Toxin-like peptides in plasma, urine and faecal samples from COVID-19 patients [version 2; peer review: 2 approved]

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Abstract

**Background:** SARS-CoV-2 that causes COVID-19 disease and led to the pandemic currently affecting the world has been broadly investigated. Different studies have been performed to understand the infection mechanism, and the involved human genes, transcripts and proteins. In parallel, numerous clinical extra-pulmonary manifestations co-occurring with COVID-19 disease have been reported and evidence of their severity and persistence is increasing. Whether these manifestations are linked to other disorders co-occurring with SARS-CoV-2 infection, is under discussion. In this work, we report the identification of toxin-like peptides in COVID-19 patients by application of the Liquid Chromatography Surface-Activated Chemical Ionization – Cloud Ion Mobility Mass Spectrometry.

**Methods:** Plasma, urine and faecal samples from COVID-19 patients and control individuals were analysed to study peptidomic toxins’ profiles. Protein precipitation preparation procedure was used for plasma, to remove high molecular weight proteins and efficiently solubilize the peptide fraction; in the case of faeces and urine, direct peptide solubilization was employed.

**Results:** Toxin-like peptides, almost identical to toxic components of venoms from animals, like conotoxins, phospholipases, phosphodiesterases, zinc metal proteinases, and bradykinins, were identified in samples from COVID-19 patients, but not in control samples.

**Conclusions:** The presence of toxin-like peptides could potentially be connected to SARS-CoV-2 infection. Their presence suggests a possible association between COVID-19 and other disorders.
COVID-19 disease and the release in the body of (oligo-)peptides almost identical to toxic components of venoms from animals. Their involvement in a large set of heterogeneous extra-pulmonary COVID-19 clinical manifestations, like neurological ones, cannot be excluded. Although the presence of each individual symptom is not selective of the disease, their combination might be related to COVID-19 by the coexistence of the panel of the here detected toxin-like peptides. The presence of these peptides opens new scenarios on the aetiology of the COVID-19 clinical symptoms observed up to now, including neurological manifestations.

**Keywords**
SARS-CoV-2, COVID-19, toxin-like peptides

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Introduction
Numerous clinical extra-pulmonary manifestations co-occurring with COVID-19 disease have been reported (e.g. neurological, haemorrhagic, and thrombotic) and evidence of their severity and persistence is increasing. Gupta et al. reviewed the extrapulmonary organ-specific pathophysiology of patients with COVID-19, ‘to aid clinicians and scientists in recognizing and monitoring the spectrum of manifestations, and in developing research priorities and therapeutic strategies for all organ systems involved’. Liotta et al. characterized the incidence of neurological manifestations in a cohort of hospitalised patients with confirmed COVID-19: the most frequent were myalgia, headache, encephalopathy, dizziness, dysgeusia, and anosmia; encephalopathy was found to be ‘associated with increased morbidity and mortality, independent of respiratory disease severity’. Whether these manifestations are linked to disorders co-occurring with SARS-CoV-2 infection is under discussion, including their concomitant occurrence, which could be strongly related COVID-19 disease. Frontera et al., by conducting a prospective, multi-centre, observational study of hospitalised adults with laboratory-confirmed SARS-CoV-2 infection, concluded that ‘neurologic disorders were detected in 13.5% of COVID-19 patients during the study timeframe. Many of these neurologic disorders occur commonly among patients with critical illness. Encephalitis, meningitis or myelitis referable to SARS-CoV-2 infection did not occur, though post-infectious Guillain-Barre syndrome was identified. Overall, neurologic disorders in the context of SARS-CoV-2 infection confer a higher risk of in-hospital mortality and reduced likelihood of discharge home’. Studies on the use of mass spectrometry in COVID-19 context focus on the search for augmented human inflammatory molecules to be used as biomarkers to assess the severity status of COVID-19 (see for example the work of Messner and colleagues). Different studies report the use of proteomic approaches to characterise SARS-CoV-2 proteins. Other studies highlight challenges in their use due to the need of enriching the protein fraction to be analysed for maximizing the technology sensitivity.

Liquid Chromatography Surface-Activated Chemical Ionization – Cloud Ion Mobility Mass Spectrometry (LC-SACI-CIMS) is reported as a high sensitivity mass spectrometry technique able to maximize the peptide signal intensity. We used LC-SACI-CIMS to reveal the presence of metabolites that could explain the clinical descriptions of neurological, coagulation and inflammatory symptoms, and here we present the results of our analyses. We found toxin-like peptides in plasma, urine, and faecal samples from COVID-19 patients, but not in control samples. As our findings do not correspond with current thinking of the aetiology related to the observed clinical manifestations in COVID-19 patients, we feel their immediate sharing with the scientific community is critical.

Methods
Rationale
Liquid Chromatography-Surface Activated Chemical Ionization – Cloud Ion Mobility Mass Spectrometry (LC-SACI-CIMS) exhibits a high selectivity in peptide detection thanks to its ability to selectively isolate peptide ions through an in-source ion mobility (IM) effect. In fact, it allows a selective regulation of the potential difference between the low voltage of the SACI surface (47 V) and the entrance lens (-50 / -600 V), and a selective focalization on solvent ion cloud species at low or high m/z ratio. By switching the entrance voltage lens between -50 and -600 V during the analysis, it is possible to separate the low m/z from the high m/z potential signal, to avoid ion trap saturation, and to maximize the number of detected compounds. The mass spectra chemical noise is also strongly reduced due to the lower amounts of solvent cluster ions that are produced in low voltage ionization conditions. Thus, the peptide detection efficiency is strongly increased by the IM selectivity and lower chemical noise with respect to the classical high voltage ionization approaches. Thanks to the specificity of the SACI-CIMS technology in focalizing the solvent ion clouds containing the high m/z (oligo-)peptide species, it was possible to increase the detection efficiency.

