



MHC genotyping from rhesus macaque exome sequences

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Abstract

Indian rhesus macaque major histocompatibility complex (MHC) variation can influence the outcomes of transplantation and infectious disease studies. Frequently, rhesus macaques are MHC genotyped to identify variants that could account for unexpected results. Since the MHC is only one region in the genome where variation could impact experimental outcomes, strategies for simultaneously profiling variation in the macaque MHC and the remainder of the protein coding genome would be useful. Here we determine MHC class I and class II genotypes using target-capture probes enriched for MHC sequences, a method we term macaque exome sequence (MES) genotyping. For a cohort of 27 Indian rhesus macaques, we describe two methods for obtaining MHC genotypes from MES data and demonstrate that the MHC class I and class II genotyping results obtained with these methods are 98.1% and 98.7% concordant, respectively, with expected MHC genotypes. In contrast, conventional MHC genotyping results obtained by deep sequencing of short multiplex PCR amplicons were only 92.6% concordant with expectations for this cohort.

Keywords *Macaca mulatta* · Exome · Major histocompatibility complex

Introduction

The major histocompatibility complex (MHC) is an intensively studied set of genes in macaques (Shiina et al. 2017; Wiseman et al. 2013). The genomic MHC region contains clusters of genes that encode the MHC class I complex and the MHC class II complex. Cells use MHC class I molecules to present intracellular peptides to immune cells like the CD8+ T cell or natural killer cells (Garcia and Adams 2005). MHC class I molecules accommodate intracellular peptides of varying specificity by having diverse amino acid sequences in the

$\alpha 1$ and $\alpha 2$ subunits, which form the peptide-binding cleft (Silver and Watkins 2017; Loffredo et al. 2009). These $\alpha 1$ and $\alpha 2$ subunits correspond to exons 2 and 3, respectively, of an MHC class I gene (Malissen et al. 1982). Most of the polymorphisms that distinguish MHC class I alleles are concentrated in exons 2 and 3 (Williams 2001). Thousands of individual MHC allelic variants have been identified in the three most widely used macaque species for biomedical research: rhesus (*Macaca mulatta*), cynomolgus (*Macaca fascicularis*), and pig-tailed macaques (*Macaca nemestrina*) (Semler et al. 2018; Karl et al. 2017; Maccari et al. 2017).

The human MHC is termed the human leukocyte antigen complex (HLA). HLA class I has a single copy of the *HLA-A*, *HLA-B*, and *HLA-C* genes on each chromosome (Daza-Vamenta et al. 2004; Shiina et al. 2017). In contrast, macaques have a variable number of genes on each chromosome that encode MHC class I MHC-A and MHC-B proteins, and macaques lack an *HLA-C* orthologue (Daza-Vamenta et al. 2004; Shiina et al. 2017; Wiseman et al. 2013). Initial approaches to genotype macaque MHC class I relied on using sequence-specific PCR oligonucleotides to test for the presence or absence of individual alleles (Kaizu et al. 2007). More recently, deep sequencing genomic DNA or complementary DNA PCR amplicons

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spanning a highly variable region of exon 2 has become commonplace (Wiseman et al. 2009; Karl et al. 2013). Amplicon sequences can be used to genotype groups of closely related MHC class I alleles, which are denoted as lineages. For example, an amplicon deep sequence that corresponds to the rhesus macaque *Mamu-A1*001* lineage demonstrates that an animal possesses *Mamu-A1*001:01*, *Mamu-A1*001:02*, or another closely related variant that has not yet been identified. This lineage-level reporting of MHC class I genotypes can be sufficient for designing experiments where specific MHC class I genotypes need to be matched between animals, balanced among experimental groups, or excluded entirely from a study (Loffredo et al. 2007, 2009; Muhl et al. 2002; Karl et al. 2013; Wiseman et al. 2009; Nomura et al. 2012; Mothe et al. 2003).

Amplicon deep sequencing can also be used for MHC class II genotyping. The human HLA class II genes *DQA1*, *DQB1*, *DPA1*, and *DPB1* have direct orthologues in macaques (Otting et al. 2017). Both macaques and humans have a variable number of MHC class II DRB genes on a single chromosome, while the DRA gene is oligomorphic (Daza-Vamenta et al. 2004; Shiina et al. 2017), and typically is not used for genotyping purposes. MHC class II molecules are members of the immunoglobulin superfamily, but they differ from MHC class I in several ways. α and β subunits comprise the MHC class II heterodimer, with separate genes encoding α and β subunits (Brown et al. 1993). Highly polymorphic regions that are diagnostic for MHC class II allele lineages can be PCR amplified in a manner that is similar to the MHC class I genotyping. Exon 2 contains the most extensive polymorphism among MHC class II alleles (Williams 2001). Each of the *DRB*, *DQA1*, *DQB1*, *DPA1*, and *DPB1* polymorphic MHC class II genes are sufficiently divergent that separate PCR amplicons are required for deep sequencing (Karl et al. 2014).

Comprehensive MHC class I and class II genotyping of macaques by amplicon deep sequencing requires preparation of six separate PCR amplicons for MHC class I and class II DRB, *DQA1*, *DQB1*, *DPA1*, and *DPB1* (Karl et al. 2014, 2017). Despite this complexity, a major advantage to amplicon deep sequencing has been its cost effectiveness. The output from a single MiSeq sequencing run can be used to determine the MHC class I and MHC class II genotypes for up to 192 macaque samples. In recent years, improved sequencing hardware and software have prompted new approaches to MHC genotyping. Instead of utilizing PCR amplicons of variable gene regions to genotype samples, researchers can use human whole exome sequencing (WES) and whole genome sequencing (WGS) datasets to determine HLA genotypes (Xie et al. 2017; Kishikawa et al. 2019; Yang et al. 2014; Posey

et al. 2016). Likewise, target capture approaches have been described for HLA genotyping with next-generation sequencing datasets (Cao et al. 2013; Wittig et al. 2015). In contrast, MHC genotyping of macaques using WGS (Xue et al. 2016; Bimber et al. 2017; de Manuel et al. 2018) or WES (Vallender 2011; Cornish et al. 2016) have not been reported to date. The macaque genome is being actively updated, but the reference database of known alleles is still incomplete (Zimin et al. 2014; Maccari et al. 2017). The unfinished state of the macaque reference database and the inherent intricacies that exist when attempting to map short sequence reads against complex, duplicated gene families both illustrate the significant challenges researchers can face with macaque genotyping.

In an era where the per-base cost of sequencing is dropping rapidly, we explored the feasibility of obtaining whole exome sequencing data while maintaining parity of results with the traditional MHC PCR amplicon approach. WES datasets provide investigators with the capability to examine sequence variants across the entire coding region of the genome. Although SIV researchers have traditionally focused on specific MHC genotypes that are associated with exceptional control of viral replication, there is also interest in evaluating variation in host restriction factor genes, such as *TRIM5*, *Tetherin*, and *APOBEC3G* receptor genes (Reynolds et al. 2011; Janaka et al. 2018; Krupp et al. 2013; Ericson et al. 2014). Exome sequencing will also provide an opportunity for investigators in multiple fields to evaluate variants from a wide variety of pathways that are important for biomedical research (Haus et al. 2014). A key advantage of WES over WGS remains to be the cost per sample of generating sequence assays. Since WES only targets the coding regions, which are 1–2% of the genome, the amount of sequencing that is required is dramatically reduced. At our institutions, currently WES can be completed for less than 20% of the cost of a comparable WGS study. The ability to evaluate at least five times more animals for the same cost opens the possibility of characterizing much larger non-human primate cohorts, such as with breeding colonies of macaques, or complete retrospective experimental studies (de Manuel et al. 2018).

