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***Basic Study***

**Dual gRNAs guided CRISPR/Cas9 system inhibits hepatitis B virus replication**

Wang J *et al*. Dual HBV-specific gRNAs inhibit HBV replication

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**Abstract**

**Aim:** To screen and investigate the effective gRNAs against hepatitis B virus (HBV) of genotypes A-D.

**Methods:** A total of 15 gRNAs against HBV of genotypes A-D were designed. 11 combinations of two above gRNAs (dual-gRNAs) covering the regulatory region of HBV were chosen. The efficiency of each gRNA and 11 dual-gRNAs on the suppression of HBV (genotypes A-D) replication was examined by the measurement of HBV surface antigen (HBsAg) or e antigen (HBeAg) in the culture supernatant. The destruction of HBV-expressing vector was examined in HuH7 cells co-transfected with dual-gRNAs and HBV-expressing vector using polymerase chain reaction (PCR) and sequencing method, and the destruction of cccDNA was examined in HepAD38 cells using KCl precipitation, plasmid-safe ATP-dependent DNase (PSAD) digestion, rolling circle amplification and quantitative PCR combined method. The cytotoxicity of these gRNAs was assessed by a mitochondrial tetrazolium assay.

**Results:** All of gRNAs could significantly reduce the HBsAg or HBeAg production in the culture supernatant, which was dependent on the region in which gRNA against. All of dual gRNAs could efficiently suppress the HBsAg and/or HBeAg production for HBV of genotypes A-D, and the efficacy of dual gRNAs in suppressing HBsAg and/or HBeAg production was significantly increased when compared to the single gRNA used alone. Furthermore, with PCR direct sequencing we confirmed that these dual gRNAs could specifically destroy HBV expressing template by removing the fragment between the cleavage sites of the two used gRNAs. Most importantly, gRNA-5 and gRNA-12 combination not only could efficiently suppressing HBsAg and/or HBeAg production, but also destroy the cccDNA reservoirs in HepAD38 cells.

**Conclusion:** These results suggested that CRISPR/Cas9 system could efficiently destroy HBV expressing templates (genotypes A-D) without apparent cytotoxicity. It may be a potential approach for eradication of persistent HBV cccDNA in chronic HBV infection patients.

**Key words**: Dual gRNAs; CRISPR/Cas9; hepatitis B; cccDNA; antiviral therapy

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**Core tip:** In this manuscript, 15 hepatitis B virus (HBV)-specific gRNAs were designed according to the HBV genome sequences of genotypes A-D. We confirmed that the CRISPR/Cas9 system with these HBV-specific gRNAs could efficiently suppress the replication of multiple genotypes HBV. Further, we demonstrated that dual gRNAs could guide the CRISPR/Cas9 system to efficiently destroy HBV cccDNA and reduce its level in HepAD38 cells. Since cccDNA, the template of HBV replication, accounts for the persistence of HBV infection. Our data suggested that CRISPR/Cas9 technique may be a useful tool to eradicate HBV of multiple genotypes.

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**INTRODUCTION**

Despite prophylactic vaccines available for many years, hepatitis B virus (HBV) infection remains an important public health problem worldwide. Current antiviral agents, including nucleos(t)ide analogues (NAs) and interferon (IFN), can control HBV production but not eliminate HBV, due to the persistence of HBV covalently closed circular DNA (cccDNA) reservoir in the nucleus of hepatocyte. NAs shown no effect on cccDNA, thus relapse of hepatitis B occurs frequently in patients who discontinued antiviral treatment[1,2]. Moreover, because the stability of HBV cccDNA was so high that it declines slowly, so life-long treatment of chronic hepatitis B (CHB) is required[1,3,4]. On the other hand, though IFN-α can degradate cccDNA followed by cytidine deamination and apurinic/apyrimidinic site formation, and can further result in virus clearance in a few patients, but its efficacy is unsatisfactorily limited[5-7]. Eradication of cccDNA, the only way to reach the clinical cure of CHB, is still an unresolved problem in the treatment of CHB.

