1 Construction of a new chromosome-scale, long-read reference

2 genome assembly for the Syrian hamster, *Mesocricetus auratus*

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24 **Abstract**

25 Background

The Syrian hamster (*Mesocricetus auratus*) has been suggested as a useful mammalian model for a variety of diseases and infections, including infection with respiratory viruses such as SARS-CoV-2. The MesAur1.0 genome assembly was generated in 2013 using whole-genome shotgun sequencing with short-read sequence data. Current more advanced sequencing technologies and assembly methods now permit the generation of near-complete genome assemblies with higher quality and greater continuity.

33 Findings

Here, we report an improved assembly of the *M. auratus* genome (BCM_Maur_2.0) using Oxford Nanopore Technologies long-read sequencing to produce a chromosomescale assembly. The total length of the new assembly is 2.46 Gbp, similar to the 2.50 Gbp length of a previous assembly of this genome, MesAur1.0. BCM_Maur_2.0 exhibits significantly improved continuity with a scaffold N50 that is 6.7 times greater than MesAur1.0. Furthermore, 21,616 protein coding genes and 10,459 noncoding genes are annotated in BCM_Maur_2.0 compared to 20,495 protein coding genes and 4,168 noncoding genes in MesAur1.0. This new assembly also improves the unresolved regions as measured by nucleotide ambiguities, where approximately 17.11% of bases in MesAur1.0 were unresolved compared to BCM_Maur_2.0 in which the number of unresolved bases is reduced to 3.00%.

45 **Conclusions**

Access to a more complete reference genome with improved accuracy and continuity
will facilitate more detailed, comprehensive, and meaningful research results for a wide
variety of future studies using Syrian hamsters as models.

49 Keywords

50 Syrian hamster, Mesocricetus auratus, genome, disease model, COVID-19

52 Data Description

53 Introduction

The Syrian hamster (*Mesocricetus auratus*, NCBI:txid10036) has been used in biomedical research for decades because it is a good model for studies of cancer [1], reproductive biology [2] and infectious diseases [3,4], including SARS-CoV-2, influenza virus, and Ebola virus [5–9]. The use of Syrian hamsters in research has declined [10], likely due to advances in the genetic and molecular tools available for other rodents, especially laboratory mice, and not to a reduction in the utility of hamsters in biomedical research [3].

61 Syrian hamsters are particularly important for COVID-19 research. They spontaneously 62 develop more severe lung disease than other animal models, such as wild-type mice, 63 macagues, marmosets, and ferrets [5,11–14]. After intranasal infection, Syrian hamsters 64 consistently show signs of respiratory distress, including labored breathing, but typically 65 recover after 2 weeks [15]. This is in stark contrast to wild-type laboratory mice that are 66 minimally susceptible to most SARS-CoV-2 strains that were circulating in 2020, though 67 laboratory mice may be more susceptible to certain variants of concern that began 68 circulating in 2021 [8,16]. Furthermore, a recent analysis has suggested that Syrian 69 hamsters fed a high-fat, high-sugar diet exhibit accelerated weight gain and pathological 70 changes in lipid metabolism, as well as more severe disease outcomes when 71 subsequently infected with SARS-CoV-2 [17]. This result has obvious parallels with 72 observations of the effects of comorbidities in humans suffering from COVID-19.

73 COVID-19 pathology in Syrian hamsters appears to be due to a dysregulated innate 74 immune response involving signal transducer and activator of transcription factor 2 75 (STAT2)-dependent type I (IFN-I) and type III interferon (IFN-III) signaling [18]. IFN-I 76 signaling can limit virus replication and dissemination and it has been shown that 77 intranasal administration of IFN-I in Syrian hamsters reduces viral load and tissue 78 damage [19]. The human angiotensin-converting enzyme 2 (ACE2) was identified as 79 the cell entry receptor of SARS-CoV-2 [20]. In addition, upon the engagement of ACE2 80 with SARS-CoV2, cellular transmembrane protease 'serine 2' (TMPRSS2) mediates the 81 priming of viral spike (S) protein by cleaving at the S1/S2 site and inducing the fusion of 82 viral and host cellular membranes, thus facilitating viral entry into the cells [21]. Human 83 ACE2 and hamster ACE2 receptors had previously been shown to share substantial 84 sequence homology, which strongly points to interaction with SARS-CoV-2 receptor 85 binding domain (RBD) structures and similar binding affinity [22]. In silico interaction 86 prediction analysis suggests that human and hamster TMPRSS2 are structurally very 87 similar. Even with slight differences in amino acid residue interactions, human and 88 hamster TMPRSS2 activity are identical for residue interactions related to SARS-CoV-2 89 infectivity [22]. As COVID-19 causes systemic disease in people, precision modeling of 90 specific aspects of pathogenesis will require carefully evaluating similarities and 91 differences across various biological processes in humans and Syrian hamsters which, 92 in turn, will require extensive genomic comparisons between the two species.