Here we introduce MHC genotyping via macaque exome sequencing (MES), which is an exome sequencing-based workflow for comprehensive MHC class I and class II genotyping in macaques. This workflow uses a commercially available human exome target-capture enrichment kit in conjunction with specialized spike-in target-capture probes to cope with the high copy number of macaque MHC genes. We show that accuracy of Indian rhesus macaque MHC class I and class II results from this workflow is comparable to conventional MiSeq genotyping when using exon 2 reference sequences.

Methods

Animals

Twenty-seven whole blood samples were collected from Indian rhesus macaques (*Macaca mulatta*). Five of these samples came from a breeding group of animals living at the Wisconsin National Primate Research Center (WNPRC). The remaining 22 samples were provided by Dr. Michele Di Mascio from the National Institutes of Health's National Institutes of Allergy and Infectious Diseases. Blood sampling was performed under anesthesia and in accordance with the regulations and guidelines outlined in the Animal Welfare Act, the Guide for the Care and Use of Laboratory Animals, and the Weatherall report (Animal Welfare Act 1966; Weatherall 2006).

Data

Exome sequence datasets have been deposited in the sequence read archive (SRA) under BioProjects PRJNA527214 and PRJNA529708. Fasta reference sequences used for MHC genotyping and sequence analysis scripts are available from <https://go.wisc.edu/jb0926>.

MHC class I and class II genotyping by amplicon deep sequencing

Genomic DNA was isolated from 250 μ L of whole blood using a Maxwell® 48 LEV Blood DNA Kit (Promega Corporation, Fitchburg, WI). Following isolation, DNA concentrations were determined with a NanoDrop 2000 and samples were normalized to 60 ng/ μ L. MHC class I and class II PCR amplicons were generated using exon 2-specific primers with adapters (CS1 and CS2) necessary for 4-primer amplicon tagging with the Fluidigm Access Array™ System (Fluidigm, San Francisco, CA, USA) by previously described methods (Karl et al. 2017, 2014). Pooled PCR products were purified using the AMPure XP beads (Agencourt Bioscience Corporation, Beverly, MA, USA) and quantified using the Quant-iT dsDNA HS Assay kit with a Qubit fluorometer (Invitrogen, Carlsbad, CA, USA), following the manufacturer's protocols. The MHC exon 2 genotyping amplicon pools were sequenced on an Illumina MiSeq instrument (San Diego, CA, USA) as previously described (Karl et al. 2017).

Analysis of the MiSeq exon 2 genotyping amplicon sequences was performed using a custom Python workflow. The workflow contained a step to remove oligonucleotide primers and sequencing adapters with `bbduk`, a step to merge reads using `bbmerge`, a step to identify unique sequences/remove chimeras with `USEARCH`, and finally mapping unique reads against a deduplicated reference database of

rhesus macaque partial MHC class I and class II exon 2 sequences with `bbmap` (Bushnell et al. 2017; Edgar 2010). The IPD-MHC NHP database on the European Bioinformatics Institute website is continuously updated as new information from researchers is submitted to it, and we elected to download the release that was available on September 8, 2018 (Maccari et al. 2017). By doing so, we created a fixed snapshot of the reference sequences for our data analyses, and this ensured consistency for us when we tested our experimental hypotheses. For this publication, we define "IPD exon 2" as this database of reference sequences. SAM output files from `bbmap` were parsed with the Python package `pandas` to enumerate the reads from each animal that were identical to IPD exon 2 reference sequences. *Mamu-A*, *-B*, *-DRB*, *-DQA1*, *-DQB1*, *-DPA1*, and *-DPB1* lineage-level haplotypes were inferred for each of the samples with a semiautomated custom workflow that identifies diagnostic alleles associated with previously defined rhesus macaque haplotypes (Karl et al. 2013; Otting et al. 2017).

MHC class I and class II genotyping by exome sequencing

Genomic DNA was isolated as described above and shipped to the Human Genome Sequencing Center at the Baylor College of Medicine. MHC and exon-containing genomic DNA fragments were selectively enriched using a custom target-capture probe set. Genome-wide exons were captured using SeqCap EZ HGSC VCRom2.1, an optimized human clinical exome probe set (Clark et al. 2013; Yang et al. 2013). SeqCap EZ HGSC VCRom2.1 contains probes designed to enrich 23,585 human genes and 189,028 non-overlapping exons. A low-coverage audit was performed to identify rhesus macaque exons inferred from the reference genome *rheMac2* that were not sufficiently enriched ($< 20\times$ coverage) with these human probes, and an additional 22,884 rhesus macaque exons were incorporated into the genotyping probe design (Prall et al. 2017). Finally, and most importantly for MHC analyses, we modified the SeqCap EZ Design: Human MHC Design to selectively enrich MHC class I and class II sequences. This previous design was prepared by the Beijing Genome Institute in collaboration with Roche/NimbleGen, and it targeted the complete 4.97-Mb HLA region with non-redundant probes designed against 8 fully sequenced HLA haplotypes (Horton et al. 2008; Cao et al. 2013). For our macaque studies, we prepared a minimal MHC target capture design using a subset of these probes that are based on all functional HLA class I (*HLA-A*, *-B*, *-C*, and *-E*) and class II (*HLA-DRA*, *-DRB1*, *-DRB3*, *-DRB4*, *-DRB5*, *-DQA1*, *-DQB1*, *-DPA1*, and *-DPB1*) genes. Probes were included to capture complete gene sequences (exons + introns +3' UTR) as well as ~ 1 kb of the 5' upstream flanking sequence. The BED file of rhesus *rheMac2* target coordinates lifted over to *rheMac8* was used

to prepare this combined minimal MHC and supplemental rhesus spike-in probe design. Because derivation of MHC results from genotyping is paramount, we used a ratio of 2.5× spike-in probes to 1× VCRom2.1 probes. The supplemental probes for MHC and rhesus macaque were a single reagent, and the MHC-specific probes only constituted 609 kb of the 37.9-Mb probes (Prall et al. 2017).

An Illumina paired-end pre-capture library was constructed with 750 ng of DNA, as described by the Baylor College of Medicine Human Genome Sequencing Center. Pre-capture libraries were pooled into 10-plex library pools for target capture according to the manufacturer's protocol. Samples were pooled in 10-plex sequence capture library pools for 151-bp paired-end sequencing in a single lane of an S4 flow cell on an Illumina NovaSeq 6000 at the Baylor College of Medicine Human Genome Sequencing Center.

Enumeration of MHC reads in exome data

The effective enrichment of MHC reads in the exome datasets was calculated by mapping the reads of each sample's exome dataset against an individual genomic reference file for each individual locus: *HLA-A* exons 2–3 (NCBI gene ID: 3105) and *HLA-E* exons 2–3 (gene ID: 3133), *HLA-DPA1* exons 2–4 (gene ID: 3113), *HLA-DPB1* exons 2–4 (gene ID: 3115), *HLA-DQA1* exons 2–4 (gene ID: 3117), *HLA-DQA2* exons 2–4 (gene ID: 3118), *HLA-DQB1* exons 2–4 (gene ID: 3119), *HLA-DQB2* exons 2–4 (gene ID: 3120), *HLA-DRB1* exons 2–4 (gene ID: 3123), *HLA-DRB3* exons 2–4 (gene ID: 3125), *HLA-DRB4* exons 2–4 (gene ID: 3126), and *HLA-DRB5* exons 2–4 (gene ID: 3127). This mapping was done by using *bbmap* with default parameters, which corresponds to a minimum alignment identity of approximately 76% (Bushnell et al. 2017). Empirically, these mapping parameters are sufficient to map macaque MHC reads to their human orthologues. Mapped reads were written to a new fastq file using *bbmap*'s *outm* = parameter. To quantify the total number of reads in a sample and the number of reads extracted with our reference file, we created a custom Python script, which is available for download.

