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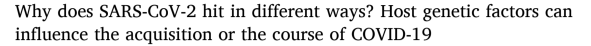
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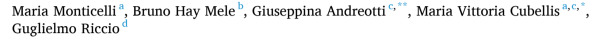
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### Review





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### ABSTRACT

The identification of high-risk factors for the infection by SARS-CoV-2 and the negative outcome of COVID-19 is crucial. The genetic background of the host might account for individual responses to SARS-CoV-2 infection besides age and comorbidities. A list of candidate polymorphisms is needed to drive targeted screens, given the existence of frequent polymorphisms in the general population.

We carried out text mining in the scientific literature to draw up a list of genes referable to the term "SARS-CoV\*". We looked for frequent mutations that are likely to affect protein function in these genes. Ten genes, mostly involved in innate immunity, and thirteen common variants were identified, for some of these the involvement in COVID-19 is supported by publicly available epidemiological data. We looked for available data on the population distribution of these variants and we demonstrated that the prevalence of five of them, Arg52Cys (rs5030737), Gly54Asp (rs1800450) and Gly57Glu (rs1800451) in MBL2, Ala59Thr (rs25680) in CD27, and Val197Met (rs12329760) in TMPRSS2, correlates with the number of cases and/or deaths of COVID-19 observed in different countries. The association of the TMPRSS2 variant provides epidemiological evidence of the usefulness of transmembrane protease serine 2 inhibitors for the cure of COVID-19. The identified genetic variants represent a basis for the design of a cost-effective assay for population screening of genetic risk factors in the COVID-19 pandemic.

### 1. Introduction

Many people affected by COVID-19 develop fever, cough, fatigue, shortness of breath, muscular pain, and headache. In the most severe cases medical conditions, such as respiratory failure, occur and eventually lead to death.

Age and comorbidities, such as cardiovascular diseases, diabetes, and chronic lung diseases, are known risk factors for worse outcomes (Fang et al., 2020). Furthermore, altered biomarkers, namely C-reactive Protein, high-sensitivity troponin T, prothrombin time, fibrin degradation products, D-Dimer, and CD4<sup>+</sup> count, help differentiate patients who will develop severe symptoms from those who will only be mildly affected (Zheng et al., 2020). These factors alone are not sufficient to

identify all subjects at high risk, and on top of that, they tend to overlap. Genetic factors, on the other hand, also influence risks associated with COVID-19 but are more likely to be independent of age-related comorbidities. Specific studies on the subject are very few and in most cases genes have been selected regardless of the existence of frequent polymorphisms in the general population, lowering their potential as markers for large scale genetic screenings (Hussain et al., 2020).

SARS-CoV-2 and SARS-CoV belong to the Betacoronavirus genus (Chen et al., 2020) and share the same cellular receptor, the angiotensin-converting enzyme 2 (ACE2), and very similar nucleotide sequences (Yan et al., 2020).

In this paper, we carried out a bioinformatic analysis to predict polymorphisms that could either influence the acquisition or the course

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of COVID-19. Text mining was carried out to extract a list of human genes that have been associated with the term "SARS-CoV\*" in the literature. We highlighted the polymorphisms of these genes that are very frequent in the general population and are likely to affect the function of the mutated protein. We hope that our prediction can guide and speed up experimental tests and help to find those high-risk subjects that have not been recognized by age, co-morbidities, or biomarkers so far.

# 2. Question: which host genetic factors are likely to influence sensitivity to covid-19?

High variability of symptoms and outcomes characterizes SARS-CoV-2 infection. The phenotypic spectrum of the disease ranges from asymptomatic individuals to patients requiring ventilation due to severe pneumonia (García, 2020). Different hypotheses have been put forward, the role of the host's genetic background is currently emerging (Godri Pollitt et al., 2020; Latini et al., 2020; Strafella et al., 2020; Zhang, S.-Y. et al., 2020). Data from a genomewide association (GWAS) study involving 1980 patients with COVID-19 and 2381 healthy control participants in Spain and Italy have been recently published (Group, 2020). Two loci were strongly associated to susceptibility to SARS-CoV-2, one on chromosome 9 covering the *ABO* blood group and another one on chromosome 3 covering six genes, *SLC6A20*, *LZTFL1*, *CCR9*, *FYCO1*, *CXCR6*, and *XCR1*.

