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(54) Title: NOVEL CRISPR ENZYMES, METHODS, SYSTEMS AND USES THEREOF

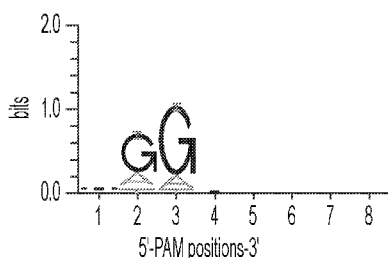


FIG. 1A

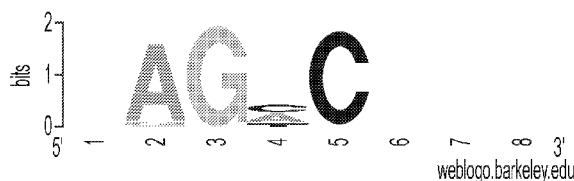


FIG. 1B

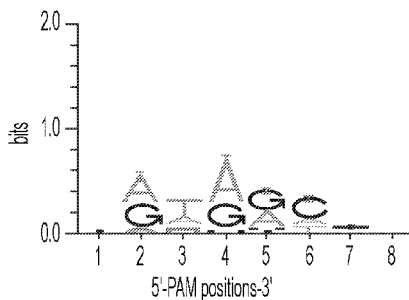


FIG. 1C

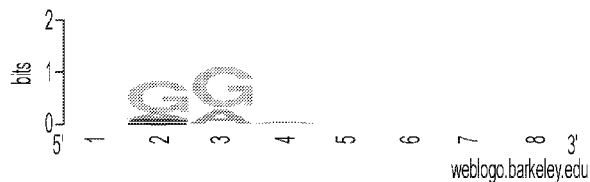


FIG. 1D

(57) Abstract: The present invention provides novel systems, methods and compositions for making and using recombinantly engineered novel Cas9 enzymes optimized for human cells, for nucleic acid targeting and manipulation. The present invention is based on the discovery of novel Cas9 enzymes from *Streptococcus constellatus*, *Sharpen spp. isolate RUG017*, *Veillonella parvula*, *Ezakiella peruensis*, *Lactobacillus fermentum strain AF15-40LB strain* and *Peptoniphilus sp. Marseille-P3761* bacteria that were codon-optimized and recombinantly produced for use in human cells. In some embodiments, novel Cas9 enzymes can be used for base editing. In some embodiments, the novel engineered Cas9 enzymes are used to treat human diseases.



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## NOVEL CRISPR ENZYMES, METHODS, SYSTEMS AND USES THEREOF

### CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority to U.S. Provisional Patent Application Serial No. 5 63/164,798, filed on March 23, 2021, which is incorporated by reference herein in its entirety for all purposes.

### BACKGROUND

Enzymes from the prokaryotic Clustered, Regularly Interspaced Short Palindromic Repeats (CRISPR) and CRISPR-associated protein (CRISPR-Cas) systems have been 10 harnessed as reprogrammable and highly specific genome editing tools for use in eukaryotes. Besides genome editing and cleavage, CRISPR-Cas9 can be used to localize effector molecules to specific sites on the genome, allowing genetic and epigenetic regulation and transcriptional modulation through a variety of mechanisms.

However, diverse genomes and genomic targets require a variety of tools for effective 15 genetic engineering, and there remains a need to expand the CRISPR toolbox through the discovery and engineering of novel Cas proteins that can recognize and target diverse sequences.

While CRISPR-Cas9 systems can be used to knock out a gene or modify the expression of a gene, certain kind of gene editing requires precise modifications to the target 20 gene, such as editing a single base within the gene. Such precise modifications remain a challenge and requires a diverse gene editing toolkit to effectuate precise genomic modifications in a wide variety of target genes.

### SUMMARY OF THE INVENTION

25 The identification of novel Cas9 enzymes with specificity for unique protospacer adjacent motifs (PAM) allows for the expansion of the available tools for gene editing. The present invention provides, among other things, engineered, non-naturally occurring novel Cas9 enzymes isolated from *Streptococcus constellatus*, *Sharpea spp. isolate RUG017*, *Veillonella parvula*, *Ezakiella peruensis*, *Lactobacillus fermentum strain AF15-40LB* and 30 *Peptoniphilus sp. Marseille-P3761* bacteria. The present invention is based, in part, on the surprising discovery that novel Cas9 enzymes discovered from different bacteria, which

recognize specific PAM sequences can be engineered for expression in eukaryotic cells (e.g., human, plant, etc.). Accordingly, the described Cas9 enzymes and their variants are functional in eukaryotes. The examples provided herewith show use of engineered, non-naturally Cas9 enzymes in human cells with diverse PAM recognition sequences to target various genomic sites. For example, Cas9 engineered from *Streptococcus constellatus*, *Ezakiella peruensis* and *Peptoniphilus sp. Marseille-P3761* recognizes the consensus PAM sequence 5'-NGG-3'. The consensus PAM sequence recognized by Cas9 isolated from *Sharpea spp. isolate RUG017* is 5'-NAGHC-3'. The consensus PAM sequence recognized by Cas9 isolated from *Veillonella parvula* was identified as 5'-NRHRRH-3'. The consensus PAM sequence recognized by Cas9 isolated from *Lactobacillus fermentum* strain AF15-40LB was identified as 5'-NNAAA-3'. (H=A, C or T; R=A or G).

In one aspect, an engineered, non-naturally occurring Cas9 protein modified from *Streptococcus constellatus* Cas9, *Sharpea* Cas9, *Veillonella parvula* Cas9, *Ezakiella peruensis* Cas9, *Lactobacillus fermentum* strain AF15-40LB Cas9 or *Peptoniphilus sp. Marseille-P3761* Cas9 is provided herein.

In some embodiments, the *Streptococcus constellatus* Cas9 protein has at least 80% sequence identity to

MGKPYSIGLDIGTNSVGVAVVTDDYKVPAAKMKVLTGNTDKQSIKKNLLGALLFDSGETAEAT  
 RLKRTARRRYTRRKNRLRYLQEI FTGEMNKVDENFFQRLDDSFLVDEDKRGEHHP I FGNIAA  
 20 EVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFK  
 DFVEEYDKTIEESHLEITVDALSILTEKVS KSSRLENLIAHYPTKKNLTFGNLIALSLDL  
 HPNFKTNFQLS EDAKLQFSKDTYEEDLEGFLGEVGDYADLFASAKNLYDAILLSGILTVDD  
 NSTKAPLSASMVKRYEEHQKDLKCLKDFIKVNAPDQYN AIFKDKNKKGYASYIESGVKQDEF  
 YKYLKGILLKINGS GDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHY PFLK  
 25 ENQDKIEKILTFRI PYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMT  
 LNDLYLP EEKVL PKHSPLYEAF TVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKY  
 TKDKLLNLYLNKEFEFRIVNLTGLDKENKAFNSSLGTYHDLRKI LDKSFLDDKAN EKTIEDI  
 IQTLTLFEDREMIRQLQKYS DIFTKAQLKKLERRHYTGWGRLSYKLINGIRNKENKKTILD  
 YLIDDDGYANRNFMLINDDALS FKEE IARAQIIDDVDDIANVVHDLPGSPA I KKGILQSVKI  
 30 VDELVKVMGHNPANII IEMARENQTTDKGRRNSQORLKL LQDSLKNLDNPVNIKNVENQQLO  
 NDRLFLLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSDD  
 VPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHV  
 AQILDERFNTEFDGNKRRIRNVKIITLKS NLVSNFRKEFELYKVREINDYHHAHDAYLNAVAV



GNALLLKY P Q L E P E F V Y G E Y P K Y N S Y R S R K S A T E K F L F Y S N I L R F F K K E D I Q T N E D G E I A W N  
 K E K H I K I L R K V L S Y P Q V N I V K K T E E Q T G G F S K E S I L P K G E S D K L I P R K T K N S Y W D P K K Y G G F  
 D S P V V A Y S I L V F A D V E K G K S K K L R K V Q D M V G I T I M E K K R F E K N P V D F L E Q R G Y R N V R L E K I I  
 K L P K Y S L F E L E N K R R R L L A S A K E L Q K G N E L V I P Q R F T T L L Y H S Y R I E K D Y E P E H R E Y V E K H K  
 5 D E F K E L L E Y I S V F S R K Y V L A D N N L T K I E M L F S K N K D A E V S S L A K S F I S L L T F T A F G A P A A F N  
 F F G E N I D R K R Y T S V T E C L N A T L I H Q S I T G L Y E T R I D L S K L G E D (SEQ ID NO: 1) .

In some embodiments, the *Sharpea* Cas9 protein has at least 80% sequence identity to

MAKNKDIRYSIGLDIGTNSVGVAVMDEHYELLKKGNNHHMWGSRLFDAAEPAATTRRASRSIRR  
 RYNKRREIRILLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQKNLGNDYKDNYNLFIEKDF  
 10 NDKTYDDKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQSFAKVYEDIEEK  
 LDNTLKKFMSLNDLDNLFVDNDINSMITVLSKIYQRSKKADDLLKIMNPTKEERAAYKEFTK  
 ALVGLKFNVS KMILAQE VKKDDKDIELDFSNDYDSTVDGLQAE LGEYIEFIEMLSINSW  
 ELQDILGNNSTISAAMVERYEEHKNDLRVLK KVI REELPKYNEVFREDNPKLHNYLGYIKY  
 PKNTPVEEFY EYIKRLLAKVD TGEAREILERIDLEK FMLKQNSRTNGSIPYQMOKDEMIQII  
 15 DNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFEDKQKERILPWNVDQIVD  
 IDATAEGFIERMQNTGTYPDPKPVMAKNSLTVSKFEVLNELNKIRINGKLIPVETKKELLS D  
 LFMKNKTI TDKKLKDWLVTHQY YDTNEELKIEGYQKDLQFSTSLAPWIDFTKIFGEINASNY  
 QLIEKIIYDISIFEDKKILKRRLK K VY Q L D D L L V D K I L K L N Y T G W S R L S E K L L T G I K S K N S K  
 ETILSILENSNMNLMEIINDESLGFKQIIEESNKKDIEGPF RYDEVK KLAGSPA I K R G I W Q A  
 20 LLVVQEITKFMKHEPSHIYIEFAREEQEKV R T E S R I A K L Q K I Y K D L N L Q T K E D Q L V Y E S L K K  
 EDAKKIDTDALYLYLQMGKSMYSGKPLDIDKLSTYHIDHILPRSLIKDDSLDNRVLVLPK  
 ENEWKLDSETVPFEIRNKMMGFWQKLHENGLMSNKKFFSLIRTD FNEKDKKRFINRQLVETR  
 QIIKNVAVIINDHYTNTNVVTVRAELSHQFRERYKIYKNRDLNDLHHAHDAYIACILGQFIH  
 QNFGNMDVNMIYGQYKKNYK K D V Q E H N N Y G F I L N S M N H I H F N D D N S V I W D P S Y I G K I K S C F C  
 25 YKDVYVTKKLEQNDAKLFDLTILPSDKNSENGVTKAKIPVNKYRKDVNKYGGFSGDAPIMLA  
 IEADKGGKHVRQVIAFPLRLKNYNDEERIKFIEKEKNLKNVKILTEVKKNQLILINHQYFFI  
 TGTNELVNATQLKLSAKNTKNL FNLVDANKHNKLESIDDANFNEVIQELICKLQEP IYSRYN  
 SIGKEFEDSYEKINAVTKQDKLYIIEYLIAIMSAKATQGYIKPELAREIGTNGKNKGRIKSF  
 TIDLNKTTFISTSVTGLFSK KYKL (SEQ ID NO: 4).

