





Thromboses and Hemostasis Disorders Associated with COVID-19: The Possible Causal Role of Cross-Reactivity and Immunological **Imprinting**

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Glob Med Genet 2021;8:162-170.

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Abstract

Keywords

- ► COVID-19
- ► SARS-CoV-2 spike qp
- cross-reactivity
- ► immunological imprinting
- thromboses-related proteins
- thromboses
- vascular diseases
- bleeding

By examining the issue of the thromboses and hemostasis disorders associated with severe acute respiratory syndrome-coronavirus-2 (SARS-CoV-2) through the lens of cross-reactivity, it was found that 60 pentapeptides are shared by SARS-CoV-2 spike qlycoprotein (qp) and human proteins that— when altered, mutated, deficient or, however, improperly functioning— cause vascular diseases, thromboembolic complications, venous thrombosis, thrombocytopenia, coaqulopathies, and bleeding, inter alia. The peptide commonality has a relevant immunological potential as almost all of the shared sequences are present in experimentally validated SARS-CoV-2 spike qpderived epitopes, thus supporting the possibility of cross-reactions between the viral qp and the thromboses-related human proteins. Moreover, many of the shared peptide sequences are also present in pathogens to which individuals have previously been exposed following natural infection or vaccinal routes, and of which the immune system has stored imprint. Such an immunological memory might rapidly trigger anamnestic secondary cross-reactive responses of extreme affinity and avidity, in this way explaining the thromboembolic adverse events that can associate with SARS-CoV-2 infection or active immunization.

Introduction

Clinical studies have shown that severe acute respiratory syndrome-coronavirus-2 (SARS-CoV-2) infection can lead to an increased incidence of disorders such as thrombosis, venous thrombosis, and pulmonary embolism. 1-3 A main conclusion of these studies is that, although it cannot be proven that the hypercoagulable state is a direct causative effect of SARS-CoV-2 infection, nonetheless it is apparent that patients with SARS-CoV-2 could have a predilection to the occurrence of thromboembolic events.¹

However, currently there are no hypotheses or data that might suggest a molecular mechanism that relates to such SARS-CoV-2-related thromboembolic events. Searching for possible mechanisms, the present study analyzes the SARS-CoV-2 spike glycoprotein (gp) for peptide sharing, that is, molecular mimicry, with human proteins, alterations of which may cause thromboses and hemostasis diseases. The underlying scientific rationale is that peptides common to a pathogen and the human host may lead to autoimmune pathologies through cross-reactivity phenomena following pathogen infection. 4-6 The results indicate that several linear

received April 13, 2021 accepted after revision April 26, 2021 published online June 26, 2021

DOI https://doi.org/ 10.1055/s-0041-1731068. ISSN 2699-9404.

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sequences shared between the SARS-CoV-2 spike gp and human proteins related to thromboembolic events can possibly generate pathogenic autoantibodies via cross-reactivity and immunologic imprinting phenomena, in this way leading to thromboses and hemostasis disorders.

Materials and Methods

Peptide sharing between spike gp (NCBI, GenBank Protein Accession, ID: QHD43416.1) from SARS-CoV-2 and human proteins related to thromboses and hemostasis disorders was analyzed as previously detailed.^{4–6} In brief, pentapeptides were used as sequence probes since a peptide grouping formed by five amino acid (aa) residues defines a minimal immune determinant that can (1) induce highly specific antibodies, and (2) determine antigen–antibody specific interaction.^{7,8} Human proteins linked to thromboses and hemostasis disorders were retrieved from UniProtKB database (www.uniprot.org).⁹ Methodologically the spike gp primary sequence was dissected into pentapeptides offset by one residue (i.e., MFVFL, FVFLV, VFLVL, FLVLL, and so forth)

and the resulting viral pentapeptides were analyzed for occurrences within the human proteins related to thromboses and hemostasis disorders. Then, the shared peptides were also controlled for occurrences in the pathogens Bordetella pertussis, Corynebacterium diphtheriae, Clostridium tetani, Haemophilus influenzae, and Neisseria meningitidis.