MHC genotyping from exome data

Two complementary data analysis strategies were employed to analyze the exome sequence data, and to verify reproducibility and confidence in the MHC genotyping results. For accuracy and quantification purposes, the expected MHC genotypes for each animal were established based on concordance among at least two out of the three described strategies and biological plausibility, e.g., no more than two alleles per *Mamu-DQA1*, *-DQB1*, *-DPA1*, or *-DPB1* locus.

Strategy 1: MHC genotyping using diagnostic subregion

The diagnostic subregion (DSR) was an intra-allelic region that encompassed polymorphisms, and these polymorphisms were distinguishable from alleles with similar sequences. Therefore, this method ensured that the DSR was captured in at least one read for each called allele. MHC class I and class II reads initially were extracted from each animal by mapping the FASTQ reads to HLA class I and class II reference sequences containing exons 2–3 and exons 2–4, respectively, plus the intervening intron(s) as described above in “Enumeration of MHC reads in exome data.” Reads were mapped to these reference sequences using *bbmap* with default parameters and the parameter (*qtrim* = *lr*) (Bushnell et al. 2017).

Following extraction of MHC reads from the total exome sequences, the MHC reads were prepared for assembly using a modified version of a data pre-processing pipeline, which included tools from the BBTools package (Bushnell et al. 2017). Briefly, optical duplicates and reads from low-quality regions of the sequencing run were removed. Next, Illumina sequencing adapters were trimmed from the ends of sequencing reads. Any residual spike-in or PhiX sequences that inadvertently survived mapping to HLA class I and class II were then removed. Three rounds of error-correction and read merging were performed to create high-confidence merged reads that were well-supported by common kmers found in the extracted MHC reads. The error-corrected reads were not merged with a minimum overlap, but instead were separately mapped against the IPD exon 2 sequences using *bbmapskimmer*. The default settings for the software tool *bbmapskimmer* were used with the following modified parameters (*semiperfectmode* = *t* *ambiguous* = *all* *ssa* = *t* *maxsites* = 50000 *maxsites2* = 50000 *expectedsites* = 50000) (Bushnell et al. 2017). The *semiperfectmode* setting accepted reads with perfect matches, as well as reads that extended off the end of contigs for no more than half of the length of the mapped read segment. The *ambiguous* setting and the “*expectedsites*” setting reported the first 50,000 matched read segments that met the “*semiperfectmode*” filtering. These settings were set exceedingly high in order to exhaustively map the reads to our IPD exon 2 reference file of approximately 874 alleles. Using the *semiperfectmode*, these were the segment sequences with the longest matching length. This output file included all mapped reads to all IPD exon 2 reference allele sequences.

We then used *samtools mpileup* with the settings (*-A -a -ff UNMAP -x -B -q 0 -Q 0*) on the *bbmapskimmer* output to calculate a depth of coverage at each position for each IPD exon 2 sequence. Next, we removed any aligned reads to the IPD exon 2 database sequences that contained less than a minimum depth of coverage of 2 across the entire reference sequence. For ambiguously mapped reads, alignments with the longest-matching region were selected for further analysis;

ties among alignments were counted multiple times. To reduce the number of false positives, we used Python pandas to only report database sequence matches that had at least one unambiguously mapped read. Based on these mapping parameters, unambiguously mapped reads must span the DSR. The MHC genotypes from this method were reported as the minimum depth of coverage for each IPD exon 2 database sequence per animal.

Strategy 2: de novo reconstruction of MHC sequences from exome reads

Most de novo sequence assemblers have been optimized for resolving long contigs, and are tolerant of small sequence mismatches that can otherwise fragment assemblies. In the case of MHC alleles, however, such closely related sequences were often biologically distinct. Because of the gene duplications in the macaque MHC, there were many valid MHC contigs that could be assembled from a single sample within the exome data. This is conceptually similar to the computational challenge of assembling viral haplotypes, where rapidly evolving viruses such as human immunodeficiency virus accumulate variants that frequently co-segregate as minor populations within an infected person (Baaijens et al. 2017). Therefore, we utilized the overlap assembly algorithm SAVAGE, originally designed to reconstruct viral haplotypes, to reconstruct MHC allele sequences from exome reads (Baaijens et al. 2017). Similar to strategy 1 above, HLA-mapped reads were pre-processed using the BBTools package to remove low-quality reads and optical duplicates. Next, adapters were trimmed, residual spike-in and PhiX sequences were removed, three steps of error-correction took place, and reads were merged. These merged reads, as well as high-confidence unmerged paired-end reads that could not be grouped into an overlapping merged read, were used for SAVAGE assembly. The following workflow was implemented in a reproducible snakemake workflow that fully documents parameter selection (Koster and Rahmann 2012), and is available upon request.

SAVAGE is designed to construct individual haplotypes from the overlap graph of individual reads. We processed the totality of sequence data in a single patch in order to maximize the sensitivity of the contiguous reconstruction with parameter “–split 1,” as well as the parameter “–revcomp” to handle reverse complement reads. The IPD exon 2 reference database that was used for MiSeq and DSR genotyping was also used to assess the quality of SAVAGE genotypes. The IPD sequences were mapped to contigs produced by SAVAGE using sensitive parameters in bbmap (minlength = 100 vslow = t subfilter = 0 indelfilter = 0 lengthtag = t ignorefrequentkmers = t

kfilter = 100) designed to identify sequences that perfectly match SAVAGE contigs. A post-processing script refined these mappings, and only retained the mappings where the length of the mapped region was the same as the length of the IPD exon 2 sequence. These mappings indicated where the reference database sequence was fully and exactly contained within a SAVAGE contig.

Results

MHC reads are efficiently enriched using target-capture probes

We designed a custom target enrichment probe set that accounts for the extensive duplication of macaque MHC genes (Prall et al. 2017). A tripartite target capture system was used in this study. The first component is SeqCap EZ HGSC VCRom2.1, an optimized human clinical exome probe set. Since macaques are closely related to humans, SeqCap EZ HGSC VCRom2.1 can also be used in macaques, though some sequences that are most divergent between macaques and humans are not efficiently captured. The second component of the target capture system is an additional 22,884 rhesus macaque exon sequences that were not effectively captured with the SeqCap EZ HGSC VCRom2.1 reagent. The third component is a collection of probes designed to specifically enrich macaque MHC class I and class II sequences. These probes span the full length of HLA class I and class II genes including introns, 3' UTRs, and approximately 1,000 bp of the flanking 5' sequence.

We obtained a median coverage of 100× for the target exon sequences across the genome with > 20× coverage for 94.97% of bases that were targeted in this study. As illustrated in Table 1, an average of 70,549,789 Illumina sequence reads per sample were obtained for the 27 animals evaluated in this study. These reads were mapped against reference files containing representative genomic HLA exons 2–3 for *HLA-A* and *HLA-E*, and exons 2–4 for *HLA-DRB1*, *-DRB3*, *-DRB4*, *-DRB5*, *-DQA1*, *-DQB1*, *-DPA1*, and *-DPB1* sequences. We identified an average of 269,057 MHC class I and class II sequence reads per sample which corresponds to an average of 0.37% of the total sequence reads evaluated per sample (Table 1). In a previous study by Ericson et al. (2014), we found that MHC sequences only accounted for an average of 0.13% of the total Illumina sequence reads that were evaluated per animal when the standard HGSC VCRom2.1 panel was used alone for target capture (Supplementary Table 1). Thus, we achieved an almost threefold increase in MHC genomic sequences after inclusion of the spike-in probes for target capture compared with use of the VCRom2.1 probe set alone.