30 In some embodiments, the *Veillonella parvula* Cas9 protein has at least 80% sequence  
 identity to

MSIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLDIGTSSVGVAVTNKAYELLKFRSHK  
 MWGSRLFDEGESAVARRGFRSMRRRLERRKLRLKLLLEELFADAMAQVDPTFFMRLRESKYHY  
 EDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHHILKYRGN  
 FLYEGATFDSNASTLDDVIKQALENITFNCFCNSAIISSIGQILMEAGKTKSDKAKAIEHLV  
 5 DTYIATDVTVDTSSTKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEEEDLKKLQITG  
 DTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPGLTISESKVKAFNKHKDDLAILK  
 SLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEY  
 ILSQIELQILLPLQRIKDNQVI PYQLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKML  
 EFRIPIYYVGPLNTHHNVDNGGFAVAVRKASGRVTPWNFDDKIDREKSAAAFIKNLTNKCTYL  
 10 LGEDVLPKSSLLYSEFMLLNELNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTKNRIEQFLK  
 DNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGESKMLRE  
 TLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGTPETIIILMRNFSYNL  
 MELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALS PAVKRAVWQALRIVDEVAHIKKA  
 LPARIFVEVTRS NKNEKKKDSRQKRLSDLYAAIKKDDVLLNGLNNEIFGELKSSLAKYDDA  
 15 ALRSKKLYLYYTQMGRCAYTGEIIELSLLNTDNYDIDHIYPRSLTKDDSFDNLVLCCKRTANA  
 QKSDAYPISEEIQKTQKPFWTF LKQOGLISERKYERLTRITPLTADDLSGFIARQLVETNQS  
 VKAATTLRRLYPGVDVVFVKAENVTD FRHDNNFIKVRSLNHHHHAKDAYLNIVVGNVYHER  
 FTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKDGKAWDVKT SMDTVKKMMSDNDVRVTKR  
 LLEQT GALADATIYKATVAGKAKDGAYIGMKTSSVFADVSKYGGMTKIKNAYSIIVQYTGK  
 20 KGEVIKEIVPLPIYLTNRNTTDQDLINIVASII PQAKDISIIYGKLCINQLVKVNGFYYYLG  
 GKTNSKFCIDNAIQVIVSNEWI PYLKVLEKFNMRKDNKDLKANVVSTRALDNKHTIEVRIV  
 EEKNIEFFDYLVSKLKMPIYQKMKGNKAAELSEKGYGLFKMSLEEQSIHLIELLNLLTNQK  
 TTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIV (SEQ ID NO: 8).

In some embodiments, the *Ezakiella peruensis* Cas9 protein has at least 80% sequence  
 25 identity to

MTKVVDYYIGLDIGTSSVGVAVTDEAYNVLKFNSSKMWGVRLFDDAKTAEERRGQRGARRRL  
 DRKKERLSLLQDFFAEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKDFKDNFHK  
 KYPTIHHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFI FEGQKFDTKSSFENSLNELKVHL  
 NDEYGLDLEFDNENLINILTPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQKLVDLFE  
 30 NPEDFDDSAIKSVDFSTTSFDDKYS DYELALGDKIALVNILKEIYDSSILENLLKEADKSKD  
 GNKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNDNYVSYTKSSISNNKRV  
 KANKFTDQEAIFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNGV  
 IPYQLKLMELNKILENQSKHHEFLNVSDYGSVCDKIASIMEFRIPIYYVGPLNPNISKYAWIK

KQKDSEITPWNFKDVVDLDS SREEFIDSLIGRCTYLKDEKVLPKASLLYNEYMVLNELNNLK  
 LNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGDFKQGLNSYND  
 FKAIVGDKVSDDDYRDKIEEIIKLVLYGDDKSYLQKKIKAGYGKYFTDSEIKKMAGLNYKD  
 WGRLSKKLLTGLEGANKITGERGSI IHFMREYNLNLMELEMSASFTFTEEIQKLN PVDDRKLS  
 5 YEMVDELYLSPSVKRWLWQSLRIVDEIKNIMGTDSKKIFIEMARGKEEVKARKESRKNQLLK  
 FYKDGKKAFFISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQLGRCMYSLEPIDISELSS  
 KNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPIPDEILNKNCYAYWKILYDKGLI  
 GQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTI CKNSEIVYSKAENASRFR  
 QEFDIVKCRAVNDLHHMHDAYINI IVGNVYNTKFTKDP MNFVKKQEKARSYNLENMFKYDVK  
 10 RGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFRMNYIESGALFNATLQRKNKGSRPLK  
 DKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLMPVEREIIYAKQKNDKKSDEIFSKY  
 LKDRFGIEDYRVVYPVKMRTLLKIDGSYFITGGSDKTLELRSALQLILPKKNEWAIKQID  
 KSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLFQDDKIENIDFKFKSMDF  
 KEKCKTLLMLVKAIRASGVRQDLKSIDLKS DYGRLSSKTNNIGNYQEFKIINQSITGLFENE  
 15 VDLLKL (SEQ ID NO: 14)

In some embodiments, the *Lactobacillus fermentum* Cas9 protein has at least 80% sequence identity to

MKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGTAI GVRLFEEGKTAAERRTFRTRRRRLKRR  
 KWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEPDAKNQAFIGKLLFPDLLKKNERGYPTL  
 20 IKMRDELPEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAVHHIVKYRGHFLNNASVDKF  
 KVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLD RQKAVAKLLEV  
 KVADKEETKRNKQIATAMSKLVLGKADFATVAMANGNEWKIDLSSETSEDEIEKFREELSD  
 AQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWTHERQLAEVKEYLATQPASARKEFD  
 QVYNKYIGQAPKEKGFDFLEKGLKILSKKENWKEIDELLKAGDFLPKQRTSANGVIPHQMHO  
 25 QELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPYYVGPLVTPEVQKATSG  
 AKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKD TYLLNEDVLPANSLLYQKYNVL  
 NELNNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMAKTRGVNKP SVEGLSDPKKF  
 NSNLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVEWLTDEQRSALV  
 KKRYKGWGRLSKKLLTGIVDENGQRIIDL MWNTDQNFMQIVNQPVFKEQIDQLNQKAITNDG  
 30 MTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSI SIEFARNEGKGEITRS  
 RRTQLQKLFEDQAHEL VKDTSLTEELEKAPDLSDRYYFYFTQGGKDMYT GDPINFDEISTKY  
 DIDHILPQS FVKDDSLDNRVLVSRAENNKSDRVP AKLYAAKMKPYWNQLLKQGLITQRKFE  
 NLTMDVDQTIKYRSLGFVKRQLVETRQVIKLTANILGSMYQEAGTDIIETRAGLTKQLREEF

DLPKVREVDYHHAVDAYLTTFAGQYLNRRYPKLRSEFFVYGEYMKFKHGS DLKLRNFNFHE  
 LMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLISNETYEETGKYFDASHESSSL  
 YLKNQNKKSCLVVPLKNKLOPEYYGAYTGITQGYMVILKLLDKKGGFGVYRIPRYAADILNK  
 CHDEVAYRNKIAEIISSDPRAPKSEFVVVPRVLKGTFLVDGEEKFILSSYRYKVNATQLILP  
 5 VSDIKLIQDNFKALKKLVNEMQTKKLEIYDNILRQVDKYYKLYDINKFRAKLHDGRSKFVE  
 LDDFGQDASKEKVIKILRGLHFGSDLQNLKEIGFGTTPLGQFQVSEAGIRLSNTAFIIFKS  
 PTGLFNRKLYLKNL (SEQ ID NO: 84).

In some embodiments, the *Peptoniphilus sp. Marseille-P3761* Cas9 protein has at least 80% sequence identity to

10 MEKKTNYTIGLDIGTDSVWAVVKDDLELVKKRMKVLGNTETNYIKKNLWGSLLFESGQTAK  
 DRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDFRRLNESFLVPEDKAFSKNPIFGTL  
 GEDKTYKYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIAINEN  
 LEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSLFKDIGRANKQSPMMA  
 FITLIVGKKAKFKAAFNLEEEISLNLTDSDYDENLEILLNTIGSDFADLFDHAQRVYNAVEL  
 15 AGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKKDLKG  
 YAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVHLAEME  
 AILDRQENYYPWLKENREKIIISLLTFRIPIYVGPLADGQSEFAWLERKSDEKIKPWNFS DVV  
 DLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIAYLDERQKRMNLS SVEK  
 KEIFETLFFKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMMD  
 20 DPDNEDLMEEIVKILTVFEDRKMRKQLEKYKERLSPEQIKELAKKHYTGWGRLSKLLVGI  
 RDKETQKTILDYLVEDDNHSGGRQHLNRNMQILINDRSLFKKTAELQ MIDPSADLYAQVQ  
 EIAGSPAIKKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTTARGKALS KRREAKIKEGL  
 AALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHII PQSFTVDN  
 SLDNLVLTNSSQNRGNKKDDVPSLEVNRQLAYWRS LKDAGLMTQRKFDNLT KAMRGGLTDK  
 25 DRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNIVLLKSAMASEFRKMFRLY  
 KVRELNDYHHAHDAYLNAAIAINLLALYPYMADDFVYGEFRYKKKPQAEKATYEKLRQWNLI  
 KRFGKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNVVKKAEERTGMLFKETINGKTNKGSR  
 IPIKKDLDP SKYGGYIEEKMAYYAVISYEDKKKKPGKTIVGISIMDKKEFEYDSISYLGKLG  
 FSNPVVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKANQIAMEQDLVNFYHLKNYD  
 30 EISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIYEEEEKKDPVDIEALVA  
 SFIELLKLTSAGAPADFI FMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMRFED (SEQ  
 ID NO: 86).

In some embodiments, the Cas9 protein comprises an amino acid sequence that is at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% identical to SEQ ID NOs: 1, 4, 8, 14, 84 or 86.

In some embodiments, the Cas9 protein further comprises a nuclear localization  
5 sequence (NLS) and/or a FLAG, HIS or HA tag.

In some embodiments, the *Streptococcus constellatus* Cas9 has an amino acid sequence at least 80% identical to

MPKKKRKVGKPYISIGLDIGTNSVGVAVVTDDYKVPKMKKVLGNTDKQSIKKNLLGALLFD  
SGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSLVDEDEKRGHEH  
10 PIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAEN  
TDVQALFKDFVEEYDKTIEESHLSSEITVDALSILTEKVS KSSRLENLIAHYPTKKNLTFGN  
LIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDYADLFASAKNLYDAILL  
SGILTVDNSTKAPLSASMVKRYEEHQDLKCLKDFIKVNAPDQYNAIFKDKNKKGYASYIE  
SGVKQDEFYKYLKGI LLKINGS GDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQ  
15 GEHY PFLKENQDKIEKILTFRIPIYVGLARKGSRFAWA EYKADEKITPWNFDDILDKEKSA  
EKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDH  
VFKENRKVTKDKLLNLYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSLDDKA  
NEKTIEDIIQTTLTFEDREMIRQLQKYS DIFTKAQLKKLERRHYTGWRLSYKLINGIRNK  
ENKKTILDYLDIDGYANRNFMLINDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPAICK  
20 GILQSVKIVDELVKVMGHNPANIIEMARENQTTDKGRRNSQQRLKLLQDSLKNLDNPNVNIK  
NVENQQQLQNDRLFLYYIQNGKDMYTGETLDINNL SQYDIDHIIIPQAFIKDNSLDNRVLRSD  
KNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGG LTEEDKAGFIKRQLVE  
TRQITKHVAQILDERFNTEFDGNKRRI RNVKIITLKS NLVSNFRKEFELYKVREINDYHHAH  
DAYLNAVVG NALLLKY PQLEPEFVYGEY PKYNSYRSRKSATEKFLFYSNILRFFKKEDIQTN  
25 EDGEIAWNKEKHIKILRKVLSYPQVNIVKKTEEQTGGFSKESILPKGESDKLI PRKTKNSYW  
DPKKYGGFDS PVVAYSILVFADVEK GKSKL RLVQDMVGITIMEKKRFEKPNVDFLEQRGYR  
NVRLEKIIKLPKYSLFELENKR RLLASAKELQKGNELVIPQRFTTLLYHSYRIEKDYEPH  
REYVEKHKDEFKELLEYISVFSRKYVLADNNLT KIEMLFSKNKDAEVSSLA KSFISLLTFTA  
FGAPAAFNFFGENIDRKRYTSVTECLNATLIHQSI TGLYETRIDL SKLGEDGKRPAATKKAG  
30 QAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 2).