We would like to contribute to gene-specific candidate-driven studies by suggesting missense variants that can cause severe symptoms or facilitate the infection and expand the list of identified by GWAS studies. We evaluated missense mutations that are frequent in the general population and occur in genes that do not cause haploinsufficiency. We selected those that are deleterious for the protein product and we excluded those associated to blood groups. A deleterious mutation on a single allele might not be enough to cause a life-threatening disease and hence it could be relatively frequent but might confer special sensitivity or resistance to viral infection. We excluded variants occurring in promoters, introns, or untranslated regions because their deleteriousness is less predictable. Hence our study will provide a subset of the genetic variants that are likely to affect the sensitivity to infection or the course of COVID-19.

### 2.1. Procedure to identify candidate variants

We obtained from the Exome Aggregation Consortium (ExAC) a list of human genes (14995) with a pLI <0.9. pLI is a value that measures the probability of being loss-of-function intolerant (pLI). Genes with a pLI <0.9 have a low probability of causing haploinsufficiency when affected by the severe mutation. We developed a simple pipeline to use such a list of genes and corresponding proteins as input and look within the Scopus database for papers containing any of these protein names and the term "SARS-CoV\*" in the article's title, abstract, or keywords. The script, which is written in R language and exploits the rscopus package, is provided in Supplementary File S1. We excluded the genes covered by the GWAS analysis (Group, 2020) obtained a "SARS-CoV\* referable genes" list (106 genes; Supplementary File S2).

We searched within GnomAD for frequent variants, missense or loss-of-function, in "SARS-CoV\* referable genes", setting the threshold to allele frequency at 1%. Lastly, we used Polyhen2 (http://genetics.bwh. harvard.edu/pph2/), to exclude benign mutations. Frequent missense variants were found in 56 genes and deleterious frequent deleterious variants in ten genes. The scheme of the search is provided in Fig. 1.

The list of identified variants with their frequencies is reported in Table 1. We observed that the candidate variants were mostly in genes involved in innate immune defense. References to databases and programs utilized in this analysis are provide as a supplementary file S3.

## 2.2. Correlation between allele frequencies and COVID-19 cases and deaths per population

Data on the prevalence in different populations were available for six of our candidate variants. The datasets relative to rs12329760 (Val197Met, *TMPRSS2*), rs25680 (Ala59Thr, *CD27*), and rs3775291 (Leu412Phe, *TLR3*) were found in ALFRED; the data relative to the *MBL2* variants rs5030737 (Arg52Cys), rs1800450 (Gly54Asp), and rs1800451 (Gly57Glu) were reported by Verdu et al. Data on the population distribution of the remaining six variants were not available. Notwithstanding the limitations associated with publicly available data on the number of cases and deaths per country that are influenced by socio-economic factors, we found interesting results for five variants, namely Arg52Cys (rs5030737), Gly54Asp (rs1800450), and Gly57Glu (rs1800451) in *MBL2*, Val197Met (rs12329760) in *TMPRSS2* and Ala59Thr (rs25680) in *CD27*.

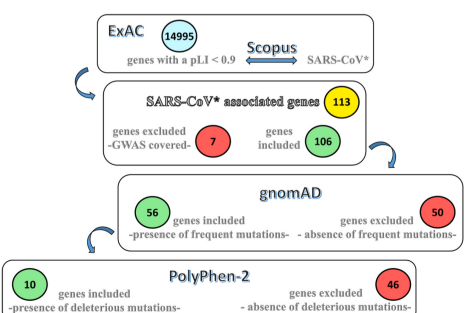


Fig. 1. Procedure to identify candidate variants The Exome Aggregation Consortium (ExAC) provided a list of human genes and annotated them with a value that measures the probability of being loss-offunction intolerant (pLI). The names of the genes were associated to the names of the corresponding proteins. Such list was used as the input to look within the Scopus database for papers containing any of the protein names and the term "SARS-CoV\* in the article's title, abstract or keywords. "SARS-CoV referable genes", encompassing frequent variants, missense or loss of function, were identified setting the threshold to allele frequency to 1%, and excluding variants that did not pass quality control or occur in non-canonical transcripts. The genes covered by the GWAS analysis (Group, 2020) were excluded. Among these genes, those encompassing frequent deleterious variants were identified.