Table 1 Fraction of total exome sequence reads corresponding to exons 2–3 of MHC class I and exons 2–4 of MHC class II genes

Animal ID	SRA accession	Total Raw Reads	MHC-I		DRB1/3/4/5	DQA1	DQB1	DPA1	DPB1 reads	MHC	MHC % of Total
			Reads Extracted	Reads Extracted	Reads Extracted	Reads Extracted	Reads Extracted	Reads Extracted	Reads Extracted	Reads Extracted	
r05029	SAMN11131055	87,345,430	79,712	215,654	24,266	39,188	29,604	112,376	500,800	0.57	
r17099	SAMN11131056	67,632,236	32,406	89,946	9,694	16,782	11,214	47,974	208,016	0.31	
r07010	SAMN11173514	81,062,856	71,694	204,352	22,528	38,382	25,392	111,626	473,974	0.58	
r17041	SAMN11131058	80,004,430	30,852	244,326	9,746	16,310	12,644	151,376	465,254	0.58	
r17061	SAMN11131059	78,428,820	70,082	204,694	20,776	41,018	19,440	95,538	451,548	0.58	
J8R	SAMN11282383	80,868,782	45,164	87,864	11,878	19,736	13,810	57,146	235,598	0.29	
J01	SAMN11282382	61,009,650	34,342	87,124	9,050	16,628	10,958	46,964	205,066	0.34	
ZC08	SAMN11282384	60,693,394	31,168	81,586	10,392	15,660	10,172	45,748	194,726	0.32	
DGKG	SAMN11282378	80,530,772	48,658	130,256	12,788	22,682	13,372	62,990	290,746	0.36	
CF18	SAMN11282368	68,593,710	37,792	96,842	10,864	18,226	13,118	51,556	228,398	0.33	
DE1AA	SAMN11282369	65,712,570	39,334	108,002	10,060	17,124	11,840	51,134	237,494	0.36	
DEG8	SAMN11282370	68,490,344	35,566	116,824	10,404	19,278	12,294	55,086	249,452	0.36	
DEXX	SAMN11282371	67,451,626	33,900	128,128	10,442	18,898	9,778	53,258	254,404	0.38	
DF24	SAMN11282372	67,768,724	41,362	96,172	10,606	18,176	10,976	50,612	227,904	0.34	
DF64	SAMN11282373	65,762,818	37,426	107,974	11,222	15,632	11,446	51,214	234,914	0.36	
DF6T	SAMN11282374	71,472,222	41,160	101,102	11,052	18,794	13,492	51,626	237,226	0.33	
DFET	SAMN11282375	77,040,660	49,068	128,862	12,588	20,952	12,586	58,238	282,294	0.37	
DFV0	SAMN11282377	73,877,184	41,924	98,866	11,630	17,690	13,952	52,394	236,456	0.32	
DJ94	SAMN11282379	66,051,412	39,616	88,196	9,910	15,892	10,978	45,972	210,564	0.32	
HIH	SAMN11282380	76,491,578	41,308	85,732	11,148	18,366	14,102	50,746	221,402	0.29	
DFJT	SAMN11282376	62,249,064	37,728	78,860	9,666	15,268	11,426	43,086	196,034	0.31	
HLJ	SAMN11282381	68,288,558	41,748	94,570	9,960	17,240	12,756	48,356	224,630	0.33	
37360	SAMN11282363	62,310,824	38,200	97,452	9,866	15,024	10,320	46,140	217,002	0.35	
OL7	SAMN11282364	64,145,834	35,302	128,778	9,310	19,992	10,686	54,196	258,264	0.40	
OR1	SAMN11282365	64,789,208	36,462	86,772	10,656	17,258	11,982	46,124	209,254	0.32	
OR8	SAMN11282366	72,836,490	38,286	118,184	10,500	22,248	13,170	56,142	258,530	0.35	
OTI	SAMN11282367	80,730,750	41,552	106,356	13,364	20,714	14,860	57,744	254,590	0.32	
Average		70,549,789	42,660	119,018	12,014	20,487	13,569	61,310	269,057	0.37	

MHC genotypes defined from target-enriched genomic DNA are comparable to those obtained by amplicon deep sequencing

We hypothesized that MHC genotypes derived from target-enriched genomic sequence would be comparable in accuracy to MHC genotypes derived from conventional amplicon deep sequencing. In order to compare the accuracy of MHC genotyping results between the methods described here, the expected genotypes for each animal were defined by concordance among at least two out of the three described strategies, as well as biological plausibility. For example, no more than two alleles should be present for each of the MHC class II *Mamu-DQA1*, *-DQB1*, *-DPA1*, or *-DPB1* loci. MHC class I and class II PCR amplicons were deep sequenced on an Illumina MiSeq from the same 27 animals that were evaluated by MES analysis. Representative MHC class I and class II

genotypes supported by the amplicon data are shown in Figs. 1 and 2, respectively. These figures illustrate genotypes from a pedigreed family of Indian rhesus macaques: a sire, a dam, and three progeny that are paternal half-siblings. Although segregation data confirmed all expected genotypes that were shared among the directly related individuals in this breeding group, this was generally not possible, since pedigree information was unavailable for 22 of 27 rhesus macaques examined in this study. Supplementary Figs. 1 and 2 show comprehensive genotyping results for all 27 animals.

The superset of alleles supported by at least two of the methods described in this manuscript (MiSeq amplicon genotyping, DSR genotyping of individual exome sequencing reads, or SAVAGE genotyping of contigs derived from exome data) was considered to be the expected MHC genotypes for these animals. A complication for this assertion was that specific genotypes were difficult for MiSeq analysis to report

