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Single-molecule analysis of interaction between p53TAD and MDM2 using aerolysin nanopores†

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Protein–protein interactions (PPIs) are regarded as important, but undruggable targets. Intrinsically disordered p53 transactivation domain (p53TAD) mediates PPI with mouse double minute 2 (MDM2), which is an attractive anticancer target for therapeutic intervention. Here, using aerolysin nanopores, we probed the p53TAD peptide/MDM2 interaction and its modulation by small-molecule PPI inhibitors or p53TAD phosphorylation. Although the p53TAD peptide showed short-lived (<100 ms) translocation, the protein complex induced the characteristic extraordinarily long-lived (0.1 s ~ tens of min) current blockage, indicating that the MDM2 recruitment by p53TAD peptide almost fully occludes the pore. Simultaneously, the protein complex formation substantially reduced the event frequency of short-lived peptide translocation. Notably, the addition of small-molecule PPI inhibitors, Nutlin-3 and AMG232, or Thr18 phosphorylation of p53TAD peptide, were able to diminish the extraordinarily long-lived events and restore the short-lived translocation of the peptide rescued from the complex. Taken together, our results elucidate a novel mechanism of single-molecule sensing for analyzing PPIs and their inhibitors using aerolysin nanopores. This novel methodology may contribute to remarkable improvements in drug discovery targeted against undruggable PPIs.

Introduction

Targeting protein–protein interactions (PPIs) for therapeutic interventions has been an emerging strategy for drugging undruggable PPI targets. With the FDA approval of a Bcl-2 inhibitor (Venetoclax), PPI inhibitors with high specificity have drawn increasing attraction in drug discovery. However, the screening of small-molecule PPI inhibitors remains still challenging because of the critical limitations of conventional biophysical techniques such as nuclear magnetic resonance (NMR), surface plasma resonance (SPR), and fluorescence polarization (FP). Thus, there is an urgent need to develop new innovative technologies to accelerate the development of PPI-targeted drugs against undruggable PPIs.

Biological nanopores have been extensively utilized for diverse analyses of biomolecules such as nucleic acid sequencing,⁴⁻⁶ characterization of structured oligonucleotides and biomolecular interactions,⁷⁻¹⁰ detection of protein interaction,^{11,12} and peptide sensing.^{13,14} Despite previous attempts at analyzing ATCR-NCBD and p53-MDM2 interactions with solid-state nanopores,¹⁵⁻¹⁷ there have been limitations in the

application of biological nanopores for PPI analysis. Most of biological nanopores used for protein detection (such as ClyA, ¹⁸ MspA, ¹⁹ and FraC²⁰) are composed of a vestibule and a channel including the constriction region. Capturing of a small protein analyte in a vestibule provides characteristic information such as protein size and conformational change. However, capturing of protein–protein complexes into vestibules is limited by the small diameter of vestibule. Previously, the engineered FhuA chimera was used as a biological nanopore sensor utilized for PPI analysis, where the binding and release of the receptor by a globular protein ligand can be measured as the transition of current between open and closed states. ¹¹ However, the FhuA pore must be specifically engineered to generate a target-fused chimera for PPI targets. In addition, an adaptor and a linker should be customized for each specific target.

Aerolysin from *Aeromonas hydrophila* is a heptameric pore without a vestibule. The narrow $(d=\sim 1.2 \text{ nm})$ and long channel $(l=\sim 10 \text{ nm})^{21-23}$ of the aerolysin pore enables excellent current separation over other pores. This allows clear discrimination at the single nucleotide and single amino acid levels. Hous, owing to its high sensitivity and reproducibility, as well as exquisite spatial resolution, aerolysin nanopores are mainly used for single-molecule sensing of oligonucleotides and DNA sequencing, the detection of peptides with different charges and lengths, analysis of protein post-translational modification (PTM), 27,28 and enzyme activity measurement. And an arrow without the matter of the matter of

p53 is a tumor suppressor protein that induces cell cycle arrest and apoptotic cell death.^{30,31} The N-terminal