**Table 1**Missense variants that might influence the acquisition or the course of COVID-19.

Gene name	Protein name/Mim phenotype	Reference SNP ID number	Amino acid change	Variant frequency	GWAS p-value
ACE	Angiotensin-converting enzyme	rs3730025	Tyr244Cys	0.0107	
AHSG	Alpha-2-HS-glycoprotein #203650	rs140827890	Ala164Thr	0.0114	0.05
CD27	CD27 antigen #615122	rs25680	Ala59Thr	0.1912	
GFAP	Glial fibrillary acidic protein #203450	rs59291670	Asp157Asn	0.11	
IFI30	Gamma-interferon-inducible lysosomal thiol reductase	rs11554159	Arg76Gln	0.2281	
MBL2	Mannose-binding protein C #614372	rs1800450	Gly54Asp	0.141	
		rs5030737	Arg52Cys	0.0558	
		rs1800451	Gly57Glu	0.0318	
MEFV	Pyrin #134610 #249100	rs3743930	Glu148Gln	0.0708	
TLR2	Toll-like receptor 2 #607948	rs5743704	Pro631His	0.0281	0.02
		rs5743708	Arg753Gln	0.0176	
TLR3	Toll-like receptor 3 #609423	rs3775291	Leu412Phe	0.2789	0.03
TMPRSS2	Transmembrane protease serine 2	rs12329760	Val197Met	0.2452	0.01528

Table 1: Likely deleterious frequent variants (allele frequency >1%) in genes referable to the term "SARS-CoV\*" in the literature. Variant frequency is referred to gnomAD total exome frequency obtained from wAnnovar annotation. For variants covered in the GWAS study, GCST90000255 Fisher p-values are provided.

MBL2 encodes a mannose-binding protein C that binds mannose, fucose, and N-acetylglucosamine on different microorganisms and activates the lectin complement pathway. The deficiency of mannosebinding protein C (MIM #614372) is associated with increased susceptibility to infection (Degn et al., 2011). MBL2 binds and activates the lectin complement pathway in viral infections such as HIV (Garred et al., 1997) and influenza (Kase et al., 1999). We tested three frequent missense mutations in MBL2: Gly54Asp (rs1800450), Arg52Cys (rs5030737), and Gly57Glu (rs1800451). All these mutations affect the ability of the protein to bind carbohydrate surfaces and MASP-2, the mannan-binding lectin serine protease 2 which activates the complement system (Turner, 2003); the lowest interaction is observed with Arg52Cys (Larsen et al., 2004). In a study conducted in China, it was proven that Gly54Asp (rs1800450) is significantly associated with susceptibility to SARS-CoV infection but not with disease severity (Ip et al., 2005; Zhang et al., 2005); unfortunately, the other two variants were not tested. We found that the allele frequency per population of Arg52Cys (rs5030737) correlates positively with the number of cases per country [p-value 0.0394] of COVID-19, but not with the number of deaths (Fig. 2, panels A and B), while the allele frequency per population of Gly54Asp (rs1800450) correlates positively both with the number of cases [p-value 0.0018] and the number of deaths [p-value 0.02] (Fig. 2, panels C and D). This finding supports the hypothesis that mannose-binding protein C plays a protective role and that MBL2 inactivation is a risk factor for SARS-CoV-2 infection, and that the immune response plays a role in the course of the disease.

Unexpectedly, the allele frequency per population of Gly57Glu (rs1800451) was found to be negatively correlated with the number of cases [*p*-value 0.0122] (Fig. 3, panel A). The correlation with the number of deaths was significant when the data from Iran were excluded [*p*-value 0.0241] (Fig. 3, panel B).

