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SHORT COMMUNICATIONS

SARS-CoV-2 spike mRNA vaccine sequences circulate in blood up to 28 days after COVID-19 vaccination

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In Denmark, vaccination against Severe Acute Respiratory Syndrome Corona Virus 2 (SARS-CoV-2) has been with the Pfizer-BioNTech (BTN162b2) or the Moderna (mRNA-1273) mRNA vaccines. Patients with chronic hepatitis C virus (HCV) infection followed in our clinic received mRNA vaccinations according to the Danish roll-out vaccination plan. To monitor HCV infection, RNA was extracted from patient plasma and RNA sequencing was performed on the Illumina platform. In 10 of 108 HCV patient samples, full-length or traces of SARS-CoV-2 spike mRNA vaccine sequences were found in blood up to 28 days after COVID-19 vaccination. Detection of mRNA vaccine sequences in blood after vaccination adds important knowledge regarding this technology and should lead to further research into the design of lipid-nanoparticles and the half-life of these and mRNA vaccines in humans.

Key words: Hepatitis C virus; SARS-CoV-2; vaccine; blood; mRNA.

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# INTRODUCTION

With the emergency approval by the FDA of two mRNA vaccines in December 2020, and subsequently very large-scale vaccine production and mass immunization programs, a breakthrough in protective measures against the global pandemic with severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) was achieved. Both the Pfizer-BioNTech (BTN162b2) and the Moderna (mRNA-1273) vaccines code for production of the fulllength SARS-CoV-2 spike protein. To ensure stability, these vaccines are composed of codonoptimized modified spike mRNA, have two stabilizing proline substitutions, and the mRNA is encapsulated in lipid nanoparticles (LNPs) [1,2]. The

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modified nucleotide sequences allow perfect identification of the vaccine sequences as being different from any coronavirus sequence. Upon intramuscular injection, the vaccine mRNA is taken up by muscle and immune cells, and transported to the regional lymph nodes and concentrated in the spleen [3]. The vaccines consist of nonreplicating mRNA and are expected to naturally decompose both within the cytosol after translation and at the injection site. The half-life of mRNA translation is estimated to be short, from hours to a day and translation is described to span up to 10 days [4-6]. The Infectious Diseases Society of America (IDSA) informs that the vaccine mRNA is degraded quickly by normal intracellular processes and states that there is no evidence for long-term detection of mRNA vaccines in vaccinated individuals by RNA-seq [1,7].

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In Denmark, the predominant vaccines against Corona Virus Disease 2019 (COVID-19) have been the two mRNA-based vaccines from Pfizer-BioNTech and Moderna. As of July 28, 2022, 80.2% of the population has received two doses and 81.6% at least one dose [8]. At Copenhagen University Hospital, Amager-Hvidovre, Denmark, patients with chronic hepatitis C virus (HCV) infection are routinely followed at Department of Infectious Diseases, while measurement of HCV viral load and genotyping by whole RNA-Seq genome sequencing of their HCV RNA, directly from plasma samples, are performed at Department of Clinical Microbiology [9]. In this paper, we describe the unexpected finding of SARS-CoV-2 vaccine mRNA sequences in plasma from 10 HCV patient samples up to 28 days after COVID-19 vaccination. These patients had recently received SARS-CoV-2 mRNA vaccinations according to the Danish rollout vaccination plan.

## **METHODS**

We analyzed five consecutive sequencing runs, from May 2021 to the end of June 2021, with 108 HCV patients, five negative controls and five HCV-positive controls consisting of HCV grown in cell culture [10]. Samples were from patients that were HCV positive and came for HCV treatment evaluation with no relation to their vaccination characteristics. The direct RNA sequencing (RNA-seq) procedure has been described previously [11]. Briefly, RNA was extracted with the ZR viral RNA kit (Zymo Research, Irvine, CA, USA) and depleted for human rRNA with the NEBNext rRNA depletion kit (New England BioLabs, Ipswich. MA, USA). Stranded RNA-seq libraries were prepared with the NEBNext Ultra II directional RNA library prep kit (New England BioLabs). Sequencing was performed with  $2 \times 150$ -bp paired-end reads on a NextSeq instrument (Illumina, San Diego, CA, USA).