Relationship	Dam			Progeny 1			Sire			Progeny 2			Progeny 3			Ambiguous Allele Groups
Animal_ID	r05029	r17099	r07010	r17041	r17061	A004	A008	A001	A008	A002a	A002a	A001	A001	A002a	A002a	
MHC-A Haplotype 1	A004	A004	A004	A008	A002a	A001	A001	A001	A008	A002a	A002a	A001	A001	A002a	A002a	
MHC-A Haplotype 2	A002a	A001	A001	A001	A001	A001	A001	A001	A001	A002a	A002a	A001	A001	A002a	A002a	
MHC-B Haplotype 1	B048	B048	B048	B001a	B001a	B001a	B001a	B001a	B001a	B001a	B001a	B001a	B001a	B001a	B001a	
MHC-B Haplotype 2	B055	B043b	B043b	B043a	B043a	B043a	B043a	B043a	B043a	B043a	B043a	B043a	B043a	B043a	B043a	
Assay	MiSeq	DSR	SAVAGE	MiSeq	DSR	SAVAGE	MiSeq	DSR	SAVAGE	MiSeq	DSR	SAVAGE	MiSeq	DSR	SAVAGE	Ambiguous Allele Groups
Mamu-A Major Alleles																
Mamu-A1*001g2				168	27	1	158	62	1				210	74	1	A1*001:01,A1*001:02
Mamu-A1*002:01	335	136	1										447	127	1	
Mamu-A1*004g1	151	137	1	126	51	1										A1*004_6_alleles
Mamu-A1*008g1							306	79	1	352	36	1				A1*008_5_alleles
Mamu-A1*023_A1*106g1										228	36	1				A1*023:01,A1*023:02,A1*106:01
Mamu-A1*059:01		45														
Mamu-A Minor Alleles																
Mamu-A2*05g1				319	25	1	712	162	1	423	15	1	438	56	1	A2*05_32_alleles
Mamu-A3*13_A4*14g1	77	79	1										91	88	1	A3*13_7_alleles,A4*14_1_allele
Mamu-A3*13g1							93	85	1	97	31	1				A3*13:03,A3*13:11
Mamu-A4*14g1	145	104	1	164	23	1				199	29	1				A4*14_14_alleles
Mamu-E Alleles																
Mamu-E*02g1	775	185	1	1923	169	1	2041	303	1	3240	125	1	914	120	1	E*02_9_alleles
Mamu-E*02g4	307	106	1										382	95	1	E*02:13,E*02:21,E*02:29
Mamu-E*02g5	1104	134	1										1583	69	1	E*02:04,E*02:06
Mamu-E*02g7							467	152	1	634	50	1				E*02:12:01,E*02:28
Mamu-E*02g8	689	116	1	1914	92	1	781	85	1				710	85	1	E*02:20,E*02:30
Mamu-B Major Alleles																
Mamu-B*001g1							161	60	1	143	28	1				B*001:01:01,B*001:04,B*001:05,B*001:06
Mamu-B*007:07							25			27						
Mamu-B*007g1							161	33	1	132	12	1				B*007_6_alleles
Mamu-B*012:01													284	60	1	
Mamu-B_022g1													171	46	1	B*022:01,B*022:02
Mamu-B*027g1										93	32	1				B*027:03,B*027:04
Mamu-B*030g1				270	12	1	303	73	-	318	11	1	671	103	1	B*030_6_alleles
Mamu-B*030g2							306	79	1	306	29	1				B*030:02,B*030:05
Mamu-B*031:03				105	12	1	139	26	1				198	35	1	
Mamu-B*031g1													127	40	1	B*031:01,B*031:02
Mamu-B*041g1	128	47	1	57	24	1										B*041:01,B*041:02
Mamu-B*043:01				223	18	1	224	49	1	263	16	1	220	41	1	
Mamu-B*048:01	275	91	1	318	26	1										
Mamu-B*052:01	54	63	1													
Mamu-B*055:01	180	65	1													
Mamu-B*058:02	177	43	1													
Mamu-B*064:01	179	56	1	44	31	1										
Mamu-B*073g1				129	19	1	140	60	1				149	32	1	B*073:01,B*073:02
Mamu-B*074g1		82	1								18	1	16	48	1	B*074:01,B*074:02,B*074:03
Mamu-B Minor Alleles																
Mamu-B*035_B*049g1	161	92	1							293	37	1	261	80	1	B*035:01,B*035:02,B*049:01
Mamu-B*053g1													114	59	1	B*053:02,B*053:03
Mamu-B*054:01	194	95	1							321	26	1				
Mamu-B*054:04																
Mamu-B*054:05													169	59	1	
Mamu-B*057:01													104	41	1	
Mamu-B*057:06				59	24	1	113	61	1				51	40	1	
Mamu-B*057g1										152	10	1				
Mamu-B*063:03	198	74	1	122	29	1	184	73	1	243	16	1				B*057:02,B*057:03,B*057:04,B*057:05
Mamu-B*063:05																
Mamu-B*063g1	121	74	1										89	68	1	B*063:01:01,B*063:01:02,B*063:02:02
Mamu-B*070g2	28	100	1													B*070:01,B*070:06
Mamu-B*072g1				218	41	1	447	161	1	365	25	1	330	74	1	B*072_8_alleles
Mamu-B*092g1				-	35	1	14	88	1	23	25	1	82	84	1	B*092:01,B*092:02,B*092:03
Mamu-B*098g1	198	108	1							324	32	1	238	60	1	B*098_9_alleles
Mamu-B*109:06	479	92	1	479	49	1										
Mamu-B*134g2	257	82	1	276	24	1										B*134:03,B*134:04,B*134:06
Mamu-B*188g1							336	42	1	535	11	1				B*188:01,B*188:02
Mamu-B11L*01:12				-	29	-	-	89	1							
Mamu-B11L*01g1	-	58	1	-	36	1	-	79	1	-	49	1	-	74	1	B11L*01:01:02,B11L*01:02,B11L*01:04,B11L*
Mamu-B16*01:01				44	34	1	72	61	1				110	29	1	
Mamu-B16*01g1	181	74	1							176	23	1				B16*01:04,B16*01:05,B16*01:07,B16*01:08
Mamu-B16*01g2							51	47	1	77	14	-				B16*01:02,B16*01:09
Mamu-B17*01g1										1221	12	1	1227	17	1	B17*01_14_alleles
Mamu-B17*01g2	1305	74	1	773	18	1	593	24	1	971	9	1				B17*01_9_alleles
Mamu-B17*01g4																B17*01:09,B17*01:15
Mamu-B20*01:02										304	43	1				
Mamu-B20*01g1	176	66	1	181	21	1	272	82	1	365	33	1				B20*01:01,B20*01:04
Mamu-I Alleles																
Mamu-I*01g1	130	63	1	129	31	1	485	151	1	532	44	1	252	68	1	I_01_66_alleles
Mamu-I*01g3										81	28	1				I*01:19,I*01:20:01

Fig. 1 Comparison of MHC class I results from MiSeq PCR amplicon versus whole exome genotyping strategies for a representative breeding group of rhesus macaques. Results for each of the three methods are provided side-by-side in the columns for each macaque. Each row indicates the detection of a specific MHC class I allele or lineage group of closely related sequences that are ambiguous because they are identical

over the IPD exon 2 database sequence. Values in the body of this figure indicate the number of sequence reads supporting each allele call for the MiSeq and DSR methods while alleles supported by a SAVAGE contig are reported with a “1”. Discrepancies between the MiSeq (pink), DSR (yellow), or SAVAGE (blue) methods are highlighted by filled cells with borders

correctly. As we have shown previously (Karl et al. 2017), the number of reads supporting each genotyping call was highly variable, ranging from tens of reads to thousands of reads per allele. A small subset of allelic variants contained nucleotide

substitutions in their sequences within the binding sites of the PCR oligonucleotides that interfere with efficient PCR amplification. Members of the *Mamu-B11L*01* allele lineage exemplify this issue; they have two nucleotide substitutions relative

Relationship Animal_ID Assay	Dam r05029			Progeny 1 r17099			Sire r07010			Progeny 2 r17041			Progeny 3 r17061			Ambiguous Allele Groups
	MS	DSR	SAV	MS	DSR	SAV	MS	DSR	SAV	MS	DSR	SAV	MS	DSR	SAV	
Mamu-DRB Alleles																
<i>DRB1*03:09</i>	198	370	1	72	131	1	170	305	1	272	107	1	156	200	1	
<i>DRB1*03g1</i>				2789	217	1	2714	497	1				3052	499	1	<i>DRB1*03:06, DRB1*03:26</i>
<i>DRB1*10:03</i>				1761	168	1	1952	358	1				2711	310	1	
<i>DRB*W2g1</i>	154	451	1	72	183	1	132	410	1	206	176	1	107	398	1	<i>DRB*W2:01, DRB*W2:05</i>
<i>DRB*W3g1</i>	1936	546	1							3097	217	1				<i>DRB*W3:03:01, DRB*W3:03:02</i>
<i>DRB*W4:01</i>	1777	414	1							2631	158	1				
Mamu-DQA Alleles																
<i>DQA1*01:02</i>				2384	98	1	2444	256	1				2964	154	1	
<i>DQA1*01g1</i>	2298	318	1	2046	96	1	2276	288	1	4317	107	1	3360	210	1	<i>DQA1*01:04:01, DQA1*01:04:02</i>
<i>DQA1*23:01</i>	2883	334	1							5079	108	1				
Mamu-DQB Alleles																
<i>DQB1*06:01</i>	3803	160	1	3980	41	1	3892	152	1	5757	33	1	3892	122	1	
<i>DQB1*06g2</i>				5229	173	1	4784	345	1				6400	282	1	<i>DQB1*06:05, DQB1*06:18</i>
<i>DQB1*18g4</i>	2656	329	1							4504	108	1				<i>DQB1*18:02, DQB1*18:27</i>
Mamu-DPA Alleles																
<i>DPA1*02g1</i>	5450	1765	1	5301	316	1	5153	840	1	8727	758	1				<i>DPA1*02_9_alleles</i>
<i>DPA1*04g1</i>				435	181	1	489	385	1				1075	350	1	<i>DPA1*04:03:01, DPA1*04:03:02, DPA1*04:05</i>
<i>DPA1*11:01</i>													-	225	1	
Mamu-DPB Alleles																
<i>DPB1*02g1</i>				106	25	1	87	73	1				342	47	1	<i>DPB1*02:01, DPB1*02:04:01, DPB1*02:04:02</i>
<i>DPB1*15g1</i>	254	217	1	286	31	1	207	129	1	3084	60	1				<i>DPB1*15:01, DPB1*15:02</i>
<i>DPB1*16:01</i>													1769	98	1	

