Proposed title:
Identification of evolutionary trajectories shared across human betacoronaviruses

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Preprint available at: https://www.biorxiv.org/content/10.1101/2021.05.24.445313v2.full

ABSTRACT (239)
Comparing the evolution of distantly related viruses can provide insights into common adaptive processes related to shared ecological niches. Phylogenetic approaches, coupled with other molecular evolution tools, can help identify mutations informative on adaptation, whilst the structural contextualization of these to functional sites of proteins may help gain insight into their biological properties. Two zoonotic betacoronaviruses capable of sustained human-to-human transmission have caused pandemics in recent times (SARS-CoV-1 and SARS-CoV-2), whilst a third virus (MERS-CoV) is responsible for sporadic outbreaks linked to animal infections.

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Moreover, two other betacoronaviruses have circulated endemically in humans for decades (HKU1 and OC43). To search for evidence of adaptive convergence between established and emerging betacoronaviruses capable of sustained human-to-human transmission (HKU1, OC43, SARS-CoV-1 and SARS-CoV-2), we developed a methodological pipeline to classify shared non-synonymous mutations as putatively denoting homoplasy (repeated mutations that do not share direct common ancestry) or stepwise evolution (sequential mutations leading towards a novel genotype). In parallel, we look for evidence of positive selection, and draw upon protein structure data to identify potential biological implications. We find 30 candidate mutations, from which four [codon sites 18121 (nsp14/residue 28), 21623 (spike/21), 21635 (spike/25) and 23948 (spike/796); SARS-CoV-2 genome numbering] further display evolution under positive selection and proximity to functional protein regions. Our findings shed light on potential mechanisms underlying betacoronavirus adaptation to the human host and pinpoint common mutational pathways that may occur during establishment of human endemcity.

KEYWORDS
Molecular Evolution, Phylogenomics, Convergence, Adaptation, Betacoronaviruses

SIGNIFICANCE STATEMENT (115)
Identifying adaptive convergence is intimately linked to the possibility of predicting evolutionary trajectories in viruses relevant to global health. In this light, we undertook a comparative approach to find evidence of adaptive convergence across betacoronaviruses capable of a sustained human-to-human transmission (HKU1, OC43, SARS-CoV-1 and SARS-CoV-2). Our methodology involved the development of a pipeline used for identifying mutations putatively denoting homoplasy and or stepwise evolution that are also evolving under positive selection, and with potential biological implications drawn from protein structural data. Coupled with future experimental data and ongoing genomic surveillance, our results raise the possibility of predicting how the evolutionary trajectory for SARS-CoV-2 may develop as the virus establishes itself as endemic to humans.

MAIN TEXT (6183)
INTRODUCTION (598)
Understanding the mutational processes that lead to adaptation in RNA viruses is crucial for developing effective control strategies. Due to their high mutation rates and small genomes, RNA viruses often display rapid evolution. However, the vast majority of mutations are either purged through purifying selection or are selectively neutral (Loewe and Hill 2010). Only a small proportion of these may contribute to adaptive evolution and be consequently fixed through positive selection (Ohta 1973; Pond et al. 2012). For most viral genomes, the mutational pathways leading to adaptation are further constrained by functional and evolutionary limitations, such as epistasis, which refers to the adaptive dependence of a given mutation on the genetic background in which it appears (Dolan et al. 2018). Therefore, viral evolutionary trajectories are often limited and may exhibit recurrent mutational patterns indicative of adaptive
convergence, especially when applied to independent virus populations that share ecological niches (Gutierrez, Escalera-Zamudio, and Pybus 2019).

The OC43 and HKU1 embecoviruses and the SARS-CoV and SARS-CoV-2 sarbecoviruses are four betacoronavirus species capable of sustained human-to-human transmission. OC43 and HKU1 were introduced into the human population through independent zoonotic events estimated to have occurred at least 50 years ago and are associated with mild respiratory disease (Su et al. 2016). In contrast, SARS-CoV and SARS-CoV-2 were independently introduced more recently, causing severe pandemic outbreaks (W. Li et al. 2005; Vijaykrishna et al. 2007; Andersen et al. 2020; Boni et al. 2020; Banerjee et al. 2021). In 2002, SARS-CoV spread to more than 20 countries, causing a short-lived outbreak characterized by sustained human-to-human transmission (Cheng et al. 2007). Although its circulation was eventually halted, the virus displayed evidence of adaptation to the human population (He et al. 2004). Almost two decades later, SARS-CoV-2 spread globally, resulting in the current pandemic, despite a low rate of adaptive change recorded during the early stages of the outbreak (van Dorp et al. 2020; MacLean et al. 2021). The continuous circulation of OC43 and HKU1 within the human population at a global scale has been accompanied by ongoing host-specific adaptation. This is now also evident for SARS-CoV-2, exemplified by the constant emergence of novel virus lineages across time and space, with sub-lineages now reflecting regional endemic patterns (O'Toole et al. 2021).

As SARS-CoV-2 becomes established in humans, it will continue to adapt to overcome the selective pressures exerted by the collective immune response of the human population (Kissler et al. 2020). We hypothesize that adaptive convergence may occur across distantly related betacoronaviruses circulating within the same ecological niche, specifically the human host. To test this, we undertook a comparative analysis to search for evidence of shared mutational pathways between established human-endemic embecoviruses and emerging sarbecoviruses, with a focus on emerging mutations observed in SARS-CoV-2. We developed a methodological pipeline that allows for the identification of non-synonymous mutations (rendering amino acid substitutions) likely associated with adaptive convergence across multiple virus species. Firstly, we detected amino acid substitutions shared across virus taxa, displaying putative evidence of homoplasy or stepwise evolution. Secondly, we assessed whether these substitutions were positively selected, and contextualized their location to functional regions of viral proteins. Following our pipeline, we initially detected 30 candidate amino acid substitutions displaying evolutionary patterns denoting putative homoplasy and/or stepwise evolution. We subsequently identified four of these (sites 18121 [nsp14/27], 21623 [spike/21], 21635...
and proximal to functional surfaces in nsp14 (Ma et al. 2015) and the spike (S) protein. Our results provide a molecular-level context for common evolutionary trajectories that betacoronaviruses may undergo during their adaptation to the human host.

