

DOI: 10.56871/RBR.2024.63.31.001

UDC [616.2+616.34]-036.21+578.834.1+579.8+577.112+547.96

SARS COV-2 DELTA VARIANT STRUCTURAL PROTEINS: HOMOLOGY WITH OPPORTUNISTIC BACTERIA

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For citation: Maryanovich AT, Kormilets DYU. SARS CoV-2 Delta variant structural proteins: homology with opportunistic bacteria. Russian Biomedical Research. 2024;9(2):5–17. DOI: <https://doi.org/10.56871/RBR.2024.63.31.001>

Received: 15.02.2024

Revised: 04.04.2024

Accepted: 20.05.2024

Abstract. The capacity of SARS CoV-2 for immune evasion can be considered universally recognized. Coronavirus and human protein homology may be one of the mechanisms of immune evasion. Delta variant necessarily has structural features that explain its specific qualities. The aim of our study is to find out whether mutations in the structural proteins of Delta variant change its homology with proteins present in the human body, i.e. human, bacterial and dietary. Using bioinformatics tools we detected homology on the heptamer level between Delta variant structural proteins and human proteins as well as some opportunistic bacteria proteins of the upper respiratory tract, lung and gut. Delta variant spike (S) and membrane (M) proteins have a large number of similarities (homologous correspondences) with the listed proteins, with the S:Δ156,157;R158G mutation having the greatest amount. The reason why SARS CoV-2 Delta variant has specific characteristics, most importantly increased lethality, is most likely to be found in a mutation at positions 156–158 of spike protein.

Keywords: SARS CoV-2, Delta variant, spike protein, opportunistic bacteria, homology

СТРУКТУРНЫЕ БЕЛКИ ДЕЛЬТА-ВАРИАНТА SARS COV-2: ГОМОЛОГИЯ С ОППОРТУНИСТИЧЕСКИМИ БАКТЕРИЯМИ

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Для цитирования: Марьянович А.Т., Кормилец Д.Ю. Структурные белки дельта-варианта SARS CoV-2: гомология с оппортунистическими бактериями // Российские биомедицинские исследования. 2024. Т. 9. № 2. С. 5–17. DOI: <https://doi.org/10.56871/RBR.2024.63.31.001>

Поступила: 15.02.2024

Одобрена: 04.04.2024

Принята к печати: 20.05.2024

Резюме. Способность SARS CoV-2 уклоняться от иммунного ответа можно считать общепризнанной. Гомология белков коронавируса и человека может быть одним из механизмов иммунного уклонения. Дельта-вариант обязательно имеет структурные особенности, которые объясняют его специфические свойства. Целью нашего исследования было выяснить, изменяют ли мутации, произошедшие в структурных белках дельта-варианта, его гомологию с белками, присутствующими в организме человека, то есть собственно человеческими, бактериальными и пищевыми. Используя инструменты биоинформатики, мы обнаружили гомологию на уровне гептамеров между структурными белками дельта-варианта и белками человека, а также белками некоторых

условно-патогенных бактерий верхних дыхательных путей, легких и кишечника. Белки шиповый (S) и мембранный (M) дельта-варианта имеют большое количество сходств (гомологических соответствий) с перечисленными белками, причем наибольшее количество — в случае мутации S:Δ156,157;R158G. Причина, по которой дельта-вариант SARS CoV-2 обладает специфическими характеристиками, и прежде всего повышенной летальностью, скорее всего, кроется в мутации в положениях 156–158 шипового белка.

Ключевые слова: SARS CoV-2, дельта-вариант, шиповидный белок, оппортунистические бактерии, гомология

INTRODUCTION

After a series of brilliant discoveries from Pasteur to Fleming and Waxman, mankind has learned to control most bacterial infections. Humans were able to create megapolises with huge population densities. In response, nature had to put forward other limiting mechanisms less humanly controllable. The COVID-19 pandemic has become and will remain one of humanity's major concerns for the near future. The very important question is why and how this CoV could cause a pandemic [1]. Some mutation-induced structural substitutions in the N-terminal domain (NTD) of the SARS-CoV-2 S-protein lead to more efficient first contact and interaction with the upper airway epithelium [2].

The extraordinary virulence of Omicron variant (B.1.1.529) is now the main focus of researchers [3]. Nevertheless, it seems to us that in order to understand the causes of SARS CoV-2 lethality, the peculiarities of Delta variant (B.1.617.2) must be studied.

Using 3D models, the researchers can determine how the spike (S) protein binds to the ACE2 receptor [4]. The peculiarity of our approach is that we seek an explanation for the properties of coronavirus in the homology (commonality of short motifs) of virus proteins with human proteins. Recently we described dozens of homologous motifs in the primary structure of SARS CoV-2 and human proteins including proteins of olfactory and taste receptors [5]. Through mutations, the virus finds a way to avoid an immune response [6].

Molecular mimicry is considered a strategy used by many viruses to subvert and regulate antiviral immunity. For example, human cytomegalovirus has hijacked or developed a number of homologous sites that mimic immunomodulatory proteins encoded by the human body. These homologues encoded by the virus can contribute to the virus' evasion of immune clearance [7].

Following Joshua Lederberg's principle [8], we took into account not only proteins synthesized by the human body, but also those that originate from other genotypes and are constantly present in the macroorganism. These are the proteins of commensal and opportunistic bacteria of the upper respiratory tract, lung, oral cavity, and GI tract. We also analyzed the most common dietary proteins that are almost constantly

present in the gut, namely those of the six world's most important cereal crops, i.e., Asian rice *Oryza sativa*, common wheat *Triticum aestivum*, maize *Zea mays*, common bean *Phaseolus vulgaris*, barley *Hordeum vulgare*, and sorghum *Sorghum bicolor*. We believed that the homology of the virus proteins with those of the named bacteria and cereals helps coronavirus to avoid or reduce the primary immune response.

THE AIM OF OUR STUDY

The aim of our study is to find out whether mutations in the structural proteins of SARS CoV-2 Delta variant change its homology with proteins present in the human body, i.e. human, bacterial and dietary.

RESULTS

Spike glycoprotein

Wuhan-Hu spike glycoprotein (S protein) molecule consists of 1273 amino acid residues. In Delta variant, as a result of two deletions (E₁₅₆Δ and F₁₅₇Δ), S protein consists of 1271 amino acid residues and contains seven substitutions in nine positions, namely T₁₉R, G₁₄₂D, R₁₅₈G, L₄₅₂R, T₄₇₈K, P₆₈₁H, and D₉₅₀N, numeration as in Wuhan-Hu variant [9].

