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Inhibition of SARS-CoV-2 spike protein entry using biologically modified polyacrylonitrile nanofibers: *in vitro* study towards specific antiviral masks[†]

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With the increase of the contagiousness rates of Coronavirus disease (COVID-19), new strategies are needed to halt virus spread. Blocking virus entry by capturing its spike (S) protein is one of the effective approaches that could help in eliminating or reducing transmission rate of viruses. Herein, we aim to develop a nanofiber-based filter for protective face masks, composed of polyacrylonitrile (PAN) nanofibers (NFs)-loaded with Angiotensin Converting Enzyme-2 (ACE-2) for capturing the spike protein of severe acute respiratory syndrome Coronavirus-2 (SARS-CoV-2) and blocking its entry. Docking simulations were performed to evaluate interactions of PAN with target proteins of both SARS-CoV-2 and Human Adenovirus type 5 (ADV-5) which was used as an in vitro model of human respiratory viruses. Scanning electron microscopy (SEM) and Fourier transformed infrared (FT-IR) spectroscopy was employed to investigate the surface morphology and to analyze the functional groups of the NFs, respectively. The mechanical properties of the electrospun NFs were investigated, according to which the tensile strengths of PAN and modified PAN NFs were 4.9 \pm 1.2 GPa and 4.5 GPa. Additionally, elongations at break were 25 \pm 2.5% to 24 \pm 1.48% for PAN and modified PAN NFs. The tensile strength test showed good mechanical characteristics of the NFs. The ACE-2-loaded NFs were shown to be safe, with promising antiviral activity towards ADV-5. Meanwhile, a binding affinity study between the spike protein and ACE-2 was performed and the dissociation constant ($K_{\rm D}$) was found to be 1.1 nM. Accordingly, the developed antiviral filters have a potential role to stand as a base for combating various human respiratory viruses

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1 Introduction

Viral outbreaks are considered as one of the primary causes of morbidity and mortality worldwide.⁴ Viral infections are associated with several complications, such as cardiovascular, respiratory, and thromboembolic diseases, which might lead to secondary bacterial infections alongside the viral infection.² The emergence of Coronavirus disease-19 (COVID-19) caused by severe acute respiratory syndrome Coronavirus-2 (SARS-CoV-2)

in Wuhan, China, led to the call of a viral emergency worldwide.^{3,4} It was reported that SARS-CoV-2 virus was discovered to be similar to severe acute respiratory syndrome Coronavirus (SARS-CoV) that caused severe acute respiratory syndrome (SARS).⁴

During the past two decades, the three pandemics caused by Coronavirus family (SARS-CoV-1, MERS-CoV and SARS-CoV-2) have affected about 29 million people around the world, according to the World Health Organization (WHO). In September 2020, it has been reported that SARS-CoV-2 virus had already taken more than 917 417 lives globally.⁵

Coronavirus infection is mediated first through binding of spike (S) glycoprotein to angiotensin-converting enzyme-2 (ACE-2) receptors on the host cells.⁶ This receptor is considered as an active site for S protein binding and hence its entry into host cells.⁶ Since ACE-2 is expressed in various human organs such as lungs, small intestine, and liver, infection can be extended to multiple human organs. It has been reported that, SARS-CoV-2 demonstrated a stronger affinity to ACE-2.⁷ This can distinguish it from SARS-CoV-1, and play a crucial role in both viral entry and transmission.⁸

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Paper

Adenoviruses (ADV-2 and ADV-5) are double-stranded DNA (dsDNA) non enveloped viruses of 90–100 nm size, with genomes of about 36 kb long.⁹ They are associated with variety of human diseases, such as respiratory infections, ocular and gastrointestinal disorders. They mainly affect children and young adults.¹⁰ No specific treatment regimen is used for adenovirus infection except only two antiviral drugs, cidofovir and ribavirin have been approved by the Food and Drug Administration (FDA).¹¹

For these reasons, development of antiviral compounds and advanced protective tools are essential to reduce morbidity and mortality among patients especially those with immunocompromised diseases, as well as to reduce economic losses.