RESULTS (2364)

Patterns of genetic variability observed in human-infecting betacoronaviruses

We performed phylogenetic analyses of human-infecting betacoronaviruses using an alignment of the Orf1ab and S viral genes (see Methods sections 1 and 2). The tree shown in Figure 1 provides a comprehensive picture of the evolutionary relationships among the four betacoronavirus species studied here, consistent with previously published phylogenies of the genus (Woo et al. 2006; Woo et al. 2010; Oong et al. 2017; Zhu et al. 2018; Bedford 2021). Our analysis confirms four well-supported clades formed by virus sequences belonging to the Embecovirus (HKU1, OC43, and related viruses) and Sarbecovirus (SARS-CoV, SARS-CoV-2, and related viruses) subgenera (ICTV et al. 2017). To further validate divergence patterns at a deeper node level, we compared individual clades (sub-trees within our trees) to species-specific phylogenies. We were also able to verify the divergence patterns described for the distinct HKU1 (A–C) and OC43 (A–H) genotypes (Woo et al. 2006; Oong et al. 2017) (Supplementary Data 1). Therefore, our phylogenetic reconstructions validate the evolutionary relationships among these four distantly related betacoronaviruses.

We then analysed the proportion of codon sites (from the total number of polymorphic sites identified), corresponding to non-synonymous mutations shared between different embeco- and sarbecovirus species (i.e., those present in any of the sarbecovirus clades, and also in HKU1 and/or OC43). Derived from the Orf1ab+S alignment (comprising a total of 8962 sites), we identify approximately 2% (205 sites) as shared. Within Orf1a region (4774 sites), 2.7% of these (129 sites) were identified as shared. Within Orf1b region (2623 sites), only 0.9% (25 sites) were further identified as shared. The Orf S region (1457 sites) displayed the highest proportion of shared mutations (3.2%, 48 sites). When analysing genetic variation patterns within single virus species, we observed a high degree of sequence conservation (>91% identity) across the Orf S of all virus species. Conserved sites were predominantly located in the membrane-proximal S2 domain, while variable sites were mostly found within the membrane-distal S1 subunit (Figure 2). The predominance of variable sites within S1 compared to S2 was most evident for embecoviruses, and less so for sarbecoviruses, suggesting for a differential
adaptation stage relative to the human host environment, evidenced by a lower degree of
genetic divergence observed in Orf S in the sarbecoviruses.

We further analysed the genetic variation across virus species, focusing on the Orf S region. As previously noted for other coronaviruses (Hulswit et al. 2016), we found that Orf S exhibited a higher proportion of variable sites relative to conserved (for definitions, see Methods section 3). Specifically, only 16% of homologous sites within the Orf S alignment were conserved, while the remaining 84% were variable (Supplementary Data 2). The S2 subunit of Orf S contained the highest proportion of conserved sites, presumably due to shared functional constraints of the viral membrane fusion machinery across coronavirus species (Li 2016). Conversely, the S1 subunit displayed a higher number of variable sites, particularly within the S1A domain (also known as the N-terminal domain, or NTD). We found that the S1B domain did not display any conserved sites across virus species, likely due to differences in receptor engagement between embecovirus- and sarbecovirus- binding mechanisms. Specifically, embecoviruses use the S1A domain to interact with sialoglycan-based receptors, while sarbecoviruses use their S1B domain to bind to angiotensin-converting enzyme 2 (ACE2) (Hulswit et al. 2019; Lan et al. 2020). Finally, we identified that the conserved R residue at site 685 corresponding to the S1/S2 cleavage site (numbering according to the SARS-CoV-2 protein, codon sites 23615-23617) is shared across and within virus species (Supplementary Data 2), reflecting a conserved proteolytic maturation mechanism of the spike protein (Millet and Whittaker 2015).

Sites displaying evidence of homoplasy and/or stepwise evolution
Although not all non-synonymous mutations putatively displaying homoplasy and/or stepwise evolution may arise from positive selection, such mutational patterns are most likely to result from adaption (Escalera-Zamudio et al. 2020; Stern et al. 2017; Gutierrez et al. 2019). Thus, amongst the non-synonymous mutations identified as shared across virus species, we further searched for those displaying putative evidence for homoplasy and/or stepwise evolution (Supplementary Text 1) using our pipeline (Methods section 3). After visual validation, we confirmed that 30 sites (representing 0.3% within the Orf1ab+S alignment) display evolutionary patterns indicative of homoplasy and/or stepwise evolution (see Supplementary Text 2, Supplementary Figure 2 and 3). Two of these were found within Orf1a, nine within Orf1b, and 19 within Orf S (Table 1). The evolutionary trajectories for different amino acid states observed for three illustrative sites (18121, 21623 and 23948, further displaying evidence of evolution under positive selection and of being proximal to regions of established protein function [see the following results sections])
are highlighted below (Figure 3). The amino acid evolution patterns observed for all other sites are available in Supplementary Data 3.

Derived from the global, expanded and the re-sampled SARS-CoV-2 trees (Methods section 1, 3 and 6, Supplementary Text 3), our results show that site 18121 (codon 18121-18123 in Orf1b, corresponding to amino acid state ‘S’ in nsp14 in SARS-CoV-2 numbering) is homoplasic between HKU1 genotype B and the sarbecoviruses (Table 1, Figure 3, Supplementary Data 3). Comparably, site 21623 (codon 21623-21626 in Orf S, corresponding to amino acid state ‘R’ in S) was identified as homoplasic between SARS-CoV-2 and OC43 genotypes D, F, G and H. This site also displayed evidence for stepwise evolution within a single virus clade (OC43), exemplified by the sequential amino acid replacement pattern of V→I→K→R (Figure 3).

For site 23948 (codon 23948-23950 in Orf S, corresponding to residue 796 in S), initial observations based on the global tree revealed that amino acid state ‘D’ was present in all virus species, except for OC43 (displaying amino acid state ‘N’). However, when replicating our analyses (expanded tree), the distribution of amino acid state ‘D’ was now found present in some embecoviruses (including OC43 but excluding HKU1) and most sarbecoviruses. These discrepancies are likely due to alignment uncertainty across genome regions of highly divergent virus taxa. Nonetheless, based on consensus protein sequences and structural comparison, the structural contextualization of amino acid 796 and adjacent sites confirmed the presence of ‘D’ in SARS-CoV-1, SARS-CoV-2 and HKU1, and ‘N’ in OC43, (Supplementary Figure 6). Thus, amino acid state ‘D’ at site 23948 shows evidence of homoplasy between the SARS-CoV-1, SARS-CoV-2 and HKU1.

