The Impact of ACE2 Polymorphisms on COVID-19 Disease: Susceptibility, Severity, and Therapy

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The coronavirus disease 2019 (COVID-19), caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), has currently spread worldwide, leading to high morbidity and mortality. As the putative receptor of SARS-CoV-2, angiotensin-converting enzyme 2 (ACE2) is widely distributed in various tissues and organs of the human body. Simultaneously, ACE2 acts as the physiological counterbalance of ACE providing homeostatic regulation of circulating angiotensin II levels. Given that some ACE2 variants are known to cause an increase in the ligand-receptor affinity, their roles in acquisition, progression and severity of COVID-19 disease have aroused widespread concerns. Therefore, we summarized the latest literature and explored how ACE2 variants and epigenetic factors influence an individual’s susceptibility to SARS-CoV-2 infection and disease outcome in aspects of ethnicity, gender and age. Meanwhile, the possible mechanisms for these phenomena were discussed. Notably, recombinant human ACE2 and ACE2-derived peptides may have special benefits for combating SARS-CoV-2 variants and further studies are warranted to confirm their effects in later stages of the disease process. As the uncertainty regarding the severity and transmissibility of disease rises, a more in-depth understanding of the host genetics and functional characteristics of ACE2 variants will not only help explain individual clinical differences of the disease, but also contribute to providing effective measures to develop solutions and manage future outbreaks of SARS-CoV-2.

Keywords: COVID-19, SARS-CoV-2, ACE2, variants, gene polymorphism, susceptibility, hrsACE2

INTRODUCTION

Coronavirus disease 2019 (COVID-19) poses a tremendous threat to human health. As of 4 August 2021, the infection has its presence around the globe with over 199 million of the world population being infected and has claimed over 4 million lives and counting (WHO, 2021). Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the virus causing COVID-19, is closely related to SARS-CoV with above 85% identity (Gralinski and Menachery, 2020). Similar to SARS-CoV, the entry of SARS-CoV-2 into the host cell is mediated by the surface-anchored spike (S) glycoprotein (Figure 1). The protein has two functional units known as "S1" and "S2", which are responsible for receptor binding and membrane fusion respectively. The receptor-binding domain (RBD) in the S1 binds to the extracellular peptidase domain (PD) of angiotensin-converting enzyme 2 (ACE2),
which leads to the exposure of S1/S2 interdomain protease site. Then, the S protein is cleaved by the proprotein convertase furin at the S1/S2 site and the transmembrane serine protease 2 (TMPRSS2) at the S2 site (Bestle et al., 2020). These processes facilitate large-scale conformational changes, triggering the fusion reaction by insertion of the fusion peptide of S2 into the host membrane (Suryamohan et al., 2021). The viral RNA is released for replication and translation upon the internalization of the SARS-CoV-2. Subsequently, new viruses are assembled and released from the cell to start another infection cycle.

As a functional receptor of SARS-CoV-2, full-length ACE2 is a 120-KDa typical zinc-metalloproteinase type I transmembrane protein, which consists of an N-terminal signal peptide of 17 amino acid residues, a PD with its HEXXH + E zinc-binding consensus sequence and a C-terminal Collectrin-like domain (Devaux et al., 2020a). It has two states, i.e. open and closed. In the open state, ACE2 opens widely from its active site, waiting for a ligand to enter. Once the ligand enters ACE2 active site, it will trigger ACE2 to close the active slot (Guo et al., 2020). ACE2 is expressed in essentially all tissues, with the highest expression in the small intestine, kidneys, testis, thyroid, heart, and adipose tissue followed by liver, lungs, colon, bladder, and adrenal gland (Li et al., 2020). Besides, there is another functional form of ACE2 with 555 amino acids that circulates in small amounts in the blood, named soluble ACE2 (sACE2), which is obtained by shedding of the full-length ACE2 through metalloproteinase 17 (ADAM17). It is worth emphasizing that binding of sACE2 to SARS-CoV-2 blocks the entry of the virus into the target cells and consequently, prevents the progression of COVID-19. Up to now, three major ACE2 functions have been described. In addition to being the viral receptor, ACE2 exhibits a protective role in the cardiovascular system and many other organs and contributes to the absorption of neutral amino acids in intestine (Kuba et al., 2010).