93 The currently available reference genome sequence for the Syrian hamster was 94 produced in 2013 using a whole-genome shotgun sequencing approach implementing 95 short read sequencing technology. The resulting MesAur1.0 reference sequence

96 (Genbank accession number GCA_000349665.1) is typical of those produced at that 97 time, containing 237,699 separate contigs with contig N50 of 22,512 bp. The quality and 98 research potential of the existing Syrian hamster genome is limited by the technology 99 that was available at the time of its development; for example, the cluster of type I 100 interferon genes was not resolvable with this technology. In this Data Note, we report 101 the production of a new Syrian hamster reference genome that was sequenced using 102 long-read methods on the Oxford Nanopore Technologies (ONT) PromethION platform 103 and assembled into highly contiguous chromosomes using a combination of Flye [23] 104 and Pilon [24] assembly software. The final assembly, BCM Maur 2.0, improves upon 105 guality and contiguity in comparison with MesAur1.0, with longer contigs and more 106 contiguous sequence, allowing for a more complete reference genome with improved 107 accuracy that will benefit a wide variety of future studies using the Syrian hamster 108 reference genome.

Methods

DNA isolation, library construction, and sequencing 110 111 All genomic DNAs for this study were isolated from a single female LVG Golden Syrian 112 hamster (SY011) that was purchased from Charles River, Inc. (Kingston, NY). All 113 procedures were performed in accordance with the guidelines set by the Institutional 114 Animal Care and Use Committee at the University of Wisconsin-Madison. The protocol 115 was approved by the Institutional Animal Care and Use Committee at the University of 116 Wisconsin-Madison (protocol number V00806). Data from this individual are available in 117 NCBI BioProject PRJNA705675, BioSamples SAMN18096087 and SAMN18096088.

Qiagen AllPrep DNA/RNA Mini kits were used to extract DNA from frozen liver while Qiagen Blood and Cell Culture DNA Midi Kits were used for extractions from frozen kidney. Ultra-high molecular weight DNA for optical mapping was purified from frozen liver using an Animal Tissue DNA Isolation Kit from Bionano Genomics, Inc. (San Diego, CA).

123 Oxford Nanopore long-read sequencing

124 We prepared three separate genomic DNA isolates from the same Syrian hamster 125 (BioSample SAMN18096087). These aliquots were sheared to distinct target fragment 126 lengths (10 kb, 20kb and 30kb) in order to assess the effect of fragment size on flowcell 127 yield and improve efficiency. The two smaller length fragment libraries were sheared 128 using Covaris gTube and the 30kb targeted size library was fragmented with Diagnode 129 Megarupter 3, all following manufacturer's recommendations. The Oxford Nanopore 130 sequencing libraries were prepared using the ONT 1D sequencing by ligation kit (SQK-131 LSK109). Briefly, 1-1.5ug of fragmented DNA was repaired with the NEB FFPE repair 132 kit, followed by end repair and A-tailing with the NEB Ultra II end-prep kit. After a clean 133 up step using AMPure beads, the prepared fragments were ligated to ONT specific 134 adapters via the NEB blunt/TA master mix kit. Each library underwent a final clean up 135 and was loaded onto a PromethION flow cell per manufacturer's instructions. One 136 library was sequenced per flow cell with standard parameters for 72 hrs. Base-calling 137 was done onboard the PromethION instrument using neuronal network based software 138 (Oxford Nanopore Technologies, UK).