Clustered regularly interspaced short palindromic repeats/Cas9 nuclease (CRISPR/Cas9) system is a novel genome editing tool derived from the adaptive immune system of bacteria and archaea[8-10]. CRISPR/Cas9 system promotes genome editing by inducing a double-strand break (DSB) at the target genomic locus. In the absence of a repair template, DSBs are re-ligated through the Non-homologous end joining (NHEJ) process, which leads to the insertion/deletion (indel) mutations[11]. CRISPR/Cas9 system has been successfully applied not only for genome editing in cells, but also for disrupting the genome of virus, including adenovirus, herpes simplex virus (HSV) and human immunodeficiency virus (HIV)[12-14]. For HBV, the CRISPR/Cas9 system has been proved to efficiently cleave the expressing templates of HBV genotype A and D[15-17]. Whereas, genotypes A, B, C, D are the predominant genotypes of HBV in East Asia and other part of the world[18-20], so it is necessary to design guide RNAs (gRNAs) specific for HBV genotypes A-D. Here, we evaluated the potential use of CRISPR/Cas9 system to clear the HBV genome of genotypes A-D.

**MATERIALS AND METHODS**

***Plasmids***

The 1.2xHBV construct (pBB4.5-HBV1.2, genotype C) was constructed using a 1.2-fold length genome of genotype C HBV DNA sequence, and was inserted into the pBB4.5-HBV1.3 (genotype D, G1896A mutation) plasmid digested with PstI and NheI enzymes. The pBB4.5-HBV1.3 (genotype D, G1896A mutation) plasmid was kindly provided by professor Locarnini SA from the Victorian Infectious Diseases Reference Laboratory, Australia[21]. The 1.2xHBV construct (pBB4.5-HBV1.2, genotype C) has been proved to efficiently produce HBV[22]. The HBV-expression vectors pGEM-HBV1.3A (genotype A) and pGEM-HBV1.3B (genotype B) were kindly provided by Professor Ningshao Xia from School of Public Health, Xiamen University, China.

Cas9 promotes genome editing by inducing a double-strand break (DSB) and re-ligating through the NHEJ process in the absence of a repair template. The gRNA/Cas9 dual expression vector pSpCas9(BB)-2A-GFP (PX458) was obtained from Addgene (Cambridge, MA). PX458 plasmid which expresses a nonsense gRNA (GGGTCTTCGAGAAGACCT) was used as a vector control in each experiment. HBV-specific gRNA/Cas9 dual expression vectors were constructed in our laboratory. The oligonucleotides sequences for the construction of HBV-specific gRNA/Cas9 dual expression vectors were listed in Supplementary Table S1.

***Transfection of cells***

Human liver cancer cell lines HuH-7[23] and HepAD38[24] (stable expression of HBV) were maintained in Dulbecco modified Eagle medium supplemented with 10% fetal bovine serum (Gibco, Carlsbad, Calif). HuH-7 and HepAD38 cells were seeded in a 12-well plate at 1.5 × 105 cells/well, HuH-7 cells were co-transfected with HBV expression vectors (HBV of genotype A, B or C) and HBV-specific gRNA/Cas9 dual expression vector [pSpCas9(BB)-2A-GFP] by lipofectamain 2000 (11668019; Life Technologies). HepAD38 cells were transfected with HBV specific gRNA/Cas9 dual expression vector by lipofectamain 3000 (L3000015; Life Technologies).

***Detection of HBsAg and HBeAg***

Seventy-two hours after transfection, cell culture supernatant were collected for detection of HBsAg and HBeAg by a time-resolved fluoroimmunoassay (TRFIA) according to manufacturer's instructions (SY60108A and SY60105A; PerkinElmer). In brief, culture supernatant (100 μl) was added into a microtiter plate coated with anti-HBsAg or anti-HBeAg and shaked for 40 min in room temperature, then washed for 4 times. Europium-labeled anti-HBsAg or anti-HBeAg was diluted 1:50 by HBsAg or HBeAg dilution buffer and added at 100 μl per well, shaked for 40 min in room temperature, then washed for 6 times. At last, after incubation with enhancement solution (100 μl) for 5 min, the plates were read using Anytest reader (SYM-BIO), and the concentrations of HBsAg and HBeAg was calculated according to the standard curve. The relative HBsAg or HBeAg level was calculated as the ratio of HBsAg or HBeAg concentration in the cell culture supernatant of gRNA treated and vector control cells.