For this same site (23948), an additional amino acid change from ‘D’ to ‘Y’ was identified as homoplasic between some SARS-CoV-1 and SARS-CoV-2 sequences (data derived from the global, expanded and the re-sampled SARS-CoV-2 trees) (Table 1). For SARS-CoV-2, amino acid state ‘Y’ emerged and was lost repeatedly during the early stage of the pandemic (represented by independent minor clusters that quickly became extinct). However, following emergence and global spread of the B.1.1.529 virus lineage (Omicron variant of concern [VOC], and descending sub-lineages), amino acid state ‘Y’ replaced amino acid state ‘D’, displaying a predominant trend associated with the dominance of the B.1.617.2 lineage (Delta VOC, and descending sub-lineages) (Table 1, Figure 3, Supplementary Data 3) (also confirmed by independently sampled SARS-CoV-2 data available up to December 2022: https://nextstrain.org/groups/neherlab/ncov/global?c=gt-S_796).
Quantifying the effects of positive selection

The \(dN/dS\) estimates we obtained across complete virus genomes and upon specific coding regions (see Methods section 4) indicates that positive selection is acting upon the Orf1ab and Orf S of SARS-CoV-2, compared to other viruses studied here. Specifically, the effect of episodic diversifying selection was detected upon 5/14 non-recombinant fragments (three in Orf1b and two in Orf S, for details see https://observablehq.com/@spond/beta-cov-analysis).

Using the Contrast-FEL method to detect the effect of a differential selection across branches separating lineages (see Methods section 4), we found 36 sites (0.4%) evolving under differential selective pressure across distinct virus clades. Furthermore, we found 0.7% of all sites (67 codons within the Orf1ab+S alignment) to be evolving under episodic diversifying positive selection (scored under MEME with a \(p\leq 0.05\) as positively-selected sites, or PSS) (Supplementary Table 1). In contrast, we found 5% of all sites (461 codons within the Orf1ab+S alignment) to be evolving under pervasive negative selection (scored under FEL with a \(p\leq 0.05\) as negatively-selected sites, or NSS). We subsequently mapped the identified PSS and NSSs onto the SARS-CoV-2 S protein structure (Methods section 5). We observe that out of a total of 22 PSSs, 18 locate within the S1 subunit (11 in S1\(^A\), 5 in S1\(^B\), 1 in S1\(^C\) and 1 in S1\(^D\) domains), whilst the remaining four mapped onto the S2 subunit. Conversely, out of a total of 82 of NSSs, 46 locate within S1 (18 in S1\(^A\), 21 in S1\(^B\), 3 in S1\(^C\) and 4 in S1\(^D\)), whilst the remaining 36 mapped onto S2 (Supplementary Figure 7).

From the 30 non-synonymous mutations we identify as displaying evolutionary patterns putatively denoting homoplasy and/or stepwise evolution (Table 1), sites 19048, 21623, 21635, 22124 and 23048 were further scored as PSS (under different methods). Sites 21623 and 21635 were inferred as PSSs along ancestral branches leading to the HKU1, OC43 and SARS-CoV-2 clades. Sites 19048 and 22124 were inferred as PSSs along the OC43 ancestral branch, whilst 23048 was inferred as a PSS along the HKU1 ancestral branch (Table 1, Supplementary Table 1). Further analysis under the branch and site model in the MEME method (Methods section 4) revealed site 18121 to be evolving under positive selection for the HKU1 clade/branch (relative to the sarbecoviruses), in agreement with our observations made on putative homoplasy detected for this site between HKU1 genotype B, SARS-CoV-1 and SARS-CoV-2 (Table 1, Figure 3). Similarly, site 23948 was also inferred to be evolving under positive selective for the SARS-CoV-1 branch, relative to other virus clades (Supplementary Table 1).

For validation, we compared our results with selection analysis available for independently sampled SARS-CoV-2 genome data available as of December 2022 (https://observablehq.com/@spond/evolutionary-annotation-of-sars-cov-2-covid-19-genomes-...
enab) (Kosakovsky Pond). Of the 30 mutations we identify, 16 of these are currently scored as PSS or NSSs, with 13 of these mapping directly onto potential T cell epitopes derived from HLA class I and HLA-DR binding peptides in SARS-CoV-2 (Nelde et al. 2021; Campbell et al. 2020) (Table 1). Additionally, up to December 2022, sites 7478, 21614, 23948, 24620 and 25166 were detected as evolving under positive selection, whilst sites 21635, 24863, and 25037 were detected as evolving under negative selection.

Contextualization of mutations using protein structural and functional information

We then mapped the 30 mutations identified onto corresponding protein structures. Below, we focus on four exemplary sites (18121, 21623, 21635, and 23948) that meet the three criteria of: displaying evidence of homoplasy and/or stepwise evolution, showing evidence of evolution under positive selection, and being proximal to regions of established protein function. A description for the other 26 identified mutations is available in the Supplementary Text 4 and Supplementary Table 2.

Site 18121 in Orf1ab

Site 18121 is located within the Orf1ab gene and corresponds to an 'S' to 'A' mutation at residue 28 within the exonuclease domain of the nsp14 protein (numbering according to the SARS-CoV protein) (Supplementary Table 2, Figure 4). Nsp14 is involved in the 5′-capping of viral mRNA and is essential for viral mRNA transcription (Ma et al. 2015). The 'S' to 'A' mutation within this region is expected to result in the loss of an intra-protein hydrogen bond and potentially modulates the protein-protein interaction (Figure 4) (assessed under PISAebi; http://www.ebi.ac.uk/pdbe/prot_int/pistart.html) (Krissinel and Henrick 2007).

