Obesity and metabolic dysfunction drive sex-associated differential disease profiles in hACE2-mice challenged with SARS-CoV-2

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Highlights
Transcriptomic analysis of infected lungs revealed unique sex-dependent differences

Obese female mice have high viral RNA burden and interferon production in the lung

Male mice have altered antibody and T cell response gene profiles after viral challenge

Metabolic dysfunction comorbidity can be studied in the hACE2 mouse model of COVID-19

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Obesity and metabolic dysfunction drive sex-associated differential disease profiles in hACE2-mice challenged with SARS-CoV-2

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SUMMARY
Severe outcomes from SARS-CoV-2 infection are highly associated with preexisting comorbid conditions like hypertension, diabetes, and obesity. We utilized the diet-induced obesity (DIO) model of metabolic dysfunction in K18-hACE2 transgenic mice to model obesity as a COVID-19 comorbidity. Female DIO, but not male DIO mice challenged with SARS-CoV-2 were observed to have shortened time to morbidity compared to controls. Increased susceptibility to SARS-CoV-2 in female DIO was associated with increased viral RNA burden and interferon production compared to males. Transcriptomic analysis of the lungs from all mouse cohorts revealed sex- and DIO-associated differential gene expression profiles. Male DIO mice after challenge had decreased expression of antibody-related genes compared to controls, suggesting antibody producing cell localization in the lung. Collectively, this study establishes a preclinical comorbidity model of COVID-19 in mice where we observed sex- and diet-specific responses that begin explaining the effects of obesity and metabolic disease on COVID-19 pathology.

INTRODUCTION
Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) continues to pose a worldwide epidemiological threat because of the emergence of novel variants with enhanced transmissibility and disease-causing capabilities. Although our understanding of the virus has increased dramatically since its emergence in 2020, questions regarding the mechanisms behind the heterogeneous nature of lethality in human subpopulations remain. Although many infections are asymptomatic or result in mild disease phenotypes, SARS-CoV-2 variants of concern (VOC) remain a considerable threat to at-risk populations where they increase host susceptibility to more severe, deadly infection (Sanyaolu et al., 2020; Djaharuddin et al., 2021; Ng et al., 2021). The most at-risk populations of concern in the COVID-19 pandemic have been suggested to include males, the elderly, pregnant individuals, and those with preexisting conditions such as obesity or metabolic disease which may cause them to be immunocompromised (Centers of Diseases and Control, 2021; Djaharuddin et al., 2021).

It has been shown that more hospitalizations and deaths from COVID-19 have occurred in men than women to-date (Capuano et al., 2020; Nguyen et al., 2021). This matches the widely accepted understanding that the immune response to pathogens is influenced by biological sex. The distribution of genes related to immunological function across the X and Y chromosomes contribute to a generally more robust immune response in females that is dampened in males, and hormonal differences are thought to boost cytokine production in females compared to males (Klein and Flanagan, 2016; Bereshchenko et al., 2018; Takahashi et al., 2020). The exact mechanisms and physiological states which predispose males to morbidity and mortality in COVID-19 are unclear. One early report pointed out that expression of the angiotensin converting enzyme 2 (ACE2) receptor is greater in men than women, which, with hormones and other behavioral tendencies like greater incidence of tobacco and alcohol consumption, may increase susceptibility to virus (Bwire, 2020; Gemmati et al., 2020). Observations that sex-differences exist in diseases contributed to a push in preclinical modeling to incorporate the evaluation of pathophysiology in both sexes, attempting to avoid the potential limitations that just one might provide (Buoncervello et al., 2017; Miller et al., 2017). Although some gaps in knowledge still remain, increased susceptibility to infections in males and...
autoimmunity in females have been modeled extensively in preclinical and clinical settings alike (Klein and Flanagan, 2016).

Obesity is another prominent comorbidity in infectious disease that has risen to “epidemic status” in the United States and other countries, occurring at an incidence above 35% in many states (Author Anonymous, 2018; Author Anonymous, n.d.). Obesity is often associated with the subsequent development of comorbidities akin to those which predispose to severe COVID-19 disease outcomes such as type 2 diabetes (T2DM) (Ekoru et al., 2019; Kulcsar et al., 2019; Nowakowska et al., 2019; Bornstein et al., 2020; Pal et al., 2021; Patone et al., 2021). Obesity is often concurrent to metabolic syndrome, a condition marked by “central” obesity with high adiposity, insulin resistance, and high blood glucose (Eckel et al., 2005; Huang, 2009; Paragh et al., 2014). Adiposity (increases in adipose tissue distribution) is accompanied by enlargement of individual adipocytes which become stressed and hypoxic at the cellular level. Chronic exposure to stress signals, hypoxic conditions, and oxidative stress causes adipocytes to produce cytokines like CRP, TNF-alpha, and IL-6 in addition to their healthy secretions intended to maintain homeostasis (Trayhurn, 2013). The resulting recruitment to and activation of proinflammatory-type macrophages cells within the adipose tissue raises basal inflammation systemically in a phenomenon known as “metabolic inflammation” (Gordon, 2003; Ouchi et al., 2011; Park et al., 2014). This inflammation contributes to metabolic dysfunction like insulin resistance in obese persons but also increases susceptibility to pathogens through cellular interactions that are deleterious over time (11-13). COVID-19 also encompasses a complex inflammatory milieu where delayed interferon responses allow virus to continue replicating, meanwhile inciting the proinflammatory actions of neutrophils and lymphocytes that drive the disease-characteristic “cytokine storm” (17). The contributions of preexisting metabolic dysfunction to this aberrant inflammatory response have yet to be mechanistically defined. In laboratory mice, severe outcomes for obese individuals during infection have been modeled for numerous agents including influenza, West Nile virus, and even the parasite Leishmania (Smith et al., 2007; Martins et al., 2020; Geerling et al., 2021). Similar studies have implicated preclinical diabetes models as comorbid conditions, but no extensive work has been done to define the comorbid outcomes of SARS-CoV-2 (Zhang et al., 2021).

T2DM is estimated to be the second most common comorbidity in patients with severe COVID-19, resulting in a 2-3 times greater likelihood to succumb compared to healthy persons (Landstra and de Koning, 2021). Over 460 million people worldwide have been diagnosed with diabetes mellitus (either T1DM or T2DM) and greater than 60% of type 2 diabetics are also clinically characterized as obese (Chan et al., 2020). SARS-CoV-2 infection combined with the metabolic dysfunction in T2DM is associated with an increased risk of pneumonia requiring ventilation, ICU admission, and “long COVID” (Unnikrishnan and Misra, 2021). Although COVID-19 vaccine implementation around the world has been a positive effort for protecting vulnerable populations, T2DM has been linked to reduced COVID-19 vaccine efficacy, with lower IgG and neutralizing antibody production (Ali et al., 2021; Pal et al., 2021). Because of their pre-disposed risk to severe outcomes, defining the immunological profile of patients with T2DM is a necessary step toward solving vaccine-established protection discrepancies.