The ACE2 gene and protein show a high degree of genetic polymorphisms including single nucleotide variation, transcriptional variation, post-transcriptional modifications and putative protein mutations (Devaux et al., 2020a), among which single nucleotide polymorphisms (SNPs) has made its way to the scientific spotlight. Recently, Suryamohan et al. (2021) found 298 unique protein-altering variants across 256 codons distributed throughout the 805 amino acid long human ACE2 (hACE2) from a number of databases (Suryamohan et al., 2021). It is noteworthy that the symptoms and severity of COVID-19 vary greatly, ranging from very mild or no symptoms to pneumonia, acute respiratory distress syndrome (ARDS), and even death (Lopera Maya et al., 2020). As SARS-CoV-2 primarily depends on ACE2 for fusion and entry, ACE2 variation is considered to be one of the causes for this difference. Thus, it’s important to systematically characterize and evaluate ACE2 polymorphism. Herein, this review provides an update on the possible role of ACE2 variants in both the susceptibility of people to SARS-CoV-2 infection and the outcome of COVID-19. Meanwhile, particular attention is given to ACE2-based potential therapeutic strategies.

**ALTED AFFINITY OF ACE2 VARIANTS FOR S PROTEIN**

Considering that close and stable contact is crucial for RBD recognition, residue changes of ACE2 in the binding interface would influence affinity that is one of the most important determinants of host susceptibility. K353 and K31 in hACE2, which have been shown to be the main hotspots that form hydrogen bonds with the main chain of N501 and Q493 in receptor-binding motif respectively, play a role in tightly binding to the S protein of SARS-CoV-2 (Sehailia and Chemat, 2020).
Due to possessing identical or similar critical residues, ACE2 molecules from pigs, ferrets, cats, orangutans and monkeys could bind to SARS-CoV-2 RBD with analogous efficiency (Wan et al., 2020). However, the presence of N30 (instead of D30) and N31 (instead of K31) in mouse ACE2 would cause the lack of salt bridges and critical H-bond in the ACE2-SARS-COV-2 RBD complex. In terms of rats, although E26 (instead of K26) in the ACE2 forms a salt bridge with K417 in RBD, it drives the relative movement of RBD over the rat ACE2 (a shift of 6.5 Å comparing to its relative position in the human complex) and contributes to a decrease in the affinity of the virus to the ACE2 receptor (Brooke and Prischi, 2020). Another hotspot residue, K353, was replaced by histidine in mice and rats, leading to unfavorable bind with the SARS-CoV-2 RBD (Gao and Zhang, 2020). Thus, ACE2 receptor polymorphism can predict species susceptibility to SARS-CoV-2 to some extent (Devaux et al., 2020b). Although pigs and ferrets can be infected with SARS-CoV-2, they generally have a large size and a long growth cycle that is difficult to process in large quantities. Given the rapid emergence of virus mutations and novel vaccine candidates, there is a continuous need for genetic-modified mouse/rat models with the high tissue-specificity of hACE2 expression and susceptibility to SARS-CoV-2 infection.

Even if a total of more than 4 billion vaccine doses have been administered at present, COVID-19 is still spreading rapidly around the world with risk factors for infection including race/ethnicity, ages and both sexes. Thus, it is particularly important to pay close attention to the genetic variation of hACE2. The majority of the ACE2 coding variants found in Genome Aggregation Database (gnomAD) are missense variants that can alter amino acid sequence of the encoded protein, and ultimately result in a change of protein structure. Ser19, situated at the beginning of the helix S19-I54, forms hydrogen bonds with Glu23 and Gln24 to stabilize the helical structure and may interact with the S477 in S protein via weak hydrophilic interaction. Its mutation to proline would destabilize the helix structure and consequently influence the recognition by SARS-COV-2 (Guo et al., 2020). Based on the OncoMX (https://oncomx.org/) and gnomAD (https://gnomad.broadinstitute.org/) datasets, we compared 23 ACE2 variants with different allele frequencies in European (non-Finnish) and East Asian (Table 1). It is shown that 14 ACE2 variants (I21V, E23K, K26R, N64K, T92I, Q102P, D206G, G211R, R219C, E329G, H378R, V447F, A501T and N720D) with enhanced susceptibility have higher allele frequencies in European (non-Finnish) populations than East Asian populations, and 2 ACE2 variants (E35K and F72V) with enhanced susceptibility have higher allele frequencies in European (non-Finnish) populations than East Asian populations while they are low or not expressed in European (non-Finnish) populations. These findings are consistent with the epidemic situation and could partially account for the differences between COVID-19 prevalence and mortality rates in Europe and East Asia.

Notably, there are still controversial issues in results of binding affinity of S protein for the same ACE2 receptor among studies. Suryamohan et al. (2021) and Wang et al. (2020) predicted that S19P, T27A and M82I enhanced ACE2 affinity whereas E37K, G326E, G352V and D355N exhibited decreased binding to RBD of S protein (Wang et al., 2020; Suryamohan et al., 2021). In contradiction to their results, another group has reported that all these variants were found to have no obvious changes in the binding energy scores (Othman et al., 2020). These discrepancies may be attributed to different criteria and methods used by various researchers. Utilizing 5 fast structure-based computational approaches, Hadi-Alijanvand and Rouhani (2020) reported the predicted affinities of 240 mutated versions of ACE2 with the SARS-CoV-2 Spike protein, and there were varying degrees of difference in these prediction results. Particularly, the predicted stabilities of 211 ACE2 variant/S1 complexes obtained using SEPAS and SAAMBE-3D methods were different, among which 101 were completely opposite (Hadi-Alijanvand and Rouhani, 2020). As these methods considered different structural aspects of protein complexes, the observed inconsistency in prediction seems natural. Therefore, to better identify the risk variant in ACE2 and develop genetic diagnostic risk profiling for COVID-19 susceptibility, it is essential not only to develop more accurate methods that consider the dynamic features of protein structures, but also to perform more validation studies in vitro.