In some embodiments, the *Sharpea* Cas9 has an amino acid sequence at least 80% identical to

MPKKKRKVGAKNKDIRYSIGLDIGTNSVGVAVMDEHYELLKKGNNHMMWGSRLFDAAEPAATR  
RASRSIRRRYNKRRERIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQKNLGN DYKDNY  
NLFIEKDFNDKTYDYKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQSFAK  
VYEDIIEKLDNTLKKFMSLNDLDNLFVDNDINSMITVLSKIYQRSKKADDLLKIMNPTKEER  
5 AAYKEFTKALVGLKFNVS KMILAQEVKKDDKDIELD FSNVDYDSTVDGLQAE LGEYIEFIEM  
LHSIN SWVELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPKYNEVFREDNPKLH  
NYLGYIKY PKNTPVEEFY EYIKRLLAKVDTGEAREILERIDLEKFMLKQNSRTNGSIPYQMQ  
KDEMIQIIDNQSVYYPQLKENREKLISILEFRIPYFGPLNTHSEFAWIKKFEDKQKERILP  
WNYDQIVDIDATAEGFIERMQNTGTYPDPKPVMAKNSLTVSKFEVLNENLKIRINGKLI PVE  
10 TKKELLSDFMKNKTITDKKLDKDWLVTHQYYDTNEELKIEGYQKDLQFSTSLAPWIDFTKIF  
GEINASNYQLIEKIIYDISIFEDKKILKRRLLKKVYQLDDLLVDKILKLN YTGWSRLSEKLLT  
GIKSKNSKETILSILENSNMNLMEIINDESLGFKQIIEESNKKDIEGPF RYDEVKKLAGSPA  
IKRGIWQALLVQEI TKFMKHEPSHIYIEFAREEQEKV RTESRIAKLQKIYKDLNLQTKEDQ  
LVYESLKKEDAKKKIDTDALYLYLQMGKSMYSGKPLDIDKLSTYHIDHILPRSLIKDDSLD  
15 NRVLVLPKENEWKLDSETVPFEIRNKMMGFWQKLHENG LMSNKKFFSLIRTD FNEKDKKRFI  
NRQLVETRQI IKNVAVIINDHYTNTNVVTVRAELSHQFRERYKIYKNRDLNDLHHAH DAYIA  
CILGQFIHQNFQNM DVNMIYGQYKKNYK KDVQEHNNYGFILNSMNH IHFNDDNSVIWDPSYI  
GKIKSCFCYKDVYVTKKLEQNDAKLFDLTI LPSDKNSENGVT KAKIPVNKYRKDVNKYGGFS  
GDAPIMLAI EADKGKKHVRQVIAFPLRLKNYNDEERIKFIEKEKNLKNVKILTEVKKNLIL  
20 INHQYFFITGTNELVNATQLKLSAKNTKNL FNLVDANKHNKLESIDDANFNEVIQELICKLO  
EPIYSRYNSIGKEFEDSYEKINAVTKQDKLYIIEYLI AIMS AKATQGYIKPELAREIGTNGK  
NKGRIKSFTIDLNKTTFISTSVTGLFSK KYKLGKRPAATKKAGQAKKKKGSYPYDVPDYAYP  
YDVPDYAYPYDVPDYA (SEQ ID NO: 5).

In some embodiments, the *Veillonella parvula* Cas9 has an amino acid sequence at  
25 least 80% identical to

MPKKKRKVGSIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLDIGTSSVGVAVTNKAYE  
LLKFRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRLKLEELFADAMAQVDPTFFMR  
LRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLF LAVH  
HILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNC FDCNSA ISSIGQILMEAGKTKSDK  
30 AKAIEHLVDTYIATD TVDTS SKTQKDQVKEDKKRLKAFANLVGLNASLIDLFGSVEELEE D  
LKKLQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILL SIKEPGLTISESKVKAFNKH  
KDDLAILKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGL  
SDKDKEYILSQIELQILLPLQRIKDNQVI PYQLHLEELKAILAKCGPKFPFLNEVADGFSV

AEKLIKMLEFRIPIYYVGPLNTHHNVDNGGFAWAVRKASGRVTPWNFDDKIDREKSAAAFIKN  
 LTNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHHKMTK  
 NRIEQFLKNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFG  
 ESKKMLRETLRKKFASCLDDEAIAKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGTPEIIIL  
 5 MRNFSYNLMELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALS PAVKRAVWQALRIVD  
 EVAHIKKALPARI FVEVTRSNKNEKKKKDSRQRLSDLYAAIKKDDVLLNGLNNEIFGELKS  
 SLAKYDDAALRSKKLYLYTQMGRCAYTGEIIELSLLNTDNYDIDHIYPRSLTKDDSFNLY  
 LCKRTANAQKSDAYPISEEIQKTQKPFWTFLKQOGLISERKYERLTRITPLTADDLSGFIAR  
 QLVETNQSVKAATTLRRLYPGVDVVFVKAENVTD FRHDNNFIKVRSLNHHHHAKDAYLNIV  
 10 VGNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKDGKAWDVKTSMDTVKKMMS  
 NDVRVTKRLLLEQTGALADATIYKATVAGKAKDGAYIGMKTSSVFADVSKYGGMTKIKNAYS  
 IIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISIYGKLCINQLVKV  
 NGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNNMRKDNKDLKANVVSTRALDNK  
 HTIEVRIVEEKNIIEFFDYLVSKLKMPIYQKMKGNKAAELSEKGYGLFKKMSLEEQSIHLIEL  
 15 LNLNLTNQKTTFEVKPLGITASRSTVSGKISNQDEFKVINESITGLYSNEVTIVGKRPAATKK  
 AGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 9).

In some embodiments, the *Ezakiella peruensis* Cas9 has an amino acid sequence at least 80% identical to

MPKKKRKVGTKVKDYIIGLDIGTSSVGWAVTDEAYNVLFNSKKMWGVRLFDDAKTAEERRG  
 20 QRGARRRLDRKKERLSLLQDFFAEVAKVDPNFFLRDLNSDLYMEDKDQKLKSKYTLFNDKD  
 FKDKNFHKYPTIHHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFI FEGQKFDTKSSFENS  
 LNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSS  
 QKLVDLFENPEDFDDSAIKSVDFSTTSFDDKYSDEYELALGDKIALVNILKEIYDSSILENLL  
 KEADKSKDGNKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEK VNDNYVS YTKS  
 25 SISNNKRVKANKFTDQEA FYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPK  
 IKSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRIPIYYVGPLNP  
 NSKYAWIKKQKDSEITPWNFKDVVDLDS SREEFIDSLIGRCTYLKDEKVL PKASLLYNEYMV  
 LNELNNLKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGDFK  
 QGLNSYNDFKAIVGDKVSDDYRDKIEEIKLIVLYGDDKSYLQKKIKAGYGYFTDSEIKK  
 30 MAGLNKYDWGRLSKKLLTGLEGANKITGERGSI IHFMREYNLNLMELEMSASFTFTEEIQKLN  
 PVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKI FIEMARGKEEVKARKE  
 SRKNQLLK FYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQLGRCMYSLEP  
 IDISELSSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPIPDEILNKNCYAYWK

ILYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTICKNSEIVYSK  
 AENASRFRQEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKDP MNFVKKQE KARSYNLE  
 NMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRK  
 NKGSRPLKDKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLPVEREIYAKQKNDKKL  
 5 SDEIFSKYLKDRFGIEDYRVVYPVVKMRTLKIDGSYFITGGSDKTLELRSALQLILPKKN  
 EWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLFQDDKIENID  
 FFKSMSDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGRSSKTNNIGNYQEFKIINQS  
 ITGLFENEVDLLKLGKRPAAATKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA  
 (SEQ ID NO: 15).

10 In some embodiments, the *Lactobacillus fermentum strain AF15-40LB* Cas9 has an amino acid sequence at least 80% identical to

MPKKKRKVGKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGTAIQVRLFEEGKTAERRTFRT  
 TRRRLKRRKWRLHYLDEIFAPHLQEVDFLRRLKQSNIHPEDEPAKNQAFIGKLLFPDLLKK  
 NERGYPTLIKMRDELPEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAVHHIVKYRGHFL  
 15 NNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLDROK  
 AVAKLLEVKVADKEETKRNKQIATAMSKLVLYGKADFATVAMANGNEWKIDLSSETSEDEIE  
 KFREEELSDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWTHERQLAEVKEYLATQP  
 ASARKEFDQVYNKYIGQAPKEKGFLEKGLKILSKKENWKEIDELLKAGDFLPKQRTSANG  
 VIPHQMHOQELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPYVYVGPLVTP  
 20 EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKDTYLLNEDVLPANSL  
 LYQKYNVLNELNNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMAKTRGVNKPVSVE  
 GLSDPKKFNENLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVEWLT  
 DEQRSALVKKRYKGGWGRSLKLLTGIVDENGQRIIDLWNTDQNFMQIVNQPVFKEQIDQLN  
 QKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSSISIEFARNEG  
 25 NKGEITRSRRTQLQKLFEDQAHELVKDTSLTEELEKAPDLSDRYFYFTQGGKDMYTGDPIIN  
 FDEISTKYDIDHILPQS FVKDDSLDNRVLVSR AENKKS DRVPAKLYAAKMPYWNQLLKQG  
 LITQRKFENLTMVDVDTIKYRSLGFVQRQLVETRQVIKLTANILGSMYQEAGTDIIETRAGL  
 TKQLREEFDL PKVREVNDYHHAVDAYLTTFAGQYLNRRYPKLRSEFFVYGEYMKFKHGS DLKL  
 RNFNFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLISNETYEETGKYFD  
 30 ASHESSSLYLKNQNKSKLVVPLKNKLQPEYYGAYTGITQGYMVLKLLDKKGGFGVYRIPR  
 YAADILNKCHDEVAYRNKIAEIISSDPRAPKSFVVVPRVLKGTFLVDGEEKFILSSYRYKV  
 NATQLILPVSDIKLIQDNFKALKKLVEMQTKKLIETIYDNILRQVDKYYKLYDINKFRAKLH  
 DGRSKFVELDDFGQDASKEKVIKILRGLHFGSDLQNLKEIGFGTTPLGQFQVSEAGIRLSN



TAFII FKSPTGLFNRKLYLKNL GKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPY  
 DVPDYA (SEQ ID NO: 85).

In some embodiments, the *Peptoniphilus sp.* Marseille-P3761 Cas9 has an amino acid sequence at least 80% identical to

5 MPKKKRKVGEKKTNYTIGLDIGTDSVGVAVVKKDDLELVKKRMKVLGNTETNYIKKNLWGSLL  
 FESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDFENFFRNLNESFLVPEDKAFS  
 KNPIFGTLGEDKTYKYTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDT  
 EHIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDIGRA  
 NKQSPMMAFITLIVGKKAKFKAAFNLEEEIISLNLTDSDYDENLEILLNTIGSDFADLFDHAQ  
 10 RVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKD  
 AQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPN  
 QVHLAEMEAILDRQENYYPWLKENREKIIISLLTFRIPIYYVGPLADGQSEFAWLERKSDEKIK  
 PWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIAYLDERQKR  
 MNLSSVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKV  
 15 PGMKSMDDPDNEDLMEEIVKILTVFEDRKMRRKQLEKYKERLSPEQIKELAKKHGTGWGRLL  
 SKKLLVGIIRDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDDRLSFKKTI AELQMI DPS  
 ADLYAQVQEIAGSPAIIKKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTTARGKALSRRR  
 EAKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHII  
 PQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEVVRQLAYWRS LKDAGLMTQRKFDNLTKA  
 20 MRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNIVLLKSAMASE  
 FRKMFRLYKVRELNDYHHAHDAYLNAATAINLLALYPYMA DDFVYGEFRYKKKPQAEKATYE  
 KLRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNVVKKAEERTGMLFKETIN  
 GKTNKGSRIPIKKDLDP SKYGGYIEEKMAYYAVISYEDKKKPGKTIVGISIMDKKEFEYDS  
 ISYLGKLGFSNPVVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKANQIAMEQDLVNF  
 25 IYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIYEEKKKDP  
 VDIEALVASFIELLKLTSAGAPADFI FMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMR  
 FEDGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 87).