Most of our knowledge regarding the positive correlation between metabolic dysfunction and SARS-CoV-2 severity comes from retrospective clinical studies, where it becomes impossible to discern the molecular mechanisms that governed severe outcomes (Rawshani et al., 2018, 2021; Barron et al., 2020; Cario et al., 2020; Chen et al., 2020; Holman et al., 2020; Muniyappa and Gubbi, 2020; Zhu et al., 2020; Deme-terco-Berggren et al., 2022). To identify and characterize the mechanisms behind increased infection and severity we developed a preclinical model of disease comorbidities by combining the K18-hACE2 transgenic mouse model and diet-induced obesity (DIO) model where metabolic disease is confirmed by the development of T2DM (Yang et al., 2007; Della Vedova et al., 2016; Avtanski et al., 2019; Bao et al., 2020; Golden et al., 2020; Jiang et al., 2020; Moreau et al., 2020; Muñoz-Fontela et al., 2020; Sun et al., 2020; Winkler et al., 2020; Yinda et al., 2021; Dong et al., 2021). K18-hACE2 mice were subjected to a 60% fat “high-fat,” diet for 8 weeks causing measurable obesity, metabolic dysfunction, and hyperglycemia. Normal diet and DIO males and females were either mock challenged or challenged with the Alpha variant of SARS-CoV-2. Female DIO mice were observed to have shorter time to morbidity than normal diet mice and exhibited higher viral RNA burden at the time of terminal euthanasia, indicating sex differences in disease pathology. RNAseq analysis was used to characterize the transcriptional responses of the lungs in all experimental cohorts. Systems based analysis revealed DIO mice have unique responses to SARS-CoV-2 challenge including lack of antibody-related gene diversity compared to normal diet K18-hACE2 mice in
addition to differential gene expression profiles. Our data illustrate how metabolic dysfunction can enhance COVID-19 disease and suggest a synergism between hyperglycemia and gene expression profile changes. This data helps to link molecular alterations with infection severity, thus constructing a profile of potential therapeutic targets for the treatment and prevention of death by COVID-19 illness.

RESULTS

K18-hACE2 mice develop obesity, metabolic dysfunction, and hyperglycemia because of high fat diet

The COVID-19 pandemic has illustrated that humans respond to infection with a great deal of heterogeneity. SARS-CoV-2 infection may be lethal in some patients but causes mild or asymptomatic disease in others. Because of this range in infection severity, it is important to understand comorbid conditions to develop therapeutic interventions that support the most at-risk populations. To understand the impact of metabolic dysfunction seen in obesity and type 2 diabetes on the outcomes of viral infection, we utilized a diet-induced model of obesity with K18-hACE2 transgenic mice (Figure 1A). Compared to normal chow mice, DIO mice gained 25% or 37% bodyweight, in females and males respectively (Figure 1B). Intraperitoneal glucose tolerance testing (IPGTT) was performed at week 6 of the diet intervention to measure glucose clearance and fasting hyperglycemia in male DIO mice (Figure 1C). One-way ANOVA with Tukey’s multiple comparisons test: p<0.0001 male ND versus male DIO; P=ns female ND versus female DIO. (D) One-way ANOVA with Tukey’s multiple comparisons test: p<0.0001 male ND versus male DIO; P=ns female ND versus female DIO; error bars represent mean with SD.

Figure 1. Development of the diet induced obesity (DIO) K18-hACE2 transgenic mouse model

(A) Experimental design: Male and female K18-hACE2 transgenic mice were provided a 60% fat “high fat” diet for 8 weeks to induce glucose impairment consistent with the diet induced obesity (DIO) model then intranasally challenged with 10^3 PFU SARS-CoV-2 Alpha variant. Control groups that were age- and sex-matched remained on normal diet for the entirety of the experiment. (n = 10 diet DIO and 10 normal diet (ND) males; 10 DIO and 10 normal diet females).

(B) Mice in each group were weighed weekly to measure the progression of obesity over 8 weeks (* indicates significance; Multiple unpaired t tests: p<0.000030 female ND versus female DIO; p<0.000001 male ND versus male DIO); error bars represent SD.

(C) Intraperitoneal glucose tolerance testing (IPGTT) was performed at week 6 of the diet intervention to measure glucose clearance and fasting hyperglycemia in male DIO mice (C) * indicates significance; Multiple unpaired t tests: p<0.000001 male ND versus male DIO; P=ns female ND versus female DIO. (D) One-way ANOVA with Tukey’s multiple comparisons test: p<0.0001 male ND versus male DIO; P=ns female ND versus female DIO; error bars represent mean with SD.
female mice presented with non-significant low to mild impairment (Figures 1C and 1D). Despite this observation, DIO females had significantly higher fasting blood glucose levels compared to normal diet females \( (p = 0.0488) \) (Figure S1). The DIO model has been previously used to study the effects of obesity and type 2 diabetes in mice, and our data here suggest that K18-hACE2-mice on the DIO diet do develop metabolic dysfunction (Della Vedova et al., 2016; Avtanski et al., 2019).