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transactivation domain of p53 (p53TAD) is well characterized as intrinsically disordered protein (IDP).32 Although IDPs such as p53TAD can offer novel opportunities for drug discovery, they are considered undruggable targets because of their inaccessibility with regard to structure determination.³³ p53 is negatively regulated through direct PPI between p53TAD and oncoprotein mouse double minute 2 (MDM2).34,35 This PPI is further controlled by post-translational modifications (PTMs) such as phosphorylation in response to apoptotic stimuli. For example, the Thr18 phosphorylation results in dissociation from MDM2 and association with the transcriptional coactivator p300/ CBP.36,37 Blocking the PPI between p53TAD and MDM2 has been an attractive strategy for cancer therapy because it can restore p53 function, resulting in cancer cell apoptosis. The minimal MDM2-binding motif in p53TAD, which is a 15-residue peptide (Ser15-Asn29, referred to as p53_{TAD1}), interacts with MDM2.³⁸ It was previously reported that the p53_{TAD1} undergoes a conformational change from disordered structure to an α-helix upon binding to MDM2.³⁵ Based on the α -helical structure of p53_{TAD1}, many p53_{TAD1}-peptidomimetic lead compounds have been identified for cancer treatment. Among them, Nutlin-3 binds specifically to the p53_{TAD1}-binding pocket of MDM2, thus potently inhibiting the PPI between p53_{TAD1} and MDM2.³⁹⁻⁴¹

In this study, we analyzed the PPI between p53 $_{\rm TAD1}$ peptide and MDM2 at the single-molecule level using aerolysin nanopores. The complex formation could be probed by monitoring the extraordinarily-long-lived events (0.1 s \sim tens of minutes) of the complex as well as the event frequency of short-lived translocation (<100 ms) of p53 $_{\rm TAD1}$ peptide using aerolysin nanopores. Furthermore, the activities of small-molecule PPI inhibitors were successfully monitored by quantifying the event frequency of the restored peptide translocation. Therefore, the aerolysin nanopore sensor may serve as a novel, robust platform for single-molecule analysis of PPI, contributing to paving a way for PPI-targeted drug discovery against undruggable PPIs.

Results and discussion

Single-molecule detection of p53TAD peptides using aerolysin nanopores

Prior to the nanopore detection of PPI, we tested the translocation of free p53_{TAD1} peptide or free MDM2 using aerolysin nanopores. The p53_{TAD1} peptide has a net charge of -2.5e at pH 8.0 (pI = 4.14). To increase the capture rate, we applied a 5-fold salt gradient between the cis and trans compartments of the nanopore detection system. Free p53_{TAD1} peptide was added to the cis compartment containing 10 mM Tris-HCl (pH 8.0), 1 mM EDTA, and 200 mM KCl (Fig. 1a and S1†). The nanopore measurements were performed at various applied potentials ranging from 80 to 160 mV. As the applied potentials increased, dwell times (t_d) of free p53_{TAD1} peptides were significantly increased, indicating that the peptides dominantly bump out of the pore (Fig. 1b). This might be because, despite being intrinsically unfolded, partial folding or local charge variation of p53_{TAD1} peptide hampers its passage through the aerolysin pore. In the scatter plot, longer dwell times (10-500 ms) for nanopore events of p53_{TAD1} peptide were observed at higher

applied voltages (Fig. 1b and d). Because of the dynamic disorder of the intrinsically unfolded $p53_{TAD1}$ peptide, its dwell time distribution is widely dispersed in the scatter plot.

Five glutamic acid tag promotes the translocation of p53_{TAD1} peptide through the aerolysin nanopore