139 Illumina sequencing

140 500ng of input genomic DNA from a kidney sample (BioSample SAMN18096088) was 141 used to generate standard PCR-free Illumina paired-end sequencing libraries. Libraries 142 were prepared using KAPA Hyper PCR-free library reagents (KK8505, KAPA Bio-143 systems) in Beckman robotic workstations (Biomek FX and FXp models). Total genomic 144 DNA was sheared into fragments of approximately 200-600 bp in a Covaris E220 145 system (96-well format) followed by purification of the fragmented DNA using AMPure 146 XP beads. A double size selection step was employed, with different ratios of AMPure 147 XP beads, to select a narrow size band of sheared DNA molecules for library 148 preparation. DNA end-repair and 3'-adenylation were then performed in the same 149 reaction followed by ligation of the barcoded adaptors to create PCR-Free libraries. The 150 resulting libraries were evaluated using the Fragment Analyzer (Advanced Analytical 151 Technologies, Ames, Iowa) to assess library size and presence of remaining adaptor 152 dimers. This was followed by gPCR assay using KAPA Library Quantification Kit and 153 their SYBR FAST qPCR Master Mix to estimate the size and quantify fragment yield.

Sequencing was performed on the NovaSeq 6000 Sequencing System using the S4 reagent kit (300 cycles) to generate 2 x 150 bp paired-end reads. The final concentration of the libraries loaded on flowcells was 400-450 pM. Briefly, the libraries were diluted in an elution buffer and denatured in sodium hydroxide. The denatured libraries were loaded into each lane of the S4 flow cell using the NovaSeq Xp Flow Cell Dock. Each lane included ~1% of a PhiX control library for run quality control.

160 Genome Assembly

161 We generated 221 gigabases of sequence data using the ONT PromethION platform 162 (NCBI BioProject PRJNA705675, SRA Experiment SRX11206953). This represents an 163 anticipated 88X coverage of the expected 2.5 Gbp Syrian hamster genome. The raw 164 sequencing reads exhibited an N50 length of 15,730 bp. We used the Flye assembler 165 v2.8.1 [23] to generate an initial de novo genome assembly. Given the potential 166 sequence error rate of PromethION reads, it is advisable to use higher quality Illumina 167 short reads mapped to an assembly to correct sequence errors in initial contigs. 168 Consequently, we used Pilon software v. 1.23 [24] with default settings and 30X 169 genome coverage of Illumina data (SRX10928323) generated from a kidney sample 170 (SAMN18096088) obtained from the same individual for this sequence polishing step. 171 Pilon sequence polishing was performed one time prior to the optical mapping analyses.

172 Optical mapping for scaffold improvement

173 Ultra-high molecular weight (UHMW) DNA was extracted following manufacturer's 174 guidelines (Bionano Prep SP Tissue and Tumor DNA Isolation protocol) from frozen 175 liver tissues obtained from the same animal used for ONT PromethION sequencing 176 (SAMN18096087). Briefly, a total of 15-20mg of liver tissue was homogenized in cell 177 buffer and digested with Proteinase K. DNA was precipitated with isopropanol and 178 bound with nanobind magnetic disk (Bionano Genomics, USA). Bound UHMW DNA 179 was resuspended in the elution buffer and quantified with Qubit dsDNA assay kits 180 (ThermoFisher Scientific). DNA labeling was performed following manufacturer's 181 protocols (Bionano Prep Direct Label and Stain protocol). Direct Labeling Enzyme 1

(DLE-1) reactions were carried out using 750 ng of purified UHMW DNA. Labeled DNA
was loaded on Saphyr chips for imaging. The fluorescently labeled DNA molecules
were imaged sequentially across nanochannel arrays (Saphyr chip) on a Saphyr
instrument (Bionano Genomics Inc, USA). Effective genome coverage of greater than
100X was achieved for all samples. All samples also met the following QC metrics:
labelling density of ~15/100 kbp; filtered (>15kbp) N50 > 230 kbp; map rate > 70%.

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189 Genome analysis of the resulting data was performed using software solutions provided 190 by Bionano Genomics Inc. Briefly, automated optical genome mapping specific 191 pipelines consisting of Bionano Access v1.4.3 and Bionano Solve v. 3.6.1 were used for 192 data processing (BioNano Access Software User Guide). Hybrid scaffolding was 193 performed using Bionano's custom software program implementing the following steps: 194 1) generate in silico maps for sequence assembly; 2) align in silico sequence maps 195 against Bionano genome maps to identify and resolve potential conflicts in either data 196 set; 3) merge the non-conflicting maps into hybrid scaffolds; 4) align sequence maps to 197 the hybrid scaffolds; and 5) generate AGP and FASTA files for the scaffolds. Pairwise 198 comparisons of all DNA molecules were made to generate the initial consensus genome 199 maps (*.cmap). Genome maps were further refined and extended with best matching 200 molecules. Optical map statistics were generated using Bionano software producing the 201 Bionano Molecule Quality Report (MQR).