### SEX AND AGE DIFFERENCES IN ACE2 POLYMORPHISMS

In the early pandemic, it was generally believed morbidity and lethality were higher in men compared with women. Although the available data indicate similar SARS-CoV-2 infection rates between the genders, males account for a greater percentage of deaths in most countries throughout the world, including China (approximately 64% male: 36% female), France (57.83%: 42.17%), Italy (56.43%: 43.57%), America (54.85%: 45.15%) and England (54.44%: 45.56%) (Global Health 50/50). As is well known, genetic variants of ACE, angiotensinogen and angiotensin II (Ang II) receptor genes show more profound effects in men, some of which are only positive in them. Therefore, apart from a higher prevalence of smoking and a deadly symbiosis of chronic systemic inflammation produced by chronic metabolic disease (such as obesity, metabolic syndrome and type 2 diabetes) in males (Mauvais-Jarvis, 2020) and a stronger innate and adaptive immune response to contagions in females, it is conceivable that ACE2 and its genetic variants also display gender-specific effects, which might be more decisive than epigenetic regulation for SARS-CoV-2 progression. For most X-chromosome genes, the dosage difference between XX females and XY males is balanced by silencing one of the two X chromosomes in females in early development. However, the ACE2 gene is located on Xp22.2, an area where genes could escape this X-inactivation, resulting in phenotypic differences between sexes. In males, the 16 residues of RBD carried by SARS-CoV-2, are in contact with 16 out of 20 residues of ACE2 protein, and there are varying degrees of difference in these prediction results. Particularly, the predicted stabilities of 211 ACE2 variant/S1 complexes obtained using SEPAS and SAAMBE-3D methods were different, among which 101 were completely opposite (Hadi-Alijanvand and Rouhani, 2020). As these methods considered different structural aspects of protein complexes, the observed inconsistency in prediction seems natural. Therefore, to better identify the risk variant in ACE2 and develop genetic diagnostic risk profiling for COVID-19 susceptibility, it is essential not only to develop more accurate methods that consider the dynamic features of protein structures, but also to perform more validation studies in vitro.

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TABLE 1 | Allele frequencies of ACE2 variants with altered affinity in European (non-Finnish) and East Asian.

| Variant | Wild aa | Position | Mutant aa | Effect on affinity | Direct or indirect | European (non-Finnish) | East Asian |
|---------|---------|----------|-----------|-------------------|--------------------|-----------------------|-------------|
| rs778030746 | I       | 21       | V         | ↑                  | Indirect           | 0.00002446            | 0.000       |
| rs756231991 | E       | 23       | K         | ↑                  | Indirect           | 0.00001222            | 0.000       |
| rs4646116   | K       | 26       | R         | ↑                  | Direct             | 0.0008688             | 0.00006738  |
| rs1348114695 | E      | 35       | K         | ↓                  | Direct             | 0.00001222            | 0.0001444   |
| rs1192192618 | Y      | 50       | F         | ↓                  | Indirect           | 0.00001227            | 0.000       |
| rs1569045690 | N      | 51       | S         | ↓                  | Indirect           | 0.00001229            | 0.000       |
| rs1325542104 | M      | 62       | V         | ↓                  | Indirect           | 0.00001266            | 0.000       |
| rs1199100713 | N      | 64       | K         | ↑                  | –                  | 0.00001081            | 0.000       |
| rs1256007252 | F      | 72       | V         | ↓                  | Indirect           | 0.000                | 0.00007243  |
| rs763956248  | T       | 92       | I         | ↑                  | Direct             | 0.00002448            | 0.000       |
| rs1395878099 | Q      | 102      | P         | ↑                  | –                  | 0.00002168            | 0.000       |
| rs142443432  | D       | 206      | G         | ↑                  | –                  | 0.00006314            | 0.000       |
| rs148771870  | G       | 211      | R         | ↑                  | –                  | 0.0001944             | 0.000       |
| rs372272603  | R       | 219      | C         | ↑                  | –                  | 0.0006950             | 0.000       |
| rs149396283  | E       | 329      | G         | ↑                  | –                  | 0.00006810            | 0.000       |
| rs707610075  | G       | 352      | V         | ↓                  | Direct             | 0.00001274            | 0.000       |
| rs961960700  | D       | 355      | N         | ↓                  | Indirect           | 0.00002591            | 0.000       |
| rs142984500  | H       | 378      | R         | ↑                  | –                  | 0.0001950             | 0.000       |
| rs762890235  | P       | 389      | H         | ↑                  | –                  | 0.00002453            | 0.000       |
| rs776328956  | V       | 447      | F         | ↑                  | –                  | 0.00007882            | 0.000       |
| rs191860450  | I       | 468      | V         | ↑                  | –                  | 0.00002198            | 0.01140     |
| rs140473595  | A       | 501      | T         | ↑                  | –                  | 0.00004746            | 0.000       |
| rs41303171   | N       | 720      | D         | ↑                  | –                  | 0.02521              | 0.000       |