All software was used with default parameters unless specified. Raw reads were trimmed for low-quality bases with fastp v 0.20.1 with minimum phred quality of 20 and minimum length of 50 [12]. Human depletion was done by mapping the trimmed reads to the human genome hg38 (GenBank accession no. GCA\_000001405.27) with Bowtie2 v 2.3.4.1 adding the parameters -k 1 -X 2000 [13]. Paired reads unmapped to human genome sequences were de novo assembled with VICUNA v 1.3 using minimum contig links of 2 and minimum identify of 90 [14]. As part of our pipeline, contigs above 1000 bp are aligned against NCBI's nt database using BLASTn v2.8.1+ with a minimum value threshold of 1 e-22 [15]. Contigs that were found to align to coronavirus sequences were subsequently aligned against two assemblies of SARS-CoV2-spikeencoding mRNA vaccines, BNT-162b2 and mRNA-1273 using BLASTn for confirmation [16]. Paired reads were also mapped against the two vaccine assemblies using BWA-mem v 0.7.16 and SAMtools v 1.2 inside the NASP pipeline v 1.1.2 [17–20]. Stats of number of reads mapping and coverage were calculated only for the SARS-CoV-2 spike gene coding region avoiding the 5' and 3' UTR as these regions share high percentage of similarity with human gene regions. Coverage was calculated with BED-Tools v 2.30.0 and plotted in R v 3.6.1 with ggplot2 [21–23]. SNP calls were generated with HaplotypeCaller from GATK v 4.2.0.0 and filtered by heterozygous genotype, mapping quality of <30 and symmetric odds ratio of >3 [24]. Consensus sequences were made in GATK and bases covered with less than 10 reads were masked with BEDTools.

The best (highest coverage) spike sequence for the Pfizer-BioNTech and similarly the two best spike sequences for the Moderna vaccine have been uploaded to NCBI (GenBank accession numbers OK120840). OK120842). Gaps in the sequences are represented by Ns. Approval by an institutional review board was not required, as this study was performed as an operational activity related to patient management.

#### RESULTS

De novo assembly of human-depleted RNA-seq reads from two patient samples produced contigs of >1000 nt with closest homology to bat coronavirus and 67% homology to the SARS-CoV-2 reference genome (NC\_045512.2). Translation of the contig nucleotide sequences to amino acid sequences revealed 100% identity to parts of the spike protein of SARS-CoV-2. From the literature we found the sequences of the two commercial SARS-CoV-2 mRNA vaccines which were used as references to map reads from all the samples [16,25]. This led to the identification of an additional eight samples with reads that matched the mRNA vaccine sequences. Both mRNA vaccine sequences have been modified and are only ~70% identical to the spike reference genome on a nucleotide level, making them distinct from circulating infectious SARS-CoV-2 sequences. Thus, of the 108 patient samples, 10 samples (9.3%) had partial or up to full sequences of the vaccine mRNA sequence (Fig. 1), identified from one to 28 days postvaccination. There was ~100% identity between the detected mRNA nucleotide sequences found in plasma and the specific mRNA vaccine given. The 10 samples had a median of 5.5 million raw read pairs available (see Table S1). Breadth and depth of coverage of the vaccine mRNA sequences ranged from completeness and >20 000, respectively, to short fragments with a depth of coverage of 100 (Fig. 1). None of the negative or the HCV-positive controls had SARS-CoV-2 matching reads.

#### DISCUSSION

We surprisingly found fragments of COVID-19 vaccine mRNA up to 28 days postvaccination in