To enhance the capture rate and promote the translocation of the peptide, we designed a new peptide construct (E₅-p53_{TAD1}), where five consecutive glutamic acids are attached to the Nterminus of the p53_{TAD1} peptide. The E₅-p53_{TAD1} peptide has a highly negative net charge of -7.5e at pH 8.0 (pI 3.67). Voltagedependent nanopore detection showed that the dwell times of the E₅-p53_{TAD1} peptide decreased with an increase in the applied voltage (Fig. 1b and S1†). This indicates that unlike the wild-type p53_{TAD1} peptide, the E₅-p53_{TAD1} peptide can pass through the aerolysin pore. 42,43 In the scatter plots, most translocation events of the E₅-p53_{TAD1} peptide have short dwell times ranging from 0.5 to 2 ms at a voltage of 160 mV (Fig. 1f and g), although the residual current mean values (0.08 and 0.21) are similar to those of $p53_{TAD1}$ peptide (0.10 and 0.19). The highly negatively charged E5-tag may strengthen the electrophoretic force on the peptide, which leads to an increase in its capture rate and subsequent short-lived translocation. Hence, this E5-tagged p53TAD1 peptide was used for PPI analysis using aerolysin pores.

p53_{TAD1} is a minimal MDM2 binding domain of intrinsically disordered p53TAD protein.⁴⁴ It was previously reported that p53_{TAD1} is characterized by partially folded and fully unfolded structures.^{45,46} The two current distributions may be caused by this structural variation of E₅-p53_{TAD1}. It is possible that, during the peptide translocation, the partially folded structure of E₅-p53_{TAD1} generates higher current blockade (mean value of $I_{\rm res}/I_0$ = \sim 0.08) than that of fully unfolded structure (mean value of $I_{\rm res}/I_0$ = \sim 0.21).

Aerolysin nanopore-based detection of PPI between $\rm E_{5}$ - $\rm p53_{TAD1}$ peptide and MDM2

To monitor the PPI between the p53_{TAD1} peptide and MDM2 (Fig. 2a), we measured nanopore events for the E_5 -p53_{TAD1} peptide, MDM2, and E₅-p53_{TAD1}/MDM2 complex (Fig. 2b). Each analyte was added to the cis compartment and derived to pass through the aerolysin pore by the applied potentials ranging from 80 to 160 mV. As shown in Fig. 2c, free E₅-p53_{TAD1} peptide showed a high frequency of nanopore events at an applied potential of 140 mV. In contrast to the E₅-p53_{TAD1} peptide, free MDM2 protein (residues 3-109, pI = 9.02) showed no translocation (Fig. 2d) because of its positive net charge (+2.2e at pH 8.0). In the nanopore measurement of the E_5 -p53_{TAD1} peptide/ MDM2 complex (at a molar ratio of 1:1), the current traces displayed notably distinct nanopore events from those of free E_5 -p53_{TAD1} peptide (Fig. 2e). Whereas free E_5 -p53_{TAD1} peptide generated short-lived (0.3-100 ms at 140 mV) translocation events (Fig. 2c), the MDM2-bound complex induced current blockages with extraordinarily prolonged dwell time (0.1 s \sim tens of minutes) (Fig. 2e). In addition, an increase in the molar ratio of E₅-p53_{TAD1} peptide to MDM2 to 1:3 enhanced the

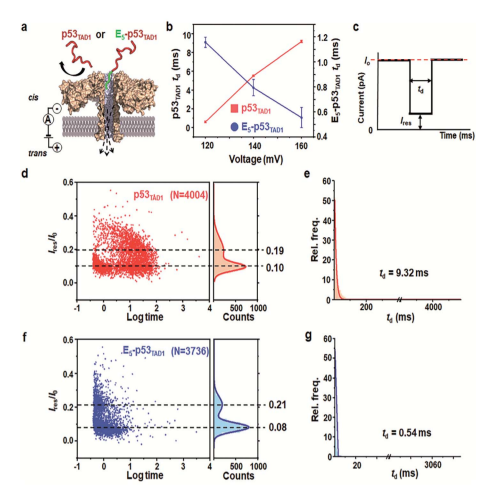


Fig. 1 Aerolysin nanopore-based detection of the intrinsically disordered p53TAD peptides. (a) The representation of the E₅-p53_{TAD1} translocation though aerolysin nanopore. (b) The voltage-dependency of dwell time in nanopore translocations of p53 $_{TAD1}$ and E_{5} -p53 $_{TAD1}$ peptides. (c) Typical nanopore event is generally characterized by the dwell time (t_d) and residual current (l_{res}/l_0) . Statistical analysis of nanopore events from the p53_{TAD1} (d and e) and E₅-p53_{TAD1} (f and g) at an applied voltage of 160 mV. The bin values used for t_d analysis are 8 and 0.5 for p53_{TAD1} and E₅-p53_{TAD1}, respectively