Reducing unregulated inflammatory lung damage. With a single X chromosome, men lack the alternative mechanisms of organ protective functions. In addition, males are hemizygous for the ACE2 gene, but in females, the rate of homozygous for variants was extremely low and 98.3% of missense variants were heterozygous, implicating that the efficacy of interaction-booster variants of ACE2 can be stronger in men than women. An analysis revealed that 7 of 10 interaction-booster variants (S19P, K26R, I21V, I21T, N64K, H378R, and T92I) were expressed in males, which means more than half of the variants have the opportunity to affect the interaction between the hACE2 and S1 protein of SARS-CoV-2 (Darbani, 2020). Nevertheless, the current data is not able to determine a clear causality, and as most of the identified variants have very low frequencies, further functional studies are needed to verify these results.

The high incidence of COVID-19 in the elderly is also associated with increased expression of ACE2 (Pruimboom, 2020). In children, the ACE2 gene in the lungs, oral tissues, and other organs is ordinarily hypermethylated and therefore actually silent (Pruimboom, 2020). Nevertheless, high-throughput, next-generation sequencing technologies and microarray revealed an increase in variability of DNA methylation with age, which caused certain genes to gradually become more active during the aging process. For example, DNA methylation at one CpG (cg08559914) near the transcription start site of the ACE2 gene in airway epithelial cells is inversely correlated with biological age (Corley and Ndhlovu, 2020). In addition, a previous study showed that ACE2 expression is also transcriptionally regulated by adenosine monophosphate kinase via sirtuin 1 (SIRT1) (Patel et al., 2016), and exogenous zinc strongly inhibits SIRT1 activity (Chen et al., 2010), hence, regulation of SIRT1 by zinc may decrease ACE2 expression and ultimately SARS-CoV-2 entry into the cell. However, some evidence indicates that zinc intakes among older adults might be marginal. An analysis of the third National Health and Nutrition Examination Survey data found that 35% to 45% of people over 60 years old had zinc intakes below the estimated average (Ervin and Kennedy-Stephenson, 2002). Notably, anosmia and ageusia, previously thought to be pathognomonic of zinc deficiency, are now also considered early signs of COVID-19, suggesting the presence of common mechanisms.

Mortality rates of COVID-19 vary dramatically across age groups. Compared with 18- to 29-year-olds, the rate of death is 95 times higher in 65- to 74-year-olds, 230 times higher in 75- to 84-year-olds, and 600 times higher in those who are 85 years and older (Centers for Disease Control and Prevention, 2021). Apart from chronic conditions and a dysregulated immune system with aging, other possible explanations for this phenomenon are as follows: (1) The vast majority of deaths result from ARDS with the histologic finding of diffuse alveolar damage (DAD). In response to DAD, type II pneumocytes enter the cell cycle and normally downregulate ACE2 expression (Baker et al., 2021). However, in the elderly suffering from COVID-19 or other diseases, the increased circulating inflammatory cytokines and age-associated epigenetic dysregulation preclude this downregulation or even lead to ACE2 upregulation, finally causing worse clinical outcomes. (2) The sACE2 levels are lower in the population over 60 years of age. In addition to the decreased Ang II/ADAM17 induced ACE2 shedding owing to the lower plasma renin concentration; some microRNAs (miRNAs) target the 3'UTR, 5'UTR or coding DNA sequence of ACE2 and influence its expression either by mediating complementary mRNA degradation or by interfering with translation. For example, miR-212-5p is significantly positively correlated with age and negatively related to sACE2 while miR-
3909, the miRNA significantly negatively correlated with age, is positively related to sACE2 (Elemam et al., 2021). (3) As the active form of vitamin D (VitD), 1,25(OH)2D3 can induce ACE2/Ang-(1-7)/MasR axis activity and suppress renin and the ACE/Ang II/AT1R axis, thereby producing a potential protective role against ARDS. Over 60 years of age, a reduction in the synthesis of VitD in response to UVB becomes apparent, which further increases when getting older (Heaney, 2006).