In some embodiments, the amino acid sequence of the Cas9 protein comprises at least  
 one, at least two, at least three, at least four, at least five, at least six, at least seven, at least  
 30 eight, at least nine, or at least 10 mutations in SEQ ID NOs: 1, 4, 8, 14, 84 or 86.

In some embodiments, the mutation is an amino acid substitution.

In some embodiments, the Cas9 protein has nickase activity.

In some embodiments, provided herein is a Cas9 protein wherein the Cas9 protein comprises a nickase mutation at an amino acid positions corresponds to one or more amino acids 10, 12, 17, 762, 840, 854, 863, 982, 983, 984, 986, 987 of wild type SpCas9.

5

In some embodiments, the at least one mutation results in an inactive Cas9 (dCas9).

In some embodiments, the Cas9 protein comprises at least one amino acid mutation in PAM Interacting, HNH and/or RuvC domain.

10 In some embodiments, provided herein is a Cas9 protein, wherein the mutation at an amino acid position corresponds to amino acid 14 in the RuvC domain of SirCas9.

In some embodiments, provided herein is a Cas9 protein, wherein the mutation at an amino acid position corresponds to amino acid 12 in the RuvC domain of EpeCas9.

In some embodiments, provided herein is a Cas9 protein, wherein the mutation at an amino acid position corresponds to amino acid 9 in the RuvC domain of LfeCas9.

15 In some embodiments, provided herein is a Cas9 protein, wherein the mutation at an amino acid position corresponds to amino acid 12 in the RuvC domain of PmaCas9.

In some embodiments, the Cas9 protein further comprises a nuclear localization sequence (NLS) and/or a FLAG, HIS or HA tag.

20 In one aspect, provided herein is an engineered, non-naturally occurring Cas9 fusion protein comprising a Cas9 protein having at least 80% identity to SEQ ID NOs: 1, 4, 8, 14, 84 or 86 and wherein the Cas9 protein is fused to a histone demethylase, a transcriptional activator, or to a deaminase.

25 In some embodiments, provided herein is an engineered, non-naturally occurring Cas9 fusion protein further comprising a nuclear localization sequence (NLS) and/or a FLAG, HIS or HA tag.

In some embodiments, provided herein is an engineered, non-naturally occurring Cas9 fusion protein having at least 80% identity to SEQ ID NOs: 2, 5, 9, 15, 85, 87, 95 or 96.

In some embodiments, the Cas9 protein is fused to a cytosine deaminase or to an adenosine deaminase.

In some embodiments, the Cas9 protein is fused to an adenosine deaminase and has an amino acid sequence at least 80% identical to

5 (a)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH  
DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA  
GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
TSESATPESSGPKKKRKGKPYSIGLAIGTNSVGVAVVTDDYKVPKAKMKVLGNTDKQSIK  
10 KNLLGALLFDSGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSF  
VDEDKRGEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHF  
LYEGDLKAENTDVQALFKDFVEEYDKTIEESHLSIEITVDALSILTEKVS KSSRLENLIAHYP  
TEKKNLTFGNLIALSLDLHPNFKTNFQLS EDAKLQFSKDTYEEDLEGFLGEVGD EYADLFAS  
AKNLYDAILLSGILTVDDNSTKAPLSASMVKRYEEHQDLKCLKDFIKVNPADQYNAIFKDK  
15 NKKGYASYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHL  
QEMHAILRRQGEHY PFLKENQDKIEKILTFRIPIYVGPLARKGSRFAWA EYKADEKITPWNF  
DDILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFD  
TNMKQEIFDHVFKENRKVTKDKLLNYLNKEFEFRIVNLTGLDKENKAFNSSLGTYHDLRKI  
LDKSFLLDDKANEKTIEDIIQTLTLFEDREMIRQLQKYSDI FTKAQLKKLERRHYTGWRLS  
20 YKLINGIRNKENKKTILDYLI DDGYANRNF MQLINDDALSFKEEIARAQIIDDVDDIANVVH  
DLPGPSPAIKKGILQSVKIVDELVKVMGHN PANII IEMARENQTTDKGRRNSQQRLKLLQDSL  
KNLDNPNVNIKNVENQQQLQNDRLFLYYIQNGKDMYTGETLDINNL SQYDIDHIIPQAFIKDNS  
LDNRVLTRS DKNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGLTEEDK  
AGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKS NLVSNFRKEFELYKV  
25 REINDYHHAHDAYLNAVVG NALLLKYPQLEPEFVYGEYPKYNSYRSRKSATEKFLFYSNILR  
FFKKEDIQTNEDEGEIAWNKEKH IKILRKVLSYPQVNI VKKTEEQTGGFSKESILPKGESDKL  
IPRKTKN SYWDPKKGFFDS PVVAYSILVFADVEK GSKKLRKVQDMVGITIMEKKRFEKNP  
VDFLEQRGYRNVRL EKI IKLPKYSLFELENKRRRL LASAKELQKGNELVIPQRFTTLLYHSY  
RIEKDYEP EHYVEKHKDEFKELLEYISVFSRKYVLADNNLT KIEMLF SKNKDAEVSSLAK  
30 SFISLLTFTAFGAPAAFNFFGENIDRKRYTSVTECLNATLIHQSI TGLYETRIDL SKLGEDG  
KRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 20);

(b)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLH  
DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA  
GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
5 TSESATPESSGPKKKRKGAKNKDIRYSIGLAIGTNSVGWAVMDEHYELLLKGNHMMWGSRL  
FDAAEPAATTRASRSIRRRYNKRERIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQK  
NLGNDYKDNYNLFIEKDFNDKTYDYKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGN  
FLKEGQSFQKVEDIEEKLDNTLKKFMSLNDLNDLFVDNDINSMITVLSKIYQRSKKADLL  
KIMNPTKEERAAYKEFTKALVGLKFNVSKMILAQEVKKDDKDIELDNVDYDSTVDGLQAE  
10 LGEYIEFIEMLHSINSWVELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPKYNE  
VFREDNPKLHNYLGYIKYPKNTPVEEFYEYIKRLLAKVDTGEAREILERIDLEKFMLKQNSR  
TNGSIPYQMQKDEMIQIIDNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKF  
EDKQKERILPWNVDQIVDIDATAEGFIERMQNTGTYPDKPVMKNSLTVSKFEVLNENLKI  
RINGKLIPVETKKELLSDLFMKNKTITDKKLDWLVTHQYYDTNEELKIEGYQKDLQFSTSL  
15 APWIDFTKIFGEINASNYQLIEKIIYDISIFEDKKILKRRLKKVYQLDDLLVDKILKLNVTG  
WSRLSEKLLTGIKSKNSKETILSILENSNMNMEIINDESLGFKQIIEESNKKDIEGPFYD  
EVKKLAGSPAIKRGIWQALLVVQEITKFMKHEPSHIYIEFAREEQEKVRTESRIAKLQKIYK  
DLNLQTKEDQLVYESLKKEDAKKKIDTDALYLYLQMGKSMYSGKPLDIDKLSTYHIDHILP  
RSLIKDDSLDNRVLVLPKENEWKLDSETVPFEIRNKMMGFQKLHENGLMSNKKFFSLIRTD  
20 FNEKDKKRFINRQLVETRQIIKNVAVIINDHYTNTNVTVRAELSHQFRERYKIYKNRDLND  
LHHAHDAYIACILGQFIHQNFGNMDVNMIYGQYKKNYKKDVQEHNNYGFILNSMNIHFND  
NSVIWDPSYIGKIKSCFCYKDVYVTKKLEQNDAKLFDLTI LPSDKNSENGVTKAKIPV NKYR  
KDVNKYGGFSGDAPIMLAIEADKGGKHVRQVIAFPLRLKNYNDEERIKFIEKEKNLKNVKIL  
TEVKKNLILINHQYFFITGTNELVNATQLKLSAKNTKNL FNLVDANKHNKLESIDDANFNE  
25 VIQELICKLQEPYISRYNSIGKEFEDSYEKINAVTKQDKLYIIIEYLIAIMSAKATQGYIKPE  
LAREIGTNGKNKGRIKSFITDLNKTTFISTSVTGLFSKKYKLGKRPAATKKAGQAKKKKGSY  
PYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 6);

(c)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLH  
30 DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA  
GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
TSESATPESSGPKKKRKGVSIIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLAIGTSSV  
GWAVTNKAYELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRLKLEELFADAM

AQVDPTFFMRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTD  
DIRKLFLAVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFCNSAISSIGQIL  
MEAGKTKSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDL  
FGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPGLTIS  
5 ESKVKA FNKHKDDLAILKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFY  
KYTKKIVEGLSDSKDKEYILSQIELQILLPLQRIKDNQVPIPYQLHLEELKAILAKCGPKFPF  
LNEVADGFSVAEKLIKMLEFRIPIYVGPLNTHHNVDNNGGFAWAVRKASGRVT PWNFDDKIDR  
EKSAAAFINKLNTNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVEHLIEAV  
FKQDHKKMTKNRIEQFLKDNQYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAE  
10 EIITDITIFGESKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAG  
DGTPETIIILMRNFSYNLMELLDGKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKR  
AVWQALRIVDEVAHIKKALPARI FVEVTRS NKNEKKKKDSRQKRLSDLYAAIKKDDVLLNGL  
NNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGRCAYTGEIIELSLLNTDNYDIDHIYPRSL  
TKDDSDNLVLCRKTANAQKSDAYPISEEIQKTQKPFWTFLKQQGLISERKYERLTRITPLT  
15 ADDLSGFIARQLVETNQSVKAATLLRRLYPGVVVFVKAENVTD FRHDNNFIKVRSLNHHH  
HAKDAYLNI VVG NVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKDGKAWDVKTS  
MDTVKMMDSNDVRVTKRLLLEQT GALADATIYKATVAGKAKDGAYIGMKT KSSVFADVSKYG  
GMTKIKNAYSIIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISIYG  
KLCINQLVKVNGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNNMRKDNKDLKAN  
20 VVSTRALDNKHTIEVRIVEEKNI EFFDYLVSKLKMPIYQKMKGNKAAELSEKGYGLFKKMSL  
EEQSIHLLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTI  
VGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 10);

(d)

MPKKKRKVSIIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLAIGTSSVGWAVTNKAYEL  
25 LKFRSHKMWGSRLFDEGESAVARRGFRSMRRLERRLKRLKLEELFADAMAQVDPTFFMRL  
RESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHH  
ILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFCNSAISSIGQILMEAGKTKSDKA  
KAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEEDL  
KKLQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPGLTISESKVKA FNKHK  
30 DDLAILKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLS  
DSKDKEYILSQIELQILLPLQRIKDNQVPIPYQLHLEELKAILAKCGPKFPFLNEVADGFSVA  
EKLIKMLEFRIPIYVGPLNTHHNVDNNGGFAWAVRKASGRVT PWNFDDKIDREKSAAAFINKL  
TNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVEHLIEAVFKQDHKKMTKN

RIEQFLKDNNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGE  
 SKKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGTPETI I ILM  
 RNFSYNLMELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVDE  
 VAHIKKALPARI FVEVTRS NKNEKKKDSRQRLSDLYAAIKKDDVLLNGLNNEIFGELKSS  
 5 LAKYDDAALRSKKLYLYYTQMGRCAYTGEI IELSLLNTDNYDIDHIYPRSLTKDDSFDNLVL  
 CKRTANAQKSDAYPISEEIQKTQKPFWTF LKQQGLISERKYERLTRITPLTADDLSGFIARQ  
 LVETNQSVKAATTLRRLYPGVDVVFVKAENVTD FRHDNNFIKVRSLNHHHHAKDAYLNIVV  
 GNVYHERFTRNFRAF FKKNGANRTYNLAKMFNYDVNCTNAKD GKAWDVKT SMDTVKKMMSDN  
 DVRVTKRLL EQT GALADATIYKATVAGKAKDGAYIGMKT KSSVFADVSKYGGMTKIKNAYS I  
 10 IVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISIIYGKLCINQLVKVN  
 GFYYLGGKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNMRKDNKDLKANVVSTRALDNKH  
 TIEVRIVEEKNI EFFDYLVSKLMP IYQMKGNKAAELSEKGYGLFKKMSLEEQSIHLIELL  
 NLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIVKRPAATKKAG  
 QAKKKKSGSETPGTSESATPESSGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNR  
 15 VIGEGWNRAIGLHDPTAHAEIMALRQGGLVMQNYRLYDATLYVT FEPCVMCAGAMIHSRIGR  
 VVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQ  
 SSTDPAAKRVKLDGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 11);

(e)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH  
 20 DPTAHAEIMALRQGGLVMQNYRLYDATLYVT FEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA  
 GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
 TSESATPESSGPKKKRKGTKVKDYIIGLAIGTSSVGWAVTDEAYNVLFKNSKKMWGVRLFD  
 DAKTAEERRGQRGARRRLDRKKERLSLLQDFFAEVAKVDPNFFLRLDNSDLYMEDKDQKLLK  
 SKYTLFNDKDFKDNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIFEGQK  
 25 FDTKSSFENSLNELKVHLNDEYGLDLEFDNENLINILTPKLNKTAKKELKSVIGDTKFLK  
 AVSAIMIGSSQKLVDLFENPEDFDDSAIKSVDFSTTSFDDKYSYELALGDKIALVNILKEI  
 YDSSILENLLKEADKSKDGNKYISNAFVKYKXKHGQDLKEFKRLVRQYHKSAYFDIFRSEKV  
 NDNYVSYTKSSI SNNKRVKANKFTDQEA FYKFAKKHLETIKYKINKVNGSKADLELIDGMLR  
 DMEFKNFMPKIKSSDNGV I PYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFR  
 30 IPYYVGPLNPN SKYAWIKKQKDSEITPWNFKDVVDLDS SREEFIDSLIGRCTYLKDEKVLPK  
 ASLLYNEYMVLNELLNKLNDLPITEEMKKKIFDQLFKTRKKVT LKAVANLLKKEFNINGEI  
 LLSGTDGDFKQGLNSYNDFKAI VGDKVDSDDYRDKIEEIKLIVLYGDDKSYLQKKIKAGYG  
 KYFTDSEIKKMAGLNYKDWGRLSKLLTGLEGANKITGERGSI IHFMREYNLNLMELEMSASF

TFTEEIQKLN PVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIF IEMAR  
 GKEEVKARKESRKNQLLKFKYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQ  
 LGRCMYSLEPIDISELSSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPI PDEI  
 LNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTI  
 5 CKNSEIVYSKAENASRFRQEFDIVKCRVNDLHHMHDAYINIIVGNVYNTKFTKDP MNFVKK  
 QEKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESG  
 ALFNATLQRKNKGSRPLKDKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLM PVEREI  
 YAKQKNDKCLSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSY YFITGGSDKTLELRS  
 ALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSF LNL  
 10 FQDDKIENIDFKFKSMD FKEKCKTLLMLVKAIRASGVRQDLKSIDLKS DYGR LSSKTNIGN  
 YQEFKIINQSITGLFENEVDLLKLGKRPAA TKKAGQAKKKKGSYPYDVPDYAYPYDVPDYA  
 PYDVPDYA (SEQ ID NO: 16);

(f)

MPKKKRKVTKVKDYYIGLAI GTSSVGWAVTDEAYNVLFNSKMMWGVRLFDDAKTAEERRGQ  
 15 RGARRRLDRKKERLSLLQDFFAEEVAKVDPNFFLRDLNSDLYMEDKDQKLKSKYTLFNDKDF  
 KDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIFEGQKFDTKSSFENSL  
 NELKVHLNDEYGLDLEFDNENLINILTPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQ  
 KLVDLFENPEDFDDSAIKSVDFSTTSFDDKYS DYELALGDKIALVNILKEIYDSSILENLLK  
 EADKSKDGNKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEK VNDNYVSYTKSS  
 20 ISNNKRVKANKFTDQEA FYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKI  
 KSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRI PYVYVGPLNPN  
 SKYAWIKKQKDSEITPWNFKDVVDLDS SREEFIDSLIGRCTYLKDEKVL PKASLLYNEYMVL  
 NELNNLKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGDFKQ  
 GLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGYFTDSEIKKM  
 25 AGLNYKDWGR LSKKLLTGLEGANKITGERGSI IHFMREYNLNLME LMSASFTFTEEIQKLN  
 PVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIF IEMARGKEEVKARKES  
 RKNQLLKFKYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQLGRCMYSLEPI  
 DISELSSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPI PDEILNKNCYAYWKI  
 LYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTICKNSEIVYSKA  
 30 ENASRFRQEFDIVKCRVNDLHHMHDAYINIIVGNVYNTKFTKDP MNFVKKQEKARSYNLEN  
 MFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRKN  
 KGSRPLKDKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLM PVEREIYAKQKNDKCLS  
 DEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSY YFITGGSDKTLELRSALQLILPKKNE

WAIKQIDKSSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLFQDDKIENIDF  
 KFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKS DYGR LSSKTNNIGNYQEFKIINQSI  
 TGLFENEVDLLKLRPAATKKAGQAKKKKSGSETPGTSESATPESGSEVEFSHEYWMRHAL  
 TLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMALRQGGLVMQNYRLYDA  
 5 TLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADE  
 CAALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPYDVPDYAYPYDVPDYAYPYDVP  
 DYA (SEQ ID NO: 17);

(g)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLH  
 10 DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA  
 GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
 TSESATPESGPKKKRKGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKKGKTAIGVRLFEEGK  
 TAAERTRFRTRRRLKRRKWRHLHYLDEIFAPHLQEVDFLRRLKQSNIHPEPKAKNQAFIG  
 KLLFPDLLKKNRGYPTLIKMRDEL PVEQRAHY PVTNIYKLREAMINEDRQFDLREVYLAVH  
 15 HIVKYRGHFLNNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLD  
 TKMRKLDKQAVAKLLEVKVADKEETKRNKQIATAMSKLVLGYKADFATVAMANGNEWKIDL  
 SSETSEDEIEKFREEELSDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWTHERQLA  
 EVKEYLATQPASARKEFDQVYNKYIGQAPKEKGFLEKGLKKILSKKENWKEIDELLKAGDF  
 LPKQRTSANGVI PHQM HQE LDR IIEKQAKYYPWLATENPATGERDRHQAKYELDQLVS FRI  
 20 PYYVGPLVTPVQKATSGAKFAWAKRKEEDGEITPWNLWDKIDRAESAEAFIKRMTVKDITYLL  
 NEDVLPANSLLYQKYNVLNELNRRVNGRRLSVGIKQDIYTELFKKT TVKAGDVASLVMK  
 TRGVNKP SVEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFA  
 DKLTEVEWLTDEQRSALVKKRYKGGWRLSKLLTGIVDENGQRIIDL MWNTDQNFMQIVNQF  
 VFKEQIDQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKS  
 25 ISIEFARNEGKGEITRSRRTQLQKLFEDQAHVELVKDTSLTEELEKAPDLSDRYFYFTQGG  
 KDMYTGDPINFDEISTKYDIDHILPQS FVKDDSLDNRVLVSRAENNKKS DRVPAKLYAAKMK  
 PYWNQLLKQGLITQRKFENLTMDVDQTIKYRSLGFVQRQLVETRQVIKLTANILGSMYQEAG  
 TDIIETRAGLTKQLREEFDLPKVREVNDYHHAVDAYLTTFAGQYLNRYPKLSRFFVYGEYM  
 KFKHGS DLKLRNFNF FHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWINLKVMLISNE  
 30 TYEETGKYFDASHES SLYLKNQNKSKLVVPLKNKLQPEYYGAYTGITQGYMVILKLLDKK  
 GFGVYRIPRYAADI LNKCHDEVAYRNKIAEIISSDPRAPKSFVVVPRVLKGTFLVDGEEK  
 FILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLVNEMQTKKLEIYDNI LRQVDKYKLY  
 DINKFRAKLHDGRSKFVELDDFGQDASKEKVIKILRGLHFGSDLQNLKEIGFGTTPLGQFQ



VSEAGIRLSNTAFII FKSPTGLFNRKLYLKNLGRPAATKKAGQAKKKKGSYPYDVPDYAYP  
YDVPDYAYPYDVPDYA (SEQ ID NO: 88);

(h)

MPKKKRKVGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAAERRTFRT  
5 TRRRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEDEPAKNQAFIGKLLFPDLLKK  
NERGYPTLIKMRDELVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAVHHIVKYRGHFL  
NNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLDROK  
AVAKLLEVKVADKEETKRNKQIATAMSKLVLGYKADFATVAMANGNEWKIDLSSETSEDEIE  
KFREEELSDAQN DILTEITSLFSQIMLNEIVPNGMSISESMMDRYWOTHERQLAEVKEYLATQP  
10 ASARKEFDQVYNKYIGQAPKEKGFLEKGLKILSKKENWKEIDELLKAGDFLPKQRTSANG  
VI PHQM HQE LDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPYVVGPLVTP  
EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKDTYLLNEDVLPANSL  
LYQKYNVLNELNNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMKTRGVNKP SVE  
GLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVEWLT  
15 DEQRSALVKKRYKGWGRLSKLLTGIVDENGQRIIDL MWNTDQNF MQIVNQPVFKEQIDQLN  
QKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVEDIVKAVGNAPKSI SIEFARNEG  
NKGEITRSRRTQLQKLFEDQAHELVKDTSLTEELEKAPDLSDRYFYFTQGGKDMYTGD PIN  
FDEISTKYDIDHILPQS FVKDDSLDNRVLVSR AENNKKS DRVPAKLYAAKMKPYWNQLLKQG  
LITQRKFENLTMDVDQTIKYRSLGFVKRQLVETRQVIKLTANILGSMYQEAGTDIIETRAGL  
20 TKQLREEFDLPKRVENDYHHAVDAYLTTFAGQYLNRRYPKLRSFFVYGEYMKFKHGS DLKL  
RNFNFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLISNETYEETGKYFD  
ASHESSLYLKNQNKSKLVVPLKNKLQPEYYGAYTGITQGYMVILKLLDKKGGFGVYRIPR  
YAADILNKCHDEVAYRNKIAEIISSDPRAPKSFVVVPRVLKGTFLVDGEEKFILSSYRYKV  
NATQLILPVSDIKLIQDNFKALKKLNEMQTKKLIEIYDNILRQVDKYYKLYDINKFRAKLH  
25 DGRSKFVELDDFGQDASKEKVIKILRGLHFGSDLQNLKEIGFGTTP LGQFQVSEAGIRLSN  
TAFII FKSPTGLFNRKLYLKNLGRPAATKKAGQAKKKKSGSETPGTSESATPESGSEVEFS  
HEYWMRHALTLAKRARDEREVPGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMALRQGGLV  
MQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVE  
ITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSPYDVPDYAYPYDVP  
30 DYAYPYDVPDYA (SEQ ID NO: 89);