The immunological potential of the peptides shared between SARS-CoV-2 spike gp and thrombosis-related proteins was analyzed by searching the Immune Epitope DataBase (IEDB [www.iedb.org/])¹⁰ for immunoreactive SARS-CoV-2 spike gp-derived epitopes hosting the shared pentaptides.

Results and Discussion

Peptide Sharing between SARS-CoV-2 Spike Glycoprotein and Thromboses-Related Human Proteins

► **Table 1** shows that 60 minimal immune determinants are shared between SARS-CoV-2 spike gp and 44 human proteins that—when altered, mutated, deficient or, however, improperly functioning—may cause diseases that include

Table 1 Pentapeptide sharing between SARS-CoV-2 spike gp and human proteins linked when altered, mutated, or deficient to blood diseases

Shared peptides		
MTKTS, NLLLQ		
TQLPP, PRTFL	ALG12: Dol-P-Man: $Man(7)GlcNAc(2)$ -PP-Dol α -1,6-mannosyltransferase Psychomotor retardation, hypotonia, coagulation disorders, and immunodeficiency	
SAIGK	ALG8: Dolichyl pyrophosphate Glc1Man9GlcNAc2 α-1,3-glucosyltransferase Pathologies: see ALG12 above	
AEIRA	ANXA6 (annexin A6) Anticoagulant protein from human placenta	
QLIRA, IRASA	AP3B1 (AP-3 complex subunit β -1) Associates with Hermansky–Pudlak syndrome. Bleeding diathesis resulting in bruising, epistaxis, gingival bleeding, postpartum hemorrhage, bleeding	
LIGAE	APLP2 (amyloid-like protein 2) The soluble form may have inhibitory properties toward coagulation factors and regulates cerebral thrombosis	
VLLPL	B3AT (band 3 anion transport protein) Involved in venous thrombosis of unknown origin	
FGGVS	B4GT1 (β-1,4-galactosyltransferase 1) Defects in the nervous system development, psychomotor retardation, dysmorphic features, hypotonia, coagulation disorders	
KGYHL	C4BPB (C4b-binding protein β chain) Controls complement activation; binds as a cofactor to C3b/C4b inactivator; possibly involved in the susceptibility to venous thrombosis	
LTVLP	CBS (cystathionine β -synthase) CBS-deficient patients are prone to vascular thrombosis	
NSVAY	CO1A1 (collagen α -1(I) chain) Connective tissue disorders characterized by fragile, bruisable skin	
PGQTG, NGLTG	CO1A2 (collagen α-2(I) chain) Pathology: see CO1A1 above	
TQSLL, GTGVL	COG1 (conserved oligomeric Golgi's complex subunit 1) Psychomotor retardation, hypotonia, coagulation disorders, and immunodeficiency	

(Continued)

 Table 1 (Continued)