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The optical map N50 (including only maps >=150 kbp and minSites >= 9) was 0.2341 Mbp and the average label density (scaffolds >= 150 kbp) was 17.40/100 kbp. This

yielded an effective molecule coverage with optical mapping information of 125.38X.
The optical mapping analysis identified 84 conflicts with the prior Flye/Pilon scaffolds
and these initial scaffolds were broken at those 84 sites. The completed assembly was
submitted to NCBI and is available under accession <u>GCA_017639785.1</u>.

Gene annotation

NCBI performed gene annotation using RNA-Seq data from multiple tissues including
lung, trachea, brain, olfactory bulb and small intestine that are targets for SARS-CoV-2
infection (NCBI BioProject <u>PRJNA675865</u>) [19].

213 **Quality assessment**

To assess the quality of our assembly compared to the previous MesAur1.0 we used Quast v5.0.2 [25] together with MUMmer v3.23 [26]. These tools provided a detailed comparison between these assemblies. In addition, the Illumina reads from the original reference (NCBI SRA <u>SRR413408</u>) were mapped to our assembly and the MesAur1.0 reference using BWA v0.7.17 [27]. Quast was used to obtain discordant pair statistics.

We next used the software Benchmarking Universal Single-Copy Orthologs (BUSCO) v5.2.2 [28] to assess the quality of the genome assembly. BUSCO is based on the concept that single-copy orthologs should be highly conserved among closely related species. BUSCO performs gene annotation on an assembly and reports the number of gene models generated. BUSCO was performed using the OrthoDB v10 (odb10) release consisting of 12,692 genes shared across the superorder Euarchontoglires [29], the appropriate test for the Syrian hamster. In addition, FRCbam [30] was used to compute Feature Response Curves (FRCurve) from the alignment of Illumina reads to the assembled contigs. FRC v1.3.0 was employed to evaluate both assemblies, using default parameters. BCM_Maur_2 was further evaluated using paired end mappings of the Illumina reads that had been used for Pilon polishing (SRX10928323). MesAur1.0 was then similarly evaluated using paired end mappings of Illumina reads used for the MesAur1.0 assembly (SRR413408).

232 **Results**

233 The initial Flye assembly consisted of 2.38 Gbp of sequence across 6,741 scaffolds with 234 a scaffold N50 of 10.56 Mbp (Table 1). Pilon polishing of the Flye assembly had little 235 effect on these metrics, but significant improvements were obtained when Bionano 236 optical mapping results were used to improve scaffolding. As shown in Table 1, the 237 optical mapping step reduced the total number of scaffolds in the final assembly by 395 (5.9%) while increasing the N50 scaffold length by more than 8-fold to 85.18 Mbp. Our 238 239 experience in comparing read lengths and total yield per flow-cell indicates that the 240 optimal target size for fragmented DNA as input into Nanopore libraries and sequencing 241 is 15-20 kb, which regularly yields 80-90 Gb of sequence data.

Of the 12,692 BUSCO gene models, 90.58% were annotated as complete genes in the initial Flye assembly (**Table 2**). Pilon polishing of this Flye-alone assembly added another 682 genes annotated completely and increased this proportion to 95.95% of the BUSCO gene model dataset. Improvements in assembly scaffolding resulting from the Bionano optical mapping step together with Pilon error correction decreased the proportions of fragmented and missing BUSCO gene models in the new assembly to

248 0.82% and 3.21% respectively, also improvements over the MesAur1.0 assembly. This

249 advance translates to an additional 1189 complete BUSCO genes identified in the new

assembly compared to MesAur1.0.