S protein Delta variant, 1271 aa

MFVFLVLLPLVSSQCVNLRTRTQLPPAYTNSFTRGVYYP
DKVFRSSVLHSTQDLFLPFFSNVTWFWHAIHVS GTNGT
KRFDPNPLPFDNGGVYFASTEKSNIIRGWIFGTTLD SKTQ
SLIVNNATNVVIVKVECFQFCNDPFLDVYHKNKNSWMES
GVYSSANNCTFEYVSQPFLLMDLEGKQGNFKNLREF
VFKNIDGYFKIYSKHTPINLVRDLPPQGSFALEPLVDLPIGINI
TRFQTLALHRSYLT PGDSSSGWTAGAAAYVGYLQPR
LLKYNENGTITDAVDCALDPLSETKCTLSFTVEKGIYQTSN
FRVQPTESIVRFPNITNLCPFGVEFNATRFASVYAWNRKRIS
NCVADYSVLNSASFSTFKCYGVSPTKLNDLCFTNVYADSF
VIRGDEVQRQIAPGQTGKIADYNYKL PDDFTGCVIWNNSNNLD
SKVGGNYNYRYRLFRKSNLKPFFERDISTEIYQAGSKPCNG
VEGFNCYFPLQSYGFQPTNGVGYQPYRVVLSFELLHAPAT
VCGPKKSTNLVKNKCVNFNFNGLTGTGVLTESNKKFLPFQQF
GRDIADTTDAVRDPQTLEILDITPCSFGGVSVITPGTNTSNQ
VAVLYQDVNCTEVPVAIHADQLTPTWRVYSTGSNVFQTRAGC
LIGAEHVNSYECDIPIGAGICASYQTQTNRRRRARSVAS

QSIAYTMSLGAENSVAYSNNISAIPTNFTISVTTEILPVSMK
 TSVDCTMYICGDSTECNLLLQYGSFCTQLNRALTGIAVEQD
 KNTQEVFAQVKQIKTPPIKDFGGFNFSQILPDPSKPSKRS
 FIEDLLFNK**VT**LADAGFIKQYGDCLGDIAARDLICAQKFN
GLTVLPLLTDEMIQYTSALLAGTITSGWTFGAGAA
 LQIPFAMQMAYRFNGIGV**TQNVLYENQKLIANQFN**SAIG
KIQDSLSSTASAL**KLQ**N**VVNQNAQALNTLVKQLSS**
NFGAISSV**LN**DIL**SRLD**KVEAEVQIDRLIT**GRLQSL**
 QTYVTQQL**LIRAAE**IRASANLAATKMSECVLGQSKRVDF
 CGKGYHLMSFPQSAPHGVVFLHVTYVPAQEKNTTAPAI
 CHDGKAHFPREGVFSNGTHWFVTQRNFYEPQIITDNT
 FVSGNCDVVIGIVNNTVYDPLQPELDSFKEE**LDKYFKN**
 HTSPDVLGDIGIN**ASV**VNI**QKEIDRLNE**VAKNLNES
 LIDLQELGKYEYIKWPWYIWLGFIAGLIAIVMVTIMLCC
 MTSCCCLKGCCSCGSCCK**DEDDSE**PVLKGVLHYT

Hereinafter, motifs homologous with human proteins [5] are highlighted in red font. Amino acid residues substituted as a result of mutations are highlighted in large letters. The N-terminal domain (NTD₁₄₋₃₀₃) is highlighted in green. Receptor-binding domain (RBD₃₁₇₋₅₃₉) is in gray italics. Receptor-binding motif RBM₄₃₆₋₅₀₆ is underlined. Heptapeptide repeat sequence 1 (HR1₉₁₀₋₉₈₂) is highlighted in blue. As a result of the double deletion $\Delta_{156,157}$, starting from G₁₅₆, the numbering of positions in Delta variant does not correspond to the numbering in Wuhan-Hu.

Delta variant, as mentioned above, has a mutation S:P₆₈₁H. The S protein motif **SPRRARS**₆₈₀₋₆₈₆ homologous with a human protein has been replaced by a heptamer **SHRRARS**₆₇₈₋₆₈₄, which has no homologues in mammals (Table 1).

Table 1

Homology of a SARS CoV-2 S protein to a human protein

Mutation	Wuhan-Hu			Delta		
P ₆₈₁ H*	S protein heptamer	Species	Homologous protein heptamer	S protein heptamer	Species	Homologous protein heptamer
	SPRRARS ₆₈₀₋₆₈₆ *	<i>Homo sapiens</i>	Hermansky-Pudlak syndrome 1 protein ₂₅₈₋₂₆₄	SHRRARS ₆₇₈₋₆₈₄	No homological heptamers in commensal	

*In Wuhan-Hu and Delta variants, the position numbering differs after position 156 as a result of the $\Delta_{156,157}$ deletions.

The heptamers of S protein that are homologous with the proteins of some commensal and opportunistic bacteria are listed in Table 2.

Table 2

The heptamers of S protein homologous with the proteins of some commensal and opportunistic bacteria

Mutation	Wuhan-Hu				Delta			
	S protein heptamer	Species	Homologous protein heptamer	Localization in the human body	S protein heptamer	Species	Homologous protein heptamer	Localization in the human body
T ₁₉ R	VNL T TRT ₁₆₋₂₂	<i>Escherichia coli</i> BCE011_MS-01	Uncharacterized protein ₂₃₋₂₉	gut	VNL R TRT ₁₆₋₂₂	<i>Streptococcus mitis</i> SK597 TnpX; <i>Streptococcus salivarius</i> (strain CCHSS3)	Site-specific recombinase ₂₇₅₋₂₈₁	nasopharynx, oral cavity, throat
	NL T TRTQ ₁₇₋₂₃	<i>Enterococcus faecalis</i>	Helicase, RecD/TraA family ₇₅₅₋₇₆₁	gut	NL R TRTQ ₁₇₋₂₃	<i>Subdoligranulum variabile</i>	Putative hydrolase ₃₄₋₄₀	gut
G ₁₄₂ D*	NDPFL G V ₁₃₇₋₁₄₃	No homological heptamers in commensal or opportunistic bacteria			NDPFL D V ₁₃₇₋₁₄₃	<i>Pasteurella multocida</i> subsp. <i>multocida</i> str	Release factor glutamine methyltransferase ₂₀₋₂₆	lung
$\Delta_{156,157}$; R158G	EF R VYSS ₁₅₆₋₁₆₂	No homological heptamers in commensal or opportunistic bacteria			ES G VYSS ₁₅₄₋₁₆₀	<i>Lachnospiraceae</i> bacterium 7_1_58FAA	Uncharacterized protein ₁₂₆₋₁₃₂	gut
						<i>Escherichia coli</i> UMEA 3609-1	Valine-tRNA ligase ₃₂₀₋₃₂₆	gut

Endind of the table 2

Mutation	Wuhan-Hu				Delta			
	S protein heptamer	Species	Homologous protein heptamer	Localization in the human body	S protein heptamer	Species	Homologous protein heptamer	Localization in the human body
	F ^R VYSSA ₁₅₇₋₁₆₃	No homological heptamers in commensal or opportunistic bacteria			s ^G VYSSA ₁₅₅₋₁₆₁	<i>Fusobacterium sp. oral taxon 370 str. F0437</i>	Hep/Hag repeat protein (Fragment) ₄₇₋₅₃	oral cavity
	R ^V YSSAN ₁₅₈₋₁₆₄	<i>Bifidobacterium animalis subsp. lactis CNCM I-2494</i>	Fibronectin-binding protein ₁₉₁₋₁₉₇	gut	G ^V YSSAN ₁₅₆₋₁₆₂	<i>Bacillus sp. NRRL B-14911</i>	Methylmalonyl-CoA mutase ₅₆₅₋₅₇₁	?
						<i>Lactobacillus faraginis JCM 14108</i>	D-alanyl-D-alanine carboxypeptidase ₁₄₉₋₁₅₅	gut
						<i>Fusobacterium nucleatum subsp. polymorphum F0401</i>	Uncharacterized protein ₂₃₄₋₂₄₀	oral cavity
						<i>Prevotella saccharolytica F0055</i>	Carbohydrate binding domain protein ₇₁₅₋₇₂₁	oral cavity, upper respiratory tract, gut
						<i>human gut metagenome</i>	Glycoside hydrolase, family 25 (Fragment) ₃₉₅₋₄₀₁	gut
L ₄₅₂ R	No homological heptamers in commensal or opportunistic bacteria							
T ₄₇₈ K	No homological heptamers in commensal or opportunistic bacteria							
P ₆₈₁ H	NS ^P RRAR ₆₇₉₋₆₈₅	No homological heptamers in commensal or opportunistic bacteria			NS ^H RRAR ₆₇₇₋₆₈₃	<i>Clostridium clostridioforme</i>	Uncharacterized protein ₁₁₆₋₁₂₂	gut
D ₉₅₀ N	KLQ ^D VVN ₉₄₇₋₉₅₃	<i>Prevotella salivae F0493</i>	Peptidase M16 inactive domain protein ₉₁₈₋₉₂₄	oral cavity, gut	KLQ ^N VVN ₉₄₅₋₉₅₁	<i>Leptotrichia buccalis (strain ATCC 14201 / DSM 1135 / JCM 12969 / NCTC 10249)</i>	GCN5-related N-acetyltransferase ₁₁₅₋₁₂₁	oral cavity
	D ^V VNQNA ₉₅₀₋₉₅₆	No homological heptamers in commensal or opportunistic bacteria			N ^V VNQNA ₉₄₈₋₉₅₄	<i>Prevotella multisaccharivorax DSM 17128</i>	Anaerobic ribonucleoside-triphosphate reductase ₁₁₄₋₁₂₀	oral cavity, gut

*The same mutation has occurred in Omicron variant.

