table S1** Genetic variants identified for traits influencing pigmentation.

Table S2 Genetics variants underlying disease and performance traits in the horse.