Sites 21623 and 21635 in S1

The S1 subunit mediates attachment of the virus to the host cell (Li 2016). Human-infecting embecoviruses bind to glycan-based cell receptors via two hydrophobic pockets within the S1^A region of the protein protein (Hulswit et al. 2019; Tortorici et al. 2019), while the receptor-binding site for human-infecting sarbecoviruses is located within the S1^B domain of the protein (Li et al. 2005; Lan et al. 2020; Shang et al. 2020). Both SARS-CoV and SARS-CoV-2 recognize the ACE2 molecule to enter the host cell, despite limited conservation amongst contact residues within the RBD of these virus species (Li et al. 2005; Lan et al. 2020). Site 21623 displays several non-synonymous mutations ('R', 'V', 'K' and 'I') mapping to residue 29 within the S1^A domain of the S1 subunit. Site 21635 also shows multiple non-synonymous mutations ('P', 'V',...
S', ‘L’ and ‘H’) mapping to residue 33 in S1\(^A\). For the OC43 S protein, this corresponds to a loop
neighbouring the hydrophobic pockets in S1\(^A\) instrumental for receptor recognition (Figure 5),
and changes within this region may potentially modulate receptor affinity (Hulswit et al. 2019).
The mutational patterns observed at these sites putatively denote homoplasy/stepwise evolution
and evidence of positive selection (Table 1), and are therefore congruent with antigenic drift
shaping the evolution of human-endemic coronaviruses (Kistler and Bedford 2021). In SARS-
CoV-2, mutations in both these sites (residue 29 and 33) have been observed for two VOCs
(B.1.351 and P.1, ‘Beta’ and ‘Gamma’) (Faria et al. 2021; Tegally et al. 2021). Even though
sarbecoviruses engage the ACE2 receptor via domain S1\(^B\), these residues locate to the ‘NTD
supersite’, serving as epitope for multiple of neutralizing antibodies (Kemp et al. 2021).

Site 23948 in S2
The S2 subunit of the betacoronavirus S protein contains the fusion machinery, responsible for
merging the viral envelope with the host cell membrane to facilitate delivery of the viral genome
into the target cell. This process is driven by the fusion peptide, which anchors the virus to the
host membrane, and requires cleavage of the S protein by host cell proteases at the S1-S2
junction (consensus RRAR|S in SARS-CoV-2) and at the S2’ cleavage site (R|S, located
immediately upstream of the fusion peptide in the S2 subunit) (Li 2016, Millet and Whittaker
2015). Site 23948 displays a non-conservative amino acid replacement ‘D’ to ‘Y’ (identified as
homoplasic between some SARS-CoV and SARS-CoV-2 sequences) at residue 796 of the S2
subunit, located immediately upstream of the S2’ cleavage site (Table 1, Supplementary Table
2). This residue locates within a loop crucial for the release of the fusion peptide, exhibiting
some variability across betacoronavirus species (Supplementary Figure 6). Our observations
suggest that the apparent relaxed local constraints at this site may facilitate cleavage activation
by securing loop accessibility. Perhaps consistently, the corresponding protein region in the
HKU1 structure remains unresolved (Kirchdoerfer et al. 2016).

DISCUSSION (1278)
In this study, we searched for signatures of adaptive convergence across distantly related
human-infecting betacoronaviruses, represented by shared non-synonymous mutations that
putatively denote homoplasy and/or stepwise evolution, further ranked according to their
selective relevance, and to their proximity to protein regions of known function. The majority of
the mutations we observe locate to the receptor binding region of the S protein (i.e., S1 subunit),
whilst a smaller proportion of these were found within non-structural proteins encoded by Orf1ab
Our *in-silico* analyses revealed four genomic sites (18121, 21623, 21635 and 23948) that display cumulative evidence of: *i*) a mutational pattern putatively denoting homoplasy and/or stepwise evolution, *ii*) evolution under positive selection, and *iii*) being structurally proximal to regions of known protein function. Below, we discuss our findings in light of three key evolutionary processes: antigenic drift, epistasis and adaptive convergence.

The host humoral immune response is an evolutionary force driving viral antigenic drift. In the case of betacoronaviruses, this is reflected by cumulative mutations in the S protein (particularly within the S1 subunit) that may allow frequent reinfections of the host population (Kistler & Bedford 2021; Yewdell 2021; Forni et al. 2021). In agreement with this observation, the emergence of some SARS-CoV-2 lineages (particularly VOC) has been associated with high levels of infection in pre-exposed human populations across different geographic regions (as an example on P.1, see Faria et al. 2021). Our results evidence antigenic drift upon the S1 subunit of distinct betacoronaviruses as a major component of the adaptation process to the human host environment, further evidenced by Orf S also being the least conserved genome region across distinct virus species (Li 2016). On the other hand, mutations found within Orf1ab could have a potential impact on viral fitness related to an enhanced replication efficacy in the human host (Menachery et al. 2017). As the evolution of Orf1ab is also driven by immune responses such as cytokine signalling cascades and antigen presentation (Wang et al. 2015; Taefehshokr et al. 2020; Hackbart et al. 2020; Yuen et al. 2020), these mutations may also be the result of concerted selective pressure(s), following that single mutational changes can have pleiotropic effects on distinct viral phenotypes and fitness components (de Wilde et al. 2018).

Identifying adaptive convergence raises the possibility of predicting mutational pathways in viruses important to global health (Gutierrez et al. 2019). When applied to SARS-CoV-2, our results reveal that some of the mutations we had initially identified as potentially relevant back in May 2021 (see (Escalera-Zamudio et al. 2021) had already been observed in other betacoronaviruses that circulate endemically in humans (Table 1), and some now display dominant trends in SARS-CoV-2 (as analysed up to December 2022). For example, amino acid state ‘R’ at residue 21 of the S protein (sites 21623) ([https://nextstrain.org/groups/neherlab/ncov/global?c=gt-S_21](https://nextstrain.org/groups/neherlab/ncov/global?c=gt-S_21)) and ‘P’ at residue 25 (site 21635) ([https://nextstrain.org/groups/neherlab/ncov/global?c=gt-S_25](https://nextstrain.org/groups/neherlab/ncov/global?c=gt-S_25)) have dominated across time. Moreover, mutation ‘D’ to ‘Y’ observed at residue 796 of the S protein (site 23948) has proven to be a successful mutational pathway, evidenced the replacement of amino acid state ‘D’ (previously observed for the B.1.617.2 lineage, Delta VOC and descending sub-lineages) by
‘Y’ (now observed for the B.1.1.529 lineage, Omicron VOC and descending sub-lineages) 
(https://nextstrain.org/groups/neherlab/ncov/global/?c=gt-S_796). Of interest, mutations at 
residue 796 of the S protein have been linked to the emergence viral variants that display 
reduced susceptibility to neutralizing antibodies (Kemp et al. 2021).