Molecular docking and computational simulations served as scientific advances in drug discovery of newer antimicrobial agents against various microbial infections.¹² Two stages are utilized for docking computations, sampling of various ligand configurations and binding pockets in protein and, calculating the binding affinity between molecular systems. Auto Dock Vina is quite possibly the most utilized docking packages in protein– ligand interactions.^{13,14} Phytochemical compounds have been given a great attention, as many of them are used as antimicrobial agents,¹⁵ where they cause minimal adverse effects to patients.

Various nano-carriers types have been employed, to a great extent, in combating infectious diseases.¹⁵ After more than 70 years of intensive research and investigation, the field of antiviral materials, both therapeutics and anti-infective, remain relatively underdeveloped. While antiviral polymeric biomaterials showed to be a promising avenue for addressing these challenges.

Currently available face masks are protective tools that can play crucial role in avoiding viral infections, due to presence of filters that can reduce inhalation of air-borne particles. However, no antiviral masks are available to provide specific protection against respiratory viruses, like SARS-CoV-2.

Nanofibers (NFs) possess unique physical and chemical characteristics, they provide a large surface area and a good potential to incorporate active chemistry,16 they have a range of advantages in terms of diameter, permeability, and, most significantly, large surface area, therefore, they have recently attracted a lot of attentions.17 The small pore size of nanofibrous mats have a significant effect on air permeability and surface porosity. Electrospinning has become one of the most common methods for fabricating nanofibrous mats, owing to its simplicity and efficiency. Poly acrylonitrile (PAN) is a synthetic thermo-plastic polymer that has good mechanical and chemical stability,18 moreover it has a relatively low cost which allows incorporation in many fields including: protective clothing, filtration media, and biomedical textiles.19 PAN NFs were previously used to fabricate protective face masks incorporating an anti-bacterial agent.¹⁷ Previously, PAN NFs were loaded with silver nanoparticles to filter air from pollutants and microbes.20 Whereas, coating PAN NFs with bovine serum albumin (BSA) protein were utilized for wound dressing applications.18

The aim of the current study is to fabricate PAN NFs loaded with anti-spike protein of corona virus, to block viral entry, thereby, specifically protects highly susceptible people, *e.g.* doctors and medical staff, against SARS-CoV-2 infection. Accordingly, *in silico* study was performed followed by *in vitro* study evaluating antiviral activity against ADV-5 as a model of human respiratory virus and against SARS-CoV-2 using binding affinity assay.

2 Materials and methods

2.1 Materials

Poly acrylonitrile (PAN, average molecular weight = 150 000 g mol⁻¹), (3-aminopropyl)triethoxysilane (APTES, \geq 98%), EDC *N*-(3-dimethylaminopropyl)-*N'*-ethylcarbodiimide hydrochloride, and *N*-hydroxy succinimide (NHS) were all purchased from Sigma-Aldrich, USA. *N*,*N*-Dimethyl formamide (DMF, \geq 99%, Laboratory reagent grade) was purchased from Fisher Chemical, USA.

Blank PAN NFs, amine-modified PAN NFs, and esterchemistry activated PAN NFs loaded with ACE-2 materials were subjected for cytotoxicity test. Vero cell line was obtained from ATCC (American tissue culture collection). Dulbecco's Modified Eagle Medium (DMEM), sodium pyruvate (100 mM), and L-glutamine (200 mM), penicillin (100 IU mL⁻¹), streptomycin (10 mg mL⁻¹), and trypsin/versine (0.05%) were all purchased from Lonza, Belgium. Fetal Bovine Serum (FBS) was purchased from Gibco, South America. MTT (3-[4,5dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide) dye was purchased from Serva Electrophoresis GmbH, Germany. Phosphate Buffer Saline tablets (PBS, pH 7.4) was purchased from Loba Chemie, India. Each tablet was dissolved in 200 mL distilled water. Dimethyl sulfoxide (DMSO) was purchased from Analysis Company, Egypt.

Blank PAN NFs, and amine-modified PAN NFs materials were subjected to anti-viral assay test. Adenovirus (VR-5) was obtained from ATCC (American tissue culture collection, USA). QIAamp viral RNA extraction kit was purchased from Qiagen, Valencia, USA. qRT-PCR kit (Power SYBR[™] Green PCR Master Mix) was purchased from Thermo-Scientific, USA. cDNA synthesis kit was purchased from Roche Diagnostics, Germany.