COMORBIDITIES AND ACE2 POLYMORPHISMS

Although the T allele of ACE2 rs2106809 has been shown to be inversely correlated with severe malaria (De et al., 2021), accumulating evidence suggests that most ACE2 polymorphisms could lead to a variety of diseases, such as atrial fibrillation, hypertension, type 2 diabetes mellitus (T2D), cardiomyopathy and so on. A summary of these results (Lieb et al., 2006; Yi et al., 2006; Wang et al., 2008; Meng et al., 2015; He et al., 2018; Kumar et al., 2018; Liu C. et al., 2018; Pan et al., 2018; Luo et al., 2019; Lozano-Gonzalez et al., 2020; Liu et al., 2021) and related research is shown in Table 2. Notably, ethnicity has been proved to alter the link between ACE2 single nucleotide polymorphisms (SNPs) and some diseases. For example, among persons of British (rs1978124, rs2074192 and rs4646188) or Finnish (rs879922) descent, there is no correlation between ACE2 SNPs and T2D, however, all the four SNPs are dramatically relevant to T2D in Xinjiang Uygur. Similarly, the sex dimorphism is observed in the relationship between ACE2 SNPs and certain diseases. The left ventricular mass index of males with the minor alleles of three SNPs (rs879922, rs4240157 and rs233575) is higher than those who do not carry these SNPs, but this phenomenon is not observed in women. On the contrary, the association of ACE2 rs2106809 with blood pressure response to ACE inhibitor was identified only in women but not in men. Specifically, after 6 weeks of ACE inhibitor therapy, female patients carrying CC or CT genotype had a greater degradation of diastolic blood pressure compared with those carrying TT genotype (Chen Y. Y. et al., 2016). Therefore, it is reasonable to assume ACE2 variant rs2106809 T allele or some other SNPs in the same linkage disequilibrium block, may be functional and could decrease the sACE2 expression and increase the susceptibility to hypertension. Considering the double effect of sACE2 on hypertension and antiviral activity, this may partly explain the high incidence rate of hypertension (58%) in critically ill patients with COVID-19 (Sulaiman et al., 2021). By the way, ACE2 rs2074192 T allele and rs2285666 G allele, which are listed in the Table 2, have been reported to be associated with more severe clinical outcomes of SARS-CoV-2 infection (Cañiero et al., 2021; Mohlendick et al., 2021). Thus, besides factors such as aging, vascular disorders and delayed virus clearance, variants-mediated risks are worth exploring in more detail. Additionally, lupus, diabetes and cancer are also comorbidities closely related to adverse outcomes or death. It is easy to understand this phenomenon because aberrant epigenetic modification of ACE2 gene is one of the characteristics of these diseases. DNA methylation defects in lupus patients (Sawalha et al., 2020), aggravated by oxidative stress generated from SARS-CoV-2 infection, could lead to ACE2 overexpression and enhance viral entry. Therefore, therapeutic strategies and drugs that efficiently target epigenetic modification should be considered in further research.

ACE2-BASED PHARMACOTHERAPY FOR COVID-19

Up to date, SARS-CoV-2 variant B.1.617 (Delta) has spread voraciously across more than 100 countries and its major mutation site, L452R, enhances infectivity by increasing the affinity of the S protein to the hACE2 (Deshpande et al., 2021). Tremendous advances have been made in vaccine development and nonpharmaceutical interventions to stop the spread of COVID-19 infection, but treatments to halt the progression of the disease are still limited. As a competitive interceptor of SARS-CoV-2 and other coronaviruses, sACE2 can be used as a protective biomarker for rapid test screening, and even one of the treatment strategies. By the way, they are not affected by common escape mutations in viral proteins. However, in view of its low concentration in plasma, it’s necessary to introduce exogenous forms. In in-vitro cell-culture experiments and engineered human organoids, an appropriate introduction of human recombinant soluble ACE2 (hrsACE2) reduced the SARS-CoV-2 growth by a factor of 1,000–5,000 (Monteil et al., 2020), directly demonstrating that this can significantly neutralize viruses. At the same time, no drug-related severe adverse events were observed in the tests of healthy volunteers and COVID-19 patients (Haschke et al., 2013). Interestingly, Khan et al. (2017) found that the infusion of hrsACE2 tends to decrease the interleukin-6 (IL-6) concentration in patients with ARDS (Khan et al., 2017). Given that the level of IL-6 has a positive correlation with the severity of the disease, hrsACE2 is considered as a promising treatment option. However, the therapeutic potential of recombinant ACE2 (rACE2) for chronic use is restricted due to its short plasma half-life of 1.8 hours. To circumvent this, rACE2-Fc is created by fusing rACE2 and an Fc fragment, with a plasma half-life of 174.2 hours and higher serum activity than rACE2 (Liu P. et al., 2018). Similarly, a C-terminally truncated hrsACE2 protein of 618 amino acids is fused with a small (5-kD) albumin-binding domain (ABD) to form a new type termed ACE2 1–618-ABD (Wysoczki et al., 2021). It offers a better way to prevent viral escape and affords more convenient dosing schedules compared with the native form of sACE2 1–740 currently undergoing clinical trials (NCT04335136). In addition, after viral binding, ACE2-ABD could direct the SARS-CoV-2–ACE2 complex to a different cell sorting pathway. When bound to albumin, it might direct the complex into the FcRn-mediated recycling pathway rather than the lysosome-endosome pathway that seems crucial for viral processing and replication.

