A series of non-enzymatic isothermal nucleic acid circuiting reactions, known as enzyme-free DNA circuits,\textsuperscript{1-4} have caught increasing attention because they are able to execute nucleic acid assembly with low cost, high flexibility, programmability, and readout comparability. Hybridization chain reaction (HCR), first reported by the Piece group,\textsuperscript{5} is one of the most important examples of enzyme-free DNA circuits.\textsuperscript{6-8} It relates to a pair of complementary DNA hairpins with sticky ends (H1, H2) and an initiator (I0). During the pathway, the pair of complementary DNA hairpins is utilized as fuel packets to achieve self-assembly, forming a long-nicked double-stranded DNA (dsDNA) polymer. Due to the reaction can provide both size and concentration amplification, HCR has been deep-studied and extensively-utilized as an intelligent sensing unit during molecular recognition, amplification, and transduction. The reactions have been engineered and designed for various different targets, such as nucleic acids,\textsuperscript{9,10} proteins,\textsuperscript{11} small molecules,\textsuperscript{12} ions,\textsuperscript{13} bacteria,\textsuperscript{14} cells,\textsuperscript{15} etc.\textsuperscript{16}

However, there are still problems in the existing HCR technology, which often limit its practical applications. For example, how to export an effective signal for an HCR reaction or products is always a serious challenge. The most-used signal is fluorescence resonance energy transfer (FRET).\textsuperscript{17} It is a common agreement that such a “signal-off” response may get easily confused with the false positives and insufficient intuitiveness. Therefore, continuous efforts have been made to develop “signal-on” responses. In many cases, classic HCR hairpin substrates have to be extended with additional oligonucleotides (named tails) to form an assembly-probe that can generate downstream signals.\textsuperscript{5-8,18-21} Taking an example from our previous publications,\textsuperscript{22} after HCR two adjacent tails can integrate into a full G-quadruplex structure. A porphyrin derivative (abbreviated as PPIX) can insert into the G-quadruplex and get much enhanced fluorescence emission. By similar concept, many other readouts, such as colorimetric, luminescent, and enzymatic signals have also been successfully realized.\textsuperscript{23,24} Even though, experimental evidence has revealed once the HCR hairpin is added with additional tail sequences, the assembly efficiency or signal-to-background ratio may be seriously exhibited, which makes the HCR difficult to control, design, and apply.\textsuperscript{25} In addition, because the HCR initiator is always the target itself or a sequence highly relevant to the target, the hairpin sequences have to be re-built, re-designed, and/or re-optimized once a new target is detected, adding the time, cost, and risk for failure.

Above issues are common for most HCR-based detection and require further advance to overcome them. With this purpose, here we report a universal and signal-on HCR based detection platform via innovatively coupling the CRISPR-Cas12a system with HCR. By using this CRISPR-HCR pathway, we can detect different targets by only changing the crRNA. The CRISPR-HCR platform coupling with an upstream amplifier can achieve a practical sensitivity as low as ~aM of ASFV gene in serum.

We report a universal and signal-on HCR based detection platform via innovatively coupling the CRISPR-Cas12a system for universal and signal-on detection\textsuperscript{†}

**Rujian Zhao,\textsuperscript{ab} Chunxu Yu,\textsuperscript{ab} Baiyang Lu\textsuperscript{*a} and Bingling Li\textsuperscript{*ab}**

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The detailed CRISPR-HCR pathway is illustrated in Scheme 1. As well-reported, Cas12a can be programmed with CRISPR RNAs (crRNAs) to specifically recognize the target DNA. Its collateral cleavage of the nonspecific single stranded DNA (ssDNA) reporter (trans-cleavage), initiated by the recognition and cleavage of the target DNA (cis-cleavage), has been used for the detection of nucleic acids (Scheme 1a).\textsuperscript{27,28} Based on our preliminary experiment, it is proved that ssDNA \( I_0 \) can be cleaved by CRISPR Cas12a in Fig. S1.\textsuperscript{†} In our CRISPR-HCR platform (Scheme 1b and c), \( H_1 \), \( H_2 \), initiator \( I_0 \) (a 24 nt ssDNA), crRNA and Cas12a are the primitive elements. A fluorophore (FAM) and a quencher is labelled on \( H_1 \) (FAM-H1) and \( H_2 \) (H2-BlackQ1) respectively. In the absence of target (Scheme 1b), the \( I_0 \) can initiate a classic HCR reaction to generate [FAM-H1:H2-BlackQ1]-polymer. Because of the adjacent position of the fluorophore and quencher, the fluorescence of FAM is quenched, getting low background signal compared with the reaction without \( I_0 \). While in presence of the targeting sequence (either ssDNA or dsDNA, Scheme 1c), the Cas12a can recognize the target through the bind of antisense guide RNA (crRNA), forming a specific hairpin scaffold structure. Then the Cas12a undergoes a series of conformational changes in its RuvC active site allowing for cis-cleavage of the target. Trans-cleavage activity is subsequently activated, and indiscriminately cleaves all ssDNA in the surrounding.\textsuperscript{29} In this way, \( I_0 \) is non-specifically cleaved and loses its ability to start the HCR reaction, the FAM and BlackQ1 molecules are still apart from each other, producing a target dependent high fluorescence intensity compared with the non-target background.