Recombinant human coronavirus SARS-CoV-2 spike glycoprotein S1 (active), (Cat. No. ab273068), recombinant anti-SARS-CoV-2 spike glycoprotein antibody H6 (chimeric), (Cat. No. ab272854), recombinant human ACE-2 protein, (Cat. No. ab151852), Goat anti-Mouse IgG H&L (Alexa Flour 647), (Cat. No. ab150115), Anti-SARS-CoV-2 spike glycoprotein antibody Coronavirus, (Cat. No. ab272504), and Goat anti-rabbit IgG Fc (Alexa Flour 647), (Cat. No. ab150091), were all purchased from Abcam, Cambridge, United Kingdom.

2.2 Methods

2.2.1 Molecular docking calculation of PAN polymer against both spike protein in SARS-CoV-2 virus and Caspid protein Hexon in ADV-5. The initial structure of the Hexon protein of adenovirus and spike protein of SARS-CoV-2 virus was fetched from the protein data bank PDB ID 6CGV and 6M17, respectively. While the structure of the PAN polymer was constructed utilizing Avogadro software and exposed to minimization calculation utilizing the quantum chemistry program ORCA. Auto-dock tools were utilized to prepare ligand and protein pdbqt files. Auto-Dock Vina was used for docking, while Protein–Ligand Interaction Profiler (PLIP) analysis was used to foresee various interactions inside the anticipated binding pockets.¹³

2.2.2 Fabrication of PAN NFs. PAN solution (8%, w/v) was completely dissolved in *N*,*N*-dimethylformamide (DMF) at 60 °C in a closed vial.¹⁸ The solution was stirred for overnight to obtained homogenous solution, which was then electrospun into NFs by electrospinner (NANON-01A, MECC, Japan). PAN solution was placed into 5 mL syringe (18 G) for electrospinning process by tubeless spinneret system. Electrospinning was performed in varying operating regimes using flow rates between (1 and 1.5 mL h⁻¹), applied voltages between (21 and 27 kV), and tip-to-target distance (13–15 cm). The NFs were collected on a static plate collector. Resultant PAN NFs were dried at 70 °C for 24 hour (h) for further testing.¹⁷ The optimized morphology of NFs was obtained at the following optimized spinning conditions: feed rate 1.5 mL h⁻¹, applied voltage 21 kV and tip-to-target distance 15 cm.

2.2.3 Chemical modification of PAN NFs. As illustrated in (Fig. 1a), PAN NFs (8%, w/v) was immersed in APTES solution (10%, w/v) that was prepared by a proportion of

(ethanol : distilled water as 9:1).^{21,22} PAN-APTES NFs were washed several times using hot alcoholic-distilled water to remove the unreacted APTES.

2.2.4 Characterization of PAN and modified PAN NFs. Surfaces of electrospun NFs were investigated by SEM model (Quattro S, Thermo-Scientific, USA). The chemical functional groups were examined by FT-IR model (Bruker Vertex 70, Germany), with IR fingerprints registered between 4000–400 cm⁻¹ using transmittance modes.

The mechanical strength of NFs was measured by uniaxial tensile test machine (Z050, ZwickRoell AG, Ulm, Germany). Tensile strength of NFs was measured by standard uniaxial tensile test. The NFs were cut into rectangular pieces with dimensions (40×10 mm) and incorporated into paper frames to protect them before the definite tensile testing. To obtain the stress–strain curves, specimens were elongated at a speed of 10 mm min⁻¹ with an initial length of 20 mm and 50 N cell load. Measurements were taken for at least six specimens of each composition to obtain average values and standard deviations.

2.2.5 Cell culture and cytotoxicity assay. Vero cells were maintained into DMEM supplemented with 10% PBS, penicillin/streptomycin, 1% of sodium pyruvate, and 1% of L-glutamine at standard conditions (37 °C and 5% CO₂, 85–95% humidity). The cytotoxic effect was investigated using MTT colorimetric assay after 48 h of cell exposure to the prepared NFs on the Vero cell line.²³ Cells were examined for any morphological changes using inverted phase contrast light



Fig. 1 Schematic showing methodology of fabricating the biologically modified PAN NFs. (a) Mechanism of modifying PAN NFs by APTES before proteins attachment, (b) antiviral assay of PAN and modified PAN NFs using adsorption mechanism, and (c) antiviral assay of PAN and modified PAN NFs using virucidal/replication mechanism.

microscope (Axio Observer 5, Zeiss, Germany) after treatment with our materials.