***Detection of HBV DNA fragments cleaved by dual gRNAs***

DNA was extracted from cells using QIAGEN DNA mini kit (51304; QIAGEN), according to the manufacturer's instruction. Specific primers listed below were used to PCR amplify the cleaved HBV DNA fragments. primer1F (nucleotide position: 1856-1877): 5'-CCTACTGTTCAAGCCTCCAAGC-3'; primer2F (nucleotide position: 321-342): 5'-CAACCTCCAATCACTCACCAAC-3'; primer1R (nucleotide position: 434-415): 5'-AGAAGATGAGGCATAGCAGC-3'; primer2R (nucleotide position: 2006-1986): 5'-CAGAGGCGGTGTCAAGGAGAT-3'; primer3R (nucleotide position: 1702-1682): 5'-GACTCAAGGTCGGTCGTTGAC-3'; primer4R (nucleotide position: 1285-1264): 5'-CTAGGAGTTCCGCAGTATGGAT-3'. Primer1F and 1R were used for detection the fragment cleaved by dual gRNAs of gRNA1+13 or gRNA2+14, primer 2F and 2R, 2F and 3R, 2F and 4R were used for detection the fragment cleaved by dual gRNAs of gRNA3+5, gRNA4+5 and gRNA5+12, respectively.

PCR reaction mixture (20 μl) contained 10 μl 2×Taq mix (Transgene), 1 μl forward primer (10 μmol/L), 1 μl reverse primer (10 μmol/L), 1 μl DNA template and 7 μl double distilled water (ddH2O). The reaction mixture was denatured at 95 °C for 5 min, followed by 35 cycles at 95 °C for 30 s, 60 °C for 30 s, 72 °C for 1 min, and at last 72 °C for 5 min. Agarose gel (1.5%) was used for separation of DNA fragments with different length. DNA markers were DL2000 and DL2000 Plus (GenStar).The DNA fragment of expected length was sequenced.

***Detection of HBV cccDNA***

Two gRNA/Cas9 dual expression vectors were co-transfected into HepAD38 cells at least 7 d after tetracycline removed. Cells were collected for DNA extraction at 72 h after transfection using QIAGEN DNA mini kit (51304; QIAGEN), according to the manufacturer's instruction. To obtain cccDNA, KCl precipitation and plasmid-safeTM ATP-dependent DNase (PSAD) (Epicentre, Madison, WI, United States) were used to remove HBV DNA integrated into cell genome, HBV rcDNA, replicative dsDNA and ssDNA. Afterwards, rolling Circle Amplification (RCA) was conducted to selectively amplify cccDNA. Finally, PCR was performed using RCA products as template, and using cccDNA specific primers which target the gap region of HBV genome[25,26].

***Mitochondrial tetrazolium assay***

Mitochondrial tetrazolium (MTT) assay is used to monitor cell viability. Cells (1500 cells per well) were plated in 96-well plates and were maintained in DMEM supplemented with 10% fetal bovine serum. 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide solution was added into the cell culture at a final concentration of 5 μg/mL and was allowed to remain in culture for 4 h before measurement. Cell viability was monitored every 24 h by measuring the absorbance in a microplate reader (Bio-Rad, Hercules, Calif).

**RESULTS**

***Construction of HBV-specific gRNAs***

Based on SpCas9 in our CRISPR/Cas9 system requires a 5’-NGG protospacer adjacent motif (PAM) sequence, HBV genome of genotypes A-D were searched for potential 18-20 base target sequences called the protospacer with 3’-downstream of PAM, which was shown as GN18-20-NGG. Finally a panel of 15 HBV-specific gRNAs targeting different regions of HBV genome were designed (Figure 1). To avoid the off-target effect, the gRNA sequences were blasted by Nucleotide Blast search (NCBI), and the difference of more than 3 nucleotides to other sequences in human genome was assured. Finally, the gRNA expression vectors were identified by sequencing method. The sequences and locations of these HBV-specific gRNAs were listed in Table 1.

***designed HBV-specific gRNAs efficiently suppress the production of HBsAg and/or HBeAg***

To examine the efficiency of each gRNA in the suppression of HBV replication, the 1.2xHBV construct (pBB4.5-HBV1.2, genotype C) was co-transfected with each gRNA/Cas9 dual expression vector in HuH-7 cells respectively. As shown in Figure 2a, all designed gRNAs could significantly suppress HBsAg level in culture supernatants, but with noticeable variation of efficacy, and gRNAs targeting the S region (gRNA-1, 2, 3, 4) and enhancer I region (gRNA-5) of HBV genome exhibited higher HBsAg suppressing efficiency than that of the gRNAs targeting either X region (gRNA-6, 7, 8, 9, 10) or preC/C (gRNA-11, 12, 13, 14, 15) region. For HBeAg, all of gRNAs could significantly reduce HBeAg levels in culture supernatant, with the exception of gRNA-6 and gRNA-8. It is noteworthy that gRNAs targeting preC/C region exhibited higher HBeAg suppressive efficiency than that of those gRNAs targeting eithther S or X regions (Figure 2b). Intriguingly, gRNA-5 against enhancer I region could efficiently suppress both HBsAg and HBeAg levels (Figure 2a and b). The off-target effect of gRNA may induce cytotoxicity, and then disturb the antiviral effect of HBV-specific gRNAs. The methods for assessing cell viability can test cytotoxicity. To exclude the possibility that the HBsAg or HBeAg suppression observed above was the result of non-specific cytotoxicity, the MTT assay which is a [colorimetric assay](http://en.wikipedia.org/wiki/Colorimetry_(chemical_method)) for assessing cell viability was conducted. The results revealed that no noticeable cytotoxicity of the gRNAs was observed in HuH-7 cells (Figure 2c).