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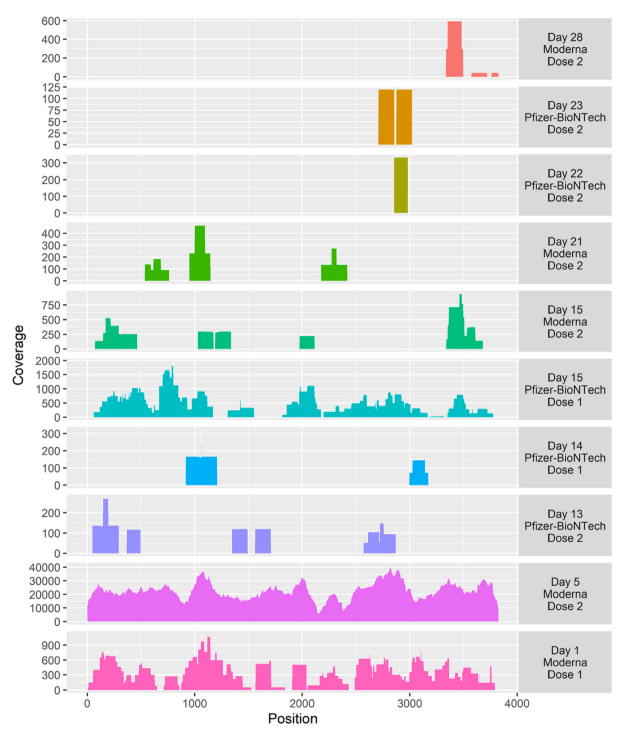


Fig. 1. Mapping of trimmed and filtered reads to the coding regions of the spike protein of the Moderna mRNA vaccine or Pfizer BioNTech vaccine. Each colored plot is a single sample. The right-hand column shows the day postvaccination, the vaccine given and found in blood and if sample was drawn after 1. vaccination or revaccination. The two bottom samples are from the same patient after first and second vaccination with the Moderna vaccine.

blood from chronic HCV patients vaccinated with mRNA vaccines from both Pfizer-BioNTech and Moderna.

Analysis of mRNA vaccine function has focused on the immune response and on protection of vaccinated individuals from SARS-CoV-2 induced

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severe COVID-19 [26,27]. The LNPs have been reported to be rapidly cleared by immune cells and mRNA is degraded by exonucleases in tissue and blood [28-30]. A recent study did not detect the vaccine mRNA by quantitative PCR in human milk after 4-48 h postdose 1 or 2 with BNT162b2 or mRNA-1273 [31].

RNA-seq is widely used in biological and medical research. We routinely use direct total RNA-seq to obtain full length HCV RNA sequences and from these infer their genotype. Our pipeline includes taxonomic analysis of contigs of more than 1000 nt which led to the discovery of coronavirus sequences that when first blasted had closest identity to a bat coronavirus.

We expect that vaccine mRNA detected in plasma is contained within LNPs and that the LNPs in plasma have been slowly released from the injection site either directly to the blood or through the lymph system. Without the LNPs protecting the mRNA, the mRNA would rapidly degrade. This allows prolonged spike protein production giving an advantage for a continuous immune response in some persons. Current studies on half-life of mRNA vaccines could have underestimated the half-life of the LNPs, primarily using results from half-life of studies of mRNA in the cytosol of human cells.

In samples where we observed only fragments of vaccine mRNA, this could indicate that the concentration of LNPs in plasma is low. This is in accordance with our findings that HCV viral load should be higher than 10 000 IU/mL to obtain a fulllength HCV RNA genome (size ~9600 nt) as genome coverage correlates with viral load. The number of SARS-CoV-2 vaccine mRNA molecules is therefore likely lower than ~4000 IU/mL, explaining the partial read coverage randomly spread over the coronavirus vaccine sequence.

To our knowledge, our study is the first to detect Pfizer-BioNTech and Moderna COVID-19 mRNA vaccine sequences in blood after vaccination, and therefore provides new knowledge regarding the timeframe in which the mRNA can be detected. This study examined a cohort of HCV-positive patients with presumed functional immune systems, as most of the patients can be cured, with negative HCV RNA 12 weeks after end of treatment with direct-acting antivirals. A future prospective study to establish the half-life of mRNA vaccines in vaccine recipients could be performed using mRNA vaccine-specific PCRs. These findings are interesting and should lead to further research into the design of LNPs and the half-life of LNPs and mRNA vaccines, but it should be emphasized that our data does not in any way change the conclusion that both mRNA vaccines are safe and effective.

#### **CONFLICT OF INTEREST**

Authors have no conflicts of interest to declare.

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## SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Table S1.** Sequencing and mapping report of the 10 patient samples. Mapping and coverage statistics are calculated for only the SARS-CoV-2 spike gene coding region of the vaccine.

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