A modified method was performed utilizing MTT (3-[4,5dimethylthiazol-2-yl]-2,5-diphenyltetrazolium bromide) dye, that is based on the reduction of the dye by mitochondrial dehydrogenases of metabolically active cells into insoluble formazan crystals according to previously published protocol.²⁴ Cell viability (%) was calculated by dividing average of 3 replicas of each material on average of the control and multiplied by 100.

2.2.6 Antiviral assay. The antiviral activity of our materials was investigated against Human adenovirus type 5 (ADV-5) propagated into normal epithelial cells derived from monkey kidney cells (Vero cells).

Viral model of human adenovirus type 5 using Real-Time PCR assay: investigating antiviral activity: This was based on the following steps.

(A) Infectivity assay. Human ADV-5 (ATCC VR-5) was propagated into Vero cells and examined every day under an inverted microscope until 80–90% of CPE (cytopathic effect) was achieved. The virus culture was then frozen and thawed for three times before being tested for viral load using a quantitative realtime PCR assay.⁹ The standard curve was generated using 10fold serial dilutions of standard VR-5 control (10–10⁶ copies per mL).⁹

(B) Titration of adenovirus-5 DNA in cell culture. Adenoviral titration was carried out by seeding 2×10^4 cells per well into a 96-well tissue culture plate. The plate was incubated for 24 h at normal conditions of 37 °C and 5% CO₂, after which the cells were infected with a two-fold serial dilution of an adenovirus stock with established viral load and incubated for 48 h until 80–90% cell lysis was observed in the undiluted viral stock, then quantitation of viral load was detected using real-time PCR assay.²⁵

(C) Quantitation of adenovirus-5 DNA in cell culture using Realtime PCR assay

(C.1) Nucleic acid extraction. QIAamp viral RNA extraction kit was used to extract adenovirus DNA from infected and uninfected cultures (Qiagen, Valencia, USA). The extraction was carried out in compliance with the manufacturer's guidelines. DNA extracts were put on ice and used for PCR right away, or they were kept at -20 °C before analysis (if needed). A Nano-Drop 2000 spectrophotometer (Thermo- Scientific/US, Canada) was used to determine the volume of viral DNA, and 100 ng of DNA template was used in the PCR assays.⁹

(C.2) Real-time PCR assay. Adenovirus DNA viral load was detected and quantified using a real-time detection system (Applied Bio systems 7500 Fast Real-time PCR). The ADV viruses were amplified using the common forward (ADV-F: ATG ACT TTT GAG GTG GAT CCC ATG GA) and the common reverse (ADV-R: GCC GAG AAG GGC GTG CGC AGG TA) primers targeting Hexon gene at nucleotide. The reaction mixture contained 12.5 μ L of Maxima SYBR Green/ROX qPCR master mix (2×) (Fermentas, California, USA), 10 μ M of each adeno primers, and 100 ng of extracted DNA in a final volume of 25 μ L distilled water. The ADV-5 cycle protocol consisted of initial

denaturation at 94.0 °C for 10 min followed by 94.0 °C for 30 s; 50.0 °C for 30 s; and at 72.0 °C for 30 s for 40 cycles then final extension at 72 °C for 7 min, finally melting curve; 95.0 °C for 15 s; 60 °C for 1 min, and 95 °C for 15 s. The PCR products were subjected to a temperature rise of up to 85 °C during the post-amplification melting temperature ($T_{\rm m}$) study, resulting in a progressive decrease in fluorescence. It was possible to differentiate between non-specific PCR products from Adeno PCR related products in this manner.⁹

(D) Antiviral activity of the anti-spike protein-loaded modified PAN NFs against adenovirus (ADV-5). The safest materials were subjected for the three antiviral mechanisms as follows:

(D.1) Adsorption mechanism. As presented in (Fig. 1b), a sixwell plate was seeded with 5 \times 10⁵ cells per mL and incubated at 37 °C and 5% CO₂ as per normal procedure. The cells were treated with our materials after 24 h. The cells were infected with IC₅₀ of adenovirus stock after 24 h and incubated for another 24 h under standard conditions.