As illustrated, the introduction of CRISPR-Cas12a system has successfully turns the HCR reaction from a “signal-off” to “signal-on” response. Also important, due to the non-specific cleavage function of Cas12a to \( I_0 \), one set HCR components (\( I_0 \), \( H_1 \), and \( H_2 \)) can be suitable to any targeting sequence without rebuilding. Therefore, here we directly select the sequences from one of the most powerful HCR reaction.\textsuperscript{24}

In initial principle verification and optimization, we use a piece of ds-DNA fragment within ASFV p72 (ref. 30) as the model target, being named dsDNA-T1. The sequence of dsDNA-T1 is rationally selected starting from a potential PAM site starting from “TTTA” at 5’ end. Its crRNA guide sequence (crRNA-T1) is designed to include antisense of dsDNA-T1 and a hairpin structure. Fig. 1a experimentally verifies the introduction of CRISPR-Cas12a system indeed turns the HCR from a “signal-off” response (Fig. 1a, left) to a “signal-on” one (Fig. 1a, right).

The experimental condition used in Fig. 1a is selected from a series of optimization. 300 nM Cas12a is used because its presence the highest signal-to-background ratio (Fig. S3†). We also investigate two critical factors that may usually affect the HCR module, the concentration ratio of \( I_0 \) to \( H_1 \) and \( H_2 \) and the concentration of \( H_1 \) (or \( H_2 \)). For convenience, the final concentrations of \( H_1 \) and \( H_2 \) are always equal. According to the 3D-bargraph shown in Fig. 1b, among all the conditions tested, the highest signal-to-background ratio appears when the concentration of \( H_1 \) (or \( H_2 \)) is 240 nM, and the concentration ratio of \( I_0 \) to \( H_1 \) (or \( H_2 \)) is 1 : 3.

![Scheme 1 Schematic illustration of the CRISPR-HCR platform. (a) Principle of CRISPR-Cas12a-mediated cis and trans cleavage of DNA. (b and c) CRISPR-HCR platform. Without the target sequence, \( I_0 \) activates the HCR assembly and produces low fluorescence background due to FRET (b). After target binds with crRNA, Cas12a cleaves \( I_0 \) and remains HCR unreacted, producing increased fluorescence targeting signal (c).](image-url)
Based on the optimized conditions, we evaluated the sensitivity of the CRISPR-HCR for detection of dsDNA-T1. A broad dynamic range (pM to nM, Fig. S4 and S5†) of more than three orders of magnitude is achieved, with the lowest detection limit around 1 pM. Furthermore, the universality of the CRISPR-HCR platform is verified by changing the crRNA guide sequence according to a new targeting sequence. Here another dsDNA fragment from Human Papilloma Virus26 (HPV, named dsDNA-T2) and a randomly designed ssDNA sequence (named ssDNA-T) are selected as the models. As shown in Fig. 1c, under the same condition the responses of all three targets (dsDNA-T1, dsDNA-T2, and ssDNA-T) are very close, presenting equally high signal-to-background ratios.

