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There are genetic risk factors that influence the outcome of COVID-19 [COVID-19 Host Genetics Initiative, *Nature* 600, 472–477 (2021)]. The major genetic risk factor for severe COIVD-19 resides on chromosome 3 and is inherited from Neandertals [H. Zeberg, S. Pääbo, *Nature* 587, 610–612 (2020)]. The risk-associated DNA segment modulates the expression of several chemokine receptors, among them CCR5, a coreceptor for HIV which is down-regulated in carriers of the COVID-19 risk haplotype. Here I show that carriers of the risk variant have an ~27% lower risk of HIV infection.

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The major genetic risk factor for severe COVID-19 was introduced into modern human populations via gene flow from Neandertals 50,000 to 70,000 y ago. Although there is no direct evidence for positive selection on the risk haplotype, it has increased in frequency since the Last Glacial Maximum (1) and is unusually common today, reaching carrier frequencies of 16% and 50% in Europe and South Asia, respectively (2). Given the prevalence of this genetic variant, it is of interest to consider whether it may offer protection against some pathogen other than severe acute respiratory syndrome coronavirus 2, either today or in the past.

The major genetic risk factor for COVID-19 severity is located on chromosome 3 in a genomic region encompassing a gene cluster encoding chemokine receptors (see Data Availability for an interactive diagram of the association in this genomic region). The chemokine genes CCR1, CCR2, CCR3, CCR5, CCR9, XCR1, and CXCR6 are all located within 0.55 megabases of the genetic variant (3) likely to confer risk for severe COVID-19 (rs17713054, chr3:45859651:G/A, hg19). Using expression data from whole blood of ~30,000 individuals (4), I find that all of the chemokine receptor genes, with the exception of XCR1, are differentially expressed in carriers of the risk variant (Table 1; P < 1e-6), with reduced expression for all genes except for CCR9. All associations pass genomewide correction for multiple comparisons ($p_{FDR} < 0.01$) and are also seen in a smaller dataset containing 1,331 individuals (Table 1).

One of the most well-studied genetic variants modulating infectious disease risk is a 32-base pair deletion that introduces a premature stop codon in *CCR5*, resulting in a nonfunctional receptor. This mutation, *CCR5*- Δ 32, which confers protection against HIV-1 infection and likely also against smallpox, has been positively selected (5). HIV-1 relies on CD4 for viral entry and commonly requires also the chemokine receptors CCR5 and/or CXCR4 as coreceptors, even if it can also utilize CCR3 (6) and CXCR6 (7).

Macrophages, dendritic cells, and memory T cells patrolling the mucosa are the first cells infected by HIV-1 (8). Indeed, a major route for HIV-1 infection among men is urethral macrophages (9), and HIV-1 can be present in macrophages in the vagina (10). Strikingly, the down-regulation of *CCR5* in carriers of the COVID-19 risk variant on chromosome 3 is primarily seen in macrophages, monocytes (which are related to dendritic cells), and memory T cells (Table 1).

Since individuals carrying the major genetic risk factor for COVID-19 have lower CCR5 as well as CCR3 and CXCR6, I hypothesized that they might have lower prevalence of HIV infection. A limiting factor for such an analysis is the low prevalence of HIV in European cohorts available for analysis. To increase statistical power, I conducted a meta-analysis of three of the largest currently available biobanks, the UK Biobank (11), the Michigan Genomics Initiative, and FinnGen. Together these cohorts contain 591 European individuals with HIV infection and 667,215 controls. In all three cohorts, carriers of the risk allele for COVID-19 have a risk ratio for HIV infection between 0.66 and 0.83 (Fig. 1). Meta-analysis of the three cohorts results in an overall risk ratio of 0.73 (95% CI: 0.59 to 0.90, P = 4.1e-3). Thus, carriers of the chromosome 3 COVID-19 risk allele has a 27% reduction in risk for HIV infection (95% CI: 9 to 40%). There was no detectable heterogeneity across the cohorts ($I^2 = 0\%$, P = 0.82).

Sequence diversity in the envelope spike of HIV-1 is known to influence coreceptor use (12). Moreover, selective pressures arising from host genetic variation affect HIV-1 sequence variation (13). I therefore investigated whether variation in the HIV-1 envelope gene Env is associated with carrier status of the COVID-19 risk haplotype. There are 34 genetic variants that cosegregate $(r^2 > 0.5)$ with rs17713054 among individuals in the 1000 Genomes Project (14). In a published dataset (13), four of these variants (rs17764980, rs17714101, rs17714228, and rs71325092) are associated (P < 7e-4) with the amino acid replacement M115L in the HIV-1 envelope protein Gp41 encoded by Env. Specifically, carriers of the risk variant for COVID-19 were more likely to be infected with a virus variant carrying a leucine residue at position 115 in Gp41 (odds ratio, OR = 1.3), a viral variant less sensitive to the fusion inhibitor enfuvirtide (15). That the association between the host genome and the viral genome involves the coreceptor and the envelope spike suggests that the protective effect of the haplotype is mediated by reduced viral entry.

The major genetic risk factor for COVID-19 rose in frequency between 20,000 to 10,000 y ago (1); since this significantly predates the HIV pandemic, it is unlikely that this increase in frequency resulted from positive selection driven by HIV. I can only speculate about the pathogen that exerted the genuine selective pressure on this allele. Variola virus emerged more than 10,000 y ago (16), making smallpox a likely candidate, whereas as *Yersina pestis* emerged later, ~7,000 y ago (17). I also note that the highest allele frequencies today (2) coincide with regions where cholera is endemic.

