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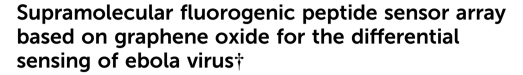


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We report on a supramolecular sensor array using fluorogenic peptide probes and graphene oxide that can target glycoproteins on a viral caspid, facilitating the differentiation of ebola virus from marburg virus and receptor-extensive vesicular stomatitis virus using principal component analysis.

Ebola virus (EBOV) is an acutely infectious pathogen towards humans. Infection with EBOV can result in a high death rate of 50–90%. The prevalence of EBOV in Africa during 2013–2016 resulted in the infection of more than twenty thousand people, and led to the death of more than ten thousand. The mechanism of EBOV invasion and the viral persistence are still not well characterized. For the effective prevention and control of the spread of EBOV, an effective diagnosis of the viral infection is a high priority in EBOV research. However, currently used biochemical methods for EBOV detection are complex and time-consuming. As a consequence, the development of new chemical tools for the effective sensing of EBOV remains challenging.

Glycoproteins (GPs) exist on the capsid of a number of viral species. They are pivotal targets for the development of virus inhibitors and vaccines.⁷ The development of anti-GP antibodies is a promising strategy to prevent EBOV infection through neutralization of GP-mediated membrane fusion with

host cells.⁸ A recent report demonstrated that peptide fragments derived from the Fab-region of an anti-GP antibody had a GP-binding affinity comparable to that of the full-length antibody.⁹ This enables the development of peptide-based biochemical assays for the detection of EBOV.

Over the past decade, rapid development of fluorescent chemosensors for a variety of applications ranging from ion detection in environmental water to imaging of disease-related biomarkers in cells and in vivo has been achieved. 10,11 While the majority of these studies use the "lock-and-key" concept, in which a ligand molecule is developed to specifically bind to its target receptor, Anslyn has developed the concept of "array sensing" in order to remove the need for specific ligand/ receptor recognition.¹² This approach, employs an array of different receptor molecules, which may only have a moderate affinity and selectivity towards the analytes. Resulting in a "fingerprint" response, when combined with an appropriate analytical method, and giving rise to characteristic patterns for each analyte. This differential sensing strategy has been successfully used for detection of ions, small molecules, proteins, cells as well as pathogens. 13-15

Guided by this elegant strategy, we developed a supramolecular sensor array consisting of graphene oxide (GO)-based fluorogenic peptide probes for the differential sensing of EBOV. GO is a prototypical two-dimensional (2D) material, which has found applications in a number of biomedical applications.¹⁶ With its strong fluorescence quenching capacity, GO-based fluorogenic sensors have been extensively developed.¹⁷ On the basis of our research on 2D material-based fluorogenic peptide probes,18 we have synthesized three 5-carboxytetramethylrhodamine (5-TAMRA)-labelled peptide probes for self-assembly with GO. Three peptide fragments, RS5, QY7 and ED17 (Fig. 1), derived from a patented anti-GP antibody, 19 were used. Marburg virus (MARV), which has similar capsid protein components to EBOV, 20 and receptor-extensive vesicular stomatitis virus (VSV), which is a common control virus for RNA-based viral investigations, were also used. For safety reasons, pseudoviruses that only contain the capsid proteins, skeletal genes

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Supramolecular peptide sensor array based on graphene oxide (GO)

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Fig. 1 (a) Structure of the 5-TAMRA-labeled peptide probes **T-RS5**, **T-QY7** and **T-ED17**. (b) Schematic illustration of the supramolecular fluorogenic peptide sensor array based on graphene oxide (GO) for differential sensing of three viruses including ebola virus (EBOV), marburg virus (MARV) and receptor-extensive vesicular stomatitis virus (VSV) using the principal component analysis (PCA).