Fig. 2 Comparison of MHC class II results from MiSeq PCR amplicon versus whole exome genotyping strategies for a representative breeding group of rhesus macaques. Results for each of the three methods are provided side by side in the columns for these five related macaques. Each row indicates detection of a specific MHC class II allele or lineage group of closely related sequences that are ambiguous because

they are identical over the IPD exon 2 database sequence. Values in the body of this figure indicate the number of sequence reads supporting each allele call for the MiSeq and DSR methods, while alleles supported by a SAVAGE contig are reported with a “1”. The *Mamu-DPA1*11:01* allele missed by the MiSeq assay due to multiple mismatches versus the amplification primers is highlighted in pink

to the 5' oligonucleotide that was used to generate MHC class I amplicons. These *Mamu-B11L*01* sequences were routinely absent in MiSeq genotypes (Fig. 1). Likewise, the *Mamu-DPA1*11:01* allele has four nucleotide substitutions relative to the oligonucleotide pair used to generate *DPA1* amplicons for MiSeq genotyping (Fig. 2).

These nucleotide substitutions do not fully account for all differences in the abundance of sequence reads for each allele. Allele lineages, such as *Mamu-B*074* and *Mamu-B*098*, exhibited significantly diminished PCR efficiency despite being perfectly matched with the oligonucleotides used for amplification (Fig. 1). False-positive genotyping calls were also noted in the MiSeq assay that resulted from intermolecular recombination during the PCR process (Ennis et al. 1990; Fichot and Norman 2013; von Wintzingerode et al. 1997). Artifacts from PCR amplification, dubbed “PCR chimeras,” arise when an incompletely extended PCR product serves as a primer with a partially mismatched template during subsequent cycles of PCR (Supplementary Fig. 3). This was exemplified by read support for the presence of *Mamu-B*007:07* in sire r07010, and progeny 2 r17041 (Fig. 1). *Mamu-B*007:07* only differs from the *Mamu-B*007g1* allele group that was determined to be present in this pair of animals by a single-nucleotide variant at the extreme 5' end of the class I genotyping amplicon. Chimeric PCR products, equivalent to the *Mamu-B*007:07* sequence, were formed between the 3' portion of the *Mamu-B*007g1* sequence and other allelic variants in these animals with this 5' SNP such as *Mamu-*

*B20*01g1*. These results illustrate that the MiSeq amplicon genotyping, while generally reflective of an animal's MHC genotype, can yield both false-positive and false-negative results. When compared with the expected genotypes for this dataset, MiSeq amplicon genotyping has 90.4% MHC class I and 97.8% MHC class II concordance (Table 2).

Two separate strategies were used to derive MHC genotypes from target-capture data using data from macaque exome sequencing. These strategies were outlined in Fig. 3, and can be described as follows. The DSR analysis was a straightforward extension of the methodology used for deriving genotypes from MiSeq amplicons. MHC sequence reads were extracted from the total exome dataset. A simplified workflow that mapped those MHC reads to reference sequences, and then assigned the genotypes based on the presence of exome reads overlapping each position of the reference sequence, was problematic. This simplified workflow was extremely vulnerable to false-positive genotypes, and the similarities among different MHC sequences often enabled those reads to map to multiple different alleles (Wiseman et al. 2013). Two or more reads each could partially match a portion of a sequence in the IPD exon 2 database, which could complement each other to provide support for an allele that was not biologically relevant. As discussed in “Methods,” the DSR encompassed polymorphisms that discriminated among closely related alleles by requiring at least one mapped read to unambiguously map to the corresponding allele within the IPD exon 2 sequences. This strategy can identify specific

Table 2 Summary of discordant results for each MHC genotyping method by animal

MHC class I					MHC class II			
Animal ID	Alleles	Discordance	Discordance	Discordance	Alleles	Discordance	Discordance	Discordance
r05029	28	2	1		10			
r17099	25	3		1	12			
r07010	26	3	1	1	12			
r17041	31	3		1	10			
r17061	29	1	2		12	1		
J8R	29	4	1		12	2		
J01	30	3	1	1	11	1		
ZC08	24	2			1	12		
DGKG	28	4	1		13			
CF18	28	3			12			
DE1AA	30	5	1		13			
DEG8	27	3	1		13			
DEXX	24	3	1	1	14			
DF24	32	4		1	12			
DF64	30	3		1	1	13		
DF6T	31	2		1	10			
DFET	30	3	1	1	13	1		1
DFV0	28	2			6			
DJ94	34	3	2	1	12			
HIH	30	3			10			
DFJT	28	2	3		11			
HLJ	29	2	1	1	11			
37360	29	2	1	1	13	1		
OL7	20	2			14			
OR1	17	1			12			
OR8	25	2			13	1		
OTI	25	2			1	12		
Total	747	72	20	13	318	7	0	1
Discordance (%)		9.6	2.7	1.7		2.2	0.0	0.3
Concordance (%)		90.4	97.3			97.8	100.0	99.7