(D.2) Viral replication mechanism. A six-well plate was seeded with 5 \times 10⁵ cells per milliliter and incubated at room temperature. After 24 h, the cells were infected with IC₅₀ of adenovirus stock and incubated for another 24 h at normal conditions.

(D.3) Virucidal mechanism. This approach was used to investigate whether the tested materials that could neutralize the viral effect and then prevent it from entering the cells. As demonstrated in (Fig. 1c), a six-well plate was seeded with 5 × 10⁵ cells per milliliter and incubated at room temperature. After incubation with IC₅₀ of virus at 4 °C for 1 h, the cells were treated with our materials after 24 h, then incubated for another 24 h at standard condition.

In each antiviral assay, two controls were run: an untreated well for cell control and an untreated infected well for viral control. All plates were then subjected for quantitative real-time PCR assay to measure viral load.²⁵

2.2.7 Attachment of anti-spike protein of coronavirus to modified NFs. The specific binding between the S protein of corona virus and its antibody was evaluated and visualized using a system composed of a monoclonal antibody and a secondary antibody conjugated with Alexa Flour 647 (Fig. 4a). This is to prove the binding between the S protein and ACE-2, thereby blocking the virus entry into the cells.

Ester chemistry was used to attach the ACE-2 protein to the electrospun NFs. The amine group was activated by a mixture of 1-ethyl-3-(30-dimethylaminopropyl)carbodiimide–*N*-hydroxysuccinimide (EDC–NHS) in 1 : 1 ratio.²⁶ The NFs were rinsed after 30 min with distilled water to remove any excess of EDC-NHS. ACE-2 protein (100 nM) was immobilized onto the modified PAN NFs by incubation for 20 min. The NFs were then rinsed with PBS to remove excess unbound proteins.

2.2.8 Binding affinity test between the spike protein and its antibody. The binding affinity between the S protein and ACE-2 was investigated to confirm that the binding is strong enough to prevent the S protein dissociation. The test was done on glass slides that were carefully cleaned and modified with APTES,

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activated with EDC-NHS to create amine groups to allow the attachment of the antibody. ACE-2 protein (10 nM) was attached to the modified glass slides by incubation for 20 min then the slide was rinsed with PBS to remove excess unbound proteins. Different concentrations of the S protein of SARS-CoV-2 (0.01, 0.1, 1, 10 and 100 nM) were incubated for 20 min and rinsed with PBS. The monoclonal antibody of S protein (10 nM) was added followed by the secondary labelled antibody (10 nM). In order to prove the specific binding, two negative controls were performed. The first control consisted of immobilized ACE-2 (100 nM) followed by the secondary labelled antibody (10 nM) that were added to the amine-activated surface, while the second one consisted of binding of the 100 nM S protein to 100 nM ACE-2, followed by 10 nM of the labelled antibody on the activated surface. The slides were placed into a 24-well microplate and fluorescence signal was measured by a microplate reader (Clariostar Plus, BMG lab tech, Germany). First, the signal was measured as a baseline, before adding the secondary labelled antibody. Second, the labelled antibody was added and the fluorescence signal was detected. Third, the proteins were rinsed with PBS for 10 min to confirm their specific binding.

3 Results and discussion

3.1 Molecular docking calculation of PAN polymer against both spike protein in SARS-CoV-2 virus and Caspid protein Hexon in ADV-5

The Docking calculations usually use different scoring functions to study the likelihood of ligand to bind to the protein target, the more negative values indicate stronger binding of the ligand.²⁷ The results demonstrate that the binding affinity of PAN against both of Hexon protein (PDB ID: 6CGV) and spike protein (PDB ID: 6M17) is about -6.3 and -5 kcal mol⁻¹, respectively. The protein–ligand interaction profiler (PLIP) analysis was employed to determine different types of interaction within the predicted binding pockets (Fig. 2). The docking results suggest that stabilization of the interaction between PAN and Hexon protein in the predicted binding site is more likely due to due to hydrophobic interactions and hydrogen bonds with the nearby residues. Where that PAN forms hydrogen bonds with amino acid residues LYS-44B, PHE-45B, ASN-47B and ASN-639C at the predicted binding pocket in Hexon protein. Where that PHE 45B and ASN-639C are shown to form shorter (stronger) hydrogen bonds with PAN molecule with average distances 2.14 Å. On the other hands, PAN forms hydrophobic interactions with residues LEU-24, LEU-28, ALA-32, LEU-41 and LYS-44 in chain B. In contrary, Hexon protein, PAN possess only hydrophobic interactions through interacting with ALA-363A in the predicted site in SARS-CoV-2 protein (Table 1).