In the use of LC-SACI-CIMS, the following strategies have been adopted:

- To reduce the presence of contamination as much as possible and to avoid the formation of acetonitrile polymers occurring in acid conditions (as reported by Eizo et al.), formic acid was not added to the CH₃CN chromatographic phase.
- To separate low from high m/z solvent ion clusters by reducing the ion trap saturation, the space/charge effect, and by increasing the detected compounds recovery, LC-SACI-CIMS entrance lens voltage was switched between -50 and -600 V every 10 ms during the analysis.
- To enhance the SACI ionization efficiency, NH₄HCO₃ was added to the samples. As reported in the literature, the peptide ionization efficiency (and consequently the sensitivity) is enhanced in SACI conditions when ionic salts are present in the sample, due to peptide ion specific coordination.
- To decrease the total run time, a shot gun chromatographic gradient was used to desalt the sample.
- To avoid sample molecular profile alteration, and to evaluate the potential biological activities of the circulating species, no enzymatic digestion was applied to samples.

Amendments from Version 1
This version contains text additions within the “Discussion” section following the comments and the suggestions made by Dr. Grumati and Dr. Arditi in the referee report on version.

Regarding the manuscript changes, please note that we added three references after reference N.20.

No changes to figures or tables.

Any further responses from the reviewers can be found at the end of the article.
• To normalize the m/z signal intensity, 5 µL of standard ESI tune mix (Agilent, USA) were added to each sample extract.

Chemicals

NH₄HCO₃, methanol, acetonitrile and formic acid were purchased from Sigma-Aldrich (Milan, Italy). Bi-distilled water was purchased from VWR (Milan, Italy).

Cohort

Samples used in the present study: plasma samples collected from 20 COVID-19 patients from different cities of Italy and from 10 control individuals (i.e. negative to SARS-CoV-2 tests and not affected by cancer or autoimmune diseases); urine samples collected from two additional COVID-19 patients and from two control individuals; stool samples from three COVID-19 patients and from three control individuals. The human biological samples used in the experimentation were collected and used with the expressed free and informed consent, of the person from whom the material was taken, according to current legislation. The study received approval from “Comitato Etiico Campania Sud” (n.36/2021, request submitted on 06-05-2020). Apart from positivity to SARS-CoV-2, no additional information (i.e. age, sex, blood serotype, severity of the disease, time of the collection, fatality, etc.) was provided.

Sample preparation

Plasma. Each plasma sample was treated as follows: 5 µL of CH₃CN were added to 50 µL of plasma and vortexed for one minute. The procedure was repeated 10 times. Then the sample was centrifuged at 1,500 g for 10 minutes and two 100 µL aliquots of supernatant were dried and resuspended in 70 µL of NH₄HCO₃ 50 mmol. The solution was analysed by LC-SACI-CIMS (see Rationale).

Urine. Each urine sample was treated as follows: an equivalent volume of bi-distilled water was added, followed by centrifugation at 1,500 g for 10 minutes. 100 µL were dried and resuspended in 70 µL of NH₄HCO₃ 50 mmol. The sample was analysed by LC-SACI-CIMS (see Rationale).

Stool. Each stool sample was treated as described by Cristoni et al. and analysed by LC-SACI-CIMS (see Rationale).

Liquid chromatography

The Ultimate 3000 LC (by ThermoFisher) was used to achieve separation of analytes for each sample prior to mass spectrometry (MS) analysis. A reversed phase Kinetex C-18 LC column (50 × 2.1 mm; particle size, 5 µm; pore size, 100 Å, by Phenomenex, USA) was used. The eluent flow was 0.25 mL/min and the injection volume was 15 µL. The mobile phases were:

A. 0.2% (v/v) formic acid (HCOOH)

B. acetonitrile (CH₃CN)

The elution gradient was: 2% (v/v) of B between 0 and 2 min; 2 to 30% between 2 and 7 min; 30 to 80% between 7 and 9 min; 80% between 9 and 12 min; 80-2% between 12 and 12.1 min. The column was rebalanced with 2% of B between 12.1 and 17 min.

Mass spectrometry

All samples were analysed for the presence of proteins with potential toxic effect by using the LC-SACI-CIMS as already described in the literature. Samples were analysed with an ORBITRAP mass spectrometer (Breme, Germany) coupled to a surface-activated chemical ionization (SACI) source and operated in positive ion mode.

The surface voltage was 47 V and the entrance lens was switched between -50 and -600 V each 10 ms. Auxiliary gas: 2 L / min; Nebulizer gas: 80 psi; Temperature: 40 °C. Full scan spectra were acquired in the 40–3,500 m/z range for non-targeted metabolomics/proteomics analyses to detect analytes. The same m/z range was used for both discovery and selective biomarker identification, and to standardize (primarily in terms of scan rate) the instrument. The software used for data elaboration is SANIST, a modified version of the Global Proteome Machine (GPM, https://www.thegpm.org/GPM/), implanted as described in 9–12. SANIST output files are available as supplementary material (see section Data availability).

SANIST software here used is freely available, upon email request to CranioMed group (dir.brogna@craniomedi.it).

Mass spectrometry on samples was performed with collision-induced dissociation using data dependent scan and helium as the collision gas. The ion trap was applied to isolate and fragment the precursor ions (windows of isolation, ± 0.3 m/z; collision energy, 30% of its maximum value, which was 5V peak to peak), and the ORBITRAP mass analyser was used to obtain fragments with an extremely accurate m/z ratio (resolution 15,000; m/z error <10 ppm).

Data elaboration

Detected high m/z peptides were used to identify toxins thanks due to the selectivity given by their long chain.

The complete UniprotKB set of manually reviewed venom proteins and toxins (UniprotKB, Animal toxin annotation project. https://www.uniprot.org/program/Toxins, Accessed October 4, 2020), mixed with a subset of non-venom proteins and toxins from UniprotKB database was used as reference protein dataset in order to give statistical significance to the results.