***Dual gRNAs exhibit synergistic effect on the suppression of HBV replication***

Since CRISPR/Cas9 system can be efficiently used for multiple genome cleavage[11], we wondering if combination of two HBV-specific gRNAs (dual gRNAs) should be more efficient than single one in suppressing HBV replication. To confirm this, pBB4.5-HBV1.2 and combinations of two different gRNA expression vectors were co-transfected into HuH-7 cells. Firstly, we chose the region modulating the expression of HBsAg and HBeAg between two gRNAs in HBV genome, including the combinations of gRNA-1+13 covering the S gene promoter and pre S1 region, gRNA-1+12 covering the S gene promoter pre S1 region and Core coding DNA sequence (CDS) region, and gRNA-8+12 covering Core promoter and Enahncer II region were tested. As expected, the synergistic effect of dual gRNAs in the suppression of HBV replication was observed. Two of the tested dual gRNAs gRNA-1+13 (gRNA1 *vs* gRNA-1+13: 0.255 *vs* 0.110, *P =* 0.0031; gRNA13 *vs* gRNA-1+13: 0.265 *vs* 0.110, *P =* 0.0017) and gRNA-8+12 (gRNA8 *vs* gRNA-8+12: 0.319 *vs* 0.110, *P =* 0.0004; gRNA12 *vs* gRNA-8+12: 0.240 *vs* 0.110, *P =* 0.0007) demonstrated significantly higher suppressive efficiency in HBsAg production, when compared to that of each gRNA used alone (Figure 3a). Consistent with HBsAg, there was synergistic effect of gRNA-1+13 (gRNA1 *vs* gRNA-1+13: 0.296 *vs* 0.124, *P =* 0.0034; gRNA13 *vs* gRNA-1+13: 0.264 *vs* 0.124, *P =* 0.0018) and gRNA-8+12 (gRNA8 *vs* gRNA-8+12: 0.889 *vs* 0.121, *P <* 0.0001; gRNA12 *vs* gRNA-8+12: 0.260 *vs* 0.121, *P =* 0.0018) in the suppression of HBeAg production (Figure 3b). While for the gRNA-1+12, such synergistic effect could only be observed in HBeAg production (gRNA1 *vs* gRNA-1+12: 0.296 *vs* 0.096, *P =* 0.0014; gRNA12 *vs* gRNA-1+12: 0.260 *vs* 0.096, *P =* 0.0006), and an antagonistic effect for HBsAg production (gRNA1 *vs* gRNA-1+12: 0.265 *vs* 1.125, *P =* 0.0001; gRNA12 *vs* gRNA-1+12: 0.240 *vs* 1.125, *P =* 0.0001) was exhibited (Figure 3c).

Following the above observation, 11 dual gRNAs (two different gRNAs at ratio of 1:1) were used for further study. The choice of gRNAs for combination was made according to the efficiency of each gRNA and its targeting regions (in promoter, enhancer or reverse transcriptional region of polymerase). Next, the efficiency of those 11 dual gRNAs in the suppression of HBsAg and HBeAg was examined, with different HBV genotypes taken into consideration. The HBV-expression vectors (genotype A, B or C constructs) and two different gRNAs expression vectors were co-transfected into HuH-7 cells. For HBV of genotype C, all of dual gRNAs could significantly suppress HBsAg production (Figure 3d), just as expected. However, though all dual gRNAs could also significantly suppress HBeAg production, the efficacy was lower than that of HBsAg, since higher HBeAg suppressive efficiency (≥ 80%) was detected in only 7 dual gRNAs (Figure 3e). Similar results were observed when HBV of genotype A and B were tested (Supplementary Figure S1a and b). To exclude the possibility that excess concentration of gRNA may interfere HBV replication, we tested the concentration effect of gRNAs in HBV replication. The result revealed that the suppression efficiency of gRNA on the HBsAg level was gRNA concentration dependent (Figure S2). Similarly, in order to exclude the interference of non-specific cytotoxicity, MTT assay was conducted. The result affirmed that there was no noticeable cytotoxicity of dual gRNAs (Figure 3f).