3.2 NFs imaging results

SEM was intensively used to investigate the accepted nanofibrous mats, as shown in (Fig. 3), SEM images show that the surface morphology of NFs demonstrated accepted morphological structure; random, uniform, filamentous and continuous shape-structure of PAN NFs with no formed beads. Meanwhile, images of activated PAN/APTES/EDC-NHS/ACE-2 protein revealed major changes after incubation of protein, where loaded protein resulted in significant swollen NFs structure, compared to blank NFs. These changes might be a result of changing the functional groups after modification by

Table 1 Residues interacting with PAN in the predicted Hexon and SARS-CoV-2 binding pockets^a

	PAN@Hexon		PAN@SARS-COV2	
Index	Resn.	Distances [Å]	Resn.	Distances [Å]
Hydrop	hobic interacti	ons		
1	24B LEU	3.84	ALA 363 A	3.74
2	28B LEU	3.95	—	—
3	32B ALA	3.93	—	—
4	41B LEU	3.93	_	_
5	44B LYS	3.75	—	—
Hydrog	en bonds			
1	44B LYS	2.65	_	_
2	45B PHE	2.17	_	_
3	47B ASN	2.12	_	_
4	639C ASN	2.26	—	—
<i>^a</i> —: no	t detected.			



Fig. 2 Docking of PAN into (a) Hexon protein of ADV-5, and (b) spike protein of SARS-CoV-2.



Fig. 3 SEM micrographs of (a) blank PAN NFs, (b) PAN NFs/APTES, and (c) PAN NFs/APTES/EDC-NHS/ACE-2/S protein (with original magnification 4000× (up) and 8000× (down) at 7 kV).

APTES and incubation of proteins as confirmed by FT-IR spectra (Fig. 4). Moreover, NFs-loaded protein, preserved their entire interior and exterior morphological structure.

3.3 NFs chemical modification

As illustrated in (Fig. 1a), PAN was modified by APTES through salinization process to modify nitrile group to amine group.²² The chemical structure of PAN NFs and modified PAN-APTES NFs were analyzed by FT-IR. As shown in (Fig. 4a), FT-IR spectra of PAN revealed appearance of amide (CO–NH) and amine (N–H) bands at ν 1657 and 1565 cm⁻¹ respectively, upon the interaction between PAN and APTES. The main characteristic peaks of Si–O–Si that appear from ν 1020 cm⁻¹ to

1200 cm⁻¹, were detected in both APTES and modified PAN, which is verifying the modification of PAN by APTES. Results show that nitrile groups of PAN were successfully modified to amide and amine groups.²⁸ The modified groups were then activated by EDC-NHS ester chemistry for protein attachment.

The results presented in (Table 2) display the effect of modification of PAN NFs with APTES through measuring specific mechanical parameters *e.g.* Young's modulus, tensile strength, and elastic modulus. Obtained data revealed that modification of PAN NFs with APTES decreases their mechanical stability insignificantly. This indicates that APTES modification affects PAN NFs surfaces. These results are consistent



Fig. 4 FT-IR spectra of (a) modified PAN NFs/APTES, and (b) loaded proteins onto amine-activated PAN NFs; PAN NFs/APTES, PAN NFs/APTES/ EDC-NHS/ACE-2, and PAN NFs/APTES/EDC-NHS/ACE-2/S protein.

with Wang *et al.*, that demonstrated minor decrease in modified NFs' strength.²²

3.4 Standard uniaxial tensile test

3.5 Cytotoxicity assay

The cytotoxic effect of tested material was performed on Vero cells was determined using MTT colorimetric assay after 48 h of cell exposure. Morphological examination (Fig. S1†) confirmed MTT results, where no changes in cell morphology were revealed upon treatment of Vero cells with PAN NFs, modified PAN/APTES NFs and activated PAN NFs loaded with ACE-2 compared to untreated cells (cell control), as shown in (Fig. 5a), where the cell viability was ranged between 80 : 100% of all tested NFs materials. The results indicate that our tested materials are safe on cells and hence can be tested for their antiviral activity.