TBLASTN was run at the National Center for Biotechnology Information (NCBI) website with default options and parameters, with the exception of the following ones: max target sequences = 1,000; expect threshold = 100; word size = 3; gap cost existence = 9; gap cost extension = 1; filter of low complexity regions = No. Searches have been performed
versus: Nucleotide collection (nr/nt); Reference RNA sequences (refseq_rna); RefSeq Genome Database (refseq_genomes); Whole-genome shotgun contigs (wgs) from metagenomic experiments; Sequence Read Archive (SRA) sequences from metagenomic experiments; Transcriptome Shotgun Assembly (TSA); Patent sequences (pat); Human RefSeqGene sequences (RefSeq_Gene); Betacoronavirus Genbank sequence dataset.

The information reported in Table 1 has been retrieved from the UniprotKB database and from the NCBI Taxonomy database20, after confirmation by BLAST sequence comparison analysis.

SANIST was set to perform the database search considering all potential protein points and post-translational modifications, and to consider proton rearrangements. No enzyme cutting rules were specified, but all the protein subsequence combinations were considered. Database search calculation was performed by means of General Processing Graphic Processing Units (GPGPU).

The MS data are available on the ZENODO platform16 (see section Data availability).

Results and discussion

The presence of (oligo-)peptides characterised as toxic components of animal venoms was observed in plasma and urine samples from SARS-CoV-2 infected patients and never in plasma, urine and faecal samples from control individuals. Examples of SACI-CIMS chromatograms are reported in Figure 1 and Figure 2 (panels a and b), showing the spectra acquired by means of the LC-SACI-CIMS technology. Figure 2c and d show the spectra obtained using ESI extracted at the same retention time. SACI-CIMS give rise to higher signal intensities probably due to the low ion trap saturation.

Several (oligo-)peptides (between 70 and 115, depending on the analysed sample) matched to different animal venom proteins and toxins like conotoxins, phospholipases A2, metalloproteinases (86% of assignments have a -log(e) higher than 25). An overview of 36 proteins covered by the toxin-like peptides found is reported in Table 1; details of -log(e) and false discovery rates are reported in Table 2. Examples of mass spectra peptide characterization together with the peptide ion fragmentation pathways are shown in Figure 3a. All the MS/MS signal were assigned to the different N-terminal y,z (blue and purple colour) and c-terminal b,c (red and yellow colour) fragmentation series (see Figure 3b for fragmentation series details). In the defined SACI-CIMS conditions, doubly charged m/z ion of medium-high molecular weight peptide species are produced, allowing high identification accuracy, in line with what is already described in the literature that high identification statistical rates are achieved analysing peptide doubly charged species with medium high molecular weight. Different fragmentation anomalies with proton rearrangements have also been detected and considered in phase of data elaboration. Only mass spectra exhibiting a statistical -log(e) score higher than 10 and a false discovery rate lower than 0.05 were considered for the identification (see Figure 3c). False discovery rate and statistical score were estimated by means of reverse sequence approach.

Some of the toxin-like peptides found mapped on the same reference protein (UniprotKB: D2DGD8), are reported in Figure 4: these peptides were found in the five plasma samples and in the three faecal samples.

The types of toxic-like peptides found resemble known conotoxins, phospholipases A2, metalloproteinases, prothrombin activators, coagulation factors, usually present in animal venoms, which are known to have high specificity and affinity towards human ion channels, receptors, and transporters of the nervous system, like the nicotinic acetylcholine receptor. Cheng et al.21 reported the discovery of a superantigen-like motif in the S1 Spike protein, as well as two other neurotoxin-like motifs that have peptide similarities to neurotoxins from *Ophiophagus* (cobra) and Bungarus genera. They conclude that neurotoxin-like motifs are present in SARS-CoV-2 protein products, acting as neurotoxin-like peptides. We checked in the full set of peptides we got (here we report only 36 examples), and we identified, in plasma and faecal samples, toxin-like peptides mapping on kappa 1a-bungarotoxin, Kappa 1b-bungarotoxin from Malayan krait, kappa-2-bungarotoxin and alpha-bungarotoxin from many-banded krait (Uniprot Accession Numbers Q8AY56, Q8AY55, P15816, and P60615, respectively), which were reported by Cheng and colleagues. Furthermore, we looked at the amino acid changes currently reported in GISAID data22, analysed by CoV-GLUE-Viz (update 15/09/2021)23, and occurring in the Y_{668}QTQTNPRRAR motif identified by these authors as homologous to neurotoxin motifs of animal venom proteins. We observed the existence of amino acid variations which makes this motif even more similar to the neurotoxin motifs of animal venom proteins (like variations Q677S and T676A observed in sequences assigned to PANGO Lineage B.1.596). Experiments to assess neurotoxicity of these peptides and of spike protein on 3D neuronal/glial model (“neurospheres”) obtained from human induced Pluripotent Stem Derived Neural Stem Cells (iPS-NSCs) are currently ongoing.

What follows is our attempt to elaborate a potential relation between their presence and extra-pulmonary COVID-19 symptomatology.

**Conotoxins**

Conotoxins are neurotoxic peptides isolated from the venom of marine (genus *Conus*) cone snails. In their mature form, they consist of 10 to 30 amino acid residues, with often one or more disulphide bonds, which are used to classify them in structural classes (μ-conotoxins, α-conotoxins, and β-conotoxins are the major classes). The mechanism of action of conotoxins is not yet fully understood24. Studies have found that they are able to modulate the activity of several receptors, including ion channels, nicotinic acetylcholine receptors...
### Table 1. Overview of candidate proteins on which toxin-like peptides have been mapped.

Thirty-six candidate protein sequences on which the identified toxin-like peptides have been mapped are here reported, together with information retrieved from UniprotKB and NCBI Taxonomy databases. The table is split in three sections according to the phylum of the reported species: **Chordata** (green), **Echinodermata** (pink), **Mollusca** (azure).