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Table 1. Assembly statistics for BCM_Maur_2.0 versus the MesAur1.0 Syrian hamsterassembly

Parameter	MesAur1.0	Flye	Flye + Pilon	Flye + Pilon + Bionano (BCM_Maur_2.0)
Assembly length (bp)	2,504,908,775	2,381,258,546	2,383,228,608	2,457,062,007
Ungapped length (bp)	2,076,159,990	2,381,254,546	2,383,226,373	2,383,228,883
Number of scaffolds	21,483	6,741	6,741	6,346
N50 scaffold length (bp)	12,753,307	10,564,357	10,573,641	85,184,847
Number of contigs	237,699	6,781	6,779	7,057
N50 contig length (bp)	22,512	10,022,145	10,097,207	9,471,653

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- Table 2. BUSCO statistics for BCM_Maur_2.0 versus the MesAur1.0 Syrian hamsterassembly

	MesAur1.0	Flye	Flye + Pilon	Flye + Pilon + Bionano (BCM_Maur_2.0)
Complete ^a	86.60%	90.58%	95.95%	95.97%
Complete and single-copy	85.75%	89.27%	94.43%	94.49%
Complete and duplicated	0.85%	1.31%	1.52%	1.47%
Fragmented	4.59%	3.23%	0.85%	0.82%
Missing	8.81%	6.19%	3.20%	3.21%

^a12,692 gene models were included in this analysis

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Assembly Comparisons

We also performed additional comparisons between the two assemblies. As background, the karyotype of *M. auratus* is diploid 2n = 44, including 14 pairs of metacentric chromosomes, 3 pairs of telocentrics and 5 pairs of acrocentrics [31]. Illumina read k-mer analyses were performed to estimate the genome size using SGA

preqc [32] (2.57 Gbp) and Jellyfish [33] (2.90 Gbp). The total length of the 268 269 BCM_Maur_2.0 assembly is 2.46 Gbp compared to the previous version's 2.50 Gbp. 270 Despite having a similar total length, BCM Maur 2.0 shows an improved continuity with 271 a scaffold N50 that is 6.7 times greater than MesAur1.0 (Table 1); the L50 (i.e. the 272 number of contigs longer than or equal to the N50 length) of BCM Maur 2.0 is 22 273 compared to MesAur1.0's 121. The longest scaffold of BCM_Maur_2.0 (187 Mb) is 2.35 274 times larger than the longest scaffold from the previous assembly. N50 is calculated in 275 the context of the assembly size rather than the genome size, so the NG50 statistic was 276 used to directly compare the different assemblies. NG50 is the same as N50 except that 277 it reports the length of the contig at which the size-ordered contigs (longest to shortest) 278 collectively reaches 50% of the known or estimated genome size [34]. Figure 1 279 illustrates the improved cumulative contig sequence length for any given NG50 value 280 that is generated from the BCM Maur 2.0 assembly as compared to MesAur1.0, based 281 on the estimated genome size of 2.57 Gb calculated using SGA-preqc. The 282 BCM_Maur_2.0 assembly further improves the unresolved regions as measured by 283 nucleotide ambiguities (i.e. number of N's included in the final contigs). Approximately 284 17.11% of bases in MesAur1.0 were unresolved. BCM_Maur_2.0 reduces the number 285 of unresolved bases to 3.00%, with only very small gaps throughout the entire genome. 286 Figure 2 displays the overall increase in continuity of the BCM Maur 2.0 assembly with 287 longer contigs than the MesAur1.0 assembly and fewer short contigs. Finally, we 288 compared feature response curves for BCM_Maur_2.0 and MesAur1.0 using FRC^{Bam} [30]. FRC^{Bam} shows that our new assembly is substantially more accurate based on the 289 290 feature response approach (Supplementary Figure 1).

To establish the correctness of the structure and completeness of BCM_Maur_2.0, we also leveraged the Illumina short-reads that were published as part of the MesAur1.0 assembly project. When mapping the MesAur1.0 Illumina reads back to the MesAur1.0 reference, only 92.19% reads mapped successfully. When the same Illumina reads were instead mapped to BCM_Maur_2.0, 97.32% mapped successfully. When considering only properly paired reads, 75.76% and 87.62% mapped to MesAur1.0 and BCM_Maur_2.0, respectively.

Alignments between the current and previous Syrian hamster assemblies performed by NCBI [35] show that BCM_Maur_2.0 covers 98.95% of MesAur1.0 while MesAur1.0 only covers 86.67% of BCM_Maur_2.0. This together with the additional 307 Mbp of ungapped sequence in BCM_Maur_2.0 indicates that BCM_Maur_2.0 is a more complete representation of the Syrian hamster genome. The percent identity in the regions aligned between the two assemblies is 99.76%.