TMPRSS2 encodes transmembrane protease serine 2. Mouse models have shown that SARS, MERS CoVs and Influenza viruses use this protease to cleave the Spike protein and achieve viral internalization (Iwata-Yoshikawa et al., 2019). A negative correlation between the allele frequency of Val197Met and the number of cases [p-value 0.0012] and deaths [p-value 0.0057] per country (Fig. 4 panels A and B) was found. We observed the number of cases and deaths as a function of prevalence clusters in two groups and the high mortality group is associated with a low prevalence of the variant, indicating that genetic TMPRSS2 inactivation is a protective factor against SARS-CoV-2 infection and progression towards severe cases. A lower frequency of

Val197Met in Italian COVID-19 patients compared to the allelic frequency of this variant in GnomAD for the EUR reference population has recently been reported (Latini et al., 2020). Val197Met is a SNP covered by the GWAS study (Group, 2020) and it is significantly associated with susceptibility to (*p*-value < 0.05). TMPRSS2 is a druggable protein that can be inhibited by camostat mesylate, a drug approved in Japan, by bromhexine and nafamostat. Clinical trials are ongoing for their usage in COVID-19 (ClinicalTrials.gov Bromexhine: NCT04355026. NCT04273763. NCT04340349: Camostat: NCT04321096. NCT04355052, NCT04353284, NCT04338906, NCT04470544; Nafamostat: NCT04352400: NCT04473053). Our data provide epidemiologic evidence of the role of the protease in SARS-CoV-2 infection and encourages the usage of protease inhibitors for the therapy (Hoffmann et al., 2020).

CD27 receptor is thought to have an essential role in connecting the CD4 and CD8 branches of the immune system. Unbalances between the two branches has been associated with a worse prognosis in SARS-CoV infections (Li et al., 2008). Known pathological mutations of CD27 exist, and they are associated with cytokine storms (Amberger et al., 2009) and hemophagocytic lymphohistiocytosis (HLH) like syndromes that frequently characterize severe SARS-CoV-2 infections (Shoenfeld, 2020). We observed a statistically significant negative correlation between the allele frequency of Ala59Thr and the number of deaths per country [p-value 0.0337] (Fig. 5, panel B), but not with the numbers of cases (Fig. 5 panel A).

# 2.3. Description of the other genes potentially related to COVID-19 response

As discussed above, the association between six inactivating variants (Table 1) and SARS-CoV-2 susceptibility could not be demonstrated because of the absence of data on population distribution. Nevertheless, literature data support a possible role of some of them in COVID-19.

The role of *ACE* in innate immunity has been reviewed (Bernstein et al., 2018). A very frequent intronic insertion/deletion I/D polymorphism influences ACE expression (Rigat et al., 1990). A recent paper has suggested the usage of *ACE* I/D polymorphisms for the identification of high-risk COVID-19 patients (Delanghe et al., 2020). The hypothesis that variants of *ACE* are associated with COVID-19 risks is very interesting and is in line with several observations. Although ACE2, rather than ACE, has been identified to be one of the receptors of the SARS-CoV-2, it has been proposed that the unbalance between ACE and