Shared peptides	Human proteins and associated functions/pathologies ^{a,b}	References 25
STNLV, GAISS	COG2. (conserved oligomeric Golgi's complex subunit 2) Pathology: as for COG1	
PINLV	COG5 (conserved oligomeric Golgi's complex subunit 5) Pathology: as for COG1	
LPFQQ, PFQQF, IGKIQ	ENTP1 (ectonucleoside triphosphate diphosphohydrolase 1) Implicated in the prevention of platelet aggregation	
YTSAL	EPHB2 (ephrin type-B receptor 2) Regulation of platelet activation and blood coagulation	
VLNDI	F13A (coagulation factor XIII A chain) Relates to hematologic disorders characterized by bleeding tendency	
DPLQP	FA5 (coagulation factor V) Central regulator of hemostasis. Parahemophilia, i.e., poor clotting; pregnancy loss, ischemic stroke, thrombophilia	
PPLLT, FVTQR	FA8 (coagulation factor VIII) Hemophilia	35
NSYEC	FA9 (coagulation factor IX) Hemophilia	35
SSANN	FIBA (fibrinogen α chain) Bleeding, amyloidosis, arterial hypertension, hepatosplenomegaly, cholestasis, petechial skin rash; thromboembolic complications	
GAGAA	GATA4 (transcription factor GATA-4) Regulates factor X, a vitamin K-dependent serine protease that functions in blood coagulation. Can predispose to dilated cardiomyopathy, and to premature death	
NDPFL	GP1BA (platelet glycoprotein lb α chain) Epistaxis; hemorrhage; menorrhagia; purpura; congenital bleeding diathesis; large platelets; thrombocytopenia; long bleeding time	
ALLAG	GPIX (platelet glycoprotein IX) Epistaxis; hemorrhage; menorrhagia; purpura; congenital bleeding diathesis; large platelets; thrombocytopenia; long bleeding time	
KLIAN	HABP2 (hyaluronan-binding protein 2) Serine protease involved in coagulation fibrinolysis and inflammatory pathways	43
TQLPP	HPS4 (Hermansky–Pudlak syndrome 4 protein) Epistaxis; reduced visual acuity; horizontal nystagmus; iris transillumination; restrictive lung disease; bruising; bleeding tendency; menorrhagia; absence of platelet dense bodies; lack of secondary aggregation response of platelets	
HTSPD	HPS5 (Hermansky–Pudlak syndrome 5 protein) As HPS4 above	
FNATR, DRLIT	HS3S5 (heparan sulfate glucosamine 3-O-sulfotransferase 5) Catalyzes a crucial step in the biosynthesis of the anticoagulant heparan sulfate	
SASFS	ITA2 (integrin α-2) Associates with increased ischemic stroke risk; thrombophilia	47,48
VRDLP	ITB3 (integrin β -3) Thrombasthenia, characterized by mucocutaneous bleeding	49
FGTTL, YDPLQ, GDISG	JAK2 (tyrosine-protein kinase JAK2) Thrombophilia, thrombocytosis	
VNLTT, GDSSS, VTYVP	MMRN1 (multimerin-1) Deficiency in multimerin-1 associates with bleeding disorder	
LLPLV	PLF4 (Platelet factor 4) Involved in thrombosis	
TFGAG	PLMN (plasminogen) may be associated with susceptibility to thrombosis	54
TVEKG, TGTGV	PROS: vitamin K-dependent protein S Anticoagulant plasma protein. Helps to prevent coagulation and stimulates fibrinolysis. Deficiency leads to impaired blood coagulation and a tendency to venous thrombosis	55,56

Table 1 (Continued)

Shared peptides	Human proteins and associated functions/pathologies ^{a,b}	References
LALHR	PROZ: vitamin K-dependent protein Z Helps hemostasis by binding thrombin and promoting its association with phospholipid vesicles. Deficiency may be a risk factor for retinal vessel occlusion	
IDRLI	PTPRJ: receptor-type tyrosine-protein phosphatase η Lack of PTPRJ leads to a bleeding tendency and defective arterial thrombosis	
VFAQV	TF (tissue factor): Initiates blood coagulation by forming a complex with circulating factor VII or VIIa	59
LFRKS	THRB: Prothrombin: Functions in blood homeostasis	60
AGAAL, GAALQ	TRBM (thrombomodulin) Relates to thrombophilia, venous thrombosis, and thromboembolic disease. TRBM administration is beneficial in sepsis-induced coagulopathy and in disseminated intravascular coagulations	61-63
TLLAL	TSP1 (thrombospondin-1): Coronary artery disease	52,64
TLLAL, SCGSC	TSP2 (thrombospondin-2): Coronary artery disease	52,64
VSSQC, LQYGS	VWF (von Willebrand factor) Von Willebrand's disease is characterized by deficiency of circulating VWF that is otherwise structurally and functionally normal. Clinical features: impaired platelet aggregation, cardiovascular diseases, mucocutaneous bleeding, epistaxis, menorrhagia	52,65-67

Abbreviations: gp, glycoprotein; SARS-CoV-2, severe acute respiratory syndrome-coronavirus-2.

coagulation disorders, bruising, bleeding, hemorrhages, retinal vessel occlusion, cerebral thrombosis, venous thrombosis, ischemic stroke, and thrombophilia, inter alia.