frequency of extraordinarily long-lived events (Fig. 2f and S2†). As shown in Fig. 2g and h, the scatter plot for the 1:3 E₅p53_{TAD1}/MDM2 complex revealed widely dispersed dwell times ranging from ~ 0.5 ms to 10^5 ms, whereas nanopore events of free E₅-p53_{TAD1} peptide were concentrated in the dwell times of <10² ms. Remarkably, the characteristic extraordinarily longlived events were detected only in the complex. Upon binding of MDM2 to E₅-p53_{TAD1} peptide, the long-lived events were significantly increased to ~4.6% of the total nanopore events. Based on these data, we chose 100 ms as the standard for the classification of long-lived nanopore events. Unlike the free E₅p53_{TAD1} peptide ($d = \sim 1$ nm), the molecular size of the complex $(>4.2 \times 3.5 \text{ nm})^{35}$ is too large to pass through a narrow channel $(d = \sim 1.2 \text{ nm})$ of aerolysin. Hence, MDM2 recruitment into the pore by the E₅-p53_{TAD1} peptide nearly completely blocked the pores, creating the characteristic extraordinarily long-lived signals.

Simultaneously, we observed a noticeable difference in the event frequencies between the free E₅-p53_{TAD1} peptide and MDM2/E₅-p53_{TAD1} complex. For quantitative analysis of event frequency, MDM2 was titrated into the E₅-p53_{TAD1} peptide at various molar ratios and then nanopore measurements were carried out. The event frequency was calculated by $f = 1/\tau_{on}$, where τ_{on} is the inter-event time between consecutive current blockades. For accuracy, 10 min-current traces without interruption of the extraordinarily long-lived events (dwell time of >5 s) were used for calculation of event frequency. As the MDM2 concentration increased, the event frequency of the complex decreased substantially (Fig. 3). The event frequency of the 1:6 E_5 -p53_{TAD1}/MDM2 complex (0.45 \pm 0.09 s⁻¹) was reduced to \sim 19.5% compared to that of free E_5 -p53 $_{TAD1}$ peptide (2.31 \pm 0.22 s^{-1}). Due to complex formation, the translocation of unbound E₅-p53_{TAD1} peptide could be impeded, which resulted in a dramatic reduction in event frequency of the peptide translocation. Based on the MDM2 concentration-dependent variation in E₅-p53_{TAD1} event frequency, we determined the binding affinity for the E₅-p53_{TAD1}/MDM2 interaction (Fig. S3 and S4, Experimental section of ESI \dagger). The determined K_d value of the E_5 -p53_{TAD1}/MDM2 interaction was 1.07 \pm 0.09 μ M (Fig. S4 \dagger), which is consistent with the previously reported K_d

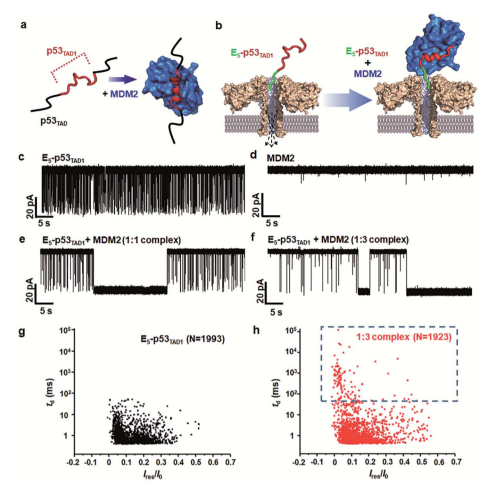


Fig. 2 Detection of PPIs between E_5 -p53_{TAD1} peptide and MDM2 by monitoring changes in current across aerolysin nanopore. (a) A schematic illustration of complex formation between intrinsically disordered p53TAD (including E_5 -p53_{TAD1}) and MDM2. The folding of E_5 -p53_{TAD1} to α -helical conformation is induced by MDM2 binding. (b) Representation of detection of the free E_5 -p53_{TAD1} and MDM2 bound form using aerolysin nanopore. (c-f) Current traces of free E_5 -p53_{TAD1}, free MDM2, and E_5 -p53_{TAD1}/MDM2 complexes (at the molar ratio of 1:1 and 1:3) at an applied voltage of 140 mV. Scatter plots of nanopore events from the E_5 -p53_{TAD1} (g) and 1:3 complex (h). The effect of the MDM2 binding with E_5 -p53_{TAD1} is represented by the dotted blue square.