Epistasis is thought to have played a central role in the emergence of human-infecting 
betacoronaviruses (Holmes & Rambaut 2004). However, inferring epistasis across diverging 
viruses is difficult given the functional differences between homologous genes and proteins. 
Through our methodological approach we cannot measure epistasis *per se*, but we can aim to 
identify adaptive convergence and subsequently discuss its possible effects. Thus, our results 
indirectly provide support for epistasis, in the sense that if the same amino acid changes are 
observed in different virus species, then associated epistatic interactions are expected to be 
shared. This is of particular importance when considering the potential role of epistasis in 
antigenic drift, where the combined effect of independent mutations could contribute to antigenic 
escape (Rochman et al 2022). In the context of our findings, sites 21623 and 21635 are 
presumed to be involved in the antigenic drift of embecoviruses. As these residues are in close 
proximity to each other (displaying a linked evolution), these could thus reflect epistatic 
interactions. Nevertheless, within the SARS-CoV-2 S1[^ACE2] interface, epistasis seems to play 
a limited role, as the effect of multiple mutations seems to be additive rather than epistatic 
(Rochman et al. 2022; Zahradník et al. 2021; Starr et al. 2022).

The mutational spectrum of SARS-CoV-2 is known to be impacted by the human host 
apolipoprotein B mRNA-editing enzyme (APOBEC) family (Di Giorgio et al. 2020). The activity 
of APOBEC induces C → U/T mutations in the viral genome through a cytidine deaminase 
activity, likely resulting in a high degree of apparent homoplasy reflected in emerging mutations 
2021). Relative to more commonly used strategies for identifying homoplasy within single virus 
species, our methodology poses an alternative approach that aims to identify homoplasy *across 
and within* virus taxa, represented by shared mutations most likely fixed under an evolutionary 
scenario driven by selection (see Supplementary Text 5). Given that candidate mutations are 
observed over longer evolutionary times, this approach represents a useful tool to decrease the 
likelihood of erroneously scoring mutations as homoplastic (such as those resulting from 
mutational biases inherent to the SARS-CoV-2 genome evolution).

However, identifying adaptive convergence faces several important limitations. First, the 
methodology we use is conservative, as it is based on strict homology. In this context, we only 
consider sites robustly identifiable as homologous that can be traced back to ancestral nodes
with confidence (consequently excluding highly divergent genes). Therefore, our approach may result in an underestimation of sites that may putatively denote adaptive convergence across highly divergent viruses. Moreover, a limited virus genome sampling across time and space (in particular for HKU1 and SARS-CoV-1), coupled with a relatively low genetic diversity observed for SARS-CoV-2 (Rausch et al. 2020), further restricts the potential to identify shared mutations across virus species (van Dorp et al. 2020). In addition, there is some uncertainty associated with the mutations identified, as (though unlikely given cumulative evidence derived from different methodological approaches) it is not possible to rule out that some of these may still derive from biological processes other than adaptation (such as founder effects, mutational hitch-hiking, linkage, and toggling at hypervariable sites) (Kosakovsky Pond et al. 2012; Delport et al. 2008; De Maio et al. 2021; Wang et al. 2020; Simmonds 2020). Finally, whilst our analysis provides insights into coronavirus evolution in humans, our approach renders us unable to identify mutations that may result from host switching events. This is due to analyses on nodes representing ancient host switching events (Corman et al. 2018) being constrained by long divergence times, differences in mutation rates across virus taxa in different animal hosts, mutational saturation, and by a considerable under-sampling of betacoronaviruses circulating in non-human hosts (Holmes & Rambaut 2004; De Maio et al. 2021).

In this sense, additional/future experimental data could help reveal the impact of mutations on viral fitness. However, performing such studies may be difficult, as these concern potential gain-of-function experiments. Alternatively, enhanced genomic surveillance of betacoronaviruses infecting the human population and of those circulating in other animal host may confirm whether the mutational pathways we identify here represent evolutionary trajectories on which betacoronaviruses converge in their adaptation process to the human host.

MATERIAL AND METHODS (1933)

1. Initial data collation

When this manuscript was first deposited as a preprint (May 2021) (Escalera-Zamudio et al. 2021), complete genomes for all HKU1, OC43 and SARS-CoV-1 viruses sampled across different geographical regions and time were downloaded from the Virus Pathogen Resource (ViPR-NCBI 2021) (Supplementary Data 4). Sequences were removed if meeting any of the following criteria: (i) being >1000nt shorter than full genome length, (ii) being identical to any other sequence, or (iii) if showing >10% of site were ambiguities (including N or X). A total of 53 HKU1, 136 OC43 and 40 SARS-CoV-1 sequences were initially retained for analyses. For SARS-CoV-2, to better reflect an early zoonotic process into the human population (MacLean et
we originally aimed to limit the genetic diversity of the sampled virus population to the first wave of infection recorded during the pandemic. For this, ~23,000 full genomes sampled worldwide before May 2021 available in the GISAID platform (GISAID 2021) were downloaded and aligned as part of an initial public dataset provided by the COG-UK consortium (COG-UK 2021) (Supplementary Data 4). To make local analyses computationally feasible, the original SARS-CoV-2 dataset was randomly subsampled to ~5% of its original size, keeping the earliest genomes, and further reducing the dataset under the quality criteria stated above. In total, 1,120 SARS-CoV-2 sequences were retained. For all virus species considered, we focused only on genomes derived from human cases, in order to reflect host-specific adaptation processes.