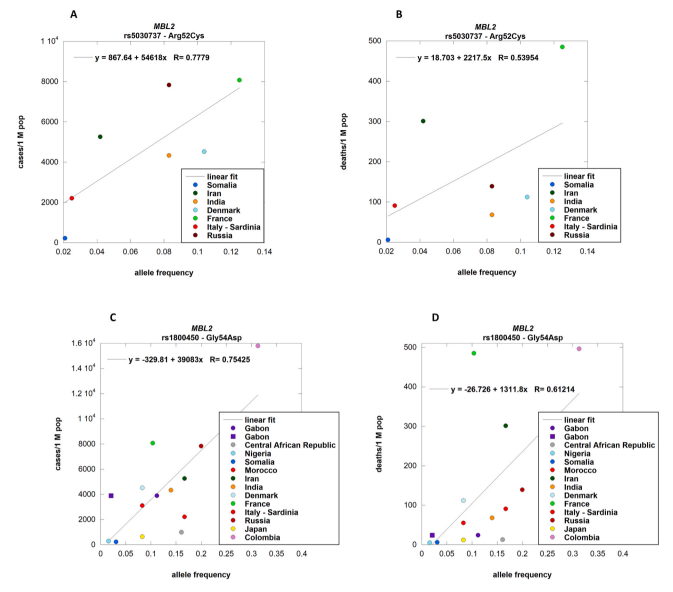


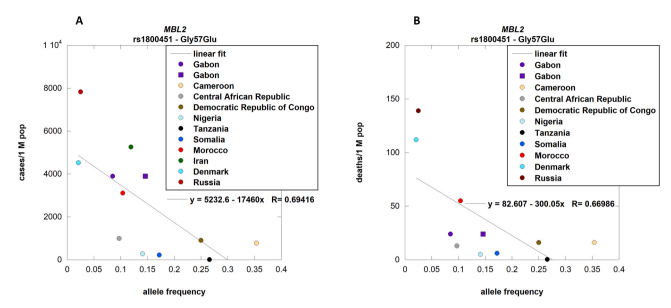
Fig. 2. *MBL2* allele frequencies per population correlation with COVID-19 cases and deaths. rs5030737 (Arg52Cys) allele frequencies in different countries from Verdu et al. were correlated with the respective number of cases (panel A; *p*-value 0.0394) and deaths (panel B; *n. s.*) of COVID-19. rs1800450 (Gly54Asp) allele frequencies were correlated both with cases (panel C; *p*-value 0.0018) and deaths (panel D; *p*-value 0.0199) of COVID-19 per country. Different symbols relate to different populations associated with the same country [Purple circle: GBP; purple square: GBB.] Details about populations and their association to countries are provided in supplementary file S1. . (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

ACE2 expression leads to lung injury (Guo et al., 2020). Children, who have higher levels of ACE than adults, are much less affected by COVID-19 (Guo et al., 2020). ACE inhibitors have been proven to upregulate ACE2, and some authors speculate a possible role of these extremely commonly used drugs in infection. Their role in COVID-19 disease, however, is very controversial, because upregulation of *ACE2* may have a protective role against lung inflammation and fibrosis (Guo et al., 2020). Publications on *ACE* I/D polymorphisms and SARS-CoV are of little help because they provide different results (Chan et al., 2005; Itoyama et al., 2004). A final word on this subject could come from a very recent paper by Richards and coworkers (Butler-Laporte et al., 2020). Although the authors did not include Tyr244Cys (rs3730025) in their study, they found that variants which decrease the expression of *ACE* do not increase COVID-19 susceptibility or severity.

AHSG encodes a protein known as human fetuin which is thought to be an immune modulator, mainly acting on cytokine release from macrophages. Macrophages play a central role in the IL1, 6, TNF alfa,

INF gamma dependent cytokine storm. Its deletion has already been linked to worse outcomes in SARS-CoV infected patients (Zhu et al., 2011). The SNP Ala164Thr is covered by the GWAS study (Group, 2020) and significantly associates with susceptibility to COVID-19 (p-value = 0.05).

The role of *GFAP* in COVID-19 might be prevalently associated to the severity of the disease and in particular to the development of neurological symptoms. *GFAP* is a TGF-beta induced gene that participates in the ERG dependent TGF-beta triggering of fibrosis consequent to ROS production. It has been proven that SARS-CoV viruses possess a deubiquitinating protein named PLpro that stimulates such pathway, inducing lung fibrosis in infected patients. Pulmonary fibrosis has been described in cases of SARS-CoV infections, but so far it has not been described to be a relevant feature of the disease (Yang et al., 2006). Very little is known so far about COVID-19 pathology, and future studies could demonstrate the importance of fibrosis since it is a central feature in uncontrolled activation of macrophage-mediated inflammation.



**Fig. 3.** *MBL2* allele frequencies per population correlation with COVID-19 cases and deaths. rs1800451 (Gly57Glu) allele frequencies were correlated with cases (panel A; *p*-value 0.0122) and deaths (panel B; *p*-value 0.0241) of COVID-19 per country. The outsider Iran was excluded to observe a significant correlation with the number of deaths.