(T-ED17)

and reporter genes without the ability to self-replicate were used for the tests. ²¹

The supramolecular peptide probes were constructed by self-assembly between GO and the three fluorescent peptide ligands (T-RS5, T-QY7 and T-ED17) (Fig. 1a). Each peptide ligand was mixed with GO, followed by gently shaking for 15 min, producing the composites (T-RS5/GO, T-QY7/GO and T-ED17/GO). To characterize the association of the peptides with GO, transmission electron microscopy (TEM) was used. As shown in Fig. 2a, small aggregates were observed in the representative TEM images of the peptides alone; similar aggregates were seen on the surface of GO flakes after the self-assembly (Fig. 2b). This is in accordance with our previous observations that fluorescent peptide probes form aggregated species on the surface of 2D materials.²²

In the UV-vis spectra, we observed a red-shifted absorption peak of TAMRA (from ca.550 nm to 558-562 nm) after assembly with GO (Fig. S1, ESI†), suggesting a strengthened conjugated system probably due to π - π stacking between the dye and GO.

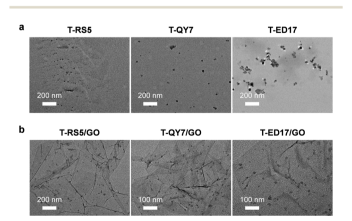


Fig. 2 Transmission electron microscopic images of (a) T-RS5, T-QY7 and T-ED17 (10 μ M), and (b) T-RS5/GO, T-QY7/GO and T-ED17/GO (peptide/GO = 10 μ M/40 μ g mL⁻¹).

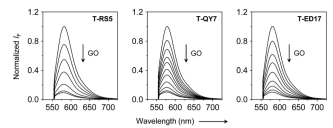


Fig. 3 Fluorescence titration of T-RS5, T-QY7 and T-ED17 (0.5 μ M) with increasing GO (from top to bottom curve: 0–12 μ g mL⁻¹ for T-RS5, 0–56 μ g mL⁻¹ for T-QY7 and 0-38 μ g mL⁻¹ for T-ED17, respectively). All measurements were carried out in PBS (0.01 M, pH 7.4, containing 1% DMSO, v/v) with an excitation of 525 nm.

Raman spectroscopy was used to characterize the self-assembly. The characteristic D band and G band centered at 1350 cm⁻¹ and 1580 cm⁻¹ were observed for GO, respectively. An increased intensity ratio of D band to G band ($I_{\rm D}/I_{\rm G}$), an increase in the disorder of carbon sp²-hybridization in the conjugated plane of GO.²4 After self-assembly with the peptide probes, we observed an increased $I_{\rm D}/I_{\rm G}$ for T-RS5/GO (1.38), T-QY7/GO (1.27) and T-ED17/GO (1.22) with respect to GO alone (0.98) (Fig. S2, ESI†). This suggests that the adsorption of peptide molecules onto the surface of GO increases the disorder in the carbon sp²-hybridization of the composite system.

Next, fluorescence spectroscopy was used to evaluate the fluorescence change of the peptide probes after assembly with GO. We observed a concentration-dependent fluorescence quenching of all the peptide probes with increasing GO (Fig. 3), which agrees with the quenching ability of the material for adsorbed photoluminescent species, ²⁵ and suggests assembly between the peptide probes and GO. Subsequently, we added the three pseudoviruses to the assembled GO-based fluorogenic peptide probes. We observed a concentration-dependent fluorescence increase with increasing concentrations of all the viruses (Fig. 4). This suggests the removal of the peptide probes from the GO surface probably by forming peptide/virus complexes results in the fluorescence recovery of TAMRA.

Similar phenomena were observed in our previous studies.²⁶ However, these results clearly suggest the inability of the probes to selectively differentiate EBOV from the other two viruses by just referring to the fluorescence recovery rate. We ascribe this observation to the week specificity of the short peptides for GP in the presence of the structurally complex viruses. This is a common problem with peptide-based sensing systems for complex biological samples.²⁷ However, we also observed that while all the GO-based peptide probes showed sensitivity to the three viruses, the extent to which the fluorescence of each peptide probe recovered varied in terms of the different viral species added. This prompted us to construct a supramolecular sensor array for differential sensing of EBOV.