Transcript and Protein Alignments and Annotation Comparisons

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NCBI annotation of BCM_Maur_2.0 [35] with Syrian hamster transcript and protein data show this assembly to be of high quality. Transcript alignments of Syrian hamster RefSeq (n=273), Genbank (n=751), and EST (n=558) data to BCM_Maur_2.0 show 99.44% or more average percent identity and 98.88% or more average percent coverage. Alignments of these same transcript datasets to MesAur1.0 show 99.13% or more average percent identity and 93.49% or more average percent coverage. Alignments of RefSeq transcripts showed a similar average percent indels in the BCM_Maur_2.0 (0.10%) and MesAur1.0 (0.11%) assemblies. Protein alignments of Syrian hamster RefSeq (n=261) and Genbank (n=485) data to BCM_Maur_2.0 show 80.95% or more average percent identity and 89.18% or more average percent coverage. Alignments of these same protein datasets to MesAur1.0 show 80.57% or more average percent identity and 84.87% or more average percent coverage.

319 NCBI annotated 21,616 protein coding genes and 10,459 noncoding genes in 320 BCM Maur 2.0 compared to 20,495 protein coding genes and 4,168 noncoding genes 321 in MesAur1.0 [36]. Only 7% of gene annotations are identical between BCM Maur 2.0 322 and MesAur1.0, suggesting that a number of previous errors have been corrected, 323 though some differences are likely to be real differences between the animals used for 324 the different assemblies. Minor changes between BCM_Maur_2.0 and MesAur1.0 were 325 made in 46% of gene annotations and major changes were made in 15% of gene 326 annotations. We further note that, based on NCBI annotation feature counts, 327 BCM Maur 2.0 has only 33 RefSeq models that were filled using transcript sequence 328 to compensate for an assembly gap [35]. This is compared to 5,050 RefSeq models 329 similarly compensated in MesAur1.0.

330 Interferon type 1 alpha gene cluster

331 Given the importance of type I interferon responses during SARS-CoV-2 infection, we 332 next compared the interferon type I alpha gene cluster in the BCM_Maur_2.0 assembly 333 relative to this genomic region in the original MesAur1.0 assembly. The MesAur1.0 334 scaffold NW_004801649.1 includes annotations for four interferon type I alpha loci but 335 this genomic sequence is riddled with numerous gaps. Of these four candidate loci, 336 only LOC101824534 appears to contain a complete interferon alpha-12-like coding 337 sequence with the ability to encode a predicted protein (XP 005074343.1). The 338 LOC101824794 gene sequence can only encode a 162 amino acid protein due to a 5' 339 remaining pair of candidate truncation. The genes (LOC101836618 and 340 LOC101836898) appear to have aberrant transcript models that have fused putative 341 exons from neighboring loci. In mice and humans, the interferon alpha gene cluster is 342 flanked by single copy interferon beta 1 (Ifnb1) and interferon epsilon (Ifne) genes. 343 Although neither of these genes are present on the MesAur1.0 scaffold 344 NW 004801649.1, this assembly does contain a *lfne* gene on a short 2,408 bp contig 345 that is predicted to code for a protein of 192 amino acids. These observations 346 emphasize the need for an improved genomic assembly for Syrian hamsters given that 347 the interferon alpha gene cluster includes more than a dozen tightly linked functional 348 genes plus multiple pseudogenes in a wide variety of species including mice and 349 humans.

350 In the BCM Maur 2.0 assembly, the interferon type I alpha gene cluster is contained on 351 the NW 024429197.1 super scaffold that spans nearly 75 Mbp. Figure 3 illustrates this 352 genomic region in comparison with the interferon type 1 alpha regions of MesAur1.0 353 (NW 004801649.1) and well-characterized the C57BL/6J mouse assembly 354 (NC 000070.7). Fourteen predicted interferon type I alpha genes as well as five 355 presumptive pseudogenes lie within a span of 196 Kbp of the new Syrian hamster 356 assembly (Figure 3 and Supplemental Table 1). This genomic organization is quite 357 comparable to that observed in the mouse genome where there are also fourteen