value for the p53 $_{TAD1}$ /MDM2 interaction (\sim 1 μ M).⁴⁷ Taken together, our results demonstrated that the aerolysin nanopore enables rapid, real-time monitoring of the PPI between the E $_5$ -p53 $_{TAD1}$ peptide and MDM2 at the single-molecule level.

Aerolysin nanopore-based detection of PPI modulation by phosphorylation

PTMs such as phosphorylation are known to govern the binding specificity for diverse partners. ^{48,49} Previous studies showed that the phosphorylation of Thr18 in p53TAD decreases binding affinity to MDM2 by 25-fold while increasing the affinity to p300 by 45-fold. ^{36,37} Mutational studies suggested that the reduction of MDM2-binding affinity results from electrostatic repulsion of negatively charged pThr18 by a proximal negative patch of the N-terminal domain of MDM2. ⁵⁰

To monitor the effect of phosphorylation on PPI between p53_{TAD1} and MDM2 using aerolysin nanopores, we generated a p53_{TAD1} construct with the E_5 -tag and Thr18 phosphorylation (referred to as E_5 -pT18-p53_{TAD1}). In the nanopore

measurements, free E_5 -pT18-p53_{TAD1} showed current traces similar to those of free E_5 -p53_{TAD1} (Fig. 4 and S5†). However, voltage-dependent nanopore detection showed that the dwell times of the E_5 -pT18-p53_{TAD1} peptide slightly increased with increase in the applied voltages (120 mV to 180 mV) (Fig. S6†), indicating that the E_5 -pT18-p53_{TAD1} peptide does not translocate through the aerolysin pore. In addition, scatter plots of E_5 -pT18-p53_{TAD1} showed slightly longer dwell times (mean dwell time = \sim 1.11 ms) than that of E_5 -p53_{TAD1} (\sim 0.8 ms) at a voltage of 140 mV. A previous study showed that the T18 phosphorylation of p53_{TAD1} induces a long-range interaction between phosphorylated T18 and K24, forming a partially distorted structure unfavourable for the MDM2 interaction. This partial distortion of E_5 -p53_{TAD1} may induce a longer dwell time than that of E_5 -p53_{TAD1}.

To monitor whether the PPI between E_5 -p53 $_{TAD1}$ and MDM2 is modulated by phosphorylation, we titrated the MDM2 protein into the *cis* compartment containing E_5 -pT18-p53 $_{TAD1}$. At an applied voltage of 140 mV, all mixtures of E_5 -pT18-p53 $_{TAD1}$ /