Through above proof-of-concept experiments, we have obtained a universal, flexible, and both dsDNA and ssDNA targets. In order to realize more practical performance which can be generalized to those real-world pathogen genes existing at very trace amounts, we further couple an ultra-isothermal nucleic acid amplification with CRISPR-HCR. LAMP reaction31,32 is selected as the model because it can amplify 103–105-fold at a constant temperature (60–65 °C) within 1–2 hours.33,34 In details shown in Fig. 2a and b, the target in above demonstration (Scheme 1c) is replaced by the LAMP amplicons for a special targeting gene. At first, the ASFV gene including the dsDNA-T1 fragment (named dsDNA-ASFV), which is selected as the model-target, has been selected as our template. The LAMP primers are self-designed. The efficiency of the LAMP reaction is verified using traditional agarose gel electrophoresis (Fig. S6†). We think that the assay detection limit depended on the LAMP detection limit, based on our experiments, the detection limit of the LAMP reaction is 1.25 copies per μL when the detection rate is 95% (Fig. S7†). To further evaluate the sensitivity of the LAMP-CRISPR-HCR platform, the LAMP products amplified from different concentrations (0.5 copies per μL to 5000 copies per μL) of synthetic dsDNA-ASFV (named ASFV-LAMP-products) are used to start the downstream CRISPR-HCR platform. The fluorescence spectrum (Fig. 2b) clearly shows ASFV-LAMP-products amplified from as low as 0.5 copies per μL (~aM) dsDNA-ASFV can already generate very high fluorescence enhancement compared with the background of water control (labelled with “0 copy”). It is notable, according to the Fig. S7,† the detection rate of 0.5 copies per μL is about 25%, and theoretically more than 5 copies per μL can achieve 100% detection rate. On contrast, the signals (Fig. 2c) for LAMP products amplified from genes of non-specific pathogens, for example, Middle East Respiratory Syndrome35 (named MERS-LAMP-products) and M13mp18 DNA32 (named M13mp18-LAMP-products), are almost same as that of water control (labelled with “NC”), which verifies the practically high sensitivity and specificity of the LAMP-CRISPR-HCR platform.

To test the feasibility of applying the LAMP-CRISPR-HCR strategy in real samples, we explore the detection of dsDNA-ASFV in pig serum, one of the most challenging matrices that contains many interfering molecules. We prepare the pig serum samples spiked with 1 × 1014 copies per μL of dsDNA-ASFV templates. After extracting the dsDNA-ASFV by the commercial extraction TIANamp genomic DNA kit, the extracts are diluted by different folds, being followed by LAMP-CRISPR-HCR detection. It is found when the sample is diluted by 1013 folds, the dsDNA-ASFV can still be detected with very high signal-to-background ratio (Fig. 2d). This result shows the components in real samples cannot affect the detection efficiency, which verifies the practicability of the platform.

In order to further evaluate the adaptability of the method, we replace the hybridization chain reaction (HCR) with catalytic hairpin assembly (CHA), which is another widely-used nucleic acid circuit.1,16 According to Fig. 3a, CHA also requires two hairpin substrates (named H3 and H4-FAM), in which H4 is FAM tagged. In the absence of the target, the CHA catalyst C1 (kind of I0 for HCR) initiates two strand-displacement (SD) reactions that forms H3:H4-FAM:C1 intermediate. Then C1 automatically dissociate from the intermediate and starts another reaction cycle, leaving H3:H4-FAM as the assembly product. At the same time, the tail of H3:H4-FAM is able to start another SD reaction to bind a probe sequence labelled with BlackQ1 (named P-BlackQ1), finally forming the H3:FAM-H4:P-BlackQ1 complex. Very similar to the CRISPR-HCR, once the targeting sequence exists, the Cas12a can recognize the target through the binding of antisense guide RNA (crRNA). Thus, C1 is non-specifically cleaved, remaining the CHA unreacted. An enhanced fluorescence signal is produced (Fig. 3b). Taking dsDNA-T1 as the model-target, the CRISPR-CHA conditions are optimized (Fig. S8–S10†). Then as low as 1.25 nM dsDNA-T1 can lead to saturated fluorescence enhancement compared with the negative background (labelled with “NC”), as shown in Fig. 3c. Further, via replacing the dsDNA-T1 with LAMP products, we
also couple the LAMP with the CRISPR-CHA strategy taking dsDNA-ASFV as the model target. As expected, Fig. 3d shows the ASFV-LAMP-products from 500 copies of dsDNA-ASFV (labelled with “PC”) has successfully produce much higher signal compared that of negative control (labelled with “NC”).

In this communication, a specific, sensitive, and universal detection platform has been established via coupling CRISPR-Cas12a system and nucleic acid signal amplification circuits. The hybridization chain reaction, HCR, has been taken as the main model circuit. With the assistance of CRISPR-Cas12a system, HCR with traditional “off” signal has been improved to an “on” response. And a well-optimized, high-performance circuit can be generalized to any targeting sequence, either ssDNA or dsDNA. By combing an upstream ultra-amplifier [i.e., LAMP], the so-called CRISPR-HCR platform can detect pathogen genes down to aM level, meeting the high requirement of real-world applications. Finally, similar platform is proven also applicable to other circuits via replacing HCR with CHA circuit. The high adaptability is further confirmed.

Conflicts of interest
There are no conflicts to declare.

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