Immunological Potential of the Viral versus Human Peptide Sharing

The data shown in **~Table 1** are quantitatively impressive and become strikingly preeminent from a pathological perspective when analyzed for their immunological potential. Indeed, exploration of the IEDB¹⁰ reveals that nearly all the shared pentapeptides described in **~Table 1** are also disseminated among SARS-CoV-2 spike gp-derived epitopes that have been experimentally validated as immunoreactive and are cataloged at the IEDB database (http://www.iedb.org).¹⁰

That is, ►Table 2 concretely supports the possibility that autoimmune cross-reactions may be triggered by SARS-CoV-2 infection/active immunization and hit human proteins related to thrombotic/thromboembolic disorders and coagulopathies, inter alia. Clinically, the vastity of the potential immunological cross-reactivity that emerges from ►Table 2 indicates that mild-to-moderate and severe forms of thrombosis and coagulopathy may unavoidably accompany SARS-CoV-2 infection/active immunization.

Autoimmunity Potential and the Immunological Memory

As already highlighted also in other infection models, ^{68–71} one has to consider that immunologic memory can powerfully enhance and amplify the autoimmune cross-reactivity potential because of interpathogen peptide sharing. Indeed, as a rule, the immune system recalls preexisting

memory responses toward past infections rather than inducing ex novo responses toward the recent ones since hallmark of the immune system is the memory for the immune determinants it has previously encountered.^{72,73}

Here, comparative sequence analyses show that 31 out of the 60 minimal immune determinants common to SARS-CoV-2 spike gp and human proteins related to thromboses are also widespread in pathogens, such as *B. pertussis*, *C. diphtheriae*, *C. tetani*, *H. influenzae*, and *N. meningitidis*, that are in pathogens with which, in general, an individual has already come into contact during his life due to infections or by vaccination (**– Table 3**).

Hence, **Table 3** indicates the possibility that a preexisting immune response to previously encountered pathogens (in the present case: *B. pertussis, C. tetani, C. diphtheriae, H. influenzae*, and/or *N. meningitidis*) might be magnified and intensified following SARS-CoV-2 infection/active immunization. That is, immunological imprinting can start a chain of events according to which followings can be measured:

- Following exposure to SARS-CoV-2, the primary response to the virus can turn into a secondary response to previously encountered pathogens of which the immune system has stored an immunological memory.
- The anamnestic secondary and, by definition, extremely powerful response against immune determinants previously encountered implies not only that a low or no immune response will be evoked against the pathogen lastly encountered, that is, SARS-CoV-2, but also entails that the anamnestic secondary reaction against the early sensitizing pathogens—in the case in point, *B. pertussis*, *C. tetani*, *C. diphtheriae*, and/or *N. meningitidis*—will fail

^aHuman proteins given by Uniprot accession and name in italics.

^bFunctions and/or pathologies: data and further references from Uniprot, PubMed, and OMIM databases

Table 2 Distribution of peptides shared between SARS-CoV-2 spike gp and human proteins related to thromboses and hemostasis disorders among 94 experimentally validated SARS-CoV-2 spike gp-derived epitopes