DIO shortens the time to morbidity in lethal SARS-CoV-2 challenge

The Alpha variant (strain B.1.1.7) of SARS-CoV-2 emerged in the UK in the spring of 2021 and moved rapidly across the globe. Alpha was observed to have enhanced virulence compared to ancestral strains of the virus in K18-hACE2-mice and other preclinical animal models (Bayarri-Olmos et al., 2021; Mok et al., 2021; O'Donnell et al., 2021; Radvak et al., 2021; Rosenke et al., 2021; Cochin et al., 2022; T. Y. Wong et al., 2022). Because of the significance of Alpha variant’s dominance among circulating strains in humans, we used this VOC to study the effects of DIO and type 2 diabetes in the K18-hACE2 mouse model of SARS-CoV-2 infection. Before the experiment, optimization of viral challenge dose was performed to determine an appropriate dose for achieving symptomatic disease (Figures 2A and 2B). The Alpha strain was intranasally administered to control K18-hACE2 mice at 10^3, 10^4, or 10^5 PFU per dose. Using a previously established disease scoring system (Lee et al., 2022; T. Y. T. Wong et al., 2022a), we observed that the 10^3 PFU dose caused lethality but postponed the time to morbidity compared to higher doses (Figure 2A) and caused disease phenotypes that increase in severity over time (Figures 2B). DIO and control diet mice were therefore challenged with 10^3 PFU of the Alpha variant to evaluate the effects of comorbidity on SARS-CoV-2 disease outcome. Male and female mice were utilized for DIO challenge studies to account for sex-based predispositions to disease severity (Kadioglu et al., 2011; Klein and Flanagan, 2013).
DIO shortened the time to morbidity following SARS-CoV-2 challenge in female DIO mice with 0% survival at 7 days after challenge (median time to death = 6 days) compared to only 20% survival in female controls (median time to death = 8 days) (Figure 2C). DIO and normal-diet male mice responded to challenge in a similar manner, with no statistical differences in survival (median time to death = 6 days in both groups). Daily disease scoring over the post-challenge period trended in a similar pattern in DIO and control challenged mice with no sex-dependent differences (Figures 2E and 2F). Collectively, these data suggest that the DIO condition has a greater impact on survival in female mice suggesting that sex specific responses may impact COVID-19 host pathogenesis.

Obese female mice experience greater viral RNA burden in the lungs

To begin identifying the factors that may contribute to the changes in survival that were observed in DIO mice, we next investigated differences in viral burden measured by qRT-PCR analysis of lung tissue for nucleocapsid transcript copy number. Total lung RNA was isolated from mice at euthanasia at their respective humane endpoints. Viral RNA burden was found to be higher in the lungs of DIO female mice than in female normal diet controls and no difference was seen in the viral RNA burden of males (Figure 3A). Normal diet female mice were observed to have approximately 1 million copies of nucleocapsid RNA transcripts on average per lung lobe whereas DIO females have 100-fold more copies, suggesting DIO enhances viral burden in females (Figure 3A). The enhanced viral burden, per PCR analysis, suggested that the differences in time to morbidity and survival in females may be related to viral burden or inability to clear virus.

Transcriptomic analysis of viral RNA confirms females have increased viral RNA due to DIO condition

As a secondary method of evaluating viral burden at morbidity, total lung RNA was used to perform bulk RNAseq analysis to measure the number of virus gene transcripts per total tissue RNA. Viral RNA reads
were mapped to the SARS-CoV-2 reference genome and represented per 50M illumina reads obtained per sample. Total viral reads mirrored the nucleocapsid qRT-PCR analysis (Figure 3B). A 100-fold increase in viral RNA was also observed for DIO females compared to normal females (Figure 3B). To identify correlations between viral burden and morbidity, total viral reads were plotted against the day post-challenge that humane euthanasia occurred (Figure 3C) or against disease score (Figure 3D). High viral reads corresponding to the mortality window 6 days post-challenge were observed for DIO mice where normal diet mice which survived longer underwent euthanasia (because of disease score or planned experimental endpoint) at lower viral burdens. Disease scoring comparisons had greater variance, but DIO mice trended toward having higher viral reads in the lung with higher disease scores (Figure 3D).

Transcriptomic analysis of mouse gene expression profiles unique to metabolic dysfunction

Bulk RNAseq analysis of infected mouse tissues allows for simultaneous pathogen and host transcriptomic analysis (Westermann and Vogel, 2018). We have previously used similar techniques to characterize mouse and bacterial gene expression during infection (Damron et al., 2016; Boehm et al., 2019; Wong et al., 2019). After evaluation of the viral transcriptomics of SARS-CoV-2 challenge in DIO mice, lung tissue RNA from non-challenged, challenged, DIO, or normal diet mice was used to characterize the host transcriptomic responses to SARS-CoV-2. Basic gene expression profiles of biological replicates in the experimental groups were compared using principal component analysis. The transcriptional profiles showed distinct patterns of gene expression between challenge and no challenge groups. Of interest, we observed a higher overlap between DIO and normal-diet in males and female mice than between challenged and non-challenged mice regardless of diet (Figures 4A and 4B). This suggested that challenge with SARS-CoV-2 has a greater impact on the lung transcriptome than diet in K18-hACE2 mice. Separation of the gene profiles into activated and repressed expression bins allowed for visualization of sex-driven differences. DIO induction increased the number of uniquely activated and repressed genes on viral challenge in females but decreased it for males (Figure 4C). Male DIO mice had smaller unique transcriptional profiles (589 DIO male-specific genes) whereas female DIO mice had a much larger transcriptional response (1974 DIO female-specific genes). When the pools of activated genes (p< 0.05) from each experimental group were compared using a Venn diagram, the unique gene profiles stemming from sex or DIO induction could be appreciated (Figure 4D). Although a core set of 835 genes were found to be activated in all SARS-CoV-2 challenged mice, DIO led to unique transcriptional profiles with differential expression of 765 unique genes in DIO females and 415 unique genes in DIO males. These data suggest that DIO and sex both influence the response to SARS-CoV-2 challenge.

Ingenuity pathway analysis identifies unique canonical pathway matches from metabolic dysfunction and infection

RNAseq analysis showed that the added variables of sex and preexisting comorbidities change the host transcriptional response to SARS-CoV-2 challenge by changing the unique differential gene expression profiles of K18-hACE2 mice. Gene expression fold change data comparing the profiles of each experimental group to no-challenge normal-diet control mice were evaluated by Ingenuity Pathway Analysis (IPA) to identify affected canonical pathways suggested by gene expression. The pathways that were associated with greater positive z-scores in the female DIO challenge datasets implicated activated inflammatory signaling pathways and immune cell activation that were not supported by the expression profiles of DIO no-challenge mice (Figure 5). Male mice’s gene expression profiles had altogether similar canonical pathway associations to females. The addition of the DIO condition to challenge groups augmented but did not significantly vary the z-scores of many pathways. Of note, we observed a decreased association of genes within the T cell receptor signaling pathway in male DIO mice. T cell responses are a major contributor to the antiviral response and are heavily implicated in the host response to COVID-19. The magnitude and polyfunctionality of the T cell response in severe cases of COVID-19 is a predictor of outcome as well as the memory response that is protective against reinfection (Peng et al., 2020; Neidleman et al., 2021). Dysregulation of the T cell response early on because of comorbidities may be partially responsible for disease outcome.