Besides rACE2, ACE2-derived peptides corresponding to the ACE2-interacting domain of SARS-CoV-2 may represent an
### TABLE 2 | Association between ACE2 polymorphisms and human diseases’ symptoms in different populations.

| Disease/Symptom | Subjects’ characteristic | Variant | Reference(s) |
|-----------------|--------------------------|---------|--------------|
| Cardiac event   | Higher risk of AF        | rs879922| Luo et al., 2019 |
|                 | Larger size of LAD       | rs2285666| Liu et al., 2021 |
|                 | Lower LV EF              | rs2074192| Liu C et al., 2018 |
| Higher parameters (LVMI and SWT) of LVH | rs233575 | Liu C et al., 2018 |
| Higher parameters (LVMI and SWT) of LVH | rs6466168 | Lieb et al., 2006 |
| DCM             | rs4646156                |         | Wang et al., 2008 |
| Blood pressure disorder | Higher levels of SBP | rs4646142 | Pan et al., 2018; Luo et al., 2019 |
|                 | Higher levels of SBP     | rs2106809| Yi et al., 2006 |
|                 | Higher levels of DBP     | rs2236306| Lozano-Gonzalez et al., 2020 |
| DCM             | rs4646188                |         | Liu et al., 2021 |
|                 | Higher levels of DBP     |         | Lozano-Gonzalez et al., 2020 |
| Dyslipidemia    | High LDL-C level         | rs2048683| Liu et al., 2018 |
|                 | High CHOL level          | rs4240157| Liu et al., 2018 |
|                 | High CHOL level          | rs4240157| Liu et al., 2018 |
|                 | High TRIG level          | rs4830542| Pan et al., 2018 |
|                 | Low HDL-C level          | rs4830542| Pan et al., 2018 |
| Ischemic stroke | Increased serum sodium level | rs50% | Pan et al., 2018 |
|                 | Low serum potassium concentration | rs50% | Pan et al., 2018 |
| CAS ≥ 50%       | EH subjects from the south Xinjiang | rs2048683 | Luo et al., 2019 |
| CAS ≥ 50%       | Uygur T2D patients       | rs2048683 | Liu et al., 2021 |
| Increased MAU   | Uygur T2D patients       | rs2048683 | Liu et al., 2021 |
| Higher level of HsCRP | Uygur T2D patients | rs2048683 | Liu et al., 2021 |
| DR and PDR      | Chinese female T2D patients | rs2048683 | Liu et al., 2021 |
| SGA             | Neonates                 | rs2048683 | Meng et al., 2015 |

13 ACE2 variants are shown in different colors and have been associated with different diseases or symptoms. AF, atrial fibrillation; LAD, left atrial end-systolic diameter; LVEF, left ventricular ejection fraction; LVMI, left ventricular mass index; SWT, septal wall thickness; LVH, left ventricular hypertrophy; DCM, dilated cardiomyopathies; EH, essential hypertension; SBP, systolic blood pressure; DBP, diastolic blood pressure; LDL-C, low-density lipoprotein cholesterol; CHOL, cholesterol; TRIG, triglyceride; HDL-C, high-density lipoprotein cholesterol; CAS, carotid artery stenosis; T2D, type 2 diabetes; MAU, microalbuminuria; HsCRP, high-sensitivity C-reactive protein; DR, diabetic retinopathy; PDR, proliferative diabetic retinopathy; SGA, small-for-gestational-age; HCM, hypertrophic cardiomyopathy.
alternative strategy to target this pathway due to their relative safety, low production complexity and remarkable curative action. Through fusing the RBD-binding proteins to the CHIPΔTPR modified E3 ubiquitin ligase domain, Chatterjee et al. (2020) designed 23-mer (A2N)-CHIPΔTPR fusion, and this ACE2-derived peptide demonstrated robust RBD degradation capabilities in human cells and reduced the infection rate of the pseudovirus by ~60% (Chatterjee et al., 2020). As for genuine SARS-CoV-2, its infection has also been reported to be inhibited by Spike-targeting ACE2-derived peptide1 (SAP1) and SAP6 in affinity precipitation assays (Larue et al., 2021).