The heptamers of S protein that are homologous with the most common cereal proteins are listed in Table 3.

Table 3

The heptamers of S protein homologous with the most common cereal proteins

Mutation	Wuhan-Hu			Delta		
	S protein heptamer	Species	Homologous protein heptamer	S protein heptamer	Species	Homologous protein heptamer
T ₁₉ R	SQC ^V NLT ₁₃₋₁₉	<i>Oryza sativa</i>	Leucine Rich Repeat family protein, expressed ₅₂₀₋₅₂₆	SQC ^V NLR ₁₃₋₁₉	No most common cereal sample	
	VNL ^T TRT ₁₆₋₂₂	<i>Oryza sativa BCE011_MS-01</i>	Uncharacterized protein ₂₃₋₂₉	VNL ^R TRT ₁₆₋₂₂	No most common cereal sample	
	L ^T TRTQL ₁₈₋₂₄	<i>Triticum aestivum</i>	Uncharacterized protein ₈₈₈₋₈₉₄	L ^R TRTQL ₁₈₋₂₄	No most common cereal sample	



End of the table 3

Mutation	Wuhan-Hu			Delta		
	S protein heptamer	Species	Homologous protein heptamer	S protein heptamer	Species	Homologous protein heptamer
L ₄₅₂ R	LYRLFRK ₄₅₂₋₄₅₈	<i>Oryza sativa</i> subsp. <i>indica</i>	Putative uncharacterized protein ₁₅₇₋₁₆₃	RYRLFRK ₄₅₀₋₄₅₆	No most common cereal sample	
		<i>Zea mays</i>	Putative NAC domain transcription factor superfamily protein (Fragment) ₁₀₀₋₁₀₆			
T ₄₇₈ K	STPCNGV ₄₇₇₋₄₈₃	No most common cereal sample		SKPCNGV ₄₇₅₋₄₈₁	<i>Phaseolus vulgaris</i>	Uncharacterized protein ₅₉₋₆₅
	SPRRARS ₆₈₀₋₆₈₆	<i>Oryza sativa</i> subsp. <i>japonica</i>	Os02g0817400 protein (Fragment) ₁₋₇	SHRRARS ₆₇₈₋₆₈₄	<i>Oryza sativa</i> subsp. <i>japonica</i>	Expressed protein ₂₉₆₋₃₀₂
		<i>Zea mays</i>	Uncharacterized protein ₅₈₋₆₄		<i>Oryza sativa</i> subsp. <i>japonica</i>	Uncharacterized protein ₆₁₆₋₆₂₂
	PRRARSV ₆₈₁₋₆₈₇	<i>Oryza sativa</i> subsp. <i>japonica</i>	Putative uncharacterized protein ₁₁₈₋₁₂₄		<i>Hordeum vulgare</i>	Predicted protein (Fragment) ₁₆₋₂₂
		<i>Zea mays</i>	Uncharacterized protein ₉₄₋₁₀₀	HRRARSV ₆₇₉₋₆₈₅	No most common cereal sample	
D ₉₅₀ N*	ALGKLQD ₈₄₄₋₉₅₀	<i>Hordeum vulgare</i> var. <i>distichum</i>	Uncharacterized protein ₁₂₃₋₁₂₉	ALGKLQN ₈₄₂₋₈₄₈	No most common cereal sample	
	LGKLQDV ₉₄₅₋₉₅₁	<i>Hordeum vulgare</i> var. <i>distichum</i>	Uncharacterized protein ₉₆₋₁₀₂	LGKLQNV ₈₄₃₋₈₄₉	No most common cereal sample	
		<i>Oryza sativa</i> subsp. <i>indica</i>	Uncharacterized protein ₂₄₈₋₂₅₄			
		<i>Zea mays</i>	Protein lap ₂₃₃₋₂₃₉			
			Golgi SNAP receptor complex member 1 ₇₅₋₈₁			
	GKLQDVV ₉₄₆₋₉₅₂	<i>Zea mays</i>	Uncharacterized protein ₃₈₈₋₃₉₄	GKLQNVV ₈₄₄₋₈₅₀	No most common cereal sample	

*The same mutation has occurred in Omicron variant.

The heptamers of S protein that are homologous with some virus proteins are listed in Table 4.

Table 4

The heptamers of S protein homologous with some virus proteins

Mutation	Wuhan-Hu			Delta			Comment
	S protein heptamer	Other virus	Homologous protein heptamer	S protein heptamer	Other viruses	Homologous protein heptamer	
P ₆₈₁ H	QTQTNSP ₆₇₅₋₆₈₁	<i>Human immunodeficiency virus 1</i>	Protease (Fragment) ₂₋₇	QTQTNSH ₆₇₃₋₆₇₉	No virus proteins homology		Homology with HIV-1 has disappeared
D ₉₅₀ N*	LQDVVNQ ₉₄₈₋₉₅₄	No virus proteins homology		LQNVVNQ ₉₄₆₋₉₅₂	<i>Human immunodeficiency virus 1</i>	Envelope glycoprotein (Fragment) ₇₁₋₇₇	Homology with HIV-1 has appeared

* The same mutation has occurred in Omicron variant.

Membrane protein

There are four mutations known in the membrane (M) protein Delta variant, namely A₂S, F₂₈L, V₇₀L, and I₈₂T [10].

M protein Delta variant, 222 aa

MSDSNGTITVEELKKLLEQWNLVIGFLLLTWICLLQFAYANR
NRFLYIIKLIFLWLLWPVTLACFVLAALYRINWITGGIATAMACLV

GLMWLSYFIASFRLFARTRSMWSFNPETNILLNVPLHGILTRP
LLESELVIGAVILRGHLRIAGHHLGRCDIKDLPEITVATSRTLSYY
 KLGASQVR**AGDSGFA**AYSRYRIGNYKLNTHSSSSDNIALLVQ

The heptamers of M protein that are homologous with the proteins of the commensal and opportunistic bacteria are listed in Table 5.