358 functional interferon alpha genes and four pseudogenes. This hamster gene cluster is 359 flanked by *lfnb1* and *lfne* genes consistent with expectations from the mouse and other 360 species. The NCBI annotations characterize twelve of these genes as interferon alpha-361 (Supplemental Table 1). The remaining pair of functional genes 12-like 362 (LOC101824794 and LOC121144100) are listed as interferon alpha-9-like and they 363 encode shorter predicted proteins. The increased length of this genomic region in the 364 mouse assembly is largely due to the presence of the interferon zeta gene family (Ifnz, 365 Gm13271, Gm13272, etc.). This *lfnz* gene family appears to be absent in Syrian 366 hamsters since the closest matches to predicted hamster protein sequences are only 367 28% identical at the amino acid level. The interferon type I alpha gene cluster in the 368 BCM_Maur_2.0 assembly lies within more than 12 Mbp of contiguous genomic 369 sequence with the nearest flanking gaps located 2.66 Mbp proximal and 9.07 Mbp distal 370 to the *lfne* and *lfnb1* genes, respectively. The availability of a contiguous hamster 371 genomic sequence and associated transcriptional regulatory elements for this complex 372 immune gene region may be helpful for investigators who are interested in unravelling 373 mechanisms that control interferon expression during infections with SARS-CoV-2 as 374 well as challenges with other viral pathogens.

375 **Conclusions**

The improved Syrian hamster assembly and annotation described here will facilitate research into this important animal model for COVID-19. Specifically, reagents for studying immune responses in hamsters have lagged behind those available for laboratory mice. BCM_Maur_2.0 will facilitate the identification of cross-reactive reagents originally developed to study immunity in other species. Additionally, a more accurate genome assembly will improve the analyses of host responses to infection byenabling more accurate interpretation of RNA-seq experiments.

383 Relative to other recent assemblies that use a combination of long-read sequencing and 384 short-read polishing, this genome assembly and annotation compares very favorably. 385 The scaffold N50 of >85 Mbp is quite consistent with other long read assemblies. The 386 contig N50 and total number of scaffolds or contigs are likewise reasonable and 387 consistent with other similar mammalian reference genomes. The number of protein 388 coding genes identified is within the expected range, although additional attention will 389 likely be needed to resolve duplicated, repetitive gene loci, potentially leveraging recent 390 advances in ultralong read sequencing.

What additional genomic resources would be needed to make hamsters a better model for COVID-19? Deep long read transcriptome analysis of multiple tissues and ages would be the best next step, in order to define not just the genes expressed but the alternative splicing of genes across tissues and developmental stages. Also, long read RNA-seq of tissues following experimental challenge with SARS-CoV-2 and other viruses would facilitate improvements in the quality of antiviral gene models.

The availability of higher accuracy sequences should lead to the development of specific reagents for monitoring immune responses. For example, epitopes that are shared between hamsters and other rodents can be used to identify monoclonal antibody reagents for flow cytometry that are predicted to be cross-reactive. Additional reagent development will be enabled by creating synthetic versions of hamster proteins that can be used as immunogens to make hamster-specific antibodies.

403 One surprising motivation for this study is that Syrian hamsters, which were quickly 404 identified as a high value model for COVID-19, did not have a higher quality reference 405 genome at the start of the pandemic. While we worked guickly to generate this data and 406 make it available to the scientific community, better preparedness will be critical for 407 future unexpected epidemics. To this end, we would encourage investment in continued 408 refinement and improvement of reference genomes for all of the rodent, bat and 409 nonhuman primate models that are commonly used to study viruses in order to prevent 410 this situation from recurring in the future. Such an investment would also yield improved 411 genomic resources that would provide broad benefit to the entire scientific community.

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Availability of Supporting Data and Materials

416 The MesAur1.0 genome assembly is available in the NCBI database under BioProject 417 PRJNA77669 (GenBank accession GCA_000349665.1). The new BCM_Maur_2.0 418 genome assembly is available in the NCBI data repository under BioProject 419 PRJNA705675 (GenBank accession GCA 017639785.1). Oxford Nanopore 420 (SRX11206953) and Illumina (SRX10928323) sequencing data are available through 421 the NCBI SRA. The Bionano data are available from the BioProject page as NCBI 422 accession SUPPF_0000004259. The Illumina RNA-Seq data from multiple tissues 423 including lung, trachea, brain, olfactory bulb and small intestine are available under 424 NCBI BioProject PRJNA675865.