***Dual gRNAs destroy HBV genome***

To confirm that the dual gRNAs mediated suppression in HBV replication was at genome level, the HBV genome region covering the cleavage sites of dual gRNAs was amplified using PCR. If dual gRNAs worked, the fragment between the two cleavage sites would be removed, consequently a relative smaller PCR product would be detected. According to the above suppressive efficiency of dual gRNAs combinations, five dual gRNAs (1+13, 5+12, 2+14, 3+5 and 4+5) were chosen for further investigating. As expected, all of dual gRNAs could destroy HBV genome, especially for the combination of gRNA-5+12, in which only the smaller fragment was detected. A up to 100% cleavage efficiency implicated the optimum combination of the two gRNAs (Figure 4a). Furthermore, direct sequencing of the PCR product showed that the smaller fragment was indeed formed by the re-ligation between the ends of two cleavage sites (Figure 4b). Then the plasmid pBB4.5-HBV1.2 were co-transfected with dual gRNA-5+12 expression vectors at different ratios to HuH-7 cells and the cleavage efficiency was assessed by PCR amplification. The result revealed that at the ratio of 1:3, almost all of HBV genome DNAs was cleft by gRNA-5+12. Surprisingly, even at the ratio of 3:1, still more than 90% of HBV genome DNAs were cleft (Figure 4c). In line with this, the efficiency in suppressing the production of HBsAg in culture supernatant was also gradually but significantly decreased from the ratio of 1:3 (reduced to 0.019% ± 0.007%) to 3:1 (reduced to 0.60% ± 0.014%), as compared to the vector control (Figure 4d). Other four dual gRNAs also showed the capability to destroy HBV genome at different extents (Figure 4a), all were confirmed by sequencing analysis (data not shown).

As shown in Figure 4e, the target genome sequence of gRNA-13 in genotype B HBV harbors a one base difference as compared to genotypes A/C/D. However, dual gRNAs composed by gRNA-1and gRNA-13 could also destroy HBV genome of genotype B (Figure 4f).

***Dual gRNAs destroy HBV cccDNA in HepAD38 cells***

Above data demonstrated that HBV-specific gRNAs could destroy HBV expressing template. Theoretically, such gRNAs could also destroy HBV cccDNA. To confirm this, the most powerful dual gRNAs (gRNA-5 and gRNA-12 expression vectors) were co-transfected into HepAD38 cells which stably express genotype D HBV and produce HBV cccDNA[24]. Firstly, the HBsAg and HBeAg levels in the culture supernatant were measured. The results showed that even with low transfection efficiency, gRNA-5 and gRNA-12 together could significantly suppress both HBsAg and HBeAg levels, particularly for HBsAg which almost dropped 50% off (Figure 5a). Besides, gRNA-5 and gRNA-12 could also significantly reduce the HBV DNA level in the HepAD38 cell culture supernatant (Figure 5b). Since cccDNA reservoirs are the cause of chronic HBV infection, next, the HBV cccDNA level in HepAD38 cells was measured. To make sure that the quantitative measurement of the HBV cccDNA was reliable, KCl precipitation, plasmid-safe ATP-dependent DNase (PSAD) digestion and rolling circle amplification (Isothermal PCR amplification for the circular HBV cccDNA rather than the linear rcDNA) were employed, followed by quantitative PCR with the cccDNA specific primers (sense primer covers the cleavage site of gRNA-12) that target the gap region of HBV genome. As expected, cccDNA level in HepAD38 cells was significantly suppressed by gRNA-5 and gRNA-12 (Figure 5c). PCR amplification confirmed that the fragment between two cleavage sites of dual gRNAs in HBV expressing template was removed (Figure 4a and c). Similarly, HBV-specific gRNA could also guide Cas9 to destroy HBV cccDNA (Figure 5d). Above results suggested that the down regulation of HBsAg and HBeAg levels was at least partially due to the destruction of HBV cccDNA.