Different symbols relate to different populations associated with the same country [Purple circle: GBP; purple square: GBB.] Details about populations and their association to countries are provided in supplementary file S1. . (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

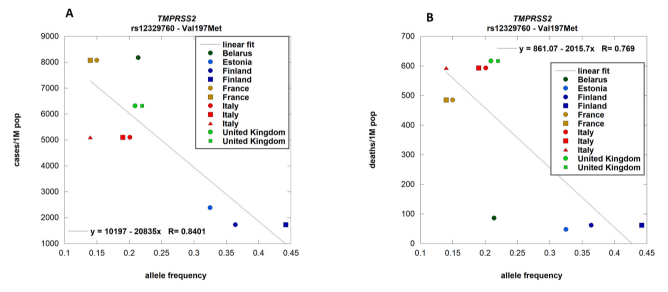


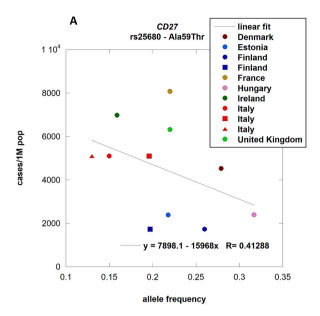
Fig. 4. TMPRSS2 allele frequencies per population correlation with COVID-19 cases and deaths. rs12329760 (Val197Met) allele frequencies in different countries from ALFRED were correlated with the respective number of cases (panel A; p-value 0.0012) and deaths (panel B; p-value 0.0057) of COVID-19. Different symbols relate to different populations associated with the same country [Blu circle: SA004049R; blu square: SA004617S. Light brown circle: SA001504K; light brown square: SA001503J. Red circle: SA004057Q; red square: SA002255O; red triangle SA001505L. Light green circle: SA004050J; light green square: SA001508O.] Details about populations and their association to countries are provided in supplementary file S1. . (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

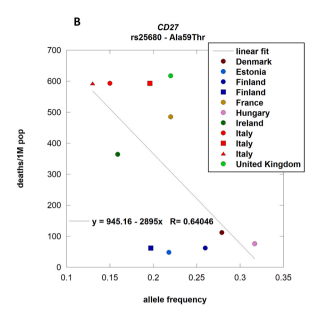
IFI30 (Gamma-interferon-inducible lysosomal thiol reductase, GILT) has a role in protecting against the internalization of SARS-CoV viruses in the lung tissue (Chen, D. et al., 2019). Polymorphisms in this gene may be correlated with a predisposition to infection by SARS-CoV-2, as well as with a worse prognosis due to higher initial viral charge. The viral charge has, in fact, already been proven to be linked to worse clinical outcomes (Chen, D. et al., 2019).

*MEFV* (Pyrin) is associated to Familial Mediterranean Fever whose symptoms, fever, abdominal/chest pain, elevation of C-reactive protein, and leucocytosis, overlap in part with those of COVID-19. Deleterious

variants in this gene might influence the severity of the infection. MEFV is thought to be involved in the NLRP3 dependant IL-1 beta modulation (Manukyan and Aminov, 2016). The capability of SARS-CoV viruses to activate the NLRP3 inflammasome has already been described (Chen, I.-Y. et al., 2019). There are some known pathological mutations of MEFV that cause autoinflammatory disease due to a IL1, 6, TNF- alfa INF-gamma cytokine storm (Manukyan and Aminov, 2016).

Toll like receptors activate Type-I interferon (alpha and beta) response that enhance innate antiviral responses such as the limitation of cell replication, the induction of apoptosis, and the synthesis of





 $\textbf{Fig. 5.} \ \, \text{CD27 allele frequencies per population correlation with COVID-19 cases and deaths.} \\ \text{rs25680 (Ala59Thr}$ 

) allele frequencies in different countries from ALFRED were correlated with the respective number of cases (panel A; n.

s.) and deaths (panel B; p-value 0.0337) of COVID-19. Different colors relate to different countries while different symbols relate to different populations associated with the same country [Blue circle: SA004377V; blue square: SA004049R. Red circle: SA002255O; red square: SA004057Q; red triangle SA001505L.] Details about populations and their association to countries are provided in supplementary file S1. . (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

enzymes with direct antiviral effect (Kawasaki and Kawai, 2014). Some viruses have been described to elude the Type I interferon response by shifting the reply towards the Type II (gamma) interferon branch, which favors viral diffusion through tissues by activating a cytokine storm (McGonagle et al., 2020).