ID ^a	Epitope ^b	ID ^a	Epitope ^b
1069137	aqYTSALLAGtitsg	1309555	qcVNLTTrTQLPPaytnsft
1069290	ctlksfTVEKGiyqt	1309558	qfnSAIGKIQdslsstasal
1071585	nlVRDLPqgfsalep	1309564	qtragcLlGAEhvnNSYECd
1071723	patvcgpkkSTNLVknkc	1309573	rLFRKSnlkpferdisteiy
1072807	skhtPINLVRDLPqg	1309595	tnftisvtteilpvsMTKTS
1072965	svtteilpvsMTKTS	1309598	tvYDPLQPeldsfkeeldky
1073281	tesnkkfLPFQQFgrdia	1309599	Tyvpaqeknfttapaichdg
1073938	vqIDRLITgrlqslq	1309600	tyvtqQLIRAAEIRASAnla
1074201	ylyrLFRKSnlkpfe	1309602	vcgpkkSTNLVknkcvnfnf
1074838	AEIRASAnlaatk	1309603	vknkcvnfnfNGLTGTGVLt
1074925	hVTYVPaqeknf	1309604	VLNDIlsrldkveaevqidr
1074969	lgaeNSVAYsnn	1309621	yskhtPINLVRDLPqgfsal
1074974	ILALHRsyl	1310254	aeNSVAYsnnsiaip
1075005	nqKLIANqf	1310281	aphgvvflhVTYVPa
1075031	rLFRKSnlk	1310303	caqkfngLTVLPpll
1075039	rqiaPGQTGkiadynykl	1310336	dskTQSLLivnnatn
1075066	sVLNDIIsrl	1310392	FGTTLdskTQSLLiv
1075079	tPINLVrdl	1310401	fkiyskhtPINLVrd
1075085	tvYDPLQPeldsfk	1310415	fngLTVLPPLLTdem
1075094	vlPPLLTdemiaqyt	1310434	GAISSVLNDIIsrld
1075125	ysvlynSASFStfk	1310444	givnntvYDPLQPel
1075131	yyvgylqPRTFLl	1310487	iginitrfqTLLALh
1087680	PINLVRDLPqgfsalepl	1310506	irgwiFGTTLdsktq
1125063	gLTVLPpll	1310513	itrfqTLLALHRsyl
1309117	ggnynylyrLFRKSn	1310592	lLALHRsyltpgdss
1309118	gpkkSTNLVknkcvn	1310611	IPPLLTdemiaqyts
1309123	khtPINLVRDLPqgf	1310633	lyenqKLIANqfnsa
1309140	tdemiaqYTSALLAG	1310787	SASFStfkcygvspt
1309147	ylqPRTFLI	1310828	svlynSASFStfkcy
1309418	AEIRASAnlaatkmsecvlg	1310852	tlvkqlssnfGAISS
1309442	ayyvgylqPRTFLlkyneng	1310865	trfqTLLALHRsylt
1309450	dplsetkctlksfTVEKGiy	1310899	VLLPLVSSQCVNLTT
1309451	dsfkeeldkyfknHTSPDvd	1310909	VNLTTrTQLPPaytn
1309461	ehvnNSYECdipigagicas	1310927	vtqnvlyenqKLIAN
1309464	esnkkfLPFQQFgrdiadtt	1310947	wTFGAGAALQipfam
1309469	fknHTSPDvdlGDISGinas	1310979	yvgylqPRTFLlkyn
1309470	fknidgyfkiyskhtPINLV	1311657	ccSCGSCckfdeddsepvlkgvkl
1309475	gccSCGSCckfdeddsepvl	1311813	rLFRKSnlkp
1309492	ilditpcsFGGVSvitpgtn	1313244	nSASFStfk
1309506	kvggnynylyrLFRKSnlkp	1313285	PINLVRDLPqgfsal
1309515	IhrsyltpGDSSSgwtagaa	1313286	PINLVRDLPqgfwal
1309516	litgrlqslqtyvtqQLIRA	1314023	ynylyrLFRKSnlkp

Table 2 (Continued)

ID ^a	Epitope ^b	ID ^a	Epitope ^b
1309523	lssnfGAISSVLNDIlsrld	1317916	gylqPRTFLl
1309524	lyenqKLIANqfnSAIGKIQ	1321084	IPPLLTdem
1309531	NGLTGTGVLtesnkkflpfq	1327418	vYDPLQPeldsf
1309532	ngLTVLPPLLTdemiaqyts	1327923	yenqKLIANqf
1309534	nitrfqTLLALHRsyltpgd	1328800	ytmslgaeNSVAY

Abbreviations: gp, glycoprotein; SARS-CoV-2, severe acute respiratory syndrome-coronavirus-2.