2. Phylogenetic analyses

Only the main viral ORFs (Orf1ab and S) were used for further phylogenetic analyses, as these are homologous amongst the four viral species studied, and encode proteins essential to certain stages of the virus life cycle (i.e., replication and entry). For each virus species, individual ORFs (codons) were extracted and aligned as translated amino acid sequences using MAFFT v7.471 (to be then reverted to codons again) (Katoh & Standley 2013). Individual alignments were concatenated to further generate species-specific concatenated Orf1ab+S alignments. The concatenated alignments were then combined to generate a global alignment comprising all virus species, that was re-aligned again at an amino acid level using a profile-to-profile approach following taxonomic relatedness (Wang & Dunbrack Jr 2004). The final alignment was reverted to codon sequences as input for all further analyses. The global alignment comprised in total 1,314 sequences and 26,883 sites.

Maximum Likelihood phylogenies were estimated for the individual and global codon alignments using RAxML v8 (Stamatakis 2015), under a general time reversible nucleotide substitution model and a gamma-distributed among-site rate variation (GTR+G). Branch support was assessed using 100 bootstrap replicates. All trees were midpoint-rooted, whilst general phylogenetic patterns observed amongst these distantly related virus species were validated by comparing to previously published phylogenies (Woo et al. 2010; Zhu et al. 2018; Lau et al. 2011; Oong et al. 2017; Woo et al. 2006; Bedford 2021). Recombination is known to be common amongst betacoronaviruses (Oong et al. 2017; Woo et al. 2006; Su et al. 2016), including SARS-CoV-2 (Gutierrez et al. 2022; Turakhia et al. 2022). However, recombinant sequences were not removed at this step, as it was important to detect potentially recombinant isolates that could display relevant mutations. Putative recombinant sequences were eventually removed for subsequent analyses (when identified, see Methods sections 6 and 7).
3. Identifying homoplasy and/or stepwise evolution

Following the pipeline described by Escalera & Golden (Escalera-Zamudio et al. 2020), variable sites across different virus taxa were identified within the global alignment as those displaying non-synonymous mutations (rendering amino acid changes) occurring in at least ≥1% of the sampled sequences. Variable sites were extracted by masking columns across the alignment showing identical sites and at least 50% gaps, followed by the 'Find Variations/SNPs' function used to compare each site to consensus sequences generated under a 95% threshold with Geneious Prime v2020.0.4 (Kearse et al. 2012). A total of 6681 variable sites were identified and used to infer ancestral amino acid state reconstructions onto the nodes/internal branches of the global tree (see Methods section 2 above). This was done using TreeTime (Sagulenko et al. 2018) under a ML approach (RAS-ML) using a time-reversible model (GTR) for state transitions. The genetic variability observed within leaves/tips of the tree was deliberately excluded, in order to only analyse changes occurring within nodes or internal branches. In parallel, conserved sites were identified as those present in ≥ 99% of the sampled virus sequences. Conserved sites were extracted by reversing the 'variable site masking', to obtain only identical sites identified across the global alignment (Supplementary Data 2).

The resulting 6681 ‘Ancestral Reconstruction Trees’ (named here ARTs) were then classified under a computational algorithm developed to sort mutational patterns based on whether or not they support homoplasy and/or stepwise evolution. Briefly, homoplasy can occur within nodes of single clade or across clades, in which the same amino acid change must be present in at least one internal node of any given clade, and in another internal node of the same/another clade. Clades with the same amino acid states must not share direct common ancestry. Conversely, stepwise evolution is represented as sequential mutations occurring at the same sites within a single clade. Any given site scored under putative ‘stepwise evolution’ must display changes between at least two different states (A→B), but without any immediate reversion (B→A). A description of the definitions used here for homoplasy and/or stepwise evolution are available as Supplementary Text 1 and Supplementary Figure 1. A description of all basic steps used in our algorithm, including a schematic representation, is available in the Supplementary Text 2, Supplementary Figure 2 and 3. Associated code is publicly available at https://github.com/nataliamv/SARS-CoV-2-ARTs-Classification.
4. Estimating \( \text{dN/dS} \)

Derived from the global alignment and tree, we estimated dN/dS (\( \omega \), the ratio between the non-
synonymous substitution rate per non-synonymous site and the synonymous substitution rate
per synonymous site) using the following site, branch and branch-site models: Mixed Effects
Model of Evolution (MEME), Fixed Effects Likelihood (FEL), and the fixed effects site-level
model (Contrast-FEL) (Kosakovsky Pond & Frost 2005; Kosakovsky Pond et al. 2021; Murrell et
al. 2012). For this, the alignment was partitioned into 14 putatively non-recombinant regions
using the Genetic Algorithm for Recombination Detection (GARD) (Kosakovsky Pond, Posada,
et al. 2006), with all subsequent analyses conducted on the partitioned data. As dN/dS models
use the GTR component for the nucleotide evolutionary rate, biased mutation rates are handled.
Further, to mitigate the inflation in dN/dS estimates that results from unresolved and/or
maladaptive evolution, testing for selection was again restricted to internal nodes/branches of
the phylogeny (Kosakovsky Pond, Frost, et al. 2006). Genome-wide comparison of dN/dS
estimates across viral genome regions was performed using the Branch-Site Unrestricted
Statistical Test for Episodic Diversification method (BUSTED) (Murrell et al. 2015). Finally, the
impact of changing biochemical properties at selected sites was further assessed under the
Property Informed Models of Evolution method (PRIME) (HyPhy 2013). Our results were further
compared to the selection analysis available for independently sampled SARS-CoV-2 genome
data available as of December 2022 (https://observablehq.com/@spond/evolutionary-

5. Mapping mutations onto betacoronavirus protein structures

To locate the non-synonymous mutations identified on viral protein regions of known function,
corresponding residues were mapped to available structural data using PyMOL v 2.4.0
(https://pymol.org/2/) (Supplementary Table 2, see Data Availability section). Mutations were
analysed in the context of their relative proximity to previously reported functional regions, and
to each other. N-linked glycosylation sites in S protein sequences were identified by searching
for the N-[not P]-[S or T] consensus sequence (Watanabe et al. 2019). None of the mutations
identified in this study resulted in generation or deletion of N-linked glycosylation sequons. In
parallel, conserved and variable sites identified (including the 30 mutations evidencing
homoplasy and/or stepwise evolution across virus species) were mapped onto published protein
structures available for the S proteins of the four human-infecting betacoronaviruses studied
here (Figure 5, Supplementary Figure 7). Finally, to compare dN/dS distributions between
specific domains of the S protein within and across virus species, sites inferred to be under
positive or negative selection (PSS, NSS) were mapped onto S protein structures (Supplementary Data 2).