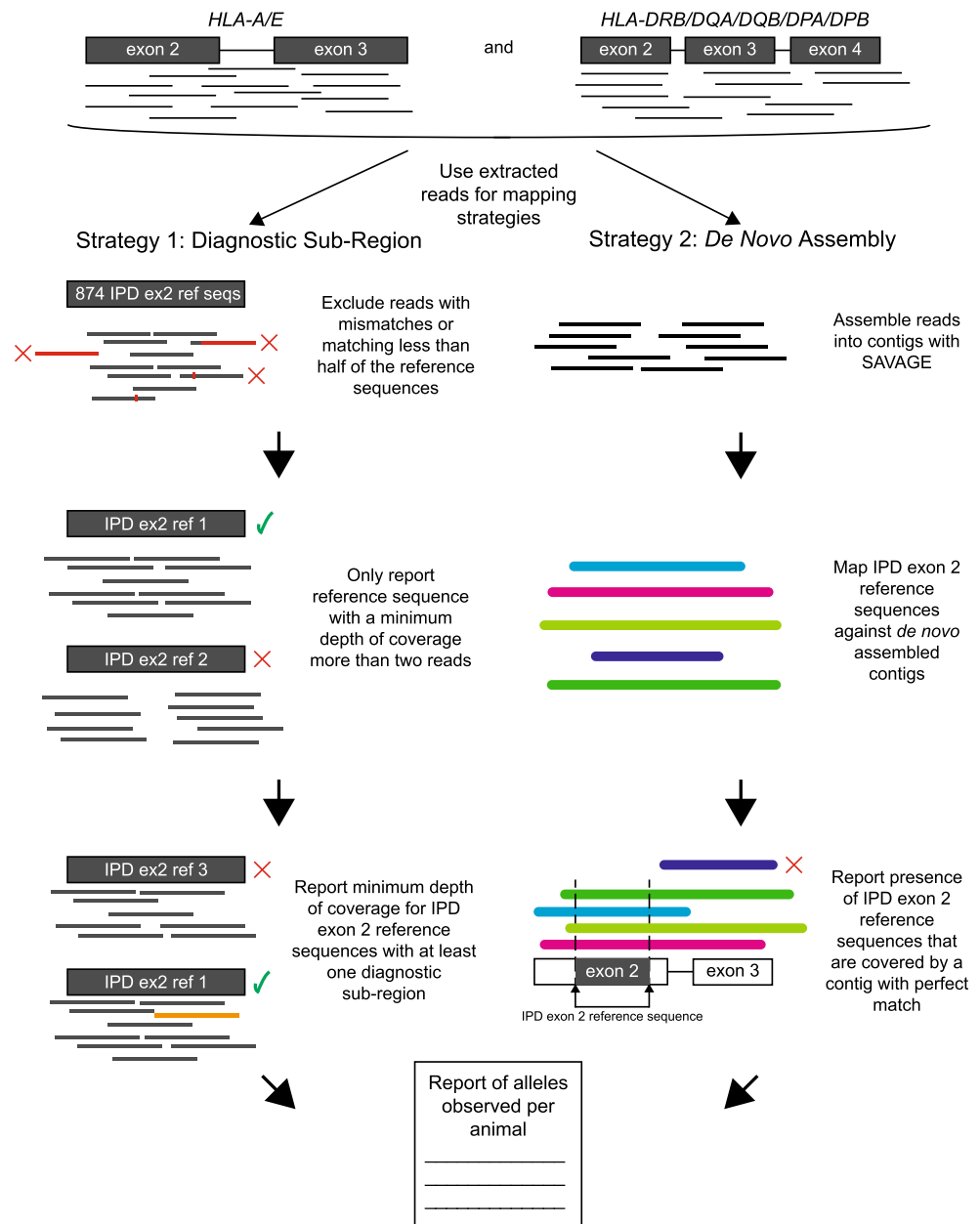
polymorphisms of interest among the IPD exon 2 sequences to produce genotypes for MHC class I and MHC class II (Figs. 1, 2, and Supplementary Figs. 1 and 2). For the 27 animals evaluated in this study, DSR was 97.3% and 100% concordant with expected MHC class I and class II genotypes, respectively (Table 2).

The DSR strategy did not consider sequences that were not among the IPD exon 2 sequences, and this lack of consideration contributes to overcalled alleles. The apparent *Mamu-A1*059:01* allele in dam r05029 was erroneously derived from reads that were from *Mamu-A1*004g1* and an unknown allele that was not among the sequences in the IPD exon 2 database. As a result of this unknown sequence not being among the known IPD exon 2 sequences, reads unambiguously mapped to *Mamu-A1*059:01*, which caused this allele to be overcalled. This type of overcalling will be mitigated with

the discovery of additional allelic variants and their inclusion in future iterations of the IPD database.

The second approach for determining genotypes performed de novo assembly on MHC class I and class II reads from each sample. The resulting assembled contigs were then mapped against IPD exon 2 reference sequences to define perfectly matching contigs. Because most assemblers were not tuned for the challenge of assembling large numbers of contigs that differ from one another by as little as 1 bp, we relied on an assembler, SAVAGE, originally designed to reconstruct viral sequencing haplotypes. As shown in Figs. 1, 2, and Supplementary Fig. 1, contigs produced by SAVAGE matched the expected genotypes. Unlike the DSR method, SAVAGE assembled sequence contigs that may be useful for downstream analyses such as higher-resolution MHC genotyping. False positives and false negatives among closely

Fig. 3 Description of strategies employed for MHC genotyping from macaque exome sequences. Exome sequencing reads are extracted by mapping to *HLA-A/E* exons 2–3 and *HLA-DRB/DQA/DQB/DPA/DPB* exons 2–4. These extracted sequences are then used for both analysis strategies. Strategy 1 maps the extracted reads to IPD exon 2 reference sequences. Only reads with perfect matches that extended off the end of contigs for no more than half of the length of the mapped read segment were retained. Next, each alignment of mapped sequences is checked to exclude those with a coverage that is less than 2 reads. The mapped alignments with this coverage are used to form a diagnostic subread, which is used to match alignments to the reference sequences. The minimum depth of coverage of the mapped alignments was reported in the output. Strategy 2 conducts a de novo assembly of the extracted reads using the computational tool SAVAGE. Then, the assembled contigs are mapped to IPD exon 2 reference sequences. To determine correct mapping, any IPD exon 2 reference sequences that fully cover and perfectly match a contig are reported in the output



related variants were mitigated by the inclusion of the filtering steps described under strategy 2. Across all 27 samples, the SAVAGE contigs are 98.3% and 99.7% accurate with respect to the expected MHC genotypes (Table 2).

While this analysis focused on genotyping using the same exon 2 reference sequences that are commonly utilized for MiSeq amplicon analyses, the SAVAGE contigs are frequently much longer than these reference sequences (Fig. 4). These extended contigs frequently contain exons 2 through 4, plus the intervening introns, and could be used to provide higher-resolution genotyping than is possible using exon 2 sequence alone. Moreover, contigs that contain complete sequences for exons 2–3 of MHC class I and exon 2 of MHC class II alleles meet the minimum criteria for obtaining formal allele

nomenclature for non-human primates from the IPD-MHC (Robinson et al. 2013).

Discussion

Here we describe MES genotyping of Indian rhesus macaques. In our estimation, this method will supersede MiSeq PCR amplicon genotyping as the most widely used macaque MHC genotyping assay in the future. This methodology has several compelling advantages. Most importantly, exome sequencing dramatically improves the overall quantity and quality of genomic information obtained from each sample. The same datasets used for MHC analyses may be used to evaluate