3.6 Antiviral assay against ADV-5 using real-time PCR assay

Viral titration was performed using quantitative PCR assay, showed that IC₅₀ (viral concentration can kill 50% of cells) determined at 10⁵ copies per mL (Fig. 5b), that was used in the antiviral assay. Results showed that PAN and modified PAN/ APTES NFs exerted antiviral activity via the two tested antiviral mechanisms, adsorption and virucidal/replication. They did not only prevent viral entry into cells, but also viral infectivity when they are incubated with the virus for 1 h. The virus was unable to replicate inside cells, as evidenced by undetected levels of viral copies using quantitative real-time PCR assay (Fig. 5b). This might be due to the affinity between the nanofiber and the viral receptors, which might lead to blocking viral path and preventing viral entry into the cells.²⁹ The second reason might be due to the negative charges on the surface of PAN,30 which when bound to viral proteins, could form a complex with the virus and hence neutralize its ability to infect the cells.31

Several approaches have been proposed to explain antiviral activity employed by anionic polymers depending on their complexation with virus proteins.³¹ This was shown to occur *via* formation of salt linkages or ion pairs between oppositely charged groups on PAN polymer and virus proteins, respectively, as previously explained in properties of anti-HIV-1 compounds.³² These results were supported by *in silico* study, where binding of PAN to Hexon protein is taking place through hydrogen bonds and hydrophobic interactions (Table 1).

3.7 Loading of anti-spike protein onto modified NFs

As shown in (Fig. 4b), FT-IR spectra of modified PAN revealed the appearance of N-H and C-H bending bands at v 1621 and 1564 cm⁻¹ respectively. Interestingly, both bands were shifted after incubation of proteins. The FT-IR spectra of (PAN NFs/ APTES/EDC-NHS/ACE-2) revealed appearance of a peak at ν 1729 cm⁻¹ after the incubation of ACE-2 protein, corresponding to -C=O group.^{33,34} The carbonyl band disappeared after incubation of S protein, owing to the interaction between S protein and its antibody, while the linkage between -COOH group of ACE-2, S proteins and -NH2 group of modified PAN NFs was confirmed by the appearance of the amide (CO-NH) stretching peak at ν 1631 cm⁻¹, that was detected in both (PAN NFs/APTES/EDC-NHS/ACE-2) and (PAN NFs/APTES/EDC-NHS/ ACE-2/S protein). This is besides, the characteristic peaks of N-H and C-H bending bands and their appearance at characteristic peaks at ν 1564 cm⁻¹ and 1456 cm⁻¹ respectively. The results verify the binding of ACE-2 and S proteins to modified PAN NFs.33,34

Regarding the stability of ACE-2-loaded PAN NFs, the samples are shown to be stable until up to 8 months at room temperature, stored under dry nitrogen. SEM images presented in (Fig. S2†) show that no visual cracks nor surface degradation. This might be due to the hydrophobic nature of PAN polymer and the storing conditions of the modified and loaded nanofibers. Moreover, the chemical structure of the protein-loaded PAN NFs, as prepared and after 8 months, doesn't indicate any structure change, as shown in the FT-IR spectra (Fig. S3†). It



Fig. 5 Schematic representation of (a) toxicity assay results showing the safety of the PAN, modified, and ACE2-loaded PAN using MTT assay, (b) antiviral assay against ADV-5 via both adsorption and virucidal/replication mechanisms.

3.8

Affinity test results

is expected that the samples can be stable up to a longer period if kept in refrigerator (4 $^{\circ}$ C).