We continued our investigation of differential gene expression profiles by narrowing in on specific IPA pathways. To gain preliminary insights into the changes in T cell responses within the different treatment groups, a heatmap of fold changes (compared to sex-matched normal-diet no-challenge groups) in the expression of genes within the T Cell Receptor Signaling pathway was generated. Of interest, male DIO
mice had higher expression of numerous T cell-related signaling genes, including many that code for the T cell receptor alpha variable region, without SARS-COV-2 challenge that were low in the male DIO challenge group and both groups of female mice (Figure 6A). Genes like RelA, which encodes the p65 transcription factor for NF-κB signaling (Ronin et al., 2019), and MAPK13, which encodes the proinflammatory p38 MAP kinase (Salvador et al., 2005) were upregulated in both male and female DIO challenge groups compared to their normal-diet counterparts. However, genes like PDK1 (Sun et al., 2021) and CARD11 (Altin et al., 2011) which support T cell proliferation were high in both normal-diet challenge mice, and comparatively low in both DIO challenge groups. These data suggest DIO resulted in changes in T cell activation.

Another pathway of interest was the coronavirus pathway, which is comprised of known biomarkers that are either activated or repressed during SARS-CoV-2 infection (Coronavirus Network Explorer, n.d.). Contrary to the most differential T cell related gene expression occurring in no-challenge DIO males, a large number of inflammatory genes appeared to be up-regulated in challenged DIO females yet repressed in normal-diet females (Figure 6B). NLRP3, encoding for the antiviral inflammasome, was up in DIO females after challenge compared to all other groups, as were genes for the proinflammatory mediators STAT3 and CCL2 (Lai et al., 2017; Chang et al., 2018). It is possible that the gene expression differences that are associated
with these pathways are directly involved in the impaired viral clearance and affected disease pathogenesis experienced because of the DIO condition.

**Metabolic dysfunction in female K18-hACE2 mice heightens host antiviral response profile**

To further refine our RNAseq data analysis, Geno Ontology (GO) term analysis was performed to identify biological processes that were affected in our experiment based on the lung tissues’ transcriptional profiles. Although DIO induction and sex resulted in different lists of suggested GO terms, a conservative list of terms was present in each analysis that was related to activation of the immune response in viral infection. The terms were graphed with their corresponding enrichment ratio to compare their relevance in the genetic profiles of each experimental condition (Figure 7A). The highest enrichment ratios were seen for the GO terms related to interferon response and were increased in the female DIO challenge sets compared to others. GO terms related to antigen processing and presentation pathways appeared to be absent in both DIO groups. To measure the inflammatory response to virus, the levels of innate cytokines and proinflammatory mediators were measured in serum and lung supernatant collected at euthanasia (Figures S2 and S3). The most striking differences in cytokine production were noted for IFN-γ in the lung (Figure 7B). In females, DIO induction caused an increase in lung supernatant concentrations of IFN-γ compared with normal diet female challenge mice (p=0.0010) and DIO male challenge mice (p=0.0234) (Figure 7B). No difference was found in the concentrations of

Figure 5. Ingenuity Pathway Analysis of differential gene expression profiles

Fold change values of genes in experimental groups compared to no-challenge were inputted to Ingenuity Pathway Analysis software and sorted into canonical pathways. The highest and lowest 20 canonical pathways by Z score in the female DIO challenge group are shown.
serum IFN-γ across the groups (Figure S3). DIO induction in males did not result in a significant change in IFN-γ production. This is an important finding because IFN-γ production is triggered during SARS-CoV2 infection and is essential for both viral clearance and resolution of the infection (Lee and Ashkar, 2018). Together, these data suggest that changes in the immune response triggered by T2DM/obesity in female mice may hinder positive disease outcomes. As shown above, challenged DIO females had 100-fold higher viral RNA burden in the lung (Figure 3A) and we speculate that the presence of higher levels of viral particles in the lung of DIO females is associated with the increased interferon response observed here.

**Sex and metabolic dysfunction influence antibody production**

In addition to T cell and interferon response, antibodies also play an important role in the immune response against SARS-CoV-2. RNAseq reads from lung RNA were analyzed to quantify antibody-related gene expression and gain insights into the effects of diet and sex on the humoral response. Overall, a trend was appreciable where antibody genes were highest expressed by male mice challenged with SARS-CoV-2 when reads were mapped and visualized by heatmap (Figure 8A). In male challenged mice, the diversity of upregulated Igh and Igk genes was greater than in the other groups, suggesting: (1) the presence of B cells in the lung, and (2) the unique activation of these B cells in male normal-diet mice after challenge compared to DIO males and both female groups. To determine if changes observed at the mRNA expression level translated into variation in protein antibody responses, quantification of anti-SARS-CoV-2 RBD and anti-nucleocapsid IgG and IgM in serum was performed via ELISA to measure virus-specific antibodies. At this early point after the challenge, only DIO challenged females produced significant levels of anti-RBD IgM antibodies compared to non-challenged female mice. We did not observe statistically significant differences in anti-nucleocapsid IgM, anti-RBD IgG or anti-nucleocapsid IgG between DIO and normal diet groups, likely because of the fact that titers were measured 11 or fewer days post-challenge. Altogether, lung mRNA and circulating serological immunoglobulin data suggest that the humoral response generated by males and females in response to SARS-CoV2 is different, with the female response characterized by higher levels of circulating IgM and the male response characterized by B cell activation and differentiation in the lung.

**DISCUSSION**

Despite vaccines, antibodies, and small molecule therapeutics, SARS-CoV-2 continues to infect individuals driving continuation of the COVID-19 pandemic. Owing to the heterogeneous nature of COVID-19 cases, it is important to consider the host factors and responses that predispose to severe infection or death. Obesity and type 2 diabetes (T2DM) have long been appreciated as comorbidities for infectious diseases, yet very little controlled experimental data has illuminated how these comorbidities interplay with COVID-19. In addition, sex-specific responses to COVID-19 are obvious in the diversity of disease manifestation (Viveiros et al., 2021). To begin the effort to explore these conditions in preclinical models, we aimed to develop a comorbidity model for COVID-19 based on the previously implicated model of diet induced obesity and T2DM (Figure 1). Utilizing our model, we designed experiments to evaluate sex and DIO as central variables. We utilized the Alpha variant of SARS-CoV-2 which we previously identified as having high virulence compared to ancestral strains (Wong et al., 2022a). K18-hACE2 mice with DIO showed that disease as well as morbidity and mortality were affected very little by the addition of DIO in males; however, the DIO condition greatly affected females (Figure 2) and resulted in 100-fold higher viral RNA burden in the lung (Figure 3). To characterize the augmented host response to viral challenge we utilized RNAseq analysis of the lung tissue to analyze the pathogen-specific airway responses to the presence of virus (Figure 4). Pathway analysis using GO term and Ingenuity Pathway analyses illuminated more sex- and diet-specific responses. Notably, it appeared that DIO non-challenged male mice develop preexisting T cell gene expression signatures in the lungs suggestive of T cell infiltration that is decreased after viral challenge (Figure 6A). DIO seemed to hinder B cell responses in the lungs of challenged male mice (Figures 6, 7, and 8) indicating that although DIO did not affect the development of morbidity in males, it did alter the host response to infection. In female mice, distinctive gene expression profiles were observed between normal diet and DIO mice. Specifically, challenged female mice have low gene expression profiles corresponding to IPA’s Coronavirus Pathogenesis Pathway, whereas the
addition of the DIO condition enhances many of the pathway’s genes (Figure 6). Our data suggest that female K18-hACE2-mice on normal diets have lower viral RNA burden and decreased inflammatory responses, and DIO impairs the clearance of virus in the lung which results in their enhanced morbidity (Figures 2 and 3). The DIO condition appeared to affect female mice more than male mice following SARS-CoV-2 challenge; however, differences were appreciable in males as well. Collectively, these data begin to shed light on the effects of DIO on COVID-19.