**DISCUSSION**

Hepatitis B is an important occupational hazard for health workers. An estimated 240 million people are chronically infected with HBV[27]. Multiple studies demonstrated that without therapy, 15%-40% of HBV infected patients would eventually develop cirrhosis, liver failure, or hepatocellular carcinoma (HCC)[28]. Because of the high HBV-related morbidity and mortality, HBV infection has been a serious public health problem worldwide. Unfortunately, none of the current therapy regimes could eradicate cccDNA in the nucleus of infected hepatocytes, which is the main cause of HBV persistent infection. In recent years, new strategies focused on targeting cccDNA are under intensive study. Three recent studies have suggested the potential use of CRISPR/Cas9 genome editing technique to cut HBV DNA and cccDNA *in vitro* and *in vivo*, as well as for chronic or *de novo* HBV infection[15-17]. However, those studies just focus on single genotype HBV, genotype A or D. Since genotypes B and C are also the major genotypes in the world, particularly in China. In this study, 15 gRNAs were designed according to the HBV genome sequences of genotypes A, B, C and D. These gRNAs covered different regions of HBV genome, including S, P, X, C, enhancer I and basic core promoter (BCP) regions. Our results here indicated that most of the gRNAs could significantly suppress the production of HBsAg and HBeAg. Interestingly, the suppressive efficiency of different gRNAs varied depending on the region in which they against.

As multiple gRNAs could be efficiently used to edit genome, here different combinations of two gRNAs were used to suppress HBV replication *in vitro*. And high efficiency against genotypes A-D HBV replication was found in different dual gRNAs combinations. However, the combination of gRNA-1 and 12 exhibited antagonistic effect in the suppression of HBsAg production. Currently we are unable to explain this phenomenon. A potential possibility is that the combination of gRNA-1 and 12 would cut off and remove the fragment between the two cleavage sites, where C, S promoter (Sp) and preS1 regions located. As a result, the enhancer II, the core promoter (Cp) and the basic core promoter (BCP) regions were re-ligated to the immediate upstream of S gene, as a result the expression of HBsAg was transcriptionally enhanced. In line with this, we found that dual gRNAs could indeed remove the fragment specifically between the two cleavage sites of gRNAs, which were confirmed by PCR sequencing. Such antagonistic effect reminded us that when multiple gRNAs were used in combination, the side effect should be taken into consideration.

In addition to genotype, due to absence of proofreading function of the viral reverse transcriptase and a high replication rate, the HBV population in an individual was a composition of genetically distinct but closely related variants known as quasispecies[29,30]. It is likely that such variants will cause the imperfect match between the designed gRNAs and their target viral genome sequence. The off-target effect of CRISPR/Cas system has been mentioned, such shortage could become an advantage when gRNAs were used to guide the cleavage of the high variety HBV genome in an infected individual. Just for this, though the target sequence in HBV genotype B harbors a one base mismatch with gRNA-13, which still could be cleft (Figure 4F). Besides, we and others have proved the oncogenicity of HBV integration[31,32]. Potentially, the multiple gRNAs can be used to remove the integrated HBV DNA to cure the HBV-related HCC. Most importantly, we found that wildtype HBV cccDNA level in HepAD38 cells was significantly down regulated by HBV-specific gRNA, which indicated that HBV cccDNA could be destroyed by CRISPR/Cas9 system. It is difficult to confirm the gRNA-induced destruction of cccDNA by southern blot for the low transfection efficiency of the PX458 palsmid. We will demonstrate the gRNA-induced destruction of cccDNA by southern blot when the CRISPR/Cas9 system was combined with the recombinant adeno-associated virus system. Above all, the indel mutation and destruction of cccDNA should be the main reason of gRNA-induced suppression of cccDNA replication.

Above all, we confirmed that CRISPR/Cas9 system had the potential to destroy cccDNA which is the template of HBV replication. Our data further demonstrated that dual gRNAs guided CRISPR/Cas9 system may be a useful tool to eradicate CHB and HBV-related disease with infection of multiple genotypes HBV. However, there is still far away for the treatment of HBV infected patients by CRISPR/Cas9 system. Next, we will validate the effect of HBV-specific gRNA on HBV replication in HBV transgenic mice by hydrodynamic injection method or use the adenovirus or adeno-associated virus system. In the future, dual gRNAs guided CRISPR/Cas9 system will be developed by adeno-associated virus system which is a potential vector for gene therapy, and be used in combination with the current NAs and/or IFN-α based antiviral therapy for the possible clinical cure of CHB.