Table 5

The heptamers of M protein homologous with the proteins of the commensal and opportunistic bacteria

Mutation	Wuhan-Hu				Delta			
	M protein heptamer	Species	Homologous protein heptamer	Localization in the human body	M protein heptamer	Species	Homologous protein heptamer	Localization in the human body
A ₂ S	M ADS NGT ₁₋₇	No homological heptamers in commensal or opportunistic bacteria			M SDS NGT ₁₋₇	No homological heptamers in commensal or opportunistic bacteria		
	A DS NGT ₁₋₈	<i>Lachnospiraceae bacterium 7_1_58FAA</i>	Uncharacterized protein ₂₅₂₋₂₅₈	gut	S DS NGT ₁₋₈			
F ₂₈ L	LVIGFL F ₂₂₋₂₈	<i>Enterococcus faecalis R508</i>	Putative ferrichrome transport system permease protein FhuG ₂₀₃₋₂₀₆	gut	LVIGFL ₂₂₋₂₈	<i>Eubacterium ventriosum</i> ATCC 27560	Putative K(+)-stimulated pyrophosphate-energized sodium pump ₅₇₃₋₅₇₉	gut
						<i>Enterococcus caccae</i> ATCC BAA-1240	Uncharacterized protein ₁₀₄₋₁₁₀	gut
						<i>Faecalibacterium</i> sp. CAG:74	Binding-protein-dependent transport systems inner membrane component ₈₆₋₉₂	gut
						<i>Prevotella histicola</i> F0411	Uncharacterized protein ₁₅₋₂₁	gut
						<i>Lachnospiraceae bacterium 2_1_58FAA</i>	Uncharacterized protein ₆₅₋₇₁	gut
						<i>Escherichia coli</i> ISC11	Putative cell envelope opacity-associated protein A ₄₂₋₄₈	gut
	VIGFL F ₂₃₋₂₉	<i>Enterococcus flavescens</i> ATCC 49996	Uncharacterized protein ₁₂₈₋₁₃₄	gut	VIGFL ₂₃₋₂₉	<i>Prevotella</i> sp. oral taxon 472 str: F0295	Uncharacterized protein ₁₇₈₋₁₈₄	gut
		<i>Lachnospiraceae bacterium COE1</i>	MATE efflux family protein ₁₁₂₋₁₁₈	gut		<i>Lactobacillus brevis</i> ATCC 14869 = DSM 20054	Potassium uptake protein, TrkH family ₂₃₉₋₂₄₅	gut
						<i>Lactobacillus antri</i> DSM 16041	Transporter, major facilitator family protein ₄₂₂₋₄₂₈	gut
						<i>Enterobacter cloacae</i> subsp. <i>cloacae</i> (strain ATCC 13047 / DSM 30054 / NBRC 13535 / NCDC 279-56)	Putative multidrug resistance protein MdtD ₁₈₃₋₁₈₉	gut
						<i>Lachnospiraceae bacterium 28-4</i>	Uncharacterized protein ₁₈₋₂₄	gut
	IGFL F _{LT} ₂₄₋₃₀	<i>Lachnospiraceae bacterium CAG:215</i>	Transporter ₄₆₈₋₄₇₄	gut	IGFL _{LT} ₂₄₋₃₀	<i>Lactobacillus paracasei</i> subsp. <i>paracasei</i> Lpp126	Oligopeptide transport system permease protein oppB ₉₋₁₅	oral cavity
						<i>Eubacterium nodatum</i> ATCC 33099	TIGR02185 family protein ₄₃₋₄₉	oral cavity
						<i>Bacteroides uniformis</i> dnLKV2	Uncharacterized protein ₇₃₇₋₇₄₃	gut
						<i>Escherichia coli</i> 2845650	Uncharacterized protein ₁₃₋₁₉	gut
						<i>Prevotella</i> sp. CAG:1320	Putative thiol:disulfide interchange protein DsbD ₈₋₁₄	gut
						<i>Enterococcus faecalis</i> 06-MB-DW-09	Putative transmembrane permease MsmF ₁₆₋₂₂	gut
	GFL F _{LTW} ₂₅₋₃₁	No homological heptamers in commensal or opportunistic bacteria			GFL _{LTW} ₂₅₋₃₁	No homological heptamers in commensal or opportunistic bacteria		

End of the table 5

Mutation	Wuhan-Hu				Delta			
	M protein heptamer	Species	Homologous protein heptamer	Localization in the human body	M protein heptamer	Species	Homologous protein heptamer	Localization in the human body
	FL ^F LTW ₂₆₋₃₂	No homological heptamers in commensal or opportunistic bacteria			FL ^L LTW ₂₆₋₃₂	No homological heptamers in commensal or opportunistic bacteria		
	L ^F LTW ₂₇₋₃₃	No homological heptamers in commensal or opportunistic bacteria			L ^L LTW ₂₇₋₃₃	<i>Peptoniphilus</i> sp. oral taxon 375 str. F0436	Na ⁺ /H ⁺ antiporter family protein ₁₀₅₋₁₁₁	gut
	F ^F LTW ₂₈₋₃₄	No homological heptamers in commensal or opportunistic bacteria			L ^L LTW ₂₈₋₃₄	No homological heptamers in commensal or opportunistic bacteria		
V ₇₀ L	CFVLA ^V ₆₄₋₇₀	<i>Enterobacter</i> sp. Ag1	Formate dehydrogenase-O subunit gamma ₂₄₋₃₀	gut	CFVLA ^L ₆₄₋₇₀	No homological heptamers in commensal or opportunistic bacteria		
	FVLA ^V ₆₅₋₇₁	No homological heptamers in commensal or opportunistic bacteria			FVLA ^L ₆₅₋₇₁	<i>Bacteroides</i> dorei CL03T12C01	HAD hydrolase, family 1A ₃₄₄₋₃₅₀	gut
	VLA ^V ₆₆₋₇₂	No homological heptamers in commensal or opportunistic bacteria			VLA ^L ₆₆₋₇₂	<i>Bifidobacterium longum</i> subsp. infantis (strain ATCC 15697 / DSM 20088 / JCM 1222 / NCTC 11817 / S12)	Putative ABC transporter permease component ₁₁₀₋₁₁₆	gut
						<i>Haemophilus parainfluenzae</i> ATCC 33392	ABC transporter, permease protein ₁₂₁₋₁₂₇	upper respiratory tract, lung
	LAA ^V ₆₇₋₇₃	<i>Lachnospiraceae</i> bacterium 3_1_57FAA_CT1	Uncharacterized protein ₁₃₀₋₁₃₆	gut	LAA ^L ₆₇₋₇₃	<i>Acinetobacter</i> sp. CIP 101966	Uncharacterized protein ₁₈₋₂₄	oral cavity
	AA ^V ₆₈₋₇₄	<i>Lautropia mirabilis</i> ATCC 51599	Selenide, water dikinase ₆₆₋₆₂	oral cavity, upper respiratory tract	AA ^L ₆₈₋₇₄	<i>Prevotella melaninogenica</i> (strain ATCC 25845 / DSM 7089 / JCM 6325 / VPI 2381 / B282) GN=HMPREF0659_A647	Hydrolase, NUDIX family ₅₄₋₆₀	upper respiratory tract
		<i>Lachnospiraceae</i> bacterium JC7	Diguanylate cyclase (GGDEF) domain-containing protein (Precursor) ₁₁₄₋₁₂₀	gut		<i>Lactobacillus ruminis</i> (strain ATCC 27782 / RF3)	Conserved hypothetical YitT family protein	gut
						<i>Bacteroides nordii</i> CL02T12C05	Uncharacterized protein ₇₀₀₋₇₀₆	gut
	AV ^V ₆₉₋₇₅	No homological heptamers in commensal or opportunistic bacteria			AL ^V ₆₉₋₇₅	No homological heptamers in commensal or opportunistic bacteria		
	VY ^V ₇₀₋₇₆	No homological heptamers in commensal or opportunistic bacteria			LY ^V ₇₀₋₇₆	No homological heptamers in commensal or opportunistic bacteria		
I ₈₂ T	ITGGIA ^I ₇₆₋₈₂	<i>Ruminococcus obeum</i> ATCC 29174	Ion channel ₁₄₃₋₁₄₉	gut	ITGGIA ^T ₇₆₋₈₂	<i>Enterococcus faecalis</i>	Dephospho-CoA kinase ₇₋₁₃	gut
		<i>Bacteroides</i> sp. 3_1_19	Putative uncharacterized protein ₁₅₈₋₁₆₄	gut		<i>Clostridium asparagiforme</i> DSM 15981	ABC transporter, permease protein ₂₆₈₋₂₇₄	gut
	TGGIA ^I ₇₇₋₈₃	No homological heptamers in commensal or opportunistic bacteria			TGGIA ^T ₇₇₋₈₃	<i>Veillonella</i> sp. oral taxon 780 str. F0422	PrpF protein ₃₁₂₋₃₁₈	oral cavity
	GGIA ^I ₇₈₋₈₄	<i>Enterobacteriaceae</i> bacterium 9_2_54FAA	Uncharacterized protein ₂₇₀₋₂₇₆	gut	GGIA ^T ₇₈₋₈₄	No homological heptamers in commensal or opportunistic bacteria		
		<i>Eubacterium sulci</i> ATCC 35585	Peptidase, M20/M25/M40 family ₁₃₆₋₁₄₂	gut				
		<i>Lactobacillus brevis</i> subsp. <i>gravesensis</i> ATCC 27305	Transporter, major facilitator family protein ₄₂₁₋₄₂₇	gut				
	GIA ^I ₇₉₋₈₅	<i>Lachnospiraceae</i> bacterium 10-1	Uncharacterized protein ₁₄₈₋₁₅₄	gut	GIA ^T ₇₉₋₈₅	<i>Enterobacter aerogenes</i> UCI 48	Uncharacterized protein ₃₂₀₋₃₂₆	gut
	IA ^I ₈₀₋₈₆	No homological heptamers in commensal or opportunistic bacteria			IA ^T ₈₀₋₈₆	No homological heptamers in commensal or opportunistic bacteria		
	AI ^I ₈₁₋₈₇	No homological heptamers in commensal or opportunistic bacteria			AI ^T ₈₁₋₈₇	<i>Lactobacillus paracasei</i> subsp. <i>paracasei</i> CNCM I-4649	Class II aldolase/adducin family protein ₁₀₁₋₁₀₇	oral cavity, gut
	I ^I AMAC ₈₂₋₈₈	No homological heptamers in commensal or opportunistic bacteria			I ^T AMAC ₈₂₋₈₈	No homological heptamers in commensal or opportunistic bacteria		