Toll-like receptor 2 interacts with TICAM2 (Stack et al., 2014) that in turn contributes to SARS-CoV pathogenesis in mice (Gralinski et al., 2017). Recent papers on TLR2 have linked the function of this gene to viral as well as to the already known bacterial targets (Carty and Bowie, 2010). TLR2 might bind SARS-CoV-2 proteins and have a key role in directing the immune response to this pathogen. Arg753Gln (rs5743708) and Pro631His (rs5743704) in TLR2 are relatively frequent in the European population, approximately 3%. A few papers confirm the role of Arg753Gln (rs5743708) in respiratory diseases (Patarčić et al., 2015) (Smelaya et al., 2016). The effects of Arg753 substitution by glutamine on TLR2 were proven in vitro. The mutation impairs agonist-induced phosphorylation hetero-dimerization with TLR6, and recruitment of myeloid differentiation primary response protein 88 (MYD88) and MyD88 adapter-like (MAL) (Xiong et al., 2012). The SNP Arg753Gln (rs5743708) is covered in the GWAS study (Group, 2020) and it is associated with COVID-19 significantly (p-value < 0.05). The role in the susceptibility to viral infection is confirmed by the finding that intranasal administration of a TLR2/6 agonist reduces upper respiratory tract viral shedding in an animal models (Proud et al., 2020).

Toll-like receptor 3 is known to be involved in the innate immune response to SARS-CoV infections (Totura et al., 2015). Its role in the new epidemics is supported by the fact that rare loss-of-function variants were found among male patients with life-threatening COVID-19 pneumonia more frequently than in mildly affected controls (Zhang, Q. et al., 2020). One frequent missense mutation in *TLR3*, Leu412Phe (rs3775291) was associated with several viral diseases (El-Bendary et al., 2018; Fischer et al., 2018; Grygorczuk et al., 2017; Studzińska et al., 2017). It has an allele frequency ranging from 25% to 35% worldwide except for African populations, where it is 6%. Although it is necessary to be extremely cautious when considering the causes of the low numbers of reported cases from Africa, nonetheless it is tempting to

associate it to the low incidence of rs3775291. Another indication in favour of the role of rs3775291 in the susceptibility to SARS-CoV-2 comes from the GWAS study (Group, 2020) where a significative difference (*p*-value < 0.03) was observed between affected people and healthy controls. When we analysed the epidemiological data of general European populations, as we did for variants in *TMPRSS2* (Fig. 4) or *CD27* (Fig. 5), we did not find significant correlation (data not shown), but it would be interesting to look in a cohort of males.

Four deleterious frequent variation were found in the genes identified by GWAS analysis (Group, 2020), although these SNPs were not covered by the study.

Thr199Met in the gene *SLC6A20* encoding Sodium- and chloridedependent transporter XTRP3 and Asp182Asn in the gene *LZTFL1* encoding Leucine zipper transcription factor-like protein 1 are relatively frequent in the European population but not in African and East Asian populations.

GluE994Lys and Arg1140Gln are in the gene *FYCO1* encoding FYVE and coiled-coil domain-containing protein 1 that participate in the transportation and maturation of autophagosomes.

### 3. Conclusion

COVID-19 pandemic upset the world. The fast spread of SARS-CoV-2 through over the world and some peculiar novelties of the virus, such as the high number of asymptomatic individuals, rendered it a difficult challenge from different aspects. A strong effort was put in the discovery of risk factors associated with a more serious phenotype. Besides this, different countries experienced different impacts caused by SARS-CoV-2. As well described by Yamamoto and Bauer (2020), many factors contribute to these differences, e.g. socio-behavioral habits, political management, co-existence with other viruses, genetic factors. Genetic factors can play a major role in defining the outcome of the infection. This correlation is hard to prove due to all the side-factors described above, that certainly influence the official data from the different countries. Moreover, the globalized world we live in certainly complicates the association between population variants and countries.

Nevertheless, our data showed that a statistically significant correlation could be proved for some common variants in three genes, namely *MBL2*, *TMPRSS7*, and *CD27*, with the number of cases or deaths observed per country.

These data suggest that the genetic background is of utmost importance in the evaluation of COVID-19 susceptibility and that the discovery of target genes could be useful in the treatment and prevention of the infection. Moreover, the evaluation of the role of these polymorphisms might be necessary for the management of very common drugs during a pandemic.

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This work is dedicated to our friend and colleague Maria Malanga.

### Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ejmg.2021.104227.

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