To the best of our knowledge this is the first study in K18-hACE2-mice to evaluate DIO and sex utilizing transcriptomic analysis to better understand SARS-CoV-2 VOC-specific host responses. A previous study utilized a mouse-adapted strain of SARS-CoV-2 to challenge DIO C57Bl6 mice to measure the protective efficacy of human convalescent serum treatment (Rathnasinghe et al., 2020, 2021). One correspondence describes a small study where DIO mice were challenged with SARS-CoV-2 and the authors reported increased lung pathology and interferon responses (Zhang et al., 2021). However, the study did not utilize sex comparisons nor did it analyze transcriptomic responses. The high-fat high-carbohydrate diet has been used to evaluate the combined effects of the “Western Diet” and COVID-19 disease in Syrian hamsters (Port et al., 2021). Comparable to what we observed in mice, Western Diet-affected hamsters had increased weight loss, lung pathology, and delayed viral clearance after challenge. Our study does have some caveats that warrant discussion. In our experiment, we only evaluated one SARS-CoV-2 challenge strain, Alpha, and there have now been three VOC strain surges (Beta, Delta, and Omicron) since Alpha was dominantly circulating. In additional studies since then, we have observed enhanced airway inflammation because of challenge with the Delta variant (Lee et al., 2022). We anticipate that different strains would result in variable host responses to what were identified using Alpha; however, additional studies will need to be performed. Another caveat is that only one challenge dose was evaluated (1,000 PFU). If lower or higher challenge doses were to be studied, we would expect to have either shorter or longer time to morbidity during which host response profiles may further develop or remain hidden because of the disease timeline. Finally, our study focused on defining transcriptomic responses to characterize the altered host responses to SARS-CoV-2 challenge. We did not analyze specific cell...
populations through cell isolation and flow cytometry, nor did we evaluate potential mechanisms responsible for this comorbidity.

This model has some limitations for the evaluation of human disease comorbidities and viral pathogenesis. The K18-hACE2 transgenic mouse was generated to act as a lethal model of SARS-CoV-2 challenge, and expresses human ACE2 in addition to mouse ACE2 (Moreau et al., 2020; Muñoz-Fontela et al., 2020; Oladunni et al., 2020; R et al., 2020; Winkler et al., 2020; Dong et al., 2021; Yinda et al., 2021). The human ACE2 gene is incorporated into the mouse genome under the human cytokeratin-18 promoter and is thus expressed across tissues and organs in a pattern unlike humans. This leads to greater tissue tropism in the mouse and a propensity for higher viral replication in the brain compared to humans, and detection of viral RNAs in the kidney, spleen, and gastrointestinal tract (Oladunni et al., 2020; Winkler et al., 2020; Kumari et al., 2021; Seehusen et al., 2022). The transgenic incorporation of hACE2 universally also detracts from the model’s ability to model comorbidities.
conditions that may alter ACE2 expression on their own. ACE2 is expressed in adipose tissues, leading to higher ACE2 expression in obesity (Emilsson et al., 2021; Gómez-zorita et al., 2021; Sarver and Wong, 2021). In our 8-week DIO mouse model, hACE2 expression in the lung did not significantly vary between the experimental groups according to RNA sequencing (data not shown). Some additional limitations of the DIO model include a level of sex-based protection from metabolic dysfunction and disease in female mice (Pettersson et al., 2012; Casimiro et al., 2021). At 6 weeks, our study’s female DIO K18-hACE2 mice were not as insulin-resistant as males according to IPGTT tests, and some studies suggest that females require a longer period of time on the high fat diet to reach metabolic dysfunction. Despite this, obese female mice in our study achieved hyperglycemia and weight gain as well as displayed worse disease outcomes than obese male mice and female controls, suggesting that some level of comorbid condition had been reached. In future studies by our lab using both female mice and the DIO model, the high fat diet will be utilized for longer periods of time with additional IPGTTs to assess the development of the T2DM-like phenotype.

SARS-CoV-2 infection in humans is generally heterogeneous in symptomology, however, increased susceptibility to severe infection requiring hospitalization are common across patients with T2DM metabolic disease and increased adiposity (obesity) (Ando et al., 2021). The role of elevated glucose and fat accumulation downstream of metabolic dysfunction has been shown in other settings to disturb cellular signaling cascades, promote cytokine synthesis and secretion (leading to hyperinflammation), and increase oxidative stress through enhancement of reactive oxygen species (Giacco and Brownlee, 2010). However, the mechanism by which elevated glucose and adiposity enhance the severity of SAR-CoV-2 infection is still unclear. As demonstrated in our findings, a proinflammatory signature exists basally in T2D hosts and is even further activated in the host response to infection supporting the hypothesis that the preexisting inflammatory state of patients with T2DM and/or obesity is a factor that contributes to the severity of infection and drives an apparent predisposition to negative disease outcomes (Roberts et al., 2021). Our data, with human RNA-Seq analysis from T2DM samples, demonstrates altered immune system activation in response to infection through changes in the antibody response. Human enrichment and GO term analysis in our mouse model compared with lung epithelium from T2DM human patients shows TNF and IL-17 signaling to be highly enriched with several other genes that are involved in the antibody response (Islam et al., 2021). We also saw that T cell populations were augmented across our study groups with male DIO challenged mice demonstrating the most noticeable decrease in T cell subtypes. In severe human COVID-19 cases, it has been demonstrated that a characteristic severe phenotype, lymphocytopenia, is likely associated with CD4+ and CD8+ T cell exhaustion (Diao et al., 2020; Mazzoni et al., 2020; Tan et al., 2020; Kusnadi et al., 2021; Rha and Shin, 2021; Xiang et al., 2021). This may be another contributor toward SARS-CoV-2’s severity in T2DM. Research surrounding T2DM and obesity pathophysiology has implicated hyperinflammatory macrophages as central agonists in the basal inflammatory state. Melvin et al., using critically ill COVID-19 patient sera, defined the role of macrophages in diabetic COVID-19, showing that coronavirus infection leads to transcription of certain enzymes in the cells that increase their proinflammatory profile including their production of IFN-γ (Melvin et al., 2021). This same phenomenon, while not yet observed in mice, ties into our findings of increased IFN-γ, albeit sex-dependent. Lastly, our RNA-Seq data in normal diet challenge mice compared with normal diet infected patient samples also demonstrates overlap in gene expression profiles relating to a heightened interferon response and cytokine storm which has been previously characterized in humans, highlighting the relevance and use of the K18-hACE2 model (Daamen et al., 2021; Jain et al., 2021). We believe that our model demonstrates some of the pathology that has been seen or suspected in human cohorts including changes in B and T cell signatures and an exaggerated proinflammatory cytokine profile (summarized in Figure 9). Unfortunately, there is still limited data on whole lung tissue samples from diabetic patients to make direct comparisons. Published reports from postmortem lung tissue genomic analysis are often non-separated based on disease presence and patient history is not described (Bass et al., 2021). However, these overlaps in non-comorbid human clinical observations and mouse data are encouraging as modeling SAR-CoV-2 in the laboratory setting is critical for understanding the molecular mechanisms at work.