Of note, although most ACE2-based pharmacotherapies exhibit an acceptable safety profile and some of them have entered clinical trials (Table 3) (Zhong et al., 2011; Zou et al., 2014; Chen L. J. et al., 2016; Gu et al., 2016; Khan et al., 2017; Liu C. et al., 2018; Rathinasabapathy et al., 2018; Chatterjee et al., 2020;...)

### TABLE 3 | A summary of the therapies based on ACE2 used until present.

| Primary drug | Condition or disease | Species | Dosage | Medication method | Results | Pros and Cons | Reference |
|--------------|----------------------|---------|--------|-------------------|---------|---------------|-----------|
| rACE2        | Pulmonary hypertension | Mice (FVB/N) | 1.2 mg/kg/day × 14 days | Mini-osmotic pumps | Significantly attenuated vascular remodeling and increased pulmonary SOD2 expression without measurable effects on pulmonary fibrosis. | – | Rathinasabapathy et al., 2018 |
| hrsACE2      | COVID-19             | Human   | 2 times/day | Intravenous injection | Significant improvement in mechanical ventilator-free Days and reduction in viral RNA load observed | – | Clinicaltrials, 2021 |
| hrsACE2      | H5N1 infection induced acute lung injury ARDS | Mice (BALB/c) | 0.1 mg/kg × 3 times | Intraperitoneal injection | Reduced acute lung injury severity and prolonged overall survival time of infected mice | – | Zou et al., 2014 |
| hrsACE2      | Respiratory syncyti al virus induced lung injury | Mice (C57BL/6) | 0.1 mg/kg × 3 times | Intravenous injection | Did not result in improvement in physiological or clinical measures of ARDS | Causes some adverse events, such as hypernatremia, pneumonia, dysphagia and rash. | Khan et al., 2017 |
| hrsACE2      | Ang II-mediated renal oxidative stress, inflammation, and fibrosis | Mice (C57BL/6) | 2 mg/kg/day × 14 days | Intraperitoneal injection | Reversed renal NADPH oxidase activation and proinflammatory changes, and attenuated tubulointerstitial fibrosis | – | Zhong et al., 2011 |
| hrsACE2      | Ang II-induced renal fibrosis | Mice (apoE-KO) | 2 mg/kg/day × 14 days | Intraperitoneal injection | Dramatically ameliorated Ang II-mediated hypertension, kidney remodeling and tubulointerstitial fibrosis | – | Chen et al., 2016a |
| rACE2-Fc     | Acute hypertension | Mice (BALB/c) | 4 kU/kg | Intravenous injection | Significantly lowered SBP following Ang II | 1. With the same dose, the peak blood concentration of rACE2-Fc was 2-fold higher than that of untagged rACE2 and could be sustained for much longer time. |
| rACE2-Fc     | Chronic hypertension, albuminuria and multiorgan fibrosis | Mice (RenTgMK) | 1 time/7 days × 42 days | Intravenous injection | 2. The levels of BP and albuminuria stayed within normal ranges. |
| rACE2-Fc     | Cardiac hypertrophy and fibrosis | Mice (C57BL/6) | 1 time/7 days × 28 days | Intravenous injection | 3. Reduced areas of fibrosis |

(Continued)
Clinical trials (2021; Larue et al., 2021; Wysocki et al., 2021), the understanding of SARS-CoV-2 pathogenesis is still in a developing stage. The possibility of virus making use of other receptors (e.g., tyrosine-protein kinase receptor UFO, kidney injury molecule-1 and cluster of differentiation 147), co-receptors/auxiliary proteins or even other mechanisms for entry into the cells cannot be ruled out. As the clinical manifestations of SARS-CoV-2 infection are very variable and can occur from asymptomatic infection to severe pneumonia, the appropriate drug concentrations in different patients should be explored in future research. Moreover, only the effect of hrsACE2 on early SARS-CoV-2 infection is understood until now, its effect in the later stages of the disease process is still unclear, and the effects on the lungs are also poorly understood. ACE2 infusion may increase Ang-(1-7) levels and reduce circulating Ang II levels to promote infectious or cardiogenic effects in later stages of the disease. Thus, the concentration in the same patient at different stages of the treatment must be precisely adjusted to remain within the therapeutic range. Furthermore, the human body's immune response to recombinant human proteins may result in the development of neutralizing antibodies and specific T cell responses, thereby compromising therapeutic potential. Therefore, before these types of peptides are approved for the clinical treatment of COVID-19, sufficient data from in vitro and in vivo studies are still needed to confirm their efficacy. Additionally, except for intravenous treatment, nasal drug administration should be paid more attention because of its fast onset of action, increased bioavailability and reduced adverse reactions.