6. Validation through resampling and by comparing mutational distributions
To validate our initial observations derived from virus genomes sampled up to May 2021, we sought to determine if the 30 mutations that had been identified initially were also present in the expanded embecov- and sarbecovirus diversity sampled up to July 25th 2022 (corresponding to the final sampling date of this study). Virus diversity now included genome sequences derived from more recently collected human isolates (only made publicly available after our initial sampling), and from other closely related embeco- and sarbecoviruses from non-human hosts. The expanded alignment comprises 1455 sequences (~700 embecovirus + SARS-CoV and ~700 SARS-CoV-2), resulting in 27503 columns that were re-aligned under a progressive profile-to-profile approach based on taxonomic relatedness to be further used to estimate an expanded 'Maximum Likelihood' tree (following Methods section 2). To additionally explore if the mutations identified were also present in a larger dataset representing an expanded SARS-CoV-2 diversity (sampled up to July 25th 2022), a set of 1400 SARS-CoV-2 genomes denoting 'evolutionary successful' virus lineages (Supplementary Table 3) was examined independently (Supplementary Text 3, Supplementary Figure 4 and 5). Both datasets were analysed following the steps described in Methods Section 2 and 3, specifically searching for the mutations listed in Table 1. Virus taxa included in both re-sampled datasets are listed in Supplementary Data 5. A full description of the sequence subsampling and methodological approach used is available as Supplementary Text 3, and Supplementary Figures 4 and 5.

We further sought to explore if the proportion of mutational patterns we classified as putatively denoting homoplasy and/or stepwise evolution were more likely to arise from an evolutionary scenario mostly driven by selection, compared to ‘random’ mutational patterns derived from evolutionary scenarios generally driven by genetic drift. For this purpose, the expanded alignment was translated to amino acid sequences and used to simulate three alignments with 'AliSim' (http://www.iqtree.org/doc/AliSim) under the 'mimick real alignment' function (mimicking a ‘real’ evolutionary process based on amino acid evolution under a LG model, and applied to the inputted original tree). To compare the corresponding proportion of sites scored under homoplasy and/or stepwise evolution, each dataset (the expanded and three simulated alignments) was analysed following the steps described in Methods section 3. The classification of mutational patterns within expanded and simulated datasets also serves the purpose of validating our algorithm, originally developed for analysing the global dataset (that
included only OC43, HKU1, SARS-CoV-1 and SARS-CoV-2 sampled from the human host). Associated results and a brief discussion are available as Supplementary Text 5.

7. Reconstruction of amino acid evolution for selected sites
To further confirm our ML-derived results (see Methods section 3), for those mutations displaying cumulative evidence of adaptive convergence (18121, 21623, 21635 and 23948, Table 1), we used the expanded dataset to infer ancestral states under a Bayesian framework. For each site, we first estimated a MCC (maximum clade credibility) tree from the resampled codon alignment using an SRD06 substitution model (Shapiro et al. 2006) and a strict molecular clock. Coded amino acid traits were then mapped onto the nodes of the MCC tree by performing reconstructions of ancestral states under an asymmetric discrete trait evolution model (DTA) in BEAST v1.8.4 (Lemey et al. 2009; Suchard et al. 2018). The DTA model was run using a Bayesian Skygrid tree prior for 100X10^6 generations and sampled every 10,000 states until all DTA-relevant parameters reached an ESS >200. For illustrative purposes, Figure 3 only shows sites 18121, 21623 and 23948. The amino acid evolution pattern observed for site 21635 is available in Supplementary Data 3.
FIGURE LEGENDS

Fig 1. Phylogenetic tree of human-infecting betacoronaviruses. The expanded tree estimated from the Orf1ab+S alignment comprising 1455 sequences (see Methods section 6), summarizing the phylogenetic pattern observed for four distantly related human-infecting betacoronaviruses: HKU1, OC43, SARS-CoV-1 and SARS-CoV-2. MERS and related virus sequences were included in the tree for rooting purposes only. Both the Embecovirus subgenus (HKU1 and OC43 and related viruses) and the Sarbecovirus subgenus (SARS-CoV-1 and SARS-CoV-2 and related viruses) are indicated, showing the positioning of the most closely related virus genome sequences derived from animal isolates (when available). The different genotypes identified for the HKU1 (A, B and C) and for the OC43 (A–H) are shown in Supplementary Data 1.

Fig 2. Distribution of conserved/variable sites with S across different virus species (a) Top-down (upper panel) and side view (bottom panel) of a cartoon representation of the multidomain architecture of the trimeric SARS-CoV-2 S ectodomain (PDB: 6VXX). The S1 subunit is coloured according to the different protein domains: S1A in cream, S1B in teal, S1C in orange, and S1D in blue, whilst the S2 subunit is shown in grey. (b) Top-down and side views of sphere-based representations of trimeric S protein ectodomains for the viruses studied: SARS-CoV-2 (PDB: 6VXX), SARS-CoV-1 (PDB: 6ACC), OC43 (PDB: 6OHW) and HKU1 (PDB: 5I08). The sphere-based representation shows conserved (shown in grey; residues present ≥99% of all sequences) and variable sites (blue; residues present in ≥1% of all sequences) across virus species. Variable sites identified as denoting homoplasry or stepwise evolutionary patterns are shown in red (see Methods section 3). The asparagine residues of N-linked glycosylation sequons are indicated in purple.

Fig 3. Reconstruction of amino acid evolution at selected sites. Maximum clade credibility (MCC) trees showing the evolutionary trajectories for different amino acid states observed for three illustrative sites (18121, 21623 and 23948) that (i) display evidence of homoplasy and/or stepwise evolution, (ii) show evidence of positive selection, and (iii) are proximal to regions of established protein function. The reconstructions of ancestral states for these sites show different amino acid states at nodes (represented by circles in different colours). The posterior probability for a given amino acid state occurring at a given node of interest is indicated. Sites 18121 display evidence of homoplasry across virus lineages, site 21623 shows evidence of both homoplasry across species and stepwise evolution within single virus species (i.e. OC43), and site 23948 shows evidence of stepwise evolution within single virus species (i.e. SARS-CoV-1), and also of homoplasry across virus species (i.e. SARS-1 and SARS-CoV-2).