(Fig. 6a) is a schematic of the system used to study the binding interaction between ACE-2 and S protein, whereas (Fig. 6b) demonstrates a rapid increase and saturation of the fluores-

cence signal occurring after adding the labelled antibody. The

samples were left for 20 min to ensure the binding of the

secondary antibody. A decrease in the fluorescence signal is

observed after the rinsing step. Despite the decrease in the

signal, it is hard to observe the dissociation owing to the fact

that it is a static type of measurement. However, after rinsing,

the signals show a variation linearly dependent on the

concentration of S protein, which is a strong proof that the

binding is specific. Specificity is further proved by results of the

two controls revealing a negligible signal demonstrating that

there is no nonspecific binding.³⁵ As illustrated in (Fig. 6c), the

affinity constant between S protein of SARS-CoV-2 and its antibody equals $0.9 \times 10^9 \text{ M}^{-1}$ and $K_{\rm D}$ equals 1.1 nM. These results

indicate a high affinity and strong specific interactions between

the S protein and ACE-2. The current findings are consistent

with the reported results of Saponaro et al., where they

demonstrated $K_{\rm D}$ values of several binding affinity studies

between S protein of SARS-CoV-2 and ACE-2.36

	PAN NFs		
Type of nanofiber	PAN	Modified PAN	
Elongation at break (%)	25 ± 2.5	24 ± 1.48	
Max. displacement (mm)	4.2	4.0	
Young's modulus (GPa)	260 ± 5.52	258 ± 1.25	
Tensile strength (GPa)	4.9 ± 1.2	4.5	
Elastic modulus (GPa)	4.5 ± 0.85	4.1 ± 0.25	

4 Conclusions

Developing broad-spectrum antiviral polymers holds enormous potential for two main reasons; to treat current viral diseases and to control transmission of viral pathogens, especially respiratory pathogens. ACE-2 protein loading onto NFs is a promising approach to develop a potential stable antiviral mask. It demonstrates a remarkable advancement to capture the virus on the nanofibrous surface and block its entry to the host cells. The loaded protein was specifically chosen considering viral pathologic features to enhance antiviral activity as evidenced by binding affinity. Molecular docking as well as physical, chemical, and biological investigations showed a promising antiviral activity against ADV-5 *via* adsorption and virucidal, and replication mechanisms. Testing the modified nanofibers over a period of 8 months showed significant stability. However, many queries are not yet answered such as



Fig. 6 (a) The system used to prove the binding between the S protein to ACE-2 on the NFs, (b) binding of the different concentrations of the S protein to the ACE2-modified slides, showing different signal level after washing, with very little change in the signal for the high concentrations, and (c) a competitive binding curve at different concentrations of the S protein revealing a dissociation constant (K_D) of 1.1 nM.

the nature of this interaction, whether the virus can be reactivated or not, and the time it takes the virus to degrade. They can also serve as microbicides or antiviral drug-delivery vehicles. Further studies are still needed to explore the exact antiviral mechanism exerted by anionic polymers.

A prototype was designed for the proposed anti-viral mask, in which ACE-2-loaded modified PAN NFs were placed on a commercial filter. For future prospective, PAN NFs should be investigated for their antiviral activity against other viruses as well. Besides, they can be loaded with many other proteins for suppressing infectious diseases. The proposed mask is inprogress to be tested as a potential commercial, specific antiviral mask.

It is worth mentioning that the fabrication of 1 mask is not expensive, taking in account the cost of the chemicals, antibody, labor, and overhead; the total cost of 1 filter with dimensions $2.5 \times 2 \text{ cm}^2$ would be 0.79 \$ and the price of the mask that will be used to implant the filter is 0.5 \$, so the total cost will be 1.28 \$. The filer can be just used and changed in the same mask.

This cost was calculated for pilot scale production; therefore, it is expected to significantly decrease by about 60% in case of mass production, the price of this mask including the modified PAN filter is expected to be: 0.768 \$. This low value makes the one-time use of PAN filter economically effective.

Author contributions

MHE performed the experimental work, conducted the graphical analysis for all the results, and initiated writing the original manuscript, under supervision of AK, EAK and SAL. HN prepared some NFs and helped in the chemical modification interpretations. FA performed the theoretical study. HE performed two antiviral experiments under the supervision and guidance of SAL. MT and RA supervised the embedding of NFs in the commercial masks. MHE and AK wrote the manuscript. All authors revised the manuscript and agreed with all its contents.

Conflicts of interest

The authors declare no competing financial interest.

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