**DISCUSSION**

In this review, we explored how ACE2 variants and epigenetic factors influence an individual’s susceptibility to SARS-CoV-2 infection and disease outcome in aspects of ethnicity, gender and age, and discussed some ACE2-based COVID-19 treatments (Figure 2). Residue changes of ACE2 in the binding interface would influence its expression and affinity with SARS-CoV-2. We compared the allele frequencies and expressions of 23 ACE2 variants in different populations, and found they could partially account for the differences in COVID-19 prevalence and mortality rates. However, in terms of affinity of the ACE2 variant to the S1 protein, current findings are controversial among studies, and the results lack validation by systems biology studies even though some variants have been believed to enhance the affinity in several reports. Thus, there is an urgent need for in vitro validation studies to assess the involvements of population-specific SNPs of ACE2 and other host factors in susceptibility toward SARS-CoV-2 infection. Moreover, we believe it is essential to promote the use of multidisciplinary tools and state-of-the-art “omics” technologies to comprehensively define the degree of inter-individual variations in susceptibility and subsequent immune responses determined by gene polymorphisms. Meanwhile, large-scale sequencing projects especially for ACE2 sequencing in the most severe patients in every population should be carried out to identify more specific susceptibility markers.

As an X-chromosome linked phenotype, the interaction-booster variants of ACE2 can have a more substantial impact on males than on females. Moreover, men lack the alternative mechanisms of organ protective functions because they have a single X chromosome, and this would be partially responsible for the gender bias towards a higher mortality rate in males. Furthermore, ethnicity and gender may alter the link between polymorphic variants of the ACE2 gene and a variety of diseases including hypertension, T2D, and cardiomyopathy, which represent risk factors for a severe prognosis of COVID-19. Notably, the high expression of ACE2 gene caused by

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### TABLE 3 | Continued

| Primary drug | Condition or disease | Species | Dosage | Medication method | Results | Pros and Cons | Reference |
|--------------|---------------------|---------|--------|-------------------|---------|-------------|-----------|
| ACE2 1–618-ABD | SARS-CoV-2 infection | Human kidney organoids | 0.2 mg/ml | – | Markedly reduced viral replication | 4. Long-acting rACE2-Fc may help to address the “Ang II escape” phenomena. | Wysocki et al., 2021 |
| ACE2 1–618-ABD | Acute hypertension | Mice (C57BL/6) | 1 mg/kg | Intraperitoneal injection | Significantly attenuated Ang II–induced hypertension | 1. Reduces the risk of immunogenicity 2. Can be readily synthesized or be efficiently packaged for delivery in a lipid nanoparticle or adeno-associated virus | Chatterjee et al., 2020 |
| ACE2 23-mer derived peptides CHIPATPR fusion | SARS-CoV-2 infection | HEK293T cells | 0.001 mg/ml | – | Reduced the infection rate of the pseudovirus by ~60% | – | – |
| SAP1, SAP6 | SARS-CoV-2 infection | HEK293T-ACE2-GFP cells | 3 mmol/L | – | Resulted in ~2-fold reduction in SARS-CoV-2 infection | – | Larue et al., 2021 |
| SAP1, SAP6 | HCoV-NL63 infection | LLC-MK2 cells | 3 mmol/L | – | Resulted in ~3-fold reduced HCoV-NL63 titers | – | Larue et al., 2021 |

ARDS, acute respiratory distress syndrome; NADPH, nicotinamide adenine dinucleotide phosphate; SBP, systolic blood pressure; BP, blood pressure.
hypomethylation and zinc deficiency may explain why the elderly are more sensitive to SARS-CoV-2 infection than younger populations and especially children. Moreover, their lower sACE2 and VitD levels cannot protect them from ARDS. Thus, it is reasonable to assume that stimulating specific DNA methylation of ACE2 gene could help to attenuate infection susceptibility and disease severity, and early supplements with zinc and VitD may have special benefits for elderly people. In addition, reducing age-related inflammation may decrease ACE2 expression in type II pneumocytes and limit infection of damaged alveolar epithelium. Indeed, dexamethasone and baricitinib (JAK1/2 inhibitor) have been shown to improve outcomes in COVID-19 patients. Whether these therapies bring benefits, at least in part, by regulating ACE2 expression may be a focus of future research.

Given the emergence of vaccine breakthrough infections, it is still necessary to explore effective treatment strategies and specific antiviral agents against the virus in addition to the development of vaccines. As a competitive interceptor of SARS-CoV-2, exogenous forms of sACE2 can be regarded as promising therapeutic options. Several rACE2 and ACE2-derived peptides have shown acceptable safety and preliminary efficacy, but there are still some challenges to overcome, especially concerning the appropriate drug concentrations in different patients and the same patient at different stages of the treatment.

AUTHOR CONTRIBUTIONS
All authors contributed to the literature review for the manuscript. The first draft of the manuscript was written by FC and all authors commented on previous versions of the manuscript. All authors contributed to the article and approved the submitted version.

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