The T2DM-COVID-19 mouse model has allowed us to observe responses to SARS-CoV-2 challenge augmented by both obesity and sex. It is apparent that DIO affects female mice and enhances viral virulence; however, it is not clear how to best ameliorate this issue using therapeutic interventions. In an earlier study (data unpublished), we evaluated treatment with Baricitinib, a JAK inhibitor that could dampen inflammation caused by SARS-CoV-2. We hypothesized that decreasing inflammation
would improve survival outcomes of SARS-CoV-2-challenged mice; however, the drug did not improve survival of SARS-CoV-2 challenged mice (data not shown). This data leads us to believe that there are still molecular events that underpin overall inflammation as well as a dynamic viral clearance timeline that need to be adjusted to improve protection. Comparing that study with our DIO model, it is clear we need to profile the cell populations that respond to each phase of viral infiltration and identify if they are ineffective at stopping disease progression. Furthermore, we propose that this DIO model can be used to evaluate differences in vaccine-induced immunity among comorbid groups which is rarely evaluated. Future work in our lab will be performed to further define the cellular mechanisms at work in this model with single cell RNA sequencing and timepoint analysis to characterize the evolving host response after challenge. We plan to continue our utilization of the novel DIO-COVID-19 mouse model to uncover therapeutic strategies to improve disease and survival outcomes in diverse persons infected with SARS-CoV-2.

STAR METHODS

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AUTHOR CONTRIBUTIONS
Studies were designed by F.H.D., H.A.C., K.S.L., B.P.R., J.R.B., and A.M.H. All authors contributed to the execution of the studies. M.T.W. and I.M. prepared and provided titred viral stocks of SARS-CoV-2 for challenge. Animal health checks, necropsy, and tissue processing were performed by F.H.D., T.Y.W., B.P.R., K.S.L., J.R.B., O.A.M., A.M.H., and H.A.C. Viral RNA qPCR was performed by H.A.C. and O.A.M. Serological analysis was executed by K.S.L., B.P.R, and N.A.R. Luminex cytokine assays were completed by BPR. Data was analyzed by K.S.L., B.P.R, H.A.C., and F.H.D. All authors contributed to the writing and revision of this manuscript.

DECLARATION OF INTERESTS
The authors declare no competing interests.

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Goat anti-Mouse IgG (H + L) Secondary Antibody [HRP] | Novus Biologicals | Cat# NBP1-75130, RRID:AB_11012784 |
| Goat anti-Mouse IgM Heavy Chain Secondary Antibody [HRP] | Novus Biologicals | Cat # NB7497, RRID:AB_524863 |
| **Bacterial and virus strains** |        |            |
| SARS-CoV-2 Alpha variant (hCoV19/England/204820464/2020) | BEI Resources | NR-54000 |
| **Chemicals, peptides, and recombinant proteins** |        |            |
| TRIReagent | Zymo Research | R2050-1 |
| Triton X-100 | Sigma-Aldrich | CAS Number: 9036-19-5 |
| Ketamine | Patterson Veterinary | 07-803-6637 |
| Xylazine | Patterson Veterinary | 07-808-1947 |
| Euthasol (pentobarbital) | Patterson Veterinary | 07-805-9296 |
| SARS-CoV-2 nucleocapsid | Sino Biological | 40588-V088 |
| SARS-CoV-2 RBD | Horspool et al., 2021 | N/A |
| **Trials** |        |            |
| TMB reagent | Biolegend | 41101 |
| Tween 20 | Sigma-Aldrich | P1379-1L |
| Dextrose (D-Glucose) Anhydrous | Fisher chemical | D16-500 |
| **Critical commercial assays** |        |            |
| Qubit Broad Range Assay Kit | Invitrogen | Q10210 |
| R&D 9-plex mouse magnetic Luminex assay | Biotechne R&D systems | LXSAMSM |
| Qubit RNA High Sensitivity Assay kit | Invitrogen | Q32852 |
| **Deposited data** |        |            |
| Gene Expression Browser | Mendeley Data | |
| Raw RNAsequencing reads | NCBI SRA | |
| **Experimental models: Organisms/strains** |        |            |
| Mouse: B6.Cg-Tg(K18-ACE2)2PrlMm/J | The Jackson Laboratory | Strain #: 034860 |
| **Oligonucleotides** |        |            |
| Nucleocapsid primers (F: ATGCTGCAATCGTGCTACAA; R: GACTGCCGCTCTTGCTC) | Winkler et al., 2020 | N/A |
| TaqMan probe (IDT:/56-FAM/TCAAGGAAC/ZEN/AAC ATTGCCAA/3IABkFQ/) | Winkler et al., 2020 | N/A |
| **Software and algorithms** |        |            |
| Prism | GraphPadSoftware | Version 9 |
| Venny 2.1 | Oliveros, 2007-2015 | https://bioinfogp.cnb.csic.es/tools/venny/ |
| BioVenn | Hulsen et al., 2008 | https://www.biovenn.nl/venndiagram.tik/create.php |
| WEB-based Gene SeT AnaLysis Toolkit | Liao et al. 2019 | http://www.webgestalt.org/ |
| Morpheus | Broad institute | https://software.broadinstitute.org/morpheus/ |
| Ingenuity Pathway Analysis | Qiagen IPA Software | N/A |
| **Other** |        |            |
| Mouse Diet, High Fat Fat Calories (60%), Soft Pellets | Bio-Serv | F3282 |
RESOURCE AVAILABILITY

Lead contact
Requests for additional information surrounding the described experiments and methods or reagents should be directed to the Lead Contact, Holly Cyphert (damron40@marshall.edu).