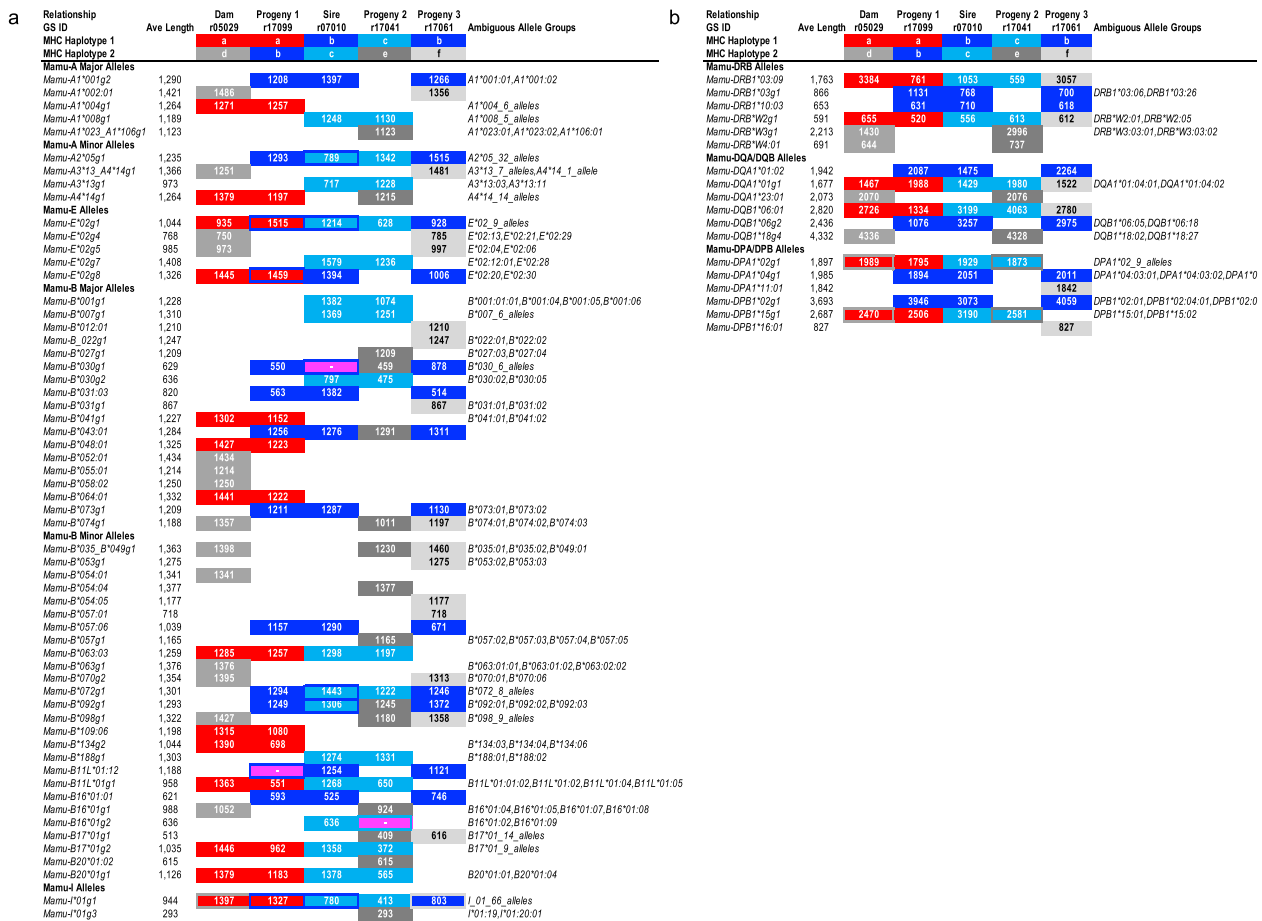


Fig. 4 Lengths of MHC sequence contigs assembled by the SAVAGE method for a representative breeding group of rhesus macaques. Values in the body of this figure indicate contig lengths generated for these MHC sequences in each animal. **a** MHC class I genotyping results are illustrated for the five animals in this breeding group. Three false-negative alleles call for SAVAGE versus the expected genotypes for these animals are highlighted in magenta, e.g., *Mamu-B*030g1* in sire r07010. Sequences highlighted in red are associated with the maternal MHC haplotype a that was inherited

by progeny 1 from its dam. Sequences in progeny 1 and progeny 3 for their MHC haplotype b that was inherited from sire r07010 are highlighted in dark blue while the light blue sequences represent the alternate paternal haplotype c that was inherited by progeny 2. Three unique extended MHC haplotypes in this breeding group are indicated with shades of gray (haplotypes d–f). MHC allele groups that are shared by both parental haplotypes are indicated by colored borders around filled cells. **b** MHC class II genotyping results are illustrated for this same breeding group

protein-coding genetic variation throughout the genome. The loss of start and stop codons exome-wide can be obtained from the same datasets by modifying the workflows in silico, as opposed to requiring the sequencing of multiple new sets of target genes (Yang et al. 2013). Exome-wide datasets offer the promise of retrospectively identifying candidate DNA sequence variants that may be responsible for unexpected experimental outcomes in studies with macaques and other non-human primates. The availability of exome-wide sequences in conjunction with MHC genotypes may also increase the rigor of prospective macaque experiments by enabling more sophisticated balancing of experimental groups, and exclusion of animals whose genetics are likely to strongly bias experimental results (Loffredo et al. 2007; Reynolds et al. 2011; Haus et al. 2014).

These same exome datasets also offer the potential to improve the quality of MHC genotyping. Full-length long-read MHC transcript sequencing offers the highest resolution, but this technology can be labor intensive and difficult to scale (Karl et al. 2017; Semler et al. 2018). The MiSeq exon 2 amplicon approach is limited in its allelic resolution, but it is the current standard deep sequencing approach for high-throughput MHC genotyping in macaques. In this report, we compare two novel strategies for MHC genotyping from MES datasets to this standard MiSeq amplicon genotyping method. The results that we obtained with all three approaches show strong concordance with expected MHC genotypes. As illustrated in Fig. 4a, MHC class I genomic contigs with an average length of approximately 1.1 kb can be assembled from

sequence reads that were initially extracted from the whole exome datasets.

Following the initial enrichment step, MHC class II genomic contigs averaging approximately 1.9 kb in length could be assembled from whole exome sequence reads that mapped to *HLA-DRB1*, *-DRB3*, *-DRB4*, *-DRB5*, *-DQA1*, *-DQB1*, *-DPA1*, and *-DPB1* reference sequences, which contained exons 2–4, and the pairs of introns (Fig. 4b). The addition of HLA spike-in probes containing both exon and intron sequences for the target capture step and the introduction of 151-bp paired end reads for the Illumina NovaSeq platform greatly facilitated our ability to assemble these extended MHC genomic contigs. Additional technological advances, including even longer sequence reads and more efficient assembly algorithms, will undoubtedly increase genomic contig lengths as well as MHC allelic resolution in future studies.

Both MHC genotyping approaches described here depend upon mapping exome sequence reads against a reference database of MHC class I and class II allele sequences. Currently, the IPD database mostly is restricted to coding regions for macaque MHC sequences, and many IPD entries are only partial transcript sequences that lack complete coding regions. It is also very challenging to correctly phase short Illumina sequence reads. The reads map to different exons of a specific allelic variant, and are separated by intronic sequences that span hundreds to thousands of base pairs in genomic DNA. Our results with the SAVAGE workflow (Fig. 4) demonstrate that exome sequence reads that have been enriched with the enhanced MHC probe design described here can be assembled into genomic contigs that span multiple exons and introns. These contigs, therefore, could also be used to improve MHC reference databases, though this requires a major effort with more animals, and is beyond the scope of this manuscript.

The current cost of genotyping is relatively high compared with amplicon deep sequencing, primarily due to two expenses. First, the amount of sequence data needed for a single sample can be high. Compared with conventional MHC genotyping, where 192 macaques' data can be collected on a single instrument run, exome data acquisition is much more expensive. On an Illumina NovaSeq instrument, which has a much higher run cost than the MiSeq, only 70 exome samples can be sequenced simultaneously per lane of an S4 flow cell. However, this cost of sequencing is rapidly decreasing, and as of early 2019, commercial providers have advertised sequencing for US\$9 per gigabyte of whole genome sequence data.¹ Second, a major expense in MHC genotyping is the production, validation, and use of target-capture arrays, as well as the development of in silico data analysis workflows. The approaches

described here, in particular MHC genotyping from SAVAGE contigs produced by de novo assembly of exome reads, is flexible and should be adaptable as sequencing approaches evolve and improve.

A major goal for future studies will be to attempt to extend these contigs to encompass full-length genomic MHC sequences using SAVAGE or other assembly software tools. The relatively compact genomic structure and consistent length of MHC class I genes increase the attainability of this goal. Establishment of comprehensive macaque MHC allele databases of extended genomic sequences will greatly facilitate mapping of exome sequence reads since they will be contiguous with the reference sequences instead of being interrupted by intervening sequences between each exon that are not included in current non-human primate IPD-MHC databases (Maccari et al. 2017).

These results demonstrate that MHC genotypes can be obtained by analyzing genomic DNA selectively enriched for MHC and protein-coding gene sequences. This represents an important advance for characterizing MHC genetics in macaques, and this suggests that analyses of whole exome and whole genome data will become the predominant method for studying macaque genetics in the coming decade.

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