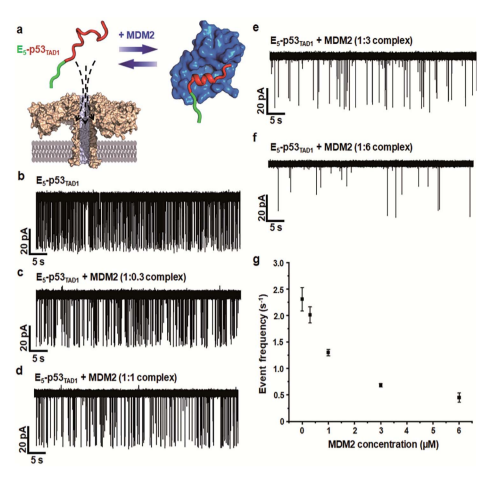


Fig. 3 Characterization of changes in the current traces obtained following E_5 -p53_{TAD1}-MDM2 interactions. (a) Representation of the nanopore detection of the E_5 -p53_{TAD1}/MDM2 complex. (b-f) The current traces of the E_5 -p53_{TAD1}/MDM2 complex at various molar ratios. The 10 min-current traces without long-lived events were selected to clearly represent variations in event frequencies. (g) Changes in nanopore event frequency of the free E_5 -p53_{TAD1} according to MDM2 concentration. The error bar indicates the standard deviation based on at least three independent experiments. The event frequencies were calculated by $f = 1/\tau_{on}$, where τ_{on} represents the inter-event interval. The value of τ_{on} at each MDM2 concentration was determined by fitting to a single exponential decay function (bin = 0.1–1.6).

MDM2 at 1:1 and 1:3 molar ratios showed negligible difference in current traces of nanopore events (Fig. 4b–d). The characteristic extraordinarily long-lived event observed in the $E_5\text{-p53}_{\mathrm{TAD1}}/\mathrm{MDM2}$ complex disappeared in the $E_5\text{-pT18-p53}_{\mathrm{TAD1}}/\mathrm{MDM2}$ mixture. Additionally, the event frequencies of $E_5\text{-pT18-p53}_{\mathrm{TAD1}}/\mathrm{MDM2}$ mixture ($f=1.71\text{--}1.81~\mathrm{s}^{-1}$) were not significantly reduced compared to those of free $E_5\text{-pT18-p53}_{\mathrm{TAD1}}$ ($f=2.19~\mathrm{s}^{-1}$) (Fig. S7†). Taken together, our results indicate that Thr18 phosphorylation of $E_5\text{-p53}_{\mathrm{TAD1}}$ inhibits the PPI between $E_5\text{-p53}_{\mathrm{TAD1}}$ and MDM2, which is consistent with previous isothermal titration calorimetry (ITC) results. 36 Thus, specific PPI modulation of p53TAD by phosphorylation could be detected at the single-molecular level using aerolysin nanopores.

Aerolysin nanopore-based detection of PPI inhibition by small-molecule drugs

Small-molecule drugs, Nutlin-3 and AMG232, has been reported to be potent p53TAD/MDM2 interaction inhibitors. The previous SPR study showed that Nutlin-3 and AMG232 have $K_{\rm d}$

values of \sim 83 nM and \sim 0.045 nM for MDM2 binding, respectively.38,52 We monitored the inhibition of the p53TAD/MDM2 interaction with the small-molecule inhibitors using aerolysin nanopores (Fig. 5a). Nanopore measurements were performed by titration of Nutlin-3 or AMG232 (at a molar ratio ranging from 1 to 5) into a 1: 3 E₅-p53_{TAD1}/MDM2 complex (Fig. 5b and S8-S11†). First, the characteristic extraordinarily long-lived events were clearly diminished upon addition of Nutlin-3 or AMG232. Second, the event frequency was gradually restored with an increase in the concentration of Nutlin-3 or AMG232, indicating that the complex of E_5 -p53_{TAD1} peptide and MDM2 was dissociated by the small-molecule inhibitors (Fig. 5b, S8 and S9†). In particular, AMG232 induced a full recovery of the event frequency of E_5 -p53_{TAD1} peptide translocation (Fig. 5b and S9†). The statistical analysis of 10 min-current traces showed a significant decrease in the number of nanopore events for the E_5 -p53_{TAD1}/MDM2 complex (N = 446), compared to that for free E_5 -p53_{TAD1} peptide (N = 1573) (Fig. 5b and S11†). However, titration of Nutlin-3 and AMG232 (at the molar ratio of 1:3:5) recovered the number of translocation events to the levels of