(i)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH  
DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA  
GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSS TDGSSGSETPG  
TSESATPESSGPKKKRKVGEKKTNYTIGLAIGTDSVGWAVVKDDLELVKKRMKVLGNTETNY  
5 IKKNLWGSLLFESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFFRLNES  
FLVPEDKAFSKNPIFGTLGEDKTYKYTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRG  
HFLIEGKLDTEHIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRI  
TSFLKDIGRANKQSPMMAFITLIVGKKAKFKA AFNLEEEISLNLTDSDYDENLEILLNTIGS  
DFADLFDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQ  
10 YDMTFVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKO  
RSFYNGVIPNQVHLAEMEAILDRQENYYPWLKENREKII SLLTFRI PYYVGPLADGQSEFAW  
LERKSDEKIKPWNFS DVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTK  
IAYLDERQKRMNLS SVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNAD  
YSTYVELAKVPGMKSMMDDPDNEDLMEEIVKILTVFEDRKMRKQLEKYKERLSPEQIKELA  
15 KKHYTGWGRLSKLLVGI RDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDDRLSFKKT  
IAELQ MIDPSADLYAQVQEIAGSPA I KKGILLGLKIVDEIIRVMGEKPENIV IEMARENQTT  
ARGKALSKRREAKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDR  
LSQYDEDHII PQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEV VNRQLAYWRS LK DAGLMT  
QRKFDNLTKAMRGG LTKDRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNI  
20 VLLKSAMASEFRKMFRLYK VRELNDYHHAHDAYLNAAIAINLLALYPYMA DDFVYGEFRYKK  
KPQAEKATYEKL RQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNVVKKAEER  
TGMLFKETINGKTNKGSRIPIK KDLDP SKYGGYIEEKMAYYAVISYEDKKKPGKTIVGISI  
MDKKEFEYDSISYLGKLGFSNPVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKANQ  
IAMEQDLVNF IYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRR  
25 IYEEEEKKDPVDIEALVASFIELLKLTSAGAPADFI FMGEAISRRRYNSMTGLFDGQVIYQS  
LTGLYETRMRFEDGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA

(SEQ ID NO: 91);

(j)

MPKKKRKVEKKTNYTIGLAIGTDSVGWAVVKDDLELVKKRMKVLGNTETNYIKKNLWGSLLF  
30 ESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFFRLNESFLVPEDKAFSK  
NPIFGTLGEDKTYKYTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTE  
HIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDIGRAN  
KQSPMMAFITLIVGKKAKFKA AFNLEEEISLNLTDSDYDENLEILLNTIGSDFADLFDHAQR

VYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDA  
 QKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQ  
 VHLAEMEAILDRQENYYPWLKENREKIISLLTFRIPIYYVGPLADGQSEFAWLERKSDEKIKP  
 WNFSDVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIAYLDERQKRM  
 5 NLSSVEKKEIFETLFFKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVP  
 GMKSMMDDPDNEDLMEEIVKILTVFEDRKMRRKQLEKYKERLSPEQIKELAKKHYTGWGRLS  
 KLLLVGIRDKETQKTILDYLVEDDNHSGGRQHNLNRNLMQLINDDRLSFKKTTAELQ MIDPSA  
 DLYAQVQEIAGSPAIIKKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTTARGKALS KRRE  
 AKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHIIP  
 10 QSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEVVRQLAYWRS LKDAGLMTQRKFDNLT KAM  
 RGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNIVLLKSAMASEF  
 RKMFRLYKVRELNDYHHAHDAYLNAAIAINLLALYPYMA DDFVYGEFRYKKKPQAEKATYEK  
 LRQWNLIKRFGEKQLFTP DHEDCWNKERDIKTIKKVMGYRQVNVVKKAEERTGMLFKETING  
 KTNKGSRIPIKKDLDP SKYGGYIEEKMAYYAVISYEDKKKKPGKTIVGISIMDKKEFEYDSI  
 15 SYLGLGFSNPVVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKANQIAMEQDLVNF I  
 YHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIYEEEEKKKDPV  
 DIEALVASFIELLKLTSAGAPADFI FMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMRF  
 EDKRPAATKKAGQAKKKKSGSETPGTSESATPES SSGSEVEFSHEYWMRHALTLAKRARDERE  
 VPGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMALRQGG LVMQNYRLYDATLYVTFEPCVM  
 20 CAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRM  
 PRRVFNAQKKAQSSTDPAAKRVKLDGSPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID  
 NO: 92).

In some embodiments, the Cas9 protein is fused to a cytosine deaminase and has an amino acid sequence at least 80% identical to

25 (a)  
 MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDV FYDPREL RKETCLLYEIKWGMSRKI WRS  
 SGKNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLV IY  
 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQY PPLWMM L  
 YALELHCIILSLPPCLKISRWRQNHLAFFRLHLQ NCHYQTIPPHILLATGLIHPSVTWRLKS  
 30 GGSSGGSSGSETPGTSESATPES SGGSSGGSPKKKRKVGKPY SIGLAIGTNSVGVAVVTDD  
 YKVPAKKMKVLGNTDKQSIKKNLLGALLFDSGETAEATRLKRTARRRYTRRKNRLRYLQEIF  
 TGEMNKVDENFFQRLDDSFLVDEDKRGEHHP IFGNIAAEVKYHDDFPTIYHLRRHLADTSKK  
 ADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFKDFVEEYDKTIEESHLS EITVDALS

ILTEKVS KSSRLENLIAHYPT EKKN TLFGNLI ALSLDLHPNF KTNFQ LSEDAKLQFSKDTYE  
 EDLEGFLGEV GDEYADLFASAKNLYDA ILLSGILTVDDNSTKAPLSASMVKRYEEHQDLKK  
 LKDFIKVNAPDQYNAIFKDKNKKG YASYIESGVKQDEFYKYLKGI LLKINGS GDFLDKI DRE  
 DFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHY PFLKENQDKIEKILTFRIPYVVGPLARK  
 5 GSRFAWA EYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTV  
 YNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTKDKLLNYLNKEFEEFRIVNLTGL  
 DKENKAFNSSLGTYHDLRKI LDKSFLDDKAN EKTI EDIIQTTLTFEDREMIRQLQKYS DIF  
 TKAQLKKLERRHYTGWGRLSYKLINGIRNKENKKTILDYLI DDGYANRNF MQLINDDALSFK  
 EEIARAQIIDDVDDIANVVHDLPGSPA I KKGILQSVKIVDELVKVMGHN PANII IEMARENQ  
 10 TTDKGRRNSQORLKL LQDSLKNLDNPVNIKNVENQQ LQNDRLFLYI IQNGKDMYTGETLDIN  
 NLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSSDDVPSIEVVHEMKS FWSKLLSVKLIT  
 QRKFDNLTKAERGG LTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRI RNVKI  
 ITLKS NLVSNFRKEFELYK VREINDYHHAH DAYLNAVVG NALLLKY PQLEPEFVYGEY PKYN  
 SYRSRKSATEKFLFY SNILRFFKKEDIQT NEDGEI AWNKEKH I KILRKVLSYPQVNIVK KTE  
 15 EQTGGFSKESILPKGESDKLIPRKT KNSYWDPKKYGGFDS PVVAYSILVFADVEKGKSKKLR  
 KVQDMVGITIMEKKRFEKNPVDFLEQRGYRNVRL EKI I KLPKYS LFELENKRRRLLASAKEL  
 QKGNELVIPQRFTTLLYHSYRIEKDYEP EHYVEKH KDEFKELLE YISVFSRKYVLADNNL  
 TKIEMLFSKNKDAEVSSLAKSFISLLTFTA FGA PAAFNFFGENIDRKRYTSVTECLNATLIH  
 QSITGLYETRIDL SKLGEDGKRPAATK KAGQAKKKKGSSGGSGGSGGSTNLSDI IEKETGKQ  
 20 LVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVM L L TSDAPEYKP WALVIQDSNG  
 ENKIKMLSGGSGGSGGSTNLSDI IEKETGKQLVIQESILMLPEEVEEVIIGNKPESDILVHTA  
 YDESTDENVM L L TSDAPEYKP WALVIQDSNGENKIKMLYPYDVPDYAYPYDVPDYAYPYDVP  
 DYA (SEQ ID NO: 21);

(b)

25 MPAARVKLDTSEKGPSTGDP TLRRIESWEFDV FYDPREL RKETCLLYEIKWGMSRKI WRS  
 SGKNTTNHVEVNF I KKF TSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVT LVIY  
 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEY YHCWRNFVNYPPGDEAHWPQY PPLWMLL  
 YALELHCII LSLPCLKISR RWQNHLA FFRHLQ NCHYQTIPPHILLATGLIHPSVTWRLKS  
 GGSSGGSSGSETPGTSESATP ESSGGSSGGS P KKKR K VGS I I N F Q R R G L M E T Q A S N Q L I S S H  
 30 LKGYPIKDYFVGLAIGTSSV GWA VTNKAYELLKFRSHKMWGSRLFDEGESAVARRGFRSMRR  
 RLERRKLRLK LLEELFADAMAQVDPTFFMRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDY  
 FKEYPTVYHLRSEL M KSGTDDIRKLFLAVHHILKYRGNFLYEGATFDSNASTLDDVIKQALE  
 NITFNCFCDCNSAISSIGQILMEAGKTKSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDK

KRLKAFANLVLGLNASLIDLFGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDDC  
 KSVYDAI ILLSIKEPGLTISESKVKAFNKHKDDLAILKSLLKSDRSIYNTMFKVDEKGLHNY  
 VHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEYIILSQIELQILLPLQRIKDNQVPIY  
 QLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKMLEFRIPIYVGPLNTHHNVDNNGFAW  
 5 AVRKASGRVT PWNFDDKIDREKSAAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNN  
 VRIDGKPLEKVVKEHLIEAVFKQDHKKMTKNRIEQFLKDNGYISETHKHEITGLDGEIKNDL  
 ASYRDMVRI LGDGFDRSMAEEIITDITIFGESKMLRET LRKKFASCLDDEAIKKLTKLRYR  
 DWGRLSQKLLNGIEGCDKAGDGPETIIILMRNFSYNLMELLGDKFSFMERIQEINAKLTEG  
 QIVNPHDIIDDLALSPAVKRAVWQALRIVDEVAHIKKALPARI FVEVTRSNKNEKKKKDSRQ  
 10 KRLSDLYAAIKKDDVLLNGLNNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGRCAYTGEII  
 ELSLLNTDNYDIDHIYPRSLTKDSDFDNLVLCRRTANAQKSDAYPISEEIQKTQKPFWTFLK  
 QQGLISERKYERLTRITPLTADDLSGFARQLVETNQSVKAATTLRRLYPGVDVVFVKAEN  
 VTDFRHDNNFIKVRSLNHHHHAKDAYLNIIVGNVYHERFTRNFRAFFKKNGANRTYNLAKMF  
 NYDVNCTNAKDGKAWDVKTSMDTVKMMDSNDVRVTKRLLLEQTGALADATIYKATVAGKAKD  
 15 GAYIGMKTKSSVFADVSKYGGMTKIKNAYSIIVQYTGKKGEVIKEIVPLPIYL TNRNTTDQD  
 LINYVASIIPQAKDISIIYGKLCINQLVKVNGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPY  
 LKVLEKFNNMRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNI EFFDYLVSKLKMPIYQKMK  
 GNKAAELSEKGYGLFKKMSLEEQS IHLIELLNLNLTNQKTTFEVKPLGITASRSTVSGKISNQ  
 DEFKVINESITGLYSNEVTIVGKRPAATKKAGQAKKKKGGSSGGSGGSGSTNLSDIEKETG  
 20 KQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTS DAPEYKPWALVIQDS  
 NGENKIKMLSGGSGGSGGSGSTNLSDIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVH  
 TAYDESTDENVMLLTS DAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAYPYDVPDYAY

(SEQ ID NO: 12);

(c)

25 MPAARVKLDTSEKGPSTGDPTLRRRIESWEFDVFDPRELRKETCLLYEIKWGMSRKIWRS  
 SGKNTTNHVEVNFIIKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIIY  
 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEY YHCWRNFVNYPPGDEAHWPQYPPLWMLL  
 YALELHCII LSLPCLKISRWRQNHLAFFRLHLQNCHYQTIPPHILLATGLIHPSVTWRLKS  
 GGSSGGSSGSETPGTSESATPESSGGSSGSPKKKRKVGTKVKDYIIGLAIGTSSVGWAVTD  
 30 EAYNVLFKFNKMMWGVRLFDDAKTAEERRGQRGARRRLDRKKERLSLLQDFFAEVAVKVDPN  
 FFLRLDNSDLYMEDKDQKLKSKYTLFNDKDFKDKNFHKKYPTIHLLMDLIEDDSKKDIRLV  
 YLACHYLLKNRGHFI FEGQKFDTKSSFENSLNELKVHLNDEYGLDLEFDNENLINILTDPKL  
 NKTAKKKELKSVIGDTKFLKAVSAIMIGSSQKLVLDLFENPEDFDDSAIKSVDFSTTSFDDKY