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<td>D2DGD8</td>
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<td>A1X8B8</td>
<td>CALCONQU</td>
<td>reviewed</td>
<td>Putative alpha-conotoxin Qc alphaL-1</td>
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<td>QcaL-1</td>
<td>68</td>
<td>101313</td>
<td>Conus quercinus</td>
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<td>P58786</td>
<td>COW_CONRA</td>
<td>reviewed</td>
<td>Contrryphan-R</td>
<td>NA</td>
<td>- . Bromocontrryphan Cleaved into 2chains</td>
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<td>Mollusca - Conidae</td>
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<td>P58811</td>
<td>CA1A_CONTU</td>
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<td>Rho-conotoxin TIA</td>
<td>NA</td>
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<td>8765</td>
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(nAChRs) and acetylcholine-degrading enzymes (acetylcholinesterases), thus resulting in the alteration of acetylcholine levels and of cholinergic transmission\textsuperscript{25–27}. Regarding cholinesterases, a potential association between cholinesterase levels and severity of pneumonia in COVID-19 patients has been proposed\textsuperscript{28}.

The presence of conotoxin peptides might explain the occurrence of many symptoms (like hyposmia, hypogeusia and the signs typical of Guillain-Barre syndrome) observed in some COVID-19 patients. Their presence can alter normal functioning of ion channels, nicotinic acetylcholine receptors and of acetylcholine levels.

**Phospholipases A2**

Phospholipases A2 (PLA\textsubscript{2}, E.C. 3.1.1.4) hydrolyse phospholipids and lead to release of lysophosphatidic acid and arachidonic acid\textsuperscript{29}. Arachidonic acid is a major precursor of many pro-inflammatory mediators like leukotriene, thromboxane and prostaglandin; as a consequence, abnormal presence of active PLA\textsubscript{2} can induce severe inflammation\textsuperscript{30}. In animal venoms, PLA\textsubscript{2} act as neurotoxic proteins: they hydrolyse membrane phospholipids of the motor nerve terminal, and the plasma membrane of skeletal muscle, thus triggering a severe inflammatory degenerative response, which in turn leads to degeneration of the nerve terminal and skeletal muscle\textsuperscript{29}. The drug dexamethasone can inhibit prostaglandins synthesis and leukotriene formation\textsuperscript{11}. As dexamethasone is still the only therapeutic shown to be effective against the novel coronavirus in patients\textsuperscript{32} with severe symptoms, it can be that the positive effect of this drug on COVID-19 patients is also due to the reduction of the here identified PLA\textsubscript{2}-like peptides.

**Metalloproteinases**

The last example of identified toxin-like peptides is those recognised as metalloproteinases present in animal venoms, zinc-dependent enzymes of varying molecular weight having multi-domain organization. These toxic enzymes cause haemorrhage,
Figure 2. Examples of full scan mass spectra, obtained by analysing a COVID-19 positive urine sample and acquired fociating solvent ion cloud species containing a) low, b) high m/z species extracted in the 5.713–5.719 min chromatographic region and ESI full scan mass spectrum obtained analysing the same sample and extracting the signal at the same retention time extracting c) low and d) high m/z ratio.
Table 2. List of proteins and the related -log(e) and false discovery ratio (FDR) expressed as p value.

<table>
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<tr>
<th>Protein</th>
<th>ID</th>
<th>Database</th>
<th>-log(e)</th>
<th>FDR p value</th>
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<td>Conotoxin Pu6.1</td>
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<td>Uniprot</td>
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local myonecrosis, skin damage, and inflammatory reaction. It has been reported that symptomatic COVID-19 patients have significantly lower zinc levels in comparison to controls and that zinc deficient patients develop more complications. The presence of this specific group of toxin-like peptides, which capture zinc, can be one of the reasons for such significantly low zinc levels in symptomatic COVID-19 patients.
Similarity searches by TBLASTN with relaxed parameters at the National Center for Biotechnology Information (NCBI) website (see Methods) revealed (in addition to mRNA sequences from the animal species reported in Table 1) almost identical short stretches (up to 10 amino acids) of these peptides in potential coding regions of many bacterial and viral sequences, but no long potential coding frame entirely covering any of them was found. Consequently, at the time of writing we have not yet identified the “genetic source” of these peptides, which could be:

- The SARS-CoV-2 RNA genome with its protein reading set, as proposed by Brogna, who reported the identification in SARS-CoV-2 RNA of many regions encoding for oligopeptides (four–five amino acids long) identical to neurotoxin peptides typical of animal venoms.

Figure 3. Examples of mass spectra peptide characterization together with the peptide ion fragmentation pathways. Example of how MS/MS signal were assigned to the different N-terminal y,z (blue and purple colour in panel a) and c-terminal b,c (red and yellow colour) fragmentation series (detailed in panel b). Only mass spectra exhibiting a statistical -log(e) score higher than 10 and a false discovery rate lower than 0.05 were considered for the identification (reported in panel c). False discovery rate and statistical score were estimated by means of reverse sequence approach.
The SARS-CoV-2 genome directly read by bacteria, assuming that the SARS-CoV-2 genome, or parts thereof, is capable of replicating with a possible ‘bacteriophage-like’ mode of action, as previously described.

Genomes of bacteria, which, as a reaction to the presence of the virus, secrete these peptides. This could happen by using still not well known and debated mechanisms, like alternative reading due to rRNA sequence heterogeneity (as described in 37, 38), or the involvement of small bacterial ncRNA (sRNAs), known to be key players of gene regulation under conditions like stress response, quorum sensing, or virulence (in this context, in 1984 Coleman et al. reported the micF non-coding RNA as a functional bacterial sRNA).

A combination of the above e.g. the ‘toxin’ genetic code is present in the bacteria and expression may be triggered by SARS-CoV-2, acting like temperate bacteriophages, which are known to interact with bacteria so that they express (or not) certain genes, as described by Carey et al.

A detailed 3D structural similarity analysis between the toxin-like peptides found and reference proteins has not yet been conducted. Accordingly, at the time of writing, we can only speculate that these toxin-like peptides are involved in the clinical extra-pulmonary manifestations in symptomatic COVID-19 patients. According to our knowledge, these toxin-like peptides have never been searched in animals considered reservoirs of SARS-CoVs.