**comments**

***Background***

The CRISPR/Cas9 system is a novel genome editing tool which leads to genome indel mutation by inducing a double-strand break (DSB) at the target genomic locus. The CRISPR/Cas9 system has been successfully applied not only for genome editing in cells, but also for disrupting the genome of virus, including adenovirus, herpes simplex virus, human immunodeficiency virus and Hepatitis B virus (HBV). It has become recognized that CRISPR/Cas9 system may be a potential tool to cure viral disease.

***Research frontiers***

Several recent studies have confirmed the potential use of CRISPR/Cas9 system to cleave HBV DNA and cccDNA *in vitro* and *in vivo*, as well as for chronic or *de novo* HBV infection.

***Innovations and breakthroughs***

Eradication of cccDNA, the only way to reach the clinical cure of chronic hepatitis B (CHB), is still an unresolved problem in the treatment of CHB. Previous studies have reported that the CRISPR/Cas9 system can efficiently cleave the expressing template of HBV. However, those studies just focus on HBV of genotype A or D. Whereas, genotypes A, B, C, D are the predominant genotypes of HBV in East Asia and other part of the world, so it is necessary to design guide RNAs (gRNAs) specific for HBV genotypes A-D. Besides, dual gRNAs can get the synergistic effect in genome editing. In this paper, we evaluated the potential use of dual gRNAs guided CRISPR/Cas9 system to clear the HBV genome of genotypes A-D.

***Applications***

This study provides a potential tool, the dual gRNAs guided CRISPR/Cas9 system, to eradicate CHB and HBV-related disease with infection of multiple genotypes HBV. In the future, the dual gRNAs guided CRISPR/Cas9 system will be developed by adeno-associated virus system which is a potential vector for gene therapy, and be used in combination with the current NAs and/or IFN-α based antiviral therapy for the possible clinical cure of CHB.

***Terminology***

The clustered regularly interspaced short palindromic repeats/Cas9 nuclease (CRISPR/Cas9) system is a novel genome editing tool derived from the adaptive immune system of bacteria and archaea. The CRISPR/Cas9 system promotes genome editing by inducing a DSB at the target genomic locus. In the absence of a repair template, DSBs are re-ligated through the Non-homologous end joining process, which leads to the insertion/deletion (indel) mutations.

***Peer-review***

The authors present a highly interesting functional study in which they showed that dual gRNA guided CRISPR/Cas9 system can suppress replication of the multiple genotypes HBV as well as promote clearance of HBV cccDNA in the cell culture. This paper confirms that the presented dual gRNA/CRISPR/Cas9 system might be a potential approach for eradication of HBV cccDNA and thus considered for a new and additional antiviral treatment option of CHB.

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**Table 1 Sequences and locations of hepatitis B virus-specific gRNAs in the hepatitis B virus genome of genotypes A-D**

|  |  |  |  |
| --- | --- | --- | --- |
| **gRNA No.** | **Nucleotide position** | **Sequence (GN18-20*NGG*, 5’-3’)** | **Genotype** |
| 1 | 56-75 | CCTGCTGGTGGCTCCAGTTC | A/B/C/D |
| 2 | 182-200 | GGACCCCTGCTCGTGTTAC | A/B/C/D |
| 3 | 415-433 | GCTGCTATGCCTCATCTTC | A/B/C/D |
| 4 | 640-658 | ATGGGAGTGGGCCTCAGTC | A/B/C |
| 5 | 1179-1197 | AGTGTTTGCTGACGCAACC | A/B/C/D |
| 6 | 1393-1410 | GCCAACTGGATCCTGCGC | B/C/D |
| 7 | 1521-1540 | GGGGCGCACCTCTCTTTACG | A/B/C/D |
| 8 | 1578-1597 | GAGGTGAAGCGAAGTGCACA | A/B/C/D |
| 9 | 1589-1608 | CTTCACCTCTGCACGTCGCA | B/C/D |
| 10 | 1775-1794 | AGGAGGCTGTAGGCATAAAT | A/B/C/D |
| 11 | 1859-1878 | AGCTTGGAGGCTTGAACAGT | A/B/C/D |
| 12 | 1865-1884 | CAAGCCTCCAAGCTGTGCCT | A/B/C/D |
| 13 | 2336-2355 | ACTACTGTTGTTAGACGACG | C/D |
| 14 | 2367-2386 | CGAGGGAGTTCTTCTTCTAG | A/B/C/D |
| 15 | 2390-2409 | GATTGAGACCTTCGTCTGCG | B/C/D |

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**Figure 1** **Illustration of the gRNA–targeted sequences located in the hepatitis B virus genome.**