SDYELALGDKIALVNI LKEIYDSSILENLLKEADKSKDGNKYISNAFVKKYNKHGQDLKEFK  
 RLVRQYHKSAYFDIFRSEKVNNDYVSYTKSSISNNKRVKANKFTDQEAFYKFAKKHLETIKY  
 KINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNGVIPPYQLKLMELNKILENQS KHHEFL  
 NVSDEYGSVCDKIASIMEFRIPIYVGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLDSREE  
 5 FIDSLIGRCTYLKDEKVLPKASLLYNEYMVLNELNNLKLNDLPITEEMKKKIFDQLFKTRKK  
 VTLKAVANLLKKEFNINGEILLSGTDGDFKQGLNSYNDFKAIVGDKVDSDDYRDKIEEIIKL  
 IVLYGDDKSYLQKKIKAGYGKYFTDSEIKKMAGLNYKDWGRLSKKLLTGLEGANKITGERGS  
 I IHFMREYNLNLMELEMSASFTFTEEIQKLN PVDDRKLSYEMVDELYLSPSVKRMLWQSLRIV  
 DEIKNIMGTDSKKIFIE MARGKEEVKARKE SRKNQLLFYKDGKKA FISEIGEERYSYLLSE  
 10 IEGEEENKFRWDNLYLYTQLGRCMYSLEPIDISELSSKNIYDQDHIYPKSKIYDSSIENRV  
 LVKKDLNSKKGNSYPI PDEILNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDELVQFIS  
 RQIVETRQATKETANLLKTI CKNSEIVYSKAENASRFRQEFDIVKCRVNDLHHMHDAYINI  
 IVGNVYNTKFTKDP MNFVKKQEKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRI  
 RKELEGTNYRFTRMNYESGALFNATLQRKNKGSRPLKDKGPKSSIEKYGGYTNINKACFAV  
 15 LDIKSKNKIERKLPVEREIIYAKQKNDKKSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLK  
 IDGSYYFITGGSDKTLELRSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNT  
 FDIIVNKFKTSVFKKSFNLNFQDDKIENIDFKFKSMDFKCKTLLMLVKAIRASGVRQDLK  
 SIDLKS DYGRLLSSKTNIGNYQEFKIINQSITGLFENEVDLLKLGKRPAAATKAGQAKKKKG  
 SSGSGSGSGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVI GNKPESDILVHTAYDESTD  
 20 ENVMLLTS DAPEYKPWALVIQDSNGENKIKMLSGSGSGSGSTNLSDIIEKETGKQLVIQES  
 ILMMLPEEVEEVI GNKPESDILVHTAYDESTDENVM L L T S D A P E Y K P W A L V I Q D S N G E N K I K M  
 LYPYDVPDYAYPYDVPDYAY (SEQ ID NO: 18);

(d)

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR  
 25 SGKNTTNHVEVNFIFKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIIY  
 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEY YHCWRNFVNYPPGDEAHWPQYPPLWMLL  
 YALELHCCIILSLPPCLKISRWRQNH LAFFRLHLQNC HYQTIPPHILLATGLIHPSVTWRLKS  
 GGSSGSSGSETPGTSESATPESGGSSGSPKKKRKVGKEYHIGLAIGTSSIGWAVTDSQF  
 KLMRIKGKTAIGVRLFEEGKTAAERRTFRTRRRRLKRRKWRLHYLDEIFAPHLQEVDENFLR  
 30 RLKQSNIHPE DPAKNQAFIGKLLFPDLLKKNERYPTLIKMRDEL PVEQRAHY PVTNIYKLR  
 EAMINEDRQFDLREVYLAVHHIVKYRGHFLNNASVDKFKVGRIDFKSFNVLNEAYEELQNG  
 EGSFTIEPSKVEKIGQLLLDTKMRKLD RQKAVAKLLEVKVADKEETKRNKQIATAMSKLVLG  
 YKADFATVAMANGNEWKIDLSSETSEDEIEKFREELSDAQNDILTEITSLSQIMLNEIVPN

GMSISESMMDRYWTHERQLAEVKEYLATQPASARKEFDQVYNKYIGQAPKEKGFDFLEKGLKK  
 ILSKKENWKEIDELLKAGDFLPKQRTSANGVI PHQMHOQELDRIIEKQAKYYPWLATENPAT  
 GERDRHQAKYELDQLVSFRIPYVYVGPLVTPEVQKATSGAKFAWAKRKEDGEITPWNLWDKID  
 RAESAEAFIKRMTVKDITYLLNEDVLPANSLLYQKYNVLNELNVRVNGRRLSVGIKQDIYTE  
 5 LFKKKKTVKAGDVASLVMAKTRGVNKPSVEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQ  
 MDLENIIEWRSVFEDGEIFADKLTEVEWLTDEQRSALVKKRYKGGWGRSLSKLLTGIVDENGQ  
 RIIDLWNTDQNFMQIVNQPVFKEQIDQLNQAITNDGMTLRERVESVLDDAYTSPQNKKAI  
 WQVVRVVEDIVKAVGNAPKSI SIEFARNEGKGEITRSRRTQLQKLFEDQAHELVKDTSLTE  
 ELEKAPDLSDRYYFYFTQGGKDMYTGDPI NFEI STKYDIDHILPQS FVKDDS LDNRVLVSR  
 10 AENNKKSDRVPKLYAAKMKPYWNQLLKQGLITQRKFENLTMDVDQTIKYRSLGFVVKRQLVE  
 TRQVIKLTANILGSMYQEAGTDIIETRAGLTKQLREEFDLPKVREVNDYHHAVDAYLTTFAG  
 QYLNRRYPKLRSEFFVYGEYMKFKHGS DLKLRNFNFHELMEGDKSQGKVVDQQTGELITTRD  
 EVADYFDWVINLKVMLISNETYEETGKYFDASHES SLYLKNQNKSKLVVPLKNKLQPEYY  
 GAYTGITQGYMILKLLDKKGGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKS  
 15 FEVVVPRVLKGTFLVDGEEKFILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLNVMQTK  
 KLIEIYDNILRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEKVI I KILRGLHFG  
 SDLQNLKEIGFGTTP LGQFQVSEAGIRLSNTAFIIFKSP TGLFNRKLYLKNLGRPAATKKA  
 GQAKKKKGS SGGSGGSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVH  
 TAYDESTDENVMLLTS DAPEYKPWALVIQDSNGENKIKMLSGGSGGSGGSTNLSDIIEKETG  
 20 KQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTS DAPEYKPWALVIQDS  
 NGENKIKMLYPYDVPDYAYPYDVPDYAY (SEQ ID NO: 90);

(e)

MPAARVKLDTSEKGPSTGDPTLRRRIESWEFDVFDYDPRELRKETCLLYEIKWGMSRKIWR  
 SGKNTTNHVEVNFIIKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIIY  
 25 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYPPLWMMML  
 YALELHCCIILSLPPCLKISRRWQNHLAFFRLHLQNCYQTI PPHILLATGLIHPSVTWRLKS  
 GGSSGGSSGSETPGTSESATPESSGGSSGSPKKKRKVGEKKTNYTIGLAIGTDSVGWAVVK  
 DDLELVKKRMKVLGNTETNYIKKNLWGSLLFESGQTAKDRRLKRVARRRYERRRNRLTELQK  
 IFAPAIDEVDENFFFRLNESFLVPEDKAFSKNPIFGTLGEDKTYKYTYPTIYHLRQHLADSE  
 30 EKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIAINENLEQFFESYNALFSEPIELRKEEL  
 IAIENILREKNSRTVKEKRITSFLKDIGRANKQSPMAFITLIVGKKAKFKAAFNLEEEISL  
 NLTDDSYDENLEILLNTIGSDFADLFDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYE  
 RHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKDLKGYAGYIDGNMSQDSFYKFVKDQLKE

VPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVHLAEMEAILDRQENYYPWLKENREKIISLL  
 TFRIPYYVGPLADGQSEFAWLERKSDEKIKPWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEY  
 VLPKKS LIYQKYMVFNELTKIAYLDERQKRMNLS SVEKKEIFETLFKKRSKVTEKQLVKFFE  
 NYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMDDPDNEDLMEEIVKILT VFEDRKMR  
 5 RKQLEKYKERLSPEQIKELAKKHGTGWGRLSKLLV GIRDKETQKTILDYLVEDDNHSGGRQ  
 HLNRLMQLINDDRLSFKKTI AELQ MIDPSADLYAQVQEIAGSPA I KKGILLGLKIVDEIIR  
 VMGEKPENIVIEMARENQTTARGKALSKRREAKIKEGLAALGSSLLKENLPGNADLSQRKIY  
 LYQTONGKDIYLDEPLDFDRLS QYDEDHII PQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSL  
 EVVNRQLAYWRS LK DAGLMTQRKFDNLT KAMRGGLTDKDRERFIQRQLVETRQITKNVAKLL  
 10 DMRLNDKKDEAGNKIRETNIVLLKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAAIAINL  
 LALYPYMA DDFVYGEFRYKKKPKAEKATYEKLRQWNLIKRFGEKQLFTP DHEDCWNKERDIK  
 TIKKVMGYRQVNVVKA EERTGMLFKETINGKTNKGSRIPIKKDLDP SKYGGYIEEKMAYYA  
 VISYEDKKKPGKTIVGISIMDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIAYPDGRRR  
 YITGATKTTKGKVELQKANQIAMEQDLVNFYHLKNYDEISHPESYAFVQSHTDYFDRLFDS  
 15 IEHYTRRFLDAETNINRLRRIYEEKKKDPVDIEALVASFIELLKLTSAGAPADFI FMGEAI  
 SRRRYNSMTGLFDGQVIYQSLTGLYETRMRFEDGKRPAATKKAGQAKKKKSGSGSGSGGS  
 TNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAP  
 EYKPWALVIQDSNGENKIKMLSGSGSGSGSTNLSDIIEKETGKQLVIQESILMLPEEVEEV  
 IGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAY  
 20 PYDVPDYAY (SEQ ID NO: 93); or

(f)

MPAKRVKLDTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDE  
 NVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGSGSGSGSTNLSDIIEKETGKQLVIQESI  
 LMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKML  
 25 SGGSGSGSGSPKKRKRVEKKTNYTIGLAIGTDSVGWAVVKDDLELVKKRMKVLGNTETNYIK  
 KNLWGSLLFESGQTAKDRRLKRVARRRYERRRNLTELQKIFAPAIDEV DENFFFRLNESFL  
 VPEDKAFSKNPIFGTLGEDKTYKYPTIYHLRQH LADSEEKADVRLIYLALAHMIKYRGHF  
 LIEGKLDTEHIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITS  
 FLKDIGRANKQSPMMAFITLIVGKKAKFKAAFNLEEEISLNLTDSDYDENLEILLNTIGSDF  
 30 ADLFDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYD  
 MTFVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRS  
 FYNGVIPNQVHLAEMEAILDRQENYYPWLKENREKIISLLTFRIPYYVGPLADGQSEFAWLE  
 RKSDEKIKPWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIA



YLDERQKRMNLS SVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYS  
 TYVELAKVPGMKSMDDPDNEDLMEEIVKILTVFEDRKMRRKQLEKYKERLSPEQIKELAKK  
 HYTGWGRLSKLLVGI RDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDRRLSFKKTIA  
 ELQMI DPSADLYAQVQEIAGSPA I KKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTAR  
 5 GKALSKRREAKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLS  
 QYDEDHIIPQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEVVNRQLAYWRS LKDAGLMTQR  
 KFDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNIVL  
 LKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAAIAINLLALYPYMA DDFVYGEFRYKKKP  
 QAEKATYEKLRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNVVKKAEERTG  
 10 MLFKETINGKTNKGSRIPIKKDLDP SKYGGYIEEKMAYYAVISYEDKKKKPGKTIVGISIMD  
 KKEFEYDSISYLGKLGFSNPV VQIILKNYSLIAYPDGR RRYITGATKTTKGKVELQKANQIA  
 MEQDLVNFIIYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRIY  
 EEEKKKDPVDIEALVASFIELLKLTSAGAPADFI FMGEAISRRRYSMTGLFDGQVIYQSLT  
 GLYETRMRFEDKRPAATKKAGQAKKKKGSSGGSSGGSSGSETPGTSESATPES SGGSSGGST  
 15 SEKGPSTGDPTLRRRIESWEFDVFDPREL RKETCLLYEIKWMSRKIWRSSGKNTTNHVEV  
 NFIKKFTSERRFHSSISCSITWFLSWSPWECSQAIREFLSQHPGVTLVIYVARLFWHMDQR  
 NRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYP PGDEAHWPQYPPLWMMMLYALELHCIILS  
 LPPCLKISR RQNH LAFFRLHLQNC HYQTI PPHILLATGLIHPSVTWRYPYDVPDYAYPYDV  
 PDYAYPYDVPDYA (SEQ ID NO: 94).