*The same mutation has occurred in Omicron variant.

Membrane protein

There are four mutations known in the membrane (M) protein Delta variant, namely A₂S, F₂₈L, V₇₀L, and I₈₂T [10].

M protein Delta variant, 222 aa
 MDSNGTITVEELKLLLEQWNLVIGFLLLTWICLLQFAYANRN
 RFLYIIKLIFLWLLWPVTLACFVLAALYRINWITGGIATAMACL
 VGLMWLSYFIASFRLFARTSRMWSFNPETNILLNVPLHGTILT

RPLLESELVIGAVILRGHLRIAGHHLGRCDIKDLPKEITVATSR
 TLSYYKLGASQQRVAGDSGFAAYSRYRIGNYKLNTHSSSSD
 NIALLVQ

The heptamers of M protein that are homologous with the proteins of the commensal and opportunistic bacteria are listed in Table 6.

Table 6

The heptamers of M protein homologous with the proteins of the commensal and opportunistic bacteria

Mutation	Wuhan-Hu				Delta			
	M protein heptamer	Species	Homologous protein heptamer	Localization in the human body	M protein heptamer	Species	Homologous protein heptamer	Localization in the human body
A ₂ S	MA _{DSNGT} ₁₋₇	No homological heptamers in commensal or opportunistic bacteria			MS _{DSNGT} ₁₋₇	No homological heptamers in commensal or opportunistic bacteria		
	A _{DSNGT} ₁₂₋₈	<i>Lachnospiraceae bacterium 7_1_58FAA</i>	Uncharacterized protein ₂₅₂₋₂₅₈	gut	S _{DSNGT} ₁₂₋₈			
F ₂₈ L	LVIGFL _F ₂₂₋₂₈	<i>Enterococcus faecalis R508</i>	Putative ferrichrome transport system permease protein PhuG ₂₀₃₋₂₀₆	gut	LVIGFL _L ₂₂₋₂₈	<i>Eubacterium ventriosum</i> ATCC 27560	Putative K(+)-stimulated pyrophosphate-energized sodium pump ₅₇₃₋₅₇₉	gut
						<i>Enterococcus caccae</i> ATCC BAA-1240	Uncharacterized protein ₁₀₄₋₁₁₀	gut
						<i>Faecalibacterium</i> sp. CAG:74	Binding-protein-dependent transport systems inner membrane component ₈₆₋₉₂	gut
						<i>Prevotella histicola</i> F0411	Uncharacterized protein ₁₅₋₂₁	gut
						<i>Lachnospiraceae bacterium 2_1_58FAA</i>	Uncharacterized protein ₆₅₋₇₁	gut
						<i>Escherichia coli</i> ISC11	Putative cell envelope opacity-associated protein A ₄₂₋₄₈	gut
	VIGFL _F ₂₃₋₂₉	<i>Enterococcus flavescens</i> ATCC 49996	Uncharacterized protein ₁₂₈₋₁₃₄	gut	VIGFL _L ₂₃₋₂₉	<i>Prevotella</i> sp. oral taxon 472 str. F0295	Uncharacterized protein ₁₇₈₋₁₈₄	gut
						<i>Lactobacillus brevis</i> ATCC 14869 = DSM 20054	Potassium uptake protein, TrkH family ₂₃₉₋₂₄₅	gut
						<i>Lactobacillus antri</i> DSM 16041	Transporter, major facilitator family protein ₄₂₂₋₄₂₈	gut
						<i>Enterobacter cloacae</i> subsp. <i>cloacae</i> (strain ATCC 13047 / DSM 30054 / NBRC 13535 / NCDC 279-56)	Putative multidrug resistance protein MdtD ₁₈₃₋₁₈₉	gut
						<i>Lachnospiraceae bacterium 28-4</i>	Uncharacterized protein ₁₈₋₂₄	gut
	IGFL _F _{LT} ₂₄₋₃₀	<i>Lachnospiraceae bacterium CAG:215</i>	Transporter ₄₆₈₋₄₇₄	gut[9]	IGFL _L _{LT} ₂₄₋₃₀	<i>Lactobacillus paracasei</i> subsp. <i>paracasei</i> Lpp126	Oligopeptide transport system permease protein oppB ₉₋₁₅	oral cavity
						<i>Eubacterium nodatum</i> ATCC 33099	TIGR02185 family protein ₄₃₋₄₉	oral cavity
						<i>Bacteroides uniformis</i> dnLKV2	Uncharacterized protein ₇₃₇₋₇₄₃	gut
						<i>Escherichia coli</i> 2845650	Uncharacterized protein ₁₃₋₁₉	gut
						<i>Prevotella</i> sp. CAG:1320	Putative thiol:disulfide interchange protein DsbD ₈₋₁₄	gut
						<i>Enterococcus faecalis</i> 06-MB-DW-09	Putative transmembrane permease MsmF ₁₆₋₂₂	gut
	GFL _F _{LTW} ₂₅₋₃₁	No homological heptamers in commensal or opportunistic bacteria			GFL _L _{LTW} ₂₅₋₃₁	No homological heptamers in commensal or opportunistic bacteria		