Table 3 Occurrence in microbial organisms of pentapeptides common to SARS-CoV-2 spike gp, SARS-CoV-2 spike gp-derived epitome, and human proteins related to thromboses, coaquiopathies, and hemostasis disorders

Organism	Shared peptides
Bordetella pertussis	AEIRA, AGAAL, ALLAG, GAALQ, GAGAA, LLPLV, PFQQF, QLIRA, SSANN, TGTGV, VLLPL, YDPLQ
Corynebacterium diphtheriae	AEIRA, AGAAL, ALLAG, DPLQP, GAALQ, GAGAA, GTGVL, LLPLV, TVEKG
Clostridium tetani	AGAAL, LQYGS
Haemophilus influenzae	AEIRA, AGAAL, FGGVS, GAALQ, GAGAA, GTGVL, KLIAN, LALHR, LLPLV, LPFQQ, LTVLP, NLLLQ, NSVAY, TLLAL, TQSLL, VLLPL, VLNDI, VNLTT, YTSAL
Neisseria meningitidis	AEIRA, AGAAL, ALLAG, DRLIT, GAALQ, GAGAA, IDRLI, KLIAN, LALHR, LTVLP, PINLV, TLLAL, VLLPL, VLNDI

Abbreviations: qp, qlycoprotein; SARS-CoV-2, severe acute respiratory syndrome-coronavirus-2.

because those early sensitizing pathogens are no more present in the organism.

 As a final result, the anamnestic, high affinity, high avidity, and extremely powerful secondary immune response triggered by the lastly encountered pathogen (SARS-CoV-2) and addressed toward past infections may find an outlet by hitting available human targets, that is, in the case in object, the human proteins related to thromboses and hemostasis diseases (~Table 1).

Conclusion

The last decades witnessed the emerging of infectious diseases and, consequently, intensive application of immunization procedures. Concomitantly, concerns about possible adverse events have increased. A recent crucial example is the immunization campaign with the dengue vaccine that highlighted the risk of enhanced disease after vaccination.⁷⁴

Today, the clinical context associated with SARS-CoV-2 infection/active immunization is no different. Actually, understanding whether undesired collateral events, such as the thrombotic manifestations and bleeding disorders discussed in this study, may causally associate with the viral infection/active immunization is a fundamental step for fighting the current pandemic. In this context, the present study:

 Analyzed the hypothesis that infectious agents can induce cross-reactive autoantibodies capable of hitting and al-

- tering human proteins that regulate hemostasis and coagulation.
- Showed that numerous peptides endowed with an immunologic potential are common to SARS-CoV-2 spike gp and human proteins, when mutated, altered, deficient or improperly functioning, are associated with thromboses and hemostasis diseases (~Tables 1 and 2).
- Documented that the peptide commonality extends to pathogens that usually have been already encountered by an individual during his life (-Table 3).

Scientifically, the data indicate that peptide sharing-associated cross-reactivity and, in conjunction, immunological imprint might help explain some of the thromboembolic events that rapidly, massively, and violently may arise following SARS-CoV-2 infection/active immunization.

Clinically, the present data warrant testing of patients' sera for autoantibodies against the peptide targets described in **-Tables 1-2** and **3**, and reiterate the suggestion advanced already in 2000⁷⁵ that immunotherapies should take advantage of the principle of peptide uniqueness, that is, of peptides present in the antigen of interest and absent in the human proteome. 71.76-81

Funding None.

Conflict of Interest None declared.

^aEpitopes listed as the Immune Epitope DataBase ID.

^bShared peptides given in capital letters.

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