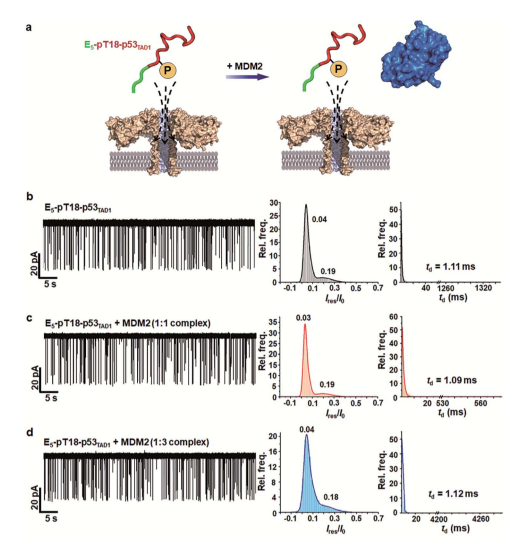


Fig. 4 Nanopore detection of inhibition of the E_5 -pT18-p53_{TAD1}/MDM2 interaction by phosphorylation. (a) A schematic illustration of nanopore translocation of E_5 -pT18-p53_{TAD1} in the absence and presence of MDM2. Thr18 phosphorylation of p53_{TAD1} inhibits interaction with MDM2. (b-d) Current traces and statistical analysis of E_5 -pT18-p53_{TAD1} in the presence of the MDM2 protein (at molar ratios of 1:1 and 1:3) applied at 140 mV. The mean t_d values were determined by fitting to single exponential decay with bin value = 1.

 \sim 61.3% (N= 965) and \sim 99.8% (N= 1570) of free E₅-p53_{TAD1} peptide, respectively (Fig. 5b). In contrast, nanopore experiment with a nonbinder control, LCL161, which is a XIAP inhibitor and does not bind to MDM2,53 did not restore the translocation of E₅-p53_{TAD1} peptide (Fig. 5b and S10†), confirming that the recovery of event frequency arises from specific PPI inhibition by drugs. Next, event frequencies (s⁻¹) based on three independent nanopore experiments (n = 3) were calculated and compared in Fig. 5c. The event frequencies of E₅-p53_{TAD1} peptide and 1 : 3 E_5 -p53 $_{TAD1}$ /MDM2 complex were 2.31 \pm 0.22 s^{-1} and 0.68 \pm 0.03 s^{-1} , respectively. The restored event frequency upon the addition of small-molecule inhibitors (at the molar ratio of 1:3:5) follows the order of AMG232 (f = 2.26 $\pm 0.14 \text{ s}^{-1}$) > Nutlin-3 ($f = 1.54 \pm 0.08 \text{ s}^{-1}$) > LCL161 ($f = 0.64 \pm 0.08 \text{ s}^{-1}$) 0.07 s⁻¹). Interestingly, this order is consistent with that of the MDM2-binding affinities reported previously AMG232 (K_d = 0.045 nM) > Nutlin-3 ($K_d = 83 \text{ nM}$) > LCL161 ($K_d = \text{n.d}$). 38,52

These results suggest that more potent PPI inhibitors induced more recovery of event frequency of E_5 -p53 $_{TAD1}$ peptide translocation. Taken together, the aerolysin nanopore sensor enables the detection of E_5 -p53 $_{TAD1}$ /MDM2 PPI and its small-molecule inhibition via single-molecule analysis of the extraordinarily long-lived events of the MDM2 complex as well as event frequency of the E_5 -p53 $_{TAD1}$ peptide translocation.

In our previous study, the PPI between p53TAD and MDM2 was probed using a solid-state nanopores. ¹⁶ Although low-noise solid-state nanopores with pyrex substrate were used for PPI detection, they showed a relatively poor $I_{\rm RMS}$ noise level (\sim 10 pA) compared to that of aerolysin nanopore (1–1.5 pA in this study). Since intrinsically disordered p53TAD showed relatively poor signal-to-noise characteristics, we engineered the protein by tagging globular-shaped GST to enhance the sensitivity.