End of the table 6

Mutation	Wuhan-Hu				Delta			
	M protein heptamer	Species	Homologous protein heptamer	Localization in the human body	M protein heptamer	Species	Homologous protein heptamer	Localization in the human body
	FLFLTWI ₂₆₋₃₂	No homological heptamers in commensal or opportunistic bacteria			FLLLTWI ₂₆₋₃₂	No homological heptamers in commensal or opportunistic bacteria		
	LFLTWIC ₂₇₋₃₃	No homological heptamers in commensal or opportunistic bacteria			LLLTWIC ₂₇₋₃₃	<i>Peptoniphilus sp. oral taxon 375 str. F0436</i>	Na ⁺ /H ⁺ antiporter family protein ₁₀₅₋₁₁₁	gut
	FLTWICL ₂₈₋₃₄	No homological heptamers in commensal or opportunistic bacteria			LLTWICL ₂₈₋₃₄	No homological heptamers in commensal or opportunistic bacteria		
V ₇₀ L	CFVLAALV ₆₄₋₇₀	<i>Enterobacter sp. Ag1</i>	Formate dehydrogenase-O subunit gamma ₂₄₋₃₀	gut	CFVLAAL ₆₄₋₇₀	No homological heptamers in commensal or opportunistic bacteria		
	FVLAALV ₆₅₋₇₁	No homological heptamers in commensal or opportunistic bacteria			FVLAALY ₆₅₋₇₁	<i>Bacteroides dorei</i> CL03T12C01	HAD hydrolase, family IA ₃₄₄₋₃₅₀	gut
	VLAALV ₆₆₋₇₂	No homological heptamers in commensal or opportunistic bacteria			VLAALY ₆₆₋₇₂	<i>Bifidobacterium longum</i> subsp. <i>infantis</i> (strain ATCC 15697 / DSM 20088 / JCM 1222 / NCTC 11817 / S12)	Putative ABC transporter permease component ₁₁₀₋₁₁₆	gut
						<i>Haemophilus parainfluenzae</i> ATCC 33392	ABC transporter, permease protein ₁₂₁₋₁₂₇	upper respiratory tract, lung
	LAAVYRI ₆₇₋₇₃	<i>Lachnospiraceae bacterium 3_1_57FAA_CT1</i>	Uncharacterized protein ₁₃₀₋₁₃₆	gut	LAALYRI ₆₇₋₇₃	<i>Acinetobacter sp. CIP 101966</i>	Uncharacterized protein ₁₈₋₂₄	oral cavity
	AAVYRIN ₆₈₋₇₄	<i>Lautropia mirabilis</i> ATCC 51599	Selenide, water dikinase ₅₆₋₆₂	oral cavity, upper respiratory tract	AALYRIN ₆₈₋₇₄	<i>Prevotella melaninogenica</i> (strain ATCC 25845 / DSM 7089 / JCM 6325 / VPI 2381 / B282) GN=HMPREF0659_A647	Hydrolase, NUDIX family ₅₄₋₆₀	upper respiratory tract
		<i>Lachnospiraceae bacterium JC7</i>	Diguanylate cyclase (GGDEF) domain-containing protein (Precursor) ₁₁₄₋₁₂₀	gut		<i>Lactobacillus ruminis</i> (strain ATCC 27782 / RF3)	Conserved hypothetical YitF family protein	gut
						<i>Bacteroides nordii</i> CL02T12C05	Uncharacterized protein ₇₀₀₋₇₀₆	gut
	AVYRINW ₆₉₋₇₅	No homological heptamers in commensal or opportunistic bacteria			ALYRINW ₆₉₋₇₅	No homological heptamers in commensal or opportunistic bacteria		
	VYRINWI ₇₀₋₇₆	No homological heptamers in commensal or opportunistic bacteria			LYRINWI ₇₀₋₇₆	No homological heptamers in commensal or opportunistic bacteria		
I ₈₂ T	ITGGIAL ₇₆₋₈₂	<i>Ruminococcus obeum</i> ATCC 29174	Ion channel ₁₄₃₋₁₄₉	gut	ITGGIAT ₇₆₋₈₂	<i>Enterococcus faecalis</i>	Dephospho-CoA kinase ₇₋₁₃	gut
		<i>Bacteroides sp. 3_1_19</i>	Putative uncharacterized protein ₁₅₈₋₁₆₄	gut		<i>Clostridium asparagiforme</i> DSM 15981	ABC transporter, permease protein ₂₆₈₋₂₇₄	gut
	TGGIAL ₇₇₋₈₃	No homological heptamers in commensal or opportunistic bacteria			TGGIAT ₇₇₋₈₃	<i>Veillonella sp. oral taxon 780 str. F0422</i>	PrpF protein ₃₁₂₋₃₁₈	oral cavity
	GGIALAM ₇₈₋₈₄	<i>Enterobacteriaceae bacterium 9_2_54FAA</i>	Uncharacterized protein ₂₇₀₋₂₇₆	gut	GGIATAM ₇₈₋₈₄	No homological heptamers in commensal or opportunistic bacteria		
		<i>Eubacterium sulci</i> ATCC 35585	Peptidase, M20/M25/M40 family ₁₃₆₋₁₄₂	gut				
		<i>Lactobacillus brevis</i> subsp. <i>gravesensis</i> ATCC 27305	Transporter, major facilitator family protein ₄₂₁₋₄₂₇	gut				
	GIALAMA ₇₉₋₈₅	<i>Lachnospiraceae bacterium 10-1</i>	Uncharacterized protein ₁₄₈₋₁₅₄	gut	GIAATAMA ₇₉₋₈₅	<i>Enterobacter aerogenes</i> UCI 48	Uncharacterized protein ₃₂₀₋₃₂₆	gut
	IALAMAC ₈₀₋₈₆	No homological heptamers in commensal or opportunistic bacteria			IATAMAC ₈₀₋₈₆	No homological heptamers in commensal or opportunistic bacteria		
	ALAMACL ₈₁₋₈₇	No homological heptamers in commensal or opportunistic bacteria			ATAMACL ₈₁₋₈₇	<i>Lactobacillus paracasei</i> subsp. <i>paracasei</i> CNCM I-4649	Class II aldolase/adducin family protein ₁₀₁₋₁₀₇	oral cavity, gut
	IAMACLV ₈₂₋₈₈	No homological heptamers in commensal or opportunistic bacteria			TAMACLV ₈₂₋₈₈	No homological heptamers in commensal or opportunistic bacteria		

Nucleocapsid protein

Two mutations are known in the Delta variant nucleocapsid (N) protein, namely R₂₀₃M and D₃₇₇Y [11].

N protein Delta variant 419 aa

MSDNGPQNQRNAPRITFGGSPDSTGSNQNGERSGARSKQR
 RPQGLPNNTASWFTALTQHGKEDLKFP**RGQGVPI**NTNS
 SPDDQIGYYRRATRRIRGGD**GKMKDLS**PRWYFYLTGT
 PEAGLPYGANKDGIWVATEGALNTPKDHIGTRNPANNA
 AIVLQLPQGTTLPGKFY**AEGRGGSQ**ASSRSSRSRNS

SRNSTPGSSMGTSPARMAGNGGDAALALLLDRLNQL
 ESKMSGKGGQQQQGQTVTKKSAEASKKPRQKRTATKA
 YNVTQAFGRRGPEQTQGNFGDQELIRQGTDYKHWPI
 AQFAPSASAFFGMSRIGMEVTPSGTWLTYTGAIKLDDK
 DPNFKDQVILLNKHIDAYKTFPTEPKDKKK**KAYETQA**
 LPQRQKKQQTVT**LLPAADLDDFSKQLQQSMSSADSTQA**

The heptamers of N protein homologous with the proteins of some opportunistic bacteria and the most common cereals are listed in Table 7.