Author contributions: H.Z. designed research, performed research, analyzed data, and wrote the paper.

The author declares no competing interest.

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Table 1.	Differential express	sion of chemokine r	receptors for carriers	s of the major g	enetic risk factor for COVID-19

	eQTLgen	eQTL cat.	NES (eQTL cat.)	P value (eQTL cat.)	P value (eQTLgen)
Whole blood					
CCR1	\downarrow	\downarrow	-0.10	1.8e-2	8.7e-14
CCR2	Ļ	Ļ	-0.05	3.4e-2	5.0e-7
CCR3	Ļ	\downarrow	-0.31	8.3e-7	8.8e-47
CCR5	\downarrow	\downarrow	-0.05	1.7e-1	1.6e-7
CCR9	Ť	↑	+0.13	2.6e-2	2.7e-7
CXCR6	Ļ	Ļ	-0.16	1.2e-3	7.9e-44
CCR5		eQTL cat.	NES (eQTL cat.)	P value	Study ref.
Macrophages		\downarrow	-1.15	7.1e-5	19
Monocytes		\downarrow	-0.59	1.7e-3	20
Memory T-follicular helper cells		\downarrow	-1.09	5.4e-3	21
Memory T helper 2 cells		\downarrow	-0.76	8.7e-3	21
Memory T helper 1 cells		\downarrow	-0.34	2.6e-2	21
Monocytes		\downarrow	-0.31	3.1e-2	22
CD8+ T cells		\downarrow	-0.69	5.3e-2	21
Macrophages		\downarrow	-0.24	5.5e-2	23
Memory T helper 1/17 cells		\downarrow	-0.24	1.5e-1	21
NK cells		\downarrow	-0.42	1.6e-1	21
Monocytes		\downarrow	-0.20	2.1e-1	21
CD16+ Monocytes		\downarrow	-0.45	3.4e-1	21
T cells		Ļ	-0.16	4.0e-1	20
Memory T regulatory cells		Ļ	-0.07	6.2e-1	21
T cells		Ļ	-0.07	6.6e-1	24
T regulatory cells		Ļ	-0.10	7.4e-1	21
Memory T helper 17 cells		ļ	-0.004	9.8e-1	21

Downward arrow and negative normalized effect sizes (NES) represent reduced expression for carriers of the COVID-19 risk allele at rs17713054. NES available from the eQTL catalog. The risk allele for COVID-19 is associated with reduced expression of all receptors except CCR9 in whole blood. Expression data from whole blood aggregated by the eQTLGen Consortium includes 26,000 to 31,569 samples, whereas the eQTL catalog comprises 1,331 whole blood samples (meta-analyzed using inverse-variance weighting of the contributing studies). CCR5 is down-regulated in all investigated immunological cell types (19–24). For CCR5 in macrophages, the NES translates to an allelic fold change of ~3.7 (log₂) based on the transcript levels (19).

The association described here highlights that gene flow from Neandertals was a double-edged sword. Whereas this genetic variant has had tragic consequences during the last 2 y in the COVID-19 pandemic, it appears to have offered considerable protection against HIV during the last 40 y. Its role in past and future pandemics remains to be seen.

Data Availability. Summary statistics for HIV infection among people of primarily European descent were obtained from FinnGen (freeze 5, https://r5. finngen.fi), Michigan Genomics Initiative (freeze 2, https://pheweb.org/MGIfreeze2), and UK Biobank (https://pan.ukbb.broadinstitute.org/). Covariates included in the analysis are age, sex, and the first four principal components (Michigan Genomics Initiative) or the first 10 principal components (UK Biobank and FinnGen). In addition, Michigan Genomics Initiative and UK Biobank controlled for genetic relatedness, and FinnGen and Michigan Genomics Initiative included genotyping batch and chip version, respectively. The HIVsummary statistics provided by these resources are on the OR scale but were here directly translated to risk ratios under the rare disease assumption. Summary statistics for expression differences were obtained from the eQTLGen Consortium (https://www.eqtlgen.org) and the eQTL Catalogue (https://www. ebi.ac.uk/eqtl/). Data were meta-analyzed using inverse-variance weighting. Linkage disequilibrium was calculated using data from the 1000 Genomes Project (15). The data on the interaction between host and viral genetic sequence variation were taken from the supplementary material of a previous study (14) which had been deposited online (https://dx.doi.org/10.5281/zenodo.

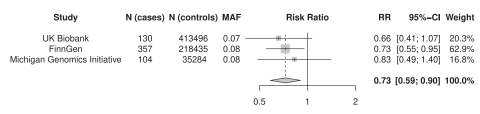


Fig. 1. Risk ratios for HIV disease for carriers of the major genetic risk factor for COVID-19. A meta-analysis by inverse-variance weighting of 591 individuals with HIV disease and 667,215 controls. Risk ratios (OR) are calculated for a linear model with effect estimates per copy of the Neandertal-derived allele at rs17713054. The MAF column shows the minor allele frequency in the cohorts. Meta-analysis was performed in R (version 4.1.2) using the package meta (version 5.1).

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7139). An interactive diagram of the association for the major risk factor for COVID-19 (COVID-19 Host Genetics Initiative, release 6) has been deposited on LocusZoom (https://my.locuszoom.org/gwas/815606/) (18).

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