As shown in Fig. 5a, we used the Principal Components Analysis (PCA) for differential sensing of the viruses. PCA is a commonly used method that relies on multivariate statistical dimensional orthogonal linear transformation for the extraction of feature signals by reducing the dimensionality of the ChemComm

1.2 1.2 T-RS5/GO T-RS5/GO T-RS5/GO 0.8 0.8 0.8 EBOV VSV MARV 0.4 0.4 0.4 0.0 0.0 0.0 550 600 650 700 550 600 650 700 550 600 650 700 T-QY7/GO T-QY7/GO T-QY7/GO 0.8 0.8 0.8 Normalized / F EBOV MARV VSV 0.4 0.4 0.4 0.0 0.0 550 600 650 700 550 600 650 700 550 600 650 700 1.2 T-ED17/GO T-ED17/GO T-ED17/GO 8.0 0.8 0.8 FBOV MARV VSV 0.4

Fig. 4 Fluorescence titration of **T-RS5/GO**, **T-QY7/GO** and **T-ED17/GO** (**T-RS5/GO** = $0.5 \, \mu \text{M}/10 \, \mu \text{g mL}^{-1}$, **T-QY7/GO** = $0.5 \, \mu \text{M}/40 \, \mu \text{g mL}^{-1}$ and **T-ED17/GO** = $0.5 \, \mu \text{M}/24 \, \mu \text{g mL}^{-1}$, respectively) with increasing EBOV (0–15 ng mL⁻¹), MARV (0–15 ng mL⁻¹) and VSV (0–15 ng mL⁻¹). The dashed curve in each panel represents the fluorescence emission of the peptide probe alone. All measurements were carried out in PBS (0.01 M, pH 7.4, containing 1% DMSO, v/v) with an excitation of 525 nm.

Wavelength (nm)

550 600 650 700

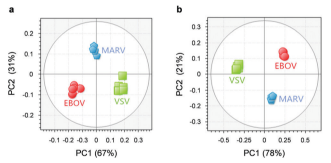


Fig. 5 Differential sensing of EBOV, MARV and VSV through transformed score of the fluorogenic response of (a) **T-RS5/GO**, **T-QY7/GO** and **T-ED17/GO** at a single composite concentration (**T-RS5/GO** = 0.5 μM/10 μg mL $^{-1}$, **T-QY7/GO** = 0.5 μM/40 μg mL $^{-1}$ and **T-ED17/GO** = 0.5 μM/24 μg mL $^{-1}$) and (b) **T-RS5/GO** (peptide/GO = 0.25 μM/5 μg mL $^{-1}$, 0.5 μM/10 μg mL $^{-1}$ and 0.1 μM/20 μg mL $^{-1}$), **T-QY7/GO** (peptide/GO = 0.25 μM/80 μg mL $^{-1}$) and **T-ED17/GO** (peptide/GO = 0.25 μM/12 μg mL $^{-1}$, 0.5 μM/24 μg mL $^{-1}$ and 0.1 μM/36 μg mL $^{-1}$) determined by PCA. The fluorogenic response expressed as histograms determined by a microplate reader of Fig. 5 is shown in Fig. S3 (ESI†).

data from multivariable problems.²⁸ The advantage of PCA is that it retains the characteristics of the data whilst reducing the dimensions, and provides more intuitive output for subsequent analysis, which promotes PCA as a method of choice for the statistical analysis of complex data.²⁹ By analyzing the different fluorescence recovery rates of the peptide sensor array in the

presence of each virus, we determined that the signals for each virus could be differentiated using the 2 top principal components (PCs) of 67% and 31%, respectively (Fig. 5a). Increasing the experimental variables for the sensor array, i.e. concentration of the peptide probes, we were able to determine a more clustered distribution for the three viruses (Fig. 5b); with the 2 top principal components (PCs) being 78% and 21%, respectively. These results suggest that, using the "sensor array" concept, it is possible to differentiate between complex viruses that express different capsid proteins using weakly specific peptide probes. Since sensing stability under varying conditions is crucial for assuring the accuracy in analyte detection, we carried out stability tests for the sensor array. The results suggest good photostability and pH stability of the GO-based fluorogenic peptide sensors (Fig. S4, ESI†). However, we note that non-specific adsorption of viruses to the exposed surface of GO could not be precluded. Therefore, in our future studies, blocking agents will be used to address this issue.

In summary, we have developed a supramolecular GO-based fluorogenic peptide sensor array for the differential sensing of viruses. Using this method, we were able to differentiate EBOV from two other viruses, MARV and VSV. We believe that our method could become a simple and general strategy for the quick differentiation of viruses of different origins without the need for specific peptide ligands.

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Conflicts of interest

No conflicts of interest.

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