425

426 Additional Files

427 Supplementary Table 1. Predicted genes in the Interferon type 1 alpha cluster of the
428 BCM_Maur_2.0 assembly.

429 Abbreviations

430 ACE2: angiotensin-converting enzyme 2; BCM: Baylor College of Medicine; bp: base 431 pairs; BUSCO: Benchmarking Universal Single-Copy Orthologs; BWA: Burrows-432 Wheeler Aligner; COVID-19: coronavirus disease 2019; EST: expressed sequence tag; 433 FFPE: formalin-fixed, paraffin-embedded; Gbp: gigabase pairs; GC: guanine-cytosine; 434 IFN: interferon; kbp: kilobase pairs; Mbp: megabase pairs; MQR: Molecule Quality 435 Report; NCBI: National Center for Biotechnology Information; NEB: New England 436 BioLabs; ng: nanogram; ONT: Oxford Nanopore Technologies; PCR: polymerase chain 437 reaction; RBD: receptor-binding domain; RNA-Seq: RNA-sequencing; SARS-CoV-2: 438 severe acute respiratory syndrome coronavirus 2; STAT2: signal transducer and 439 activator of transcription factor 2; TMPRSS2: transmembrane protease serine 2

440 **Competing interests**

441 The authors declare that they have no competing interests.

442 Funding

This research was supported by contract HHSN272201600007C awarded to DHO from theNational Institute of Allergy and Infectious Diseases of the National Institutes of Health. The

445 content of this publication is solely the responsibility of the authors and does not446 necessarily represent the official views of the National Institutes of Health.

447 Authors' Contributions

448 R.A.H. performed genome assembly and quality assessment, data and metadata 449 submission, and contributed to manuscript preparation. F.S. and M.M. performed 450 assembly assessment and comparison analyses. T.M.P. and R.W.W. performed 451 transcript and annotation comparisons. D.H.O. managed experimental design and 452 oversight and coordinated manuscript preparation. H.D., Q.M. and Y.H. developed, 453 optimized and implemented protocols for ONT PromethION sequencing. M.R., D.M., J.A.K. and J.R. performed project and/or data management. R.A.H., D.H.O., D.T.L., 454 455 T.M.P., R.W.W., M.M., F.S. and J.R. wrote the manuscript. All authors approved the 456 manuscript.

457 Acknowledgements

We are extremely grateful to Dr. Tadashi Maemura for collecting the Syrian hamster tissues that were used for the sequence analyses described here. We also thank Dr. Benjamin tenOever for sharing Syrian hamster RNA-Seq datasets generated by his group prior to publication. And we also wish to thank two reviewers for their helpful comments.

464

Figure 1: Cumulative length and continuity comparison of MesAur1.0 and BCM_Maur_2.0. This summarizes the length of contigs/scaffolds across the assemblies. Given the length of contigs, the NG50 (mid x-axis) summarizes the sequence length of the shortest contig/scaffold at 50% of the total genome length. For genome length, the SGA preqc estimate of 2.57 Gbp was used.

Figure 2: Contig length and count comparison between BCM_Maur_2.0 and
MesAur1.0. Log length of contigs on the X axis and normalized count on the Y axis
comparing BCM_Maur_2.0 assembly and the previous assembly. Contigs from
BCM_Maur_2.0 are shown red and contigs for MesAur1.0 are shown in gray.

474 Figure 3: Comparison of interferon type 1 alpha gene cluster between MesAur1.0, 475 BCM_Maur_2.0 and GCRm39 mouse genome assembly. The genomic intervals 476 illustrated here are defined by the flanking interferon beta 1 and interferon epsilon 477 genes except for MesAur1.0 which does not include an interferon epsilon or beta 1 gene 478 in a continuous sequence with interferon type 1 alpha genes. White space within each 479 scaffold represents gaps in the MesAur1.0 assembly. Accession numbers for each 480 genomic sequence are indicated on the right with genomic coordinates for the extracted 481 intervals shown below their respective accession numbers. Predicted interferon type 1 482 alpha genes are highlighted in blue while putative pseudogenes are depicted with open 483 symbols and labelled below each assembly.

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