Table 7

The heptamers of N protein homologous with the proteins of some opportunistic bacteria and the most common cereals

Mutation	Wuhan-Hu				Delta			
	N protein heptamer	Species	Homologous protein heptamer	Localization in the human body	N protein heptamer	Species	Homologous protein heptamer	Localization in the human body
R ₂₀₃ M	STPGSSR ₁₉₇₋₂₀₃	<i>Prevotella buccalis</i> ATCC 35310	NHL repeat protein ₃₀₆₋₃₁₂	oral cavity	STPGSSM ₁₉₇₋₂₀₃	No bacterial or cereal sample		
	TPGSSRG ₁₉₈₋₂₀₄	No bacterial or cereal sample			TPGSSMG ₁₉₈₋₂₀₄	<i>Bacteroides uniformis</i> CAG:3	Uncharacterized protein ₁₂₈₋₁₃₄	gut
	PGSSRGT ₁₉₉₋₂₀₅	<i>Zea mays</i>	Putative WRKY DNA-binding domain superfamily protein ₇₈₋₈₄	gut	PGSSMGT ₁₉₉₋₂₀₅	<i>Oryza sativa</i> subsp. <i>indica</i>	Putative uncharacterized protein ₅₅₈₋₅₆₄	gut
		<i>Sorghum bicolor</i>	Putative uncharacterized protein Sb07g002490 ₂₇₋₃₃	gut				
	GSSRGTS ₂₀₀₋₂₀₆	<i>Sorghum bicolor</i>	Putative uncharacterized protein Sb08g014350 ₁₇₆₋₁₈₂	gut	GSSMGTS ₂₀₀₋₂₀₆	<i>Fusobacterium</i> sp. CM21	Permease family protein ₂₉₄₋₃₀₀	oral cavity
	SSRGTSP ₂₀₁₋₂₀₇	<i>Hordeum vulgare</i> var. <i>distichum</i>	Uncharacterized protein ₂₆₇₋₂₇₃	gut	SSMGTS ₂₀₁₋₂₀₇	No bacterial or cereal sample		
		<i>Oryza sativa</i> subsp. <i>japonica</i>	Expressed protein ₂₁₆₋₂₂₂	gut				
	SRGTSPA ₂₀₂₋₂₀₈	No bacterial or cereal sample			SMGTSPA ₂₀₂₋₂₀₈	No bacterial or cereal sample		
D ₃₇₇ Y	RGTSRAR ₂₀₃₋₂₀₉	<i>Oryza sativa</i> subsp. <i>japonica</i>	Os06g0523800 protein ₁₁₈₋₁₂₄	gut	MGTSRAR ₂₀₃₋₂₀₉	No bacterial or cereal sample		
	DKKKKA ₃₇₁₋₃₇₇	<i>Prevotella</i> sp. oral taxon 473 str. F0040	Pseudouridine synthase, RluA family ₂₉₅₋₃₀₁	oral cavity	DKKKKAY ₃₇₁₋₃₇₇	<i>Lachnospiraceae</i> bacterium 3-1	Oligoendopeptidase F ₄₃₉₋₄₄₅	gut
		<i>Prevotella</i> sp. oral taxon 473 str. F0040	Pseudouridine synthase, RluA family ₂₉₆₋₃₀₂	oral cavity				
		<i>Enterococcus faecalis</i>	Uncharacterized protein ₃₉₆₋₄₀₂	gut				
	KKKA ₃₇₂₋₃₇₇	No significant sample			KKKAY ₃₇₂₋₃₇₇	<i>Oryza sativa</i> subsp. <i>indica</i>	Putative uncharacterized protein ₁₀₉₀₋₁₀₉₆	gut
	KKKADET ₃₇₃₋₃₇₉	No bacterial or cereal sample			KKKAYET ₃₇₃₋₃₇₉	<i>Bacillus infantis</i> NRRL B-14911	GntR family transcriptional regulator ₂₋₈	?
	KKADETQ ₃₇₄₋₃₈₀	No bacterial or cereal sample			KKAYETQ ₃₇₄₋₃₈₀	No bacterial or cereal sample		
	KADETQA ₃₇₅₋₃₈₁	<i>Homo sapiens</i>	Myopalladin ₉₀₋₉₆	?	KAYETQA ₃₇₅₋₃₈₁	No bacterial or cereal sample		
	ADETQAL ₃₇₆₋₃₈₂	<i>Oryza glaberrima</i>	Uncharacterized protein (Fragment) ₄₇₄₋₄₈₀	gut	AYETQAL ₃₇₆₋₃₈₂	<i>Lachnospiraceae</i> bacterium M18-1	Uncharacterized protein ₂₄₄₋₂₅₀	gut
	DETQALP ₃₇₇₋₃₈₃	No bacterial or cereal sample			YETQALP ₃₇₇₋₃₈₃	<i>Lachnospiraceae</i> bacterium M18-1	Uncharacterized protein ₂₄₅₋₂₅₁	gut



As shown above, some of the mutations that occurred in the Delta variant increased the homology of its structural proteins with those of the opportunistic and some other bacteria. These data are summarized in Table 8.

Information about the effects that mutations in SARS CoV-2 Delta variant have on the homology between its structural proteins and human opportunistic bacteria proteins are summarized in Figure 1.

DISCUSSION

In Wunan-Hu variant, the S protein molecule contains dozens of heptamers homologous to human proteins. Their total length is 169 amino acid residues, or 13.3% of the S protein molecule total length [5]. For the sake of brevity, we suggest calling **homologous motifs homots**. For example, a SARS CoV-2 S protein human homot means a motif common for the S protein and any human protein. The same way “in SARS CoV-2 S protein, the motif SPRRARS is a human homot” means that motif SPRRARS is present in the S protein of coronavirus as well as in some human protein. The term *mimics*, proposed by Damoiseaux et al. [12], is close in meaning but less specific.

We assumed that the reason for the special qualities of SARS CoV-2 Delta variant should be sought in the greater homology of its proteins with those of the human body. However, we did not find any significant differences between Wuhan-Hu variant and Delta variant in their homology to human proteins.

Delta variant stays on the nasal mucosal surface significantly longer than Wuhan-Hu variant (14 vs. 8 days) [13].

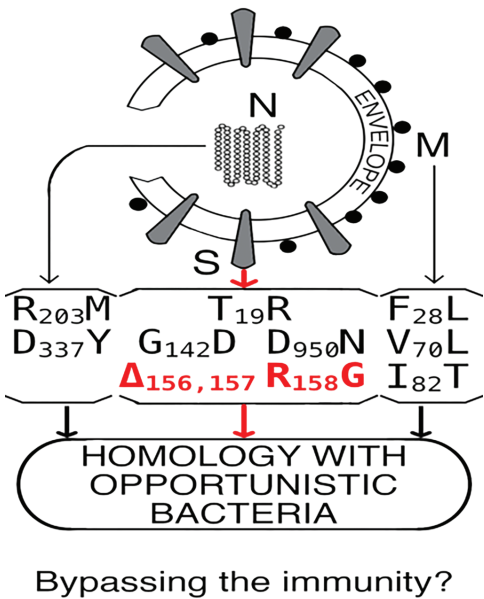


Fig. 1. The effect of mutations in SARS CoV-2 Delta variant structural proteins S, M, and N on their homology with human opportunistic bacteria. The most important mutation, in our opinion, is highlighted in red font

As has been already mentioned, we considered the human proteome in general as a set of proteins synthesized by the macroorganism itself, proteins of commensal and opportunistic bacteria, and the most common digestive proteins, therefore studying the homology of SARS CoV-2 Delta variant with all the listed types of proteins.