Owing to unique long, narrow geometry of channel and lack of vestibule, ^{21,22} aerolysin nanopores make it possible to analyze

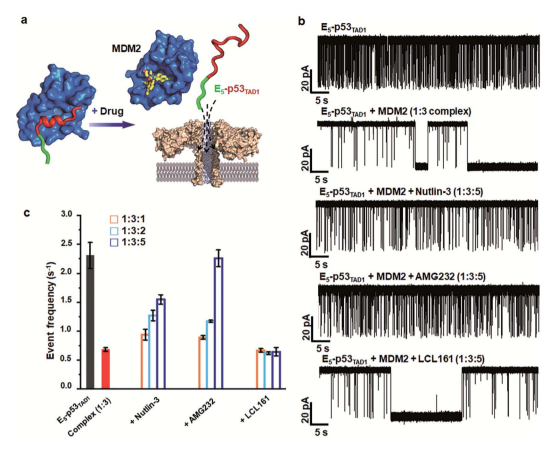


Fig. 5 Nanopore detection of inhibition of E₅-p53_{TAD1}/MDM2 interaction by small-molecule inhibitors. (a) Representation of the nanopore detection of the inhibition of the interaction of E_5 -p53 $_{TAD1}$ and MDM2 by small-molecule inhibitors. (b) Current traces obtained from free E_5 p53_{TAD1} and E₅-p53_{TAD1}/MDM2 complex in the presence of small-molecule PPI inhibitors. (c) PPI inhibition effects of small-molecule PPI inhibitors as monitoring the events frequency. The 10 min-current trace of each nanopore experiment was analyzed for the determination of event frequency. The error bar indicates the standard deviation based on at least three independent nanopore experiments

unfolded peptide or IDPs at the single-molecule level with high sensitivity and excellent resolution. Here, we provide a novel sensing mechanism for single-molecule PPI analysis using aerolysin nanopores. In the aerolysin nanopores, the PPI generated two noticeable changes of current signal, one from (p53TAD peptide) and the other from its globular protein partner (MDM2): (i) upon the complex formation, intrinsically disordered E₅-p53_{TAD1} peptide hijacks a large-sized, globular MDM2 to the pore entrance, leading to characteristic extraordinarily long (0.1 s \sim tens of min) current blockage. (ii) Simultaneously, the protein complex formation sequesters free p53TAD peptide and thereby prevents its translocation through the pore, which eventually reduces the event frequency of shortlived peptide translocation. Moreover, upon addition of smallmolecule PPI inhibitors (Nutlin-3 and AMG232) or Thr18 phosphorylation of the p53TAD peptide, the observed current signal changes were reversed; extraordinarily long-lived events diminished and the event frequency of the p53TAD peptide translocation was restored. Taken together, the activities of small-molecule PPI inhibitors can be simply and quantitatively monitored using an aerolysin nanopore.

Conclusions

Drugging the intractable PPI targets is a key challenge in drug discovery that requires innovation and the development of new technologies.54,55 Most current biophysical approaches for PPI detection, such as NMR,56,57 SPR, and FP,58 require large amounts of samples, highly expensive instruments, and complicated time-consuming procedures. However, owing to its single-molecule detection, the aerolysin nanopore sensor provides strong advantages for PPI analysis and screening of small-molecule PPI inhibitors: ultrahigh sensitivity, real-time detection, low-cost, and no immobilization. Particularly, our method has two remarkable advantages viz. (i) using aerolysin nanopores, the PPI between p53TAD and MDM2 could be detected in a \sim 10 picomole range, which shows \sim 4000-fold higher sensitivity than NMR; (ii) the association/dissociation of the protein complex can be monitored in real-time through only a \sim 10 min-current trace analysis. Taken together, these notable merits suggest that aerolysin nanopores can be applied for single-molecule-based structure-activity-relationship (SAR) analysis or drug screening of small-molecule PPI inhibitors, which can be further accelerated through integration into a multi-array and nanofluidic platform.59

In summary, we demonstrated a novel principle of single-molecule sensing for PPI of p53TAD and PPI inhibition by protein phosphorylation or small molecule inhibitors using aerolysin nanopores. Our results provide a proof-of-concept that aerolysin can be a novel, valuable platform for single-molecule-based PPI analysis and ultrasensitive, rapid screening of PPI inhibitors. This might open up new possibility for application of protein nanopore sensor to facilitate PPI-targeted drug discovery against undruggable PPIs.

Author contributions

S. O. and M.-K. L. performed experimental research. M.-K. L. and S.-W. C. designed and supervised the research, analysed the data, and wrote the paper.

Conflicts of interest

The authors declare no competing financial interest.

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