Dear Editor,

Today, the need for time-effective and cost-effective nucleic acid detection methods is still growing in fields such as human genotyping and pathogen detection. Using synthetic biomolecular components, many methods have been developed for fast nucleic acid detection\(^1\)–\(^3\); however, they may not be able to satisfy specificity, sensitivity, speed, cost and simplicity at the same time. Recently, a very promising CRISPR-based diagnostic (CRISPR-Dx) (namely SHERLOCK) was established, which was based on the collateral effect of an RNA-guided and RNA-targeting CRISPR effector, Cas13a\(^4\). SHERLOCK is of high sensitivity and specificity, and is very convenient in detection of target RNA. However, to detect DNA sequences, in vitro transcription of DNA to RNA must be conducted prior to the SHERLOCK test, which could be inconvenient.

In a recent study, we found that Cas12a, which belongs to the class 2 type V-A CRISPR-Cas system\(^5\), performed collateral cleavage on non-targeted ssDNAs upon the formation of the Cas12a/crRNA/target DNA ternary complex\(^6\). Here, with the employment of this feature, we used a quenched fluorescent ssDNA reporter (e.g., HEX-N12-BHQ1 in Supplementary Table S1) as the probe, and developed HOLMES (an one-HOURLow-cost Multipurpose highly Efficient System), which could be used for fast detection of target DNA as well as target RNA. In HOLMES, if a target DNA exists in the reaction system, the Cas12a/crRNA binary complex forms a ternary complex with the target DNA, which will then trans-cleave non-targeted ssDNA reporter in the system, illuminating the HEX fluorescence (or any other fluorescence) (Fig. 1a).

We ever purified ten Cas12a proteins (Supplementary Table S3) and found all showed the ssDNA trans-cleave activity\(^6\). To find the most suitable Cas12a for HOLMES (i.e., with high signal-to-noise ratios), we tested all ten Cas12a proteins and found Lachnospiraceae bacterium ND2006 Cas12a (LbCas12a), Orbacterium sp. NK2B42 Cas12a (OsCas12a), Lachnospiraceae bacterium NC2008 Cas12a (Lb5Cas12a) and Francisella tularensis Cas12a (FnCas12a) showed good performance, among which LbCas12a was chosen for the following studies (Fig. 1b). To determine the sensitivity of HOLMES, we titrated target DNA, and found the minimum detectable concentration for Cas12a-crRNA was approximately 0.1 nM; however, when combined with PCR, the detectable concentration could be as low as 10 aM (Fig. 1c), which was comparable to the SHERLOCK system\(^4\) and was better than PCR alone or quantitative PCR using the SYBR Green method (Supplementary Figure S1). Therefore, to achieve higher sensitivity, PCR amplification was employed in the HOLMES test thereafter.

To test whether HOLMES could discriminate single-base differences, we made point mutations at different positions in the target DNA sequence, including both the PAM region and the guide sequences (Supplementary Figure S2a). When a full length of crRNA guide sequence (24-nt crRNA, Supplementary Table S2) was used, we found mutations in either the PAM sequences or the region of the 1st–7th bases of the guide sequence resulted in clear decline of the fluorescence signal; however, no significant difference was observed when the mutation was within the region of the 8th–18th bases (Supplementary Figure S2b), which was highly consistent with the previous report that the 5′-end seed region in the crRNA

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Fig. 1 (See legend on next page.)
guide sequence was extremely important for Cas12a recognition⁶. In addition, based on our previous findings⁸, Cas12a with a reduced length of crRNA guide sequence showed higher cleavage specificity. Therefore, we then tested shorter guide sequences, and found point mutations within a larger region (1st−16th bases) resulted in more than 2-fold difference in fluorescence signals for both 16-nt and 17-nt crRNA guide sequences (Supplementary Figure S2b), suggesting that shorter guide sequences might be used in HOLMES. Furthermore, considering the fact that there might exist no suitable PAM sequence nearby the SNP site, primers for PCR amplification were specially designed to introduce the PAM sequence (Supplementary Figure S3), which therefore allowed for sequence-independent detection of any single nucleotide polymorphism (SNP) sites.

We then chose a dozen of SNP loci that are related to human health and personal characteristics (Supplementary Table S4). We either extracted genomic DNA from cultured human 293T cells or collected saliva from human individuals, and then PCR amplification targeted the SNP regions, followed by the HOLMES assay to distinguish alleles (Fig. 1d). The results clearly showed that HOLMES had sufficiently high specificity to determine both homozygous and heterozygous genotypes (Fig. 1e and Supplementary Figure S4a). We also collected nineteen volunteers’ saliva samples to detect the SNP rs1014290, which is related to gout risk, and proved that HOLMES could be used to rapidly and easily detect human SNP genotypes (Supplementary Figure S4b).

Moreover, HOLMES could also be used to detect DNA viruses (e.g., pseudorabies virus (PRV), Supplementary Figure S5a) and RNA viruses (e.g., Japanese encephalitis virus (JEV), Supplementary Figure S6a), and the sensitivity for both could be as low as 1–10 aM (Supplementary Figures S5b and S6b). For JEV, total RNA was first extracted and then reverse transcribed into cDNA before being detected by HOLMES. Because of the high sensitivity, HOLMES successfully detected PRV virus in both the PRV-infected cells and the culture supernatant (Supplementary Figure S5c). In addition, the high specificity of HOLMES also enabled it to distinguish between virus strains. For example, the PRV Ra classical strain, the cmz variant strain and the Bartha-K61 vaccine strain were easily discriminated by the gE46 site (Supplementary Figure S5d and S5e). Similarly, the JEV NJ2008 strain and the live-attenuated vaccine strain SA14–14–2 were well differentiated by the site E138 (Supplementary Figure S6c and S6d).

The “SHERLOCK” nucleic acid detection system was recently established with the employment of the “RNA collateral effect” of Cas13a and an isothermal amplification method⁹. Although both HOLMES and SHERLOCK show attomolar detection sensitivity and can be used to detect both DNA and RNA targets, this study indicates that HOLMES may have advantages in DNA detection, while SHERLOCK is more convenient for RNA detection. In addition, isothermal amplification methods (e.g., the recombinase polymerase amplification (RPA) and loop-mediated isothermal amplification (LAMP)) can also be used although rapid PCR amplification was used in HOLMES in this study. Similar to SHERLOCK, HOLMES requires no expensive reagents and no special instruments, making it low cost and easily accessible for nucleic acid detection. In addition to the medical applications described above, HOLMES may also be used for a variety of applications that require rapid detection of nucleic acids, including monitoring foods and the environment.

(While this manuscript has been ready to submit to Cell Discovery, two pieces of work were published on Science, both of which described the use of the Cas12a trans-cleavage activity on ssDNAs for nucleic acid detection⁹, ¹⁰.)
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S.Y.L., G.P.Z. and J.W. designed the experiments. Q.X.C. and X.Y.L. conducted most of the experiments. J.K.L. purified the Cas12a proteins. J.M.W., Z.L.Z., S.G. and R.B.C. cultured cells and performed viral transfection experiments. S.Y.L. and J.W. wrote the paper. All authors read and approved the final version of the manuscript. J.W. supervised the whole project.

Conflict of interest
Q.X.C., S.Y.L., J.W and X.Y.L. are co-inventors on patent applications filed by Shanghai Tolo Biotechnology Company Limited and Shanghai Institutes for Biological Sciences relating to the work in this manuscript. The remaining authors declare that they have no conflict of interest.

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