**Conclusions**

The presence of (oligo-)peptides almost identical to toxic components of venoms from animals has been observed. Data and results reported here suggest an association between COVID-19 disease and the release in the body of these, and raise a series of questions:

- Are these findings in line with what was proposed by Tizabi et al., i.e. a potential therapeutic role for nicotinic agonists, or positive allosteric modulators of nicotinic cholinergic receptors in COVID-19?
- If induced by SARS-CoV-2, can the production of toxin-like peptides be involved in the neurological disorders and injuries observed in hospitalized COVID-19 patients?
- If induced by SARS-CoV-2, can the production of toxin-like peptides influence complex diseases apparently triggered or enhanced by COVID-19, like e.g. Guillain-Barré Syndrome or Parkinson’s disease?
- Are toxin-like peptides associated with SARS-CoV-2 infection or to other viral infections or, more in general, is their presence related to sickness condition?
- Are our findings supporting the suggestion made by the iVAMP Consortium on the relationships between animal venom glands and microorganisms’ microenvironments?

We consider that the immediate sharing of these results can contribute to the untangling of the multifaceted set of clinical manifestations in symptomatic COVID-19 patients, and to the further understanding of the mechanisms involved.

**Data availability**

**Underlying data**

Uniprot: Kunitz-type serine protease inhibitor homolog beta-bungarotoxin B1 chain [Bungarus candidus (Malayan krait)]. Accession number Q8AY46, [https://identifiers.org/uniprot:Q8AY46](https://identifiers.org/uniprot:Q8AY46)
Uniprot: Basic phospholipase A2 BFPA, svPLA2, EC 3.1.1.4 (Antimicrobial phospholipase A2) (Phosphatidylcholine 2-acylhydrolase) [Bungarus fasciatus (Banded krait) (Pseudoboa fasciata)]. Accession number A6MEY4, https://identifiers.org/Uniprot:A6MEY4

Uniprot: Phospholipase A2 MALT0035C, svPLA2, EC 3.1.1.4 [Micrurus alitostris (Uruguayan coral snake) (Elaps alitostris)]. Accession number F5CPF1, https://identifiers.org/Uniprot:F5CPF1

Uniprot: Zinc metalloproteinase-disintegrin-like NaMP, EC 3.4.24.- (Snake venom metalloproteinase, SVMP) [Naja atra (Chinese cobra)]. Accession number A8QL59, https://identifiers.org/Uniprot:A8QL59

Uniprot: Acidic phospholipase A2 D, svPLA2, EC 3.1.1.4 (APLA) (Phosphatidylcholine 2-acylhydrolase) [Naja sputatrix (Malayan spitting cobra) (Naja naja sputatrix)]. Accession number Q9F900, https://identifiers.org/Uniprot:A9F900

Uniprot: Venom prothrombin activator omicarin-C non-catalytic subunit, vPA (Venom coagulation factor Va-like protein) [Cleaved into: Omericarin-C non-catalytic subunit heavy chain; Omericarin-C non-catalytic subunit light chain] [Oxyuranus microlepidotus (Inland taipan) (Dienema microlepidota)]. Accession number A58L90, https://identifiers.org/Uniprot:A58L90

Uniprot: Venom prothrombin activator oscurarin-C non-catalytic subunit, vPA (Venom coagulation factor Va-like protein) [Cleaved into: Osciurarin-C non-catalytic subunit heavy chain; Osciurarin-C non-catalytic subunit light chain] [Oxyuranus scutellatus (Coastal taipan)]. Accession number Q58L91, https://identifiers.org/Uniprot:Q58L91

Uniprot: Short neurotoxin 4, SNTX4 (Alpha-neurotoxin 4) [Pseudonaja textilis (Eastern brown snake)]. Accession number Q9W7J9, https://identifiers.org/Uniprot:Q9W7J9


Uniprot: Coagulation factor V [Cleaved into: Coagulation factor V heavy chain; Coagulation factor V light chain] [Pseudonaja textilis (Eastern brown snake)]. Accession number Q593B6, https://identifiers.org/Uniprot:Q593B6

Uniprot: Venom prothrombin activator pseutarin-C non-catalytic subunit, PCNS, vPA (Venom coagulation factor Va-like protein) [Cleaved into: Pseutarin-C non-catalytic subunit heavy chain; Pseutarin-C non-catalytic subunit light chain] [Pseutarin-C non-catalytic subunit light chain] [Pseudonaja textilis (Eastern brown snake)]. Accession number Q7SZN0, https://identifiers.org/Uniprot:Q7SZN0

Uniprot: Cysteine-rich venom protein ENH1, CRVP (Cysteine-rich secretory protein ENH1, CRISP-ENH1) [Pseudoferania polypleis (Macleay's water snake) (Enhydris polylepis)]. Accession number Q2XXQ3, https://identifiers.org/Uniprot:Q2XXQ3

Uniprot: Bradykinin-potentiating and C-type natriuretic peptides (Brain BPP-CNP, bBPP-CNP) (Evasin-CNP) [Cleaved into 12 chains] [Bothrops jararaca (Jaraaraca)]. Accession number Q9PW56, https://identifiers.org/Uniprot:Q9PW56

Uniprot: Snake venom metalloprotease inhibitor 02D01 (02E11) (10F07) (Svmi-Eoc7) [Cleaved into 15 chains] [Echis ocellatus (Ocellated saw-scaled viper)]. Accession identifier A8YPR6, https://identifiers.org/Uniprot:A8YPR6

Uniprot: Zinc metalloproteinase-disintegrin [Cleaved into: Snake venom metalloproteinase brevilsin L4, SVMP (Snake venom metalloproteinase hx1-1, EC 3.4.24.-); Disintegrin brevicudin-1a; Disintegrin brevicudin-1b (Disintegrin adinbitor) (Disintegrin halystatin)] [Gloydius brevicaudus (Korean slam osa snake) (Agkistrodon haly haly)]. Accession number Q698K8, https://identifiers.org/Uniprot:Q698K8