Table 8

Mutational changes of homology SARS CoV-2 structural proteins with proteins of opportunistic bacteria and some other functionally significant proteins

Protein	Mutation	Increases homology with proteins of commensal or opportunistic bacteria, inhabitants of the oral cavity, upper respiratory tract or lung	Increases homology with proteins of gut commensal or opportunistic bacteria and/or the most common cereals	Increases homology with some other proteins
S (Table 2)	T ₁₉ R	+	+	
	G ₁₄₂ D*	+	-	
	Δ _{156,157} , R ₁₅₈ G	+++	+++	Homology with a protein of <i>Bacillus</i> sp. NRRL B-14911 that can provoke autoimmune damage to the heart
	L ₄₅₂ R	-	-	
	T ₄₇₈ K*	-	-	
	P ₆₈₁ H	-	-	
	D ₉₅₀ N*	+	+	Homology with a protein of <i>Human immunodeficiency virus 1</i> (Table 4)
M (Table 6)	A ₂ S	-	-	
	F ₂₈ L	+	++	
	V ₇₀ L	++	++	
	I ₈₂ T	+	-	
N (Table 7)	R ₂₀₃ M	-	-	
	D ₃₇₇ Y	-	+	

*The same mutation has occurred in Omicron variant.

In S protein, mutations at the positions 19, 142, 156-158, and 950 created a number of heptamers homologous to proteins of bacteria, that are always present in the human nasopharynx, mouth, throat, upper respiratory tract, and lung (Table 2). It is possible that the presence of such homologous motifs allows Delta to bypass the innate immunity protection more successfully.

Mutations S:G₁₄₂D and S:D₉₅₀N are also found in Omicron variant, while the mutations S:T₁₉R and S:Δ_{156,157};R₁₅₈G are only present in Delta variant. These exclusive Delta variant mutations especially the ones at the positions 156-158 may be the reason for its specific qualities.

The L₄₅₂R and T₄₇₈K mutations did not affect the homology of S protein with proteins of opportunistic bacteria (Table 2).

In Delta variant, the positions where the most significant increase in homology occurred — S:Δ_{156,157};R₁₅₈G — are located in the N-terminus domain (NTD₁₄₋₃₀₃). So far, researchers have paid less attention to this domain than to the Receptor-binding domain (RBD₃₁₇₋₅₃₉). It is logically consistent to assume that in the S protein molecule one domain is responsible for binding to the receptor and other for structural mimicry and evasion.

The delta variant differs from the other SARS COV-2 variants in 14 positions. According to our data (Fig. 1), six of these alterations involved in the increase in the homology of coronavirus proteins with those of opportunistic bacteria. None of these six alterations are common to the Delta and non-VOC variants. This suggests that the increase in homology with proteins of opportunistic infections is specific to the Delta variant.

We are not yet able to analyze homology data for SARS CoV-2 S protein and the HIV-1 C protein (Table 4).

In M protein, the F₂₈L, V₇₀L, and I₈₂T mutations resulted in the emergence of heptamers homologous to proteins of numerous commensal and opportunistic upper respiratory and gut bacteria (Table 6). M protein is located on the outer side of the virion envelope [5], and these heptamers can participate in immune evasion.

In N protein (Table 7), the mutation N:R₂₀₃M resulted in the motif GSSMGTS₂₀₀₋₂₀₆ which is homologous to the Permease family protein₂₉₄₋₃₀₀ of *Fusobacterium nucleatum*, an opportunistic periodontal pathogen of the oral cavity [14]. The mutation M:D₃₇₇Y caused the following effects: (a) disappearance of the heptamer KADETQA₃₇₅₋₃₈₁, homologous to the human protein Myopalladin (MYPN₉₀₋₉₆), which is involved in communication between the sarcomere and the nucleus in cardiac and skeletal muscles [15]; and (b) emergence of KKKAYET₃₇₃₋₃₇₉, homologous to the heptamer GntR family transcriptional regulator₂₋₈ *Bacillus infantis*, which is involved in the provocation of immune myocardial disorder [16].

A recent review of the available evidence for immune mechanisms of cardiovascular damage COVID-19 has been presented [17]. N protein, located inside of the virion,

should act at the later stages of the infectious process, for example, provoking an autoimmune response.

Of all the Delta variant mutations we studied, none caused an increase in the homology of the SARS CoV-2 S protein with proteins with the most common cereals (Table 3).

Natural selection fixes some substitutions in the primary structure of the protein molecules of viruses and eliminates others. One of the “aims” of selection might be immune evasion. A virus can achieve this by making the most functionally important parts of the protein molecule as similar as possible to the proteins permanently present in the host. Microorganisms, due to their genetic diversity and the huge size of their combined genome, provide more opportunities for viral mimicry than the macroorganism itself. Delta variant has increased homology of S and M proteins with proteins already familiar to human immunity, namely with opportunistic bacteria proteins.

The capacity of SARS CoV-2 for immune evasion can be considered universally acknowledged [3]. Coronavirus and human protein homology may be one of the mechanisms of immune evasion [5]. Delta variant necessarily has structural features that explain its specific qualities. Perhaps the reason is the homology of its proteins with those of commensal bacteria and opportunistic infections of the upper respiratory tract and lung. In this case, the S:Δ_{156,157};R₁₅₈G mutation deserves special attention. The reason why SARS CoV-2 Delta variant has these specific qualities, most importantly increased lethality, is most likely to be found in a mutation at positions 156-158 of spike protein. It has not yet been concluded whether the homology of Delta variant proteins with gut bacteria proteins and dietary protein is of any significance.

We hope that this preliminary study will open the door to further research into the immunology and bioinformatics.

METHODS

We used our original way of presenting the text search. The data were obtained from the Uniprot open-access protein database, in which the amino acid sequences of proteins are encoded by a one-letter code. We cut the primary structures of the coronavirus proteins into heptamers using the frame-shift method and searched a separate database of 75777 molecules of human proteins [18]. This number is about three times the real number of all human proteins because of repetition and minor differences in the records. We looked for a full match of the 7-mer amino acid sequences in SARS CoV-2 proteins [19] with proteins of other organisms throughout the taxonomic range of evolution from bacteria and plants to humans. Heptamers were chosen as a criterion for homology because of the lack of matches in octamers and tens of thousands of matches in hexamers. In the case of matching heptamers, an alignment was performed on the matching site.

ДОПОЛНИТЕЛЬНАЯ ИНФОРМАЦИЯ

Вклад авторов. А.Т. Марьянович и Д.К. Кормилец написали основной текст рукописи. А.Т. Марьянович и Д.К. Кормилец подготовили анализ данных. Авторы прочли и одобрили финальную версию перед публикацией.

Конфликт интересов. Авторы декларируют отсутствие явных и потенциальных конфликтов интересов, связанных с публикацией настоящей статьи.

Источник финансирования. Данное исследование представляет собой инициативный проект авторов, финансируемый исключительно из их личных источников.

Заявление о доступности данных. Источником базы данных по 75 777 строкам белков человека является [18]. Источник базы данных объемом ок. 33 млн нитей всех видов белков [19].

Иллюстрации. Для создания наших иллюстраций мы использовали GIMP (версия 2.10.22). Рисунок полностью оригинальный и нигде не публиковался.

ADDITIONAL INFORMATION

Author contributions. A.T. Maryanovich and D.Yu. Kormilets wrote the main manuscript text. A.T. Maryanovich and D.Yu. Kormilets prepared data analysis. The authors read and approved the final version before publication.

Competing interests. The authors declare that they have no competing interests.

Funding: This research is an authors' initiative project funded exclusively from their personal sources.

Funding source. This study was not supported by any external sources of funding.

Data Availability Statement: The source of database of 75777 strings of human proteins is [18]. The source of database of approx. 33 mln strings of all species proteins is [19].

Artwork. We used GIMP (Version 2.10.22) to create our artwork. The figure is completely original and have not been published anywhere.

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