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**Figure 2 Fifteen hepatitis B virus-specific gRNAs could efficiently suppress the production of hepatitis B virus surface antigen or hepatitis B virus e antigen.** A: The plasmid pBB4.5-HBV1.2 (0.5 μg) was co-transfected with each individual gRNA/Cas9 dual expression vector (1.5 μg) to HuH-7 cells. HBsAg level in culture supernatant was measured at 72 h post transfection using a time-resolved fluoroimmunoassay analysis; b: HBeAg level in culture supernatant was measured using a time-resolved fluoroimmunoassay analysis as above; c: The cytotoxicity of each HBV-specific gRNA was examined using an MTT assay. Data was shown as mean ± SE of 5 independent experiments. All *P* value is student’s *t*-test. HBV: hepatitis B virus; HBsAg: hepatitis B virus surface antigen; HBeAg: hepatitis B virus e antigen; MTT: Mitochondrial tetrazolium.

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**Figure 3 Eleven dual gRNAs could efficiently suppress the production of hepatitis B virus surface antigen or hepatitis B virus e antigen.** The plasmid pBB4.5-HBV1.2 was co-transfected with gRNA-1 and 13 (a), gRNA-8 and 12 (b), or gRNA-1 and 13 (c) expression vectors to HuH-7 cells, alone or in combination. HBsAg and HBeAg levels in cell culture supernatant were measured at 72 h post transfection using a time-resolved fluoroimmunoassay; d: The plasmid pBB4.5-HBV1.2 (0.5 μg) was co-transfected with different combinations of two gRNAs expression vectors (each 0.75 μg) to HuH-7 cells. HBsAg level in culture supernatant was measured at 72 h post transfection using a time-resolved fluoroimmunoassay; e: HBeAg level in culture supernatant was measured using time-resolved fluoroimmunoassay as above; f: The cytotoxicity of dual gRNAs was examined using an MTT assay. Data was shown as mean ± SE of 3 independent experiments. All *P* value is student’s *t*-test. HBV: hepatitis B virus; HBsAg: hepatitis B virus surface antigen; HBeAg: hepatitis B virus e antigen; MTT: Mitochondrial tetrazolium.

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**Figure 4 Dual gRNAs could destroy hepatitis B virus genome.** a: The plasmid pBB4.5-HBV1.2 (0.5 μg) was co-transfected with gRNA-1 and 13, gRNA-5 and 12, gRNA-2 and 14, gRNA-3 and 5, or gRNA-4 and 5 expression vectors (each 0.75 μg) to HuH-7 cells. Cellular DNA was extracted at 72 h post transfection, and PCR amplifications were performed using the primers beyond the cleavage sites of each dual gRNAs; b: Sequencing analysis of the smaller fragment formed by gRNA-5 and 12; c: The plasmid pBB4.5-HBV1.2 was co-transfected with the gRNA-5 and 12 expression vectors at different ratios to HuH-7 cells. The smaller fragment was amplified at 72 h post transfection; d: HBsAg level in culture supernatant was measured using a time-resolved fluoroimmunoassay. Data was shown as mean ± SE of 3 independent experiments (student’s *t*-test); e: Comparative analysis for the sequences of gRNA13 and HBV genome of genotypes A-D; f: The plasmid pGEM-HBV1.3A or pGEM-HBV1.3B and gRNA-1 and 13 expression vectors were co-transfected into HuH-7 cells, and PCR amplifications were performed at 72 h post transfection as above. HBV: hepatitis B virus; HBsAg: hepatitis B virus surface antigen.

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**Figure 5 hepatitis B virus-specific gRNA could suppress the genotype D hepatitis B virus replication and destroy cccDNA.** a: HepAD38 cells were seeded into a 6 well plate. Then, gRNA-5 and 12 expression vectors (each 2 µg) were co-transfected into HepAD38 cells. HBsAg and HBeAg levels in the culture supernatant of HepAD38 cells were measured at 72 h post transfection using a time-resolved fluoroimmunoassay; b: HBV DNA levels in the culture supernatant of HepAD38 cells were measured at 72 h post transfection using real-time quantitative PCR; c: HBV cccDNA levels in HepAD38 cells were measured using KCl precipitation, plasmid-safe ATP-dependent DNase (PSAD) digestion, rolling circle amplification and quantitative PCR combined method; d: PCR amplification was performed using the primers beyond the cleavage sites of dual gRNAs following KCl precipitation, plasmid-safe ATP-dependent DNase (PSAD) digestion and rolling circle amplification. HBV: hepatitis B virus; HBsAg: hepatitis B virus surface antigen.