Uniprot: Alpha-elapitoxin-Oh2b, Alpha-EPXT-Oh2b (Alpha-neurotoxin) (LNTX3) (Long neurotoxin OH-6A/OH-6B) (OH-3) [Ophiophagus hannah (King cobra) (Naja hannah)]. Accession number P82662, https://identifiers.org/Uniprot:P82662


Uniprot: Bradykinin-potentiating and C-type natriuretic peptides (BPP-CNP) [Cleaved into six chains] [Protopothebrops flavoviridis (Habu) (Trimeresurus flavoviridis)]. Accession number P0C7P5, https://identifiers.org/Uniprot:P0C7P5


Uniprot: Conotoxin C9.6 [Californiconus californicus (California cone) (Conus californicus)]. Accession number D6C4M3, https://identifiers.org/Uniprot:D6C4M3

Uniprot: Kunitz-type serine protease inhibitor conotoxin Cal9.1a [Californiconus californicus (California cone) (Conus
Californiconus californicus]. Accession number D2Y488, https://identifiers.org/Uniprot:D2Y488

Uniprot: Conotoxin Cl14.9 [Californiconus californicus (California cone) (Conus californicus)]. Accession number D6C4I8, https://identifiers.org/Uniprot:D6C4I8

Uniprot: Alpha-conotoxin CIB (Cl1.2) [Conus catus (Cat cone)]. Accession number P0DPT2, https://identifiers.org/Uniprot:P0DPT2

Uniprot: Conotoxin Fla16d (Conotoxin Fla16.1) [Cleaved into: Conotoxin fla16a; Conotoxin fla16b; Conotoxin fla16c] [Conus flavus (Yellow Pacific cone)]. Accession number V5V893, https://identifiers.org/Uniprot:V5V893

Uniprot: Sigma-conotoxin GVIIIA [Conus geographus (Geography cone) (Nubecula geographus)]. Accession number P58924, https://identifiers.org/Uniprot:P58924

Uniprot: Conotoxin Mr15.2 (Mr094) [Conus marmoreus (Marble cone)]. Accession number P0DM19, https://identifiers.org/Uniprot:P0DM19

Uniprot: Conotoxin m3g (Mr3.6) [Conus marmoreus (Marble cone)]. Accession number P0C1N5, https://identifiers.org/Uniprot:P0C1N5

Uniprot: Conotoxin Pu6.1 [Conus pulicarius (Flea-bitten cone)]. Accession number D2DGD8, https://identifiers.org/Uniprot:D2DGD8

Uniprot: Alpha-conotoxin-like Pu1.5 [Conus pulicarius (Flea-bitten cone)]. Accession number P0C8U9, https://identifiers.org/Uniprot:P0C8U9

Uniprot: Putative alpha-conotoxin Qc alphaL-1, QcαL-1 [Conus quercinus (Oak cone)]. Accession number A1X8B8, https://identifiers.org/Uniprot:A1X8B8

Uniprot: Contryphan-R (Bromocontryphan) [Cleaved into: [Des-Gly1]-contryphan-R] [Conus radiatus (Rayed cone)]. Accession number P58786, https://identifiers.org/Uniprot:P58786


Uniprot: Conotoxin 10 [Conus virgo (Virgin cone)]. Accession number Q5K0C5, https://identifiers.org/Uniprot:Q5K0C5

Uniprot: Conotoxin Vi15a (Vi15.1) [Conus virgo (Virgin cone)]. Accession number B3FIA5, https://identifiers.org/Uniprot:B3FIA5


This project contains the following underlying data:
- Data file 1: Toxins.fasta
- Data file 2: Toxins.mgf

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY4.0)

Consent
The human biological samples used in the experimentation were collected and used with the expressed free and informed written consent of the person from whom the material was taken, according to current legislation.

Acknowledgements
The authors thank Martina Larini and Simone Madama for paper revision.

Declarations
The scientific output expressed does not imply a policy position of the European Commission. Neither the European Commission nor any person acting on behalf of the Commission is responsible for the use that might be made of this publication.

References


15. Prasatty V, Radhar M, Iystyontso E: Natural Peptides in Drug Discovery


Open Peer Review

Current Peer Review Status: ✔️ ✔️

Version 2

Reviewer Report 29 October 2021

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Moshe Arditi
Department of Pediatrics, Division of Pediatric Infectious Diseases and Immunology, Cedars-Sinai Medical Center, Los Angeles, CA, USA

I approve the paper as it is now – the paper is excellent now.

Competing Interests: No competing interests were disclosed.

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 18 October 2021

https://doi.org/10.5256/f1000research.78337.r96936

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Paolo Grumati
Telethon Institute of Genetics and Medicine (TIGEM), Pozzuoli, Italy

Author addressed the requested points.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: biology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.
Moshe Arditi

Department of Pediatrics, Division of Pediatric Infectious Diseases and Immunology, Cedars-Sinai Medical Center, Los Angeles, CA, USA

The investigators report the identification of toxin-like peptides in COVID-19 patients samples (Plasma, urine and stool samples) by using Liquid Chromatography Surface-Activated Chemical Ionization-Cloud Ion Mobility Mass Spectrometry. The investigators used a study cohort (for plasma) of 15 COVID 19 patients from different cities of Italy and from 5 control uninfected individuals. They collected urine samples from 2 COVID19 patients and 2 controls, and stool samples from 3 COVID19 patients and 3 controls. They report that toxin-like peptides, almost identical to toxic components such as conotoxins, phospholipases, phosphodiesterases etc. were identified from COVID19 patients, but not in any control samples. They report an overview of 36 proteins covered by the toxin-like peptides they have found in plasma of COVID1- patients. These toxin-like peptides they discovered are very much like various neurotoxins, such as alpha Conotoxins, alpha Cobratoxins or similar to Bungarotoxins, all known to be neurotoxins.