Fig 4. Residue Ser28 of nsp14 is situated near the nsp14-nsp10 interface. Cartoon representation of the SARS-CoV-1 nsp14-nsp10 protein complex (PDB: 5C8S) with Ser28 (corresponding to site 18121 in SARS-CoV-2 genome coordinates) shown as a red sphere. This residue is located within the nsp14 ExoN domain (cream) and is approximately 9 Å from the interface with nsp10 (light blue, the proximal nsp10 residue Cys41 was used to calculate the distance and is indicated as a sphere). The distance between nsp14’s Ser28 and nsp10’s Cys41 is annotated and indicated by a dashed black line. Zoomed-in panel: detailed representation of the intra-nsp14 hydrogen-bond between the side chain of Ser28 and the main chain of Thr25 (identified with the PISAebi server). The side chain of Ser28 is indicated as a red stick and Thr25 is indicated in sticks and coloured according to atom (C, cream; O, red; N, blue). The hydrogen-bond is indicated as a dashed black line.

Fig 5. Mapping of mutations exhibiting homoplasy onto the S protein structure of SARS-CoV-2. Top-down (left) and side view (right) of a cartoon representation of the multidomain architecture of the trimeric SARS-CoV-2 S ectodomain (PDB: 6VXX). The S2 subunit is highlighted in grey and the S1 ectodomain is divided into S1A (highlighted in cream), S1B (teal), S1C (orange), and S1D (blue) domains, following the colour scheme in Figure 3. Homoplastic mutations that co-localize to known functional surfaces (see Supplementary Table 4) are indicated in the structure and coloured in groups: Arg21 (corresponding to site 21623 in SARS-CoV-2 genome coordinates, in green), Pro25 (site 21635, in green), Asp796 (site 23948, in yellow), Ile1418 (site 24614, in red), Ala1020 (site 24620, in red) and Leu1024 (site 24632, in red). All representations are shown with a transparent protein surface for clarity.
ACKNOWLEDGEMENTS
MEZ was supported by Leverhulme Trust ECR Fellowship (ECF-2019-542). RJGH is supported by the European Molecular Biology Organisation (ALTF 869-2019). RJGH and TAB are supported by the Medical Research Council (MR/S007555/1). The Wellcome Centre for Human Genetics is supported by Wellcome Trust grant 203141/Z/16/Z. SKP is supported in part by the NIH (grants AI134384, AI140970, GM110749) and the NSF (grant 2027196). JT and RPDI were supported by European Union’s Horizon 2020 project MOOD (874850). LvD is supported by a UCL Excellence Fellowship. HGCS is supported by funding through the “Vigilancia Genómica del Virus SARS-CoV-2 en México” grant from the National Council for Science and Technology-México (CONACyT). Author contributions: MEZ and OGP designed research. MEZ and RJGH performed research. MEZ, RJGH, BG, SKP, LvD, JT, RPDI and HGCS analysed data. NM developed the code for implementing the computational pipeline. OGP and TAB supervised data analysis. MEZ and RJGH wrote the manuscript, with comments from all authors. We thank Dr. Louis Du Plessis for his help with the phylogenomic analysis and Dr. Nicola De Maio for constructive comments on our analytical approach.

COMPETING INTERESTS
The authors declare no competing interests.

DATA AVAILABILITY
Taxa IDs and accession numbers for virus sequences used in this study are provided in the Supplementary Data 4 and 5 files. All SARS-CoV-2 genome sequences and associated metadata used in this study are published in GISAID’s EpiCoV database under the EPI SET GISAID Identifier: EPI_SET_230131zy. To view the contributors of each individual sequence with details such as accession number, virus name, collection date, originating and submitting lab, as well as the list of all authors, visit 10.55876/gis8.230131zy. PDB files used are listed as follows: S protein (HKU1 PDB:5I08, OC43 PDB:6OHW, SARS-CoV-1 PDB:6ACC and SARS-CoV-2 PDB:6VXX, 6ZGI). Orf1a (SARS-CoV-1 nsp3 PDB:2W2G). Orf1b (SARS-CoV-2 nsp13 PDB:6XEZ, SARS-CoV-1 nsp14 PDB:5C8S and SARS-CoV-2 nsp15 PDB:6WLC). Full code for our algorithm is available as open source: https://github.com/nataliamv/SARS-CoV-2-ARTs-Classification. An interactive notebook with our full selection analysis results is available at https://observablehq.com/@spond/beta-cov-analysis.


### Table 1. Potentially relevant sites across human-infecting betacoronaviruses

<table>
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<th>SARS-CoV-2 genome coordinates</th>
<th>ORF</th>
<th>Protein/Residue</th>
<th>Ancestral LNA</th>
<th>OC43</th>
<th>HKU1</th>
<th>Ancestral LNB</th>
<th>SARS-CoV-1</th>
<th>SARS-CoV-2</th>
<th>Expanded</th>
<th>1400 SARS-CoV-2</th>
<th>Homoplaspy (H)/Stepwise Evolution (SWE)</th>
<th>Selection across species (Method, p-value)**</th>
<th>Selection in SARS-CoV-2 (recent amino acid changes)</th>
<th>Epitope</th>
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† Positions indicate the start of the codon for reference genome Wuhan-Hu-1 (NC_045512.2). Sites in bold refer to those highlighted in the results section.
§ Sites/branches scored under MEME/FEL and Contrast-FEL (CF); CF tests for differences in selective pressures between clades.
\[\]

Representing virus diversity sampled as of May 2021
\[\]

Representing viral diversity sampled as of December 2022 available from: https://observablehq.com/@spond/sars_cov_2/sites
\[\]

* Potential T cell epitopes derived from HLA class I and HLA-DR SARS-CoV-2 binding peptides (Campbell et al. 2020; Satpute et al. 2021)
Figure 2
159x74 mm (x DPI)
Figure 3
159x54 mm (x DPI)
Figure 4
77x65 mm (x DPI)
Figure 5
125x62 mm (x DPI)