Materials availability
- This study did not generate any new unique reagents.
- The RNAseq datasets (gene expression browsers) generated in this study have been deposited to and are available at: https://doi.org/10.17632/zds9bng2yc.1.

Data and code availability
RNA sequencing data have been deposited at NCBI SRA: PRJNA875205 and are publicly available as of the date of publication. Accession numbers are listed in the Key resources table.

Any additional information required to reanalyze the data reported in this article is available from the lead contact on request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animal ethics statement
SARS-CoV-2 research experiments in mice were conducted in compliance with West Virginia University IACUC protocol #2009036460. K18-hACE2 transgenic mice were obtained for use from The Jackson Laboratory (B6.Cg-Tg(K18-ACE2)2Prlmn/J; JAX strain number #034860). West Virginia University’s Biosafety Level 3 Laboratory and Animal Housing facility were used for the cultivation of SARS-CoV-2 stocks and mouse challenge studies under IBC protocol #20-09-03.

Biosafety statement
Additional work in BSL2 conditions were performed after mouse tissue samples containing SARS-CoV-2 were treated with either 1% Triton (Sigma-Aldrich T8787) by volume (serum and supernatants) or TRIzol reagent (Zymo R2050-1) at a 1:1 ratio or greater (homogenates) to inactivate virus.

High fat diet induced K18-hACE2 mouse model of obesity and type 2 diabetes
Six-week-old male and female K18-hACE2 mice were provided a 60% fat diet (Bio-serv) ad libitum for 8 weeks to induce obesity and metabolic dysfunction. Age- and sex-matched control mice were concurrently provided a standard chow diet. Mouse weights were monitored weekly. At week 6, intraperitoneal glucose tolerance testing (IPGTT) was performed to assess insulin sensitivity. Mice were fasted for 6 h then injected with a solution of glucose in PBS (2% w/v) intraperitoneally (2 mg/g bodyweight). At 0, 15, and 60 min, blood glucose was measured from tail snips using a hand-held glucometer.

SARS-CoV-2 challenge strain cultivation
The Alpha variant of SARS-CoV-2 was obtained first from BEI: hCoV19/England/204820464/2020 (Alpha; NR-54000)(GISAID: EPI_ISL_683466). Vero E6 cells were grown to 90% confluency in a T150 flask before culture media was replaced with 20mL of viral stock in infection media. After 72 h of incubation at 37°C and monitoring for cytopathic effects, virus-containing cell media was pipetted off and transferred to two 15mL conical tubes. Media was centrifuged at 450 × g for 5 min at 4°C to pellet any cellular debris. Supernatant was then aliquoted into 1mL O-ring cryovials and stored as stocks at −80°C until use. Viral titers were confirmed by plaque assay.

METHOD DETAILS

K18-hACE2 mouse challenge with SARS-CoV-2
High fat diet and control male and female 6-week-old K18-hACE2 mice were health checked by West Virginia University Office of Laboratory Animal Resources vet staff before they were moved to the ABSL3 facility for SARS-CoV-2 challenge experiments. On the day of challenge, mice were anesthetized using an intraperitoneal injection of ketamine (Patterson Veterinary 07-803-6637) + xylazine (Patterson Veterinary 07-808-1947) (80 mg/kg). A suspension of the SARS-CoV-2 Alpha variant (10^3 PFU) was then administered
intranasally to each mouse by pipetting 25μL of the stock into each nare of the anesthetized mice (50 μL total).

**K18-hACE2 mouse disease scoring**

Beginning on the day of challenge, the overall health of the mice was evaluated using in-person assessments of rectal temperature, weight, appearance, and behavior. In addition to in-person checks, mice were monitored using the SwifTAG Systems video monitoring system. Mice received daily scores for activity (0-3), appearance (0-2), changes in respiratory phenotype (0-2), eye closure/conjunctivitis (0-2) and weight loss (0-4). Each score was based on a scale where 0 indicated a normal mouse phenotype and the numerical scale represents increasingly severe phenotypes. The combined total of each category’s scores were awarded to each mouse as its daily disease score. Mice that reached 20% weight loss experienced a significant drop in temperature and/or a disease score of 5 were euthanized in compliance with predetermined humane endpoints. Daily health checks were performed until day 11 post-challenge.

**Mouse necropsy and tissue collection**

On the day of euthanasia, mice were administered an intraperitoneal injection (390 mg/kg) of Euthasol (pentobarbital) (Patterson Veterinary 07-249 805-9296) diluted in 0.9% sterile NaCl. Once mice were unresponsive, blood was collected via cardiac puncture in serum separator tubes (BD, 365,967). To separate the serum, each tube was centrifuged for 5 min at 15,000 × g. Mice were dissected to collect the lung and brain tissues as well as nasal wash fluid for downstream analysis. Nasal wash fluid was collected by pushing 1mL sterile PBS by catheter through the nasal pharynx. Right lobes of the lung were homogenized in 1mL sterile PBS using gentleMACS C tubes (Miltenyi Biotec, 130-096-334) and the m_Lung_02 program on the gentleMACS Dissociator (Miltenyi Biotec, 130-093-235). Whole brains were homogenized similarly. For RNA analysis 300μL of lung homogenate, or 500μL of brain homogenate was added to 1mL TRizol Reagent (Zymo R2050-1). For nasal wash RNA analysis, 500μL of nasal wash was added to 500μL of TRizol Reagent. For cytokine analysis, 300μL of lung homogenate was centrifuged at 15,0000 × g for 5 min and the supernatant was collected.