These are very important observations, the authors are asking the question is the COVID-19 infection is somehow inducing these toxin-like peptides in the host and if so, if these neurotoxin-like peptides maybe playing a functional role of the neurologic findings that are frequently associated with COVID-19 infection. One very important and potentially critical paper that must be mentioned in the discussion and that authors have missed was recently published by Mary Hongying Cheng et al. (2020) where the investigators discovered a Superantigen-like motif in the S1 Spike protein, as well as two other neurotoxins that have peptide similarities to alpha cobra toxin and alpha bungarotoxin, alpha cobra toxin etc. Given this PNAS paper, it is now clear that the SARS-CoV2 virus contains neurotoxin-like peptides already. It would be incredibly interesting to see if the neurotoxin like peptides described and discovered in this PNAS paper are present in the toxin-like peptides described in this specific study. It will make this paper and its discussion much more impactful. At a minimum the PNAS paper should be discussed and cited.

References

Is the work clearly and accurately presented and does it cite the current literature?
Partly
Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Partly

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Immunology, Infectious Diseases, Innate Immune Responses

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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**Author Response 05 Oct 2021**

Mauro Petrillo, European Commission, Joint Research Centre (JRC), Ispra, Italy

Dear Dr Arditi,

thanks a lot for the valuable comments and suggestions that you have provided in the report.
We will address all of them in order to provide a fully revised version of the manuscript.
Best regards,

Mauro Petrillo, on behalf of the authors.

**Competing Interests:** No competing interests were disclosed.

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**Author Response 09 Oct 2021**

Mauro Petrillo, European Commission, Joint Research Centre (JRC), Ispra, Italy

Dear Dr Arditi,

Thanks a lot for your valuable comments and suggestions that you have provided in the report. As anticipated, we have addressed all your points, and provided a new version of the manuscript:

- One very important and potentially critical paper that must be mentioned in the discussion and that authors have missed was recently published by Mary Hongying Cheng et al.(2020)
where the investigators discovered a Superantigen-like motif in the S1 Spike protein, as well as two other neurotoxins that have peptide similarities to alpha cobratoxin and alpha bungarotoxin, alpha cobra toxin etc. Given this PNAS paper, it is now clear that the SARS-CoV2 virus contains neurotoxin-like peptides already. It would be incredibly interesting to see if the neurotoxin like peptides described and discovered in this PNAS paper are present in the toxin-like peptides described in this specific study. It will make this paper and its discussion much more impactful. At a minimum the PNAS paper should be discussed and cited.

**Response:** We thank the Dr. Arditi for the comment and we fully agree with his suggestion to cite the Mary Hongying Cheng et al. (2020) PNAS paper. With respect to this point, we would like to highlight that the peptides reported in the paper as examples are a subset of more than 100 peptides identified by MS. We checked in the full set of peptides, and we can confirm that we identified, in plasma and faecal samples, toxin-like peptides mapping on kappa 1a-bungarotoxin, Kappa 1b-bungarotoxin from Malayan krait, kappa-2-bungarotoxin and alpha-bungarotoxin from many-banded krait (Uniprot Accession Numbers Q8AY56, Q8AY55, P15816, and P60615, respectively). To address this point, we added the following paragraph in the section “Results and discussion”: Cheng et al. [REF] reported the discovery of a superantigen-like motif in the S1 Spike protein, as well as two other neurotoxin-like motifs that have peptide similarities to neurotoxins from Ophiophagus (cobra) and Bungarus genera. They conclude that neurotoxin-like motifs are present in SARS-CoV-2 protein products, acting as neurotoxin-like peptides. We checked in the full set of peptides we got (here we report only 36 examples), and we identified, in plasma and faecal samples, toxin-like peptides mapping on kappa 1a-bungarotoxin, Kappa 1b-bungarotoxin from Malayan krait, kappa-2-bungarotoxin and alpha-bungarotoxin from many-banded krait (Uniprot Accession Numbers Q8AY56, Q8AY55, P15816, and P60615, respectively), which were reported by Cheng and colleagues. Furthermore, we looked at the amino acid changes currently reported in GISAID data [REF], analysed by CoV-GLUE-Viz (update 15/09/2021) [REF], and occurring in the Y674QTQTNSPRRAR685 motif identified by these authors as homologous to neurotoxin motifs of animal venom proteins. We observed the existence of amino acid variations which makes this motif even more similar to the neurotoxin motifs of animal venom proteins (like variations Q677S and T676A observed in sequences assigned to PANGO Lineage B.1.596). Experiments to assess neurotoxicity of these peptides and of spike protein on 3D neuronal/glial model (“neurospheres”) obtained from human induced Pluripotent Stem Derived Neural Stem Cells (iPS-NSCs) are currently ongoing. What follows is our attempt to elaborate a potential relation between their presence and extra-pulmonary COVID-19 symptomatology.

We hope that the quality of the manuscript, thanks to your comments, has been improved and you consider it suitable for indexing.

Best regards,
Mauro Petrillo, on behalf of the authors.

**Competing Interests:** No competing interests were disclosed.
In the present manuscript the authors proposed an interesting consequence of COVID 19 infection. The idea is that COVID infection induces the production of toxins that are responsible for the specific clinical manifestations. Despite the fact that the origin of the toxins is not clear. The authors identified via mass spectrometry peptides that match the sequences of toxin components of venoms from animals. These observations are surprising and provocative. However, there are several points that the authors should consider:

1. COVID-19 outbreak is a pandemic therefore the number of affected people is extremely high. The number of samples analysed should be much higher. Authors should consider to have at least three different groups. Non-infected control, infected without clinical symptoms, affected with severe symptoms. For each group at least 10 samples for each analysis (blood, urine) should be analysed in order to have a more reliable statistic analysis. Moreover, does the amount of toxins correlate with the severity of the phenotype?

2. Authors should provide some biological data that the toxins they identified are responsible for the clinical phenotype. They should perform some in vitro experiments infecting cells (CALU are the most used) with COVID or treating them with toxins. The outcome should be similar.