**Quantification of SARS-CoV-2 nucleocapsid RNA**

The Direct-zol RNA miniprep kit (Zymo, R2050) was used according to the manufacturer’s protocol to purify RNA from the nasal wash, lung, and brain homogenates of SARS-CoV-2 challenged mice. Resulting RNA concentrations were quantified using the Qubit 3.0 Flurometer and Qubit Broad Range Assay Kit (Invitrogen, Q10210). SARS-CoV-2 nucleocapsid RNA was quantified by qPCR using the Applied Biosystems TaqMan RNA-to-CT One-Step Kit (Thermo Fisher, 4,392,938). qPCR reactions for each sample and tissue were prepared in triplicate in Micro-Amp Fast optical 96 well reaction plates (Applied Biosystems, 4,306,737) and contained: 2XTaqMan RT-PCR Mix, 900nM Forward and reverse primers, 250nM TaqMan probe, 40X TaqMan RT enzyme mix and 100ng RNA template. RNA template from nasal wash samples was added at a set volume of 2 μL because of low RNA concentrations. Nucleocapsid primers (F: ATGCTGCAATCGTGCTACAA; R: GACTGCCGCCTTGCTC); and TaqMan probe (IDT:56-FAM/TCAAGGAAC/ZEN/AACATTGCCCAA/3IABkFQ/) synthesized according to Winkler et al. (2020) (Winkler et al., 2020). qPCR reactions were performed on the StepOnePlus Real-Time System machine using the following parameters: Reverse transcription for 15 min at 48°C, activation of AmpliTaq Gold DNA polymerase for 10 min at 95°C, and 50 cycles of denaturing for 15 s at 95°C and annealing at 60°C for 1 min.

**Serological analysis**

Anti-nucleocapsid and anti-RBD specific IgG as well as IgM antibodies were measured in the serum of SARS-CoV-2 challenged mice by ELISA. High binding 96-well plates (Pierce, 15,041) were coated overnight (4°C) with RBD or nucleocapsid at a concentration of 2 μg/μL. On the day of, plates were washed 3× with PBS-0.1% Tween 20, then incubated shaking for 1 h with 200μL of 3% non-fat milk in PBS-0.1% Tween 20 per well to block. After blocking, the plates were washed 4× with PBS-0.1% Tween 20 before adding sample. Mouse serum was diluted 1:20 in the wells of row A with 1% non-fat milk in PBS-0.1% Tween 20, then serially diluted 1:2 into 1% non-fat milk in PBS-0.1% Tween 20 across the concurrent wells of two plates (total of 15 dilutions). Plates were then incubated shaking for 1 h with sample before being washed again 4× with PBS-0.1% Tween 20. Secondary antibody diluted 1:2,000 for IgG (Goat anti-mouse IgG (H + L) HRP (Novus biological, NB101-75130)) or 1:10,000 for IgM (Goat anti-mouse IgM HRP (Novus biological, NB7497)) in 1% non-fat milk in PBS-0.1% Tween 20 was then added to all wells at a volume of 100μL, and plates were
incubated shaking for an additional 1 h. After incubation, unbound secondary was washed away with 5 ×
3 washes of PBS-0.1% Tween 20. TMB reagent (Biolegend, 421,101) was added to each well at a volume of
100μL to develop the reaction. After 15 min in the dark, the reaction was stopped using 50μL 2N Sulfuric
acid. Assays were analyzed at 450nm using the Synergy H1 plate reader. Antibody titers were quantified
using area under the curve analysis in GraphPad Prism v.9.0.0.

Cytokine analysis
To measure IFN-γ as well as other cytokines, samples of lung supernatant and serum from each mouse were
run on the R&D 9-plex mouse magnetic Luminex assay (Ref LXSAMSM). The manufacturer’s protocols were
followed to prepare and run samples. The plate was analyzed on the Luminex Magpix to calculate concen-
trations (pg/mL) based on of the individual standard curves for each cytokine. MSD assay plates were
analyzed using the Meso Scale Discovery Sector 2400.

Illumina library preparation
RNA concentrations from SARS-CoV-2 challenged mouse lung homogenates were measured with the Qu-
bit 3.0 Fluorometer using the RNA High Sensitivity kit (Invitrogen, Q32852) and RNA integrity was assessed
using an Agilent TapeStation. RNA was DNAsed before library preparation. Illumina sequencing libraries
were created with the KAPA RNA Hyper-Prep Kit with RiboErase (Basel, Switzerland). Resulting libraries
passed standard Illumina quality control PCR and were sequenced on an Illumina NovaSeq s4 4000 at Ad-
mera Health (South Plainfield, NJ).

QUANTIFICATION AND STATISTICAL ANALYSIS
RNA sequencing and in silico bioinformatics analysis
A total of ~100 million 150 base pair reads were acquired per SARS-CoV-2 challenged mouse lung sample.
Sequencing data will be deposited to the Sequence Read Archive. The reads were trimmed for quality and
mapped to the Mus musculus reference genome using CLC Genomics Version 21.0.5. An exported gene
expression browser table is provided as supplemental materials (Table S1). Statistical analysis was per-
formed with the Differential Expression for RNA Seq tool and genes were annotated with the reference
mouse gene ontology terms. PCA plots were formed in CLC Genomics Version 21.0.5. Quantification of
the number of activated or repressed genes unique to each experimental group was performed using
Venny 2.1 (Olivers, n.d.) and visually modeled using the WEB-based BioVenn (Hulsen et al., 2008). Genes
from each experimental comparison with significant fold changes compared to no-challenge (Bonferroni
≤0.04) were submitted to the WEB-based Gene SeT AnaLysis Toolkit’s OverRepresentation Analysis
(ORA) software compared to the reference set “affy mg u74a” to determine GO terms from gene ontology
and biological process databases (FDR ≤0.05) (Liao et al., 2019). GO Term heat maps were generated us-
ing Morpheus (Morpheus, n.d.). To analyze the expression of the hACE2 transgene, the RNA reads were
mapped to the human ACE2 gene (GRCh38). SARS-CoV-2 reads were analyzed by mapping the reads to
the SARS-CoV-2 WA-1 reference genome. hACE2 and viral reads were normalized by dividing counts
per 50M total reads in each sample.

Ingenuity pathway analysis
RNAseq fold change gene expression data was submitted to Ingenuity Pathway analysis using a cut off
value of p=0.05. Pathways that were statistically enriched were exported and plotted into heat maps using
Morpheus as described above.

Statistical analysis
Statistical tests were performed in GraphPad Prism v.9.0.0. K18-hACE2 mouse studies were performed with
an n = 5 per experimental group. In preluding experiments to determine the challenge dose, n ≥ 3. Kaplan
Meier survival curves were analyzed using Mantel-Cox log rank tests. Student’s t-tests were used for com-
parisons made between two groups. When three or four groups were being compared, statistical differ-
ences were assessed using one-way ANOVA with Dunnett’s multiple comparisons test or two-way
ANOVA with Tukey’s multiple comparisons test for parametric data. For any non-parametric data,
Kruskal-Wallis tests with Dunn’s multiple comparisons tests were used.