3. It is difficult to access the original data. It would be interesting to see the peptide sequences identified from the mass spectrometry. In Fig.4 the authors reported an example but it is unlike that a secreted toxin contains the leader region pro-peptide.

4. The text needs some editing. Citation of other research papers should be conform to the standard.

Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Partly

Are sufficient details of methods and analysis provided to allow replication by others?
Partly
If applicable, is the statistical analysis and its interpretation appropriate?
No

Are all the source data underlying the results available to ensure full reproducibility?
Partly

Are the conclusions drawn adequately supported by the results?
No

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** biology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

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**Author Response 12 Aug 2021**

**Mauro Petrillo**, European Commission, Joint Research Centre (JRC), Ispra, Italy

Dear Dr Grumati,

Thanks a lot for your valuable comments and suggestions that you have provided in the report.

We will address all of them, and wait for those of other reviewers, in order to provide a fully revised version of the manuscript.

Best regards,
Mauro Petrillo, on behalf of the authors.

**Competing Interests:** No competing interests were disclosed.

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**Author Response 09 Oct 2021**

**Mauro Petrillo**, European Commission, Joint Research Centre (JRC), Ispra, Italy

Dear Dr Grumati,

Thanks a lot for your valuable comments and suggestions that you have provided in the report. As anticipated, we have addressed all your points, and provided a new version of the manuscript:

- **COVID-19 outbreak is a pandemic therefore the number of affected people is extremely high.** The number of samples analysed should be much higher. Authors should consider to have at least three different groups. **Non-infected control, infected without clinical symptoms, affected with severe symptoms.** For each group at least 10 samples for each analysis (blood, urine) should be analysed in order to have a more reliable statistic.
analysis. Moreover, does the amount of toxins correlate with the severity of the phenotype?

- **Response:** We thank the Dr. Grumati for the comment. We increased the number of analysed plasma samples to address this point: in addition to the 5 control cases and 15 hospitalised cases (already mentioned in the section “Methods: Cohort”), we have analysed 5 plasma samples of infected individuals with mild or no symptoms, together with 5 additional controls. Thus, for plasma, there are now 30 cases (20 positive to SARS-CoV-2 tests plus 10 controls). In this new added group, we observe the presence of toxin-like peptides, apparently in lower amounts (in terms of -loge) with respect to the samples from hospitalised individuals. However, as we have no additional information about the grade of severity of the hospitalised subjects, we prefer to not infer any correlation between the amount of identified peptides and the severity of the phenotype. To better clarify this point, we have added the following sentence in the section “Methods:Cohort”: *Apart from positivity to SARS-CoV-2, no additional information (i.e. age, sex, blood serotype, severity of the disease, time of the collection, fatality, etc.) was provided.*

- Authors should provide some biological data that the toxins they identified are responsible for the clinical phenotype. They should perform some in vitro experiments infecting cells (CALU are the most used) with COVID or treating them with toxins. The outcome should be similar.

- **Response:** We thank the Dr. Grumati for the comment. Experiments to assess neurotoxicity of these peptides on 3D neuronal/glial model (“neurospheres”) obtained from human induced Pluripotent Stem Derived Neural Stem Cells (iPS-NSCs) are ongoing and will be part of an additional publication. We added a sentence in the section “Results and discussion” to address this point: *Experiments to assess neurotoxicity of these peptides on 3D neuronal/glial model (“neurospheres”) obtained from human induced Pluripotent Stem Derived Neural Stem Cells (iPS-NSCs) are ongoing.*

- **It is difficult to access the original data. It would be interesting to see the peptide sequences identified from the mass spectrometry. In Fig.4 the authors reported an example but it is unlike that a secreted toxin contains the leader region pro-peptide.**

- **Response:** We thank the Dr. Grumati for these comments. Regarding the data, we followed the journal policy: all data produced and here presented are freely and publicly available on the Zenodo platform (http://www.doi.org/10.5281/zenodo.4903154, as reported in the section “Data availability” and in Reference 16, including MS files in MGF format and sequence files in FastA format). Regarding the alignment in Fig.4, the shown peptides correspond to the longest peptides we observed (LC-SACI-CIMS is particularly able in detect long peptides), aligned with respect to the precursor of the protein. We did not make any specific selection for secreted proteins; thus, precursors are expected to be present in our samples. To address this point, we modified the legend of Fig.4 accordingly.

- **The text needs some editing. Citation of other research papers should be conform to the standard.**

- **Response:** Thanks a lot for spotting these inconsistencies. We revised citations according to the journal specifications. We hope that the quality of the manuscript, thanks to your comments, has been improved and you consider it suitable for publication.

Best regards,
Comments on this article

Author Response 22 Jul 2021

Mauro Petrillo, European Commission, Joint Research Centre (JRC), Ispra, Italy

Dear Dr. de Bernardis,

Thanks a lot for your valuable comment.

You are perfectly right: it can't be excluded that the findings aren't specific to COVID and that might be common to other conditions.

And in fact, one of the questions of the Conclusions section of the manuscript is "Are toxin-like peptides associated with SARS-CoV-2 infection or to other viral infections or, more in general, is their presence related to sickness condition?"

The aim of our manuscript is to immediately share these observations with the scientific community as they are (together with a series of other observations which we have recently reported in https://doi.org/10.12688/f1000research.52540.3) quite unexpected, at least to us.

Thanks again for your time and interest. I am happy to further discuss, also privately.

Best regards,
Mauro Petrillo

Competing Interests: None

Reader Comment 19 Jul 2021

Ernesto de Bernardis, ASP SR, Italy

I don't understand why the Authors' hypotheses about the origin of these peptides don't include the host response during severe inflammation or ARDS, or during a pharmacological therapy similar to those given to COVID patients.

The paper compares peptides from COVID patients with those from healthy controls, but doesn't
include controls with other diseases, e.g. inflammatory diseases, other viral diseases, or people treated with the same medications that were administered to their sample of COVID patients.

So, I guess it can't be excluded that the findings aren't specific to COVID and that might be common to other conditions.

**Competing Interests:** No competing interests were disclosed.