20 In some embodiments, the *Streptococcus constellatus* Cas9 protein recognizes a PAM sequence comprising 5'-NGG-3'.

In some embodiments, the *Streptococcus constellatus* Cas9 protein recognizes a PAM sequence comprising 5'-NGC-3'.

25 In some embodiments, a Cas9 protein disclosed herein (e.g., SirCas9, VapCas9, EpeCas9, LfeCas9, or PmaCas9) recognizes a PAM sequence comprising 5'-NGC-3'.

In some embodiments, the *Sharpea* Cas9 protein recognizes a PAM sequence comprising 5' – NAGHC – 3' wherein H=A, C or T.

30 In some embodiments, the *Veillonella parvula* Cas9 protein recognizes a PAM sequence comprising 5' – NRHRRH – 3', wherein H is adenine, cytosine or thymine, and R is adenine or guanine.

In some embodiments, the *Ezakiella peruensis* Cas9 protein recognizes a PAM sequence comprising 5'-NGG-3'.

In some embodiments, the *Lactobacillus fermentum* strain AF15-40LB Cas9 protein recognizes a PAM sequence comprising 5'-NGG-3'.

5 In some embodiments, the *Peptoniphilus sp. Marseille-P3761* Cas9 protein recognizes a PAM sequence comprising 5'-NNAAA-3'

In some embodiments, a nucleic acid encoding the Cas9 protein is provided.

In some embodiments, the nucleic acid is codon-optimized for expression in mammalian cells.

10 In some embodiments, the nucleic acid is codon-optimized for expression in human cells.

In some embodiments, a eukaryotic cell comprising the Cas9 protein is provided.

In some embodiments, the cell is a human cell. In some embodiments, the cell is a plant cell.

15 In one aspect, a method of cleaving a target nucleic acid in a eukaryotic cell is provided comprising: contacting the cell with a Cas9 as described herein, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and wherein the Cas9 protein is capable of binding to the RNA guide and of causing a break in the target  
20 nucleic acid sequence complementary to the RNA guide.

In one aspect, a method of altering expression of a target nucleic acid in a eukaryotic cell is provided comprising: contacting the cell with a Cas9 as described herein, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and  
25 wherein the Cas9 protein is capable of binding to the RNA guide and of causing a break in the target nucleic acid sequence complementary to the RNA guide.

In one aspect, a method of altering expression of a target nucleic acid in a eukaryotic cell is provided comprising: contacting the cell with a Cas9 as described herein, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct  
30 repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and

wherein the Cas9 protein is capable of binding to the RNA guide and editing the target nucleic acid sequence complementary to the RNA guide.

In one aspect, a method of modifying a target nucleic acid in a eukaryotic cell is provided comprising: contacting the cell with a Cas9 as described herein, and an RNA guide  
 5 or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and wherein the Cas9 protein is capable of binding to the RNA guide and editing the target nucleic acid sequence complementary to the RNA guide.

In some embodiments, the Cas9 protein is an inactive Cas9 (dCas9).

10 In some embodiments, the dCas9 is fused to a deaminase.

In some embodiments, the RNA guide comprises a crRNA and a tracrRNA.

In some embodiments, the RNA guide comprises a sgRNA.

In some embodiments, the sgRNA for use with *Streptococcus constellatus* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to  
 15 5' –  
 GUUUUAGAGCUGUGCUGUUUAAACAACACAGCAAGUUAAAUAAGGCUUUGUCCGUACUCAA  
 GCUUGCAAAAGCGUGCACCGAUUCGGUGCU–3' (SEQ ID NO: 3).

In some embodiments, the sgRNA for use with *Sharpea* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to  
 20 5' –  
 GUUUUAGAGUUGUGUUUUAUUGAAAAUAACACAACGAGUUAAAUAAGCUUAUGCUUAAAUG  
 CCAGCUUUGCUGGUGUCAUUUAGAUGACUUUACUAAGGUUGCUUCGGCAACCUUUUU–3'  
 (SEQ ID NO: 7).

In some embodiments, the sgRNA for use with *Veillonella parvula* Cas9 comprises a  
 25 scaffold comprising a sequence having at least about 80% identity to  
 5' –  
 GUUUGAGAGUAGUGUGAAAACAUUACGAGUUCAAAUACAAAUAAUUUACAAUGCCUUCGGG  
 CUGCCCGACGUAGGGCACCUACUCUCAAUUCUUCGGAAUUGAGUU–3' (SEQ ID NO: 13).

In some embodiments, the sgRNA for use with *Ezakiella peruensis* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

5' –  
 GUUUGAGAGUUAUGUAAUUGAAAAUUACAUGACGAGUUCAAAUAAAAUUUAUCAAACCG  
 5 CCUAUUUAUAGGCCGCAGAUUGUUCUGCAUUAUGCUUGCUAUUGCAAGCUU–3' (SEQ ID  
 NO: 19).

In some embodiments, the sgRNA for use with *Lactobacillus fermentum strain AF15-40LB* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

5' –  
 10 GUCUUUGAUGAGUGUGAAAACACUCAUAGUCAAGAUCAAACGAGUGGUUUUCCACGAGUUUAU  
 UACUUUUUGAGGUCUUUAUUGGCCCAUACAUAAAAGGAGUCGGAAUUUCCGGCUCUUUUUCU  
 U–3' (SEQ ID NO: 95).

In some embodiments, the sgRNA for use with *Peptoniphilus sp. Marseille-P3761* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

15 5' –  
 GUUUUAGAGCCAUGUAGAAUACAUUGCAAGUUAAAAUAAGGCUUUGUCCGUAUAUCAACUUG  
 AAAAAGUGGCGCUGUUUCGGCGCUUU–3' (SEQ ID NO: 96).

In some embodiments, the crRNA comprises a guide sequence of between about 16 and 26 nucleotides long.

20 In some embodiments, the crRNA comprises a guide sequence between 18 and 24 nucleotides long.

In some embodiments, the break in the target nucleic acid is a single-stranded or double-stranded break.

In some embodiments, the break in the target nucleic acid is a single-stranded break.

25 In some embodiments, the Cas9 protein is a nuclease that cleaves both strands of the target nucleic acid sequence. In some embodiments, the Cas9 is a nickase that cleaves one strand of the target nucleic acid sequence.

In some embodiments, the target nucleic acid is 5' to a protospacer adjacent motif (PAM) sequence.

In some embodiments, the Cas9 is operably linked to a promoter sequence for expression in a eukaryotic cell, and wherein the guide RNA is operably linked to a promoter  
5 sequence for expression in a eukaryotic cell.

In some embodiments, the eukaryotic cell is a human cell.

In some embodiments, the promoter sequence is a eukaryotic or viral promoter.

In one aspect, provided herein is an engineered, non-naturally occurring CRISPR-Cas system comprising: an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA  
10 guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to a target nucleic acid; and a codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to SEQ ID NOs: 1, 4, 8, 14, 84 or 86 and wherein the Cas protein is capable of binding to the RNA guide and of causing a break in the target nucleic acid sequence complementary to the RNA guide.

15 In some embodiments, provided herein is an engineered, non-naturally occurring CRISPR-Cas system comprising a codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to SEQ ID NOs: 2, 5, 9, 15, 85, 87, 95 or 96, and wherein the Cas protein is capable of binding to the RNA guide and of causing a break in the target nucleic acid sequence complementary to the RNA guide.

20 In one aspect, provided herein is an engineered, non-naturally occurring CRISPR-Cas system comprising: an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to a target nucleic acid; and a codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to SEQ ID NOs: 1, 4, 8, 14, 84 or 86; wherein the Cas protein is  
25 fused to a deaminase, and wherein the Cas protein fusion is capable of binding to the RNA guide and of editing the target nucleic acid sequence complementary to the RNA guide.

In some embodiments, the engineered, non-naturally occurring CRISPR-Cas system comprises a codon-optimized CRISPR-associated (Cas) protein further comprising a nuclear localization sequence (NLS) and/or a FLAG, HIS or HA tag.

In some embodiments, the engineered, non-naturally occurring CRISPR-Cas system comprises a codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to SEQ ID NOS: 2, 5, 9, 15, 85, 87, 95 or 96, wherein the Cas protein is fused to a deaminase, and wherein the Cas protein fusion is capable of binding to the RNA guide and of editing the target nucleic acid sequence complementary to the RNA guide.

In one embodiment, the Cas9 protein is an inactive Cas9 (dCas9).

In one embodiment, the RNA guide comprises a crRNA and a tracrRNA.

In one embodiment, the RNA guide comprises an sgRNA.

In one embodiment, the Cas protein is operably linked to a promoter sequence for expression in a eukaryotic cell, and wherein the guide RNA is operably linked to a promoter sequence for expression in a eukaryotic cell.

In one embodiment, the eukaryotic cell is a human cell.

In one embodiment, the promoter sequence is a eukaryotic promoter sequence.

In one embodiment, a nucleic acid encoding the system described herein is provided.

In one embodiment, a vector comprising the system described herein is provided.

In one embodiment, the vector is a plasmid vector or a viral vector.

In one embodiment, the viral vector is an adeno associated virus (AAV) vector or a lentiviral vector.

In one embodiment, the viral vector is an AAV vector.

In one embodiment, more than one AAV vector is used for packaging the system.

In one embodiment, a method of treating a disorder or a disease in a subject in need thereof comprises administering to the subject the system described herein, wherein the guide RNA is complementary to at least 10 nucleotides of a target nucleic acid associated with the condition or disease; wherein the Cas protein associates with the guide RNA; wherein the guide RNA binds to the target nucleic acid; wherein the Cas protein causes a break in the target nucleic acid, optionally wherein the Cas9 is an inactive Cas9 (dCas9) fused to a

deaminase and results in one or more base edits in the target nucleic acid, thereby treating the disorder or disease.

In some embodiments, the guide RNA is complementary to about 18-24 nucleotides.

In some embodiments, the guide RNA is complementary to 20 nucleotides.

5 In some embodiments, the base editor comprises a fusion protein.

In some embodiments, the base editor comprises an adenosine deaminase domain or a cytidine deaminase domain.

In some embodiments, provided herein is a method of editing a nucleobase of a polynucleotide, the method comprising contacting the polynucleotide with a base in complex  
10 with one or more guide RNAs, wherein the base editor comprises an adenosine deaminase domain, and wherein the one or more guide RNAs target the base editor to effect an A•T to G•C alteration in the polynucleotide.

In some embodiments, provided herein is a method of editing a nucleobase of a polynucleotide, the method comprising contacting the polynucleotide with a base editor in  
15 complex with one or more guide RNAs, wherein the base editor comprises a cytidine deaminase domain, and wherein the one or more guide RNAs target the base editor to effect a C•G to T•A alteration in the polynucleotide.

In some embodiments, the editing results in less than 50 % indel formation in the target polynucleotide sequence.

20 In some embodiments, the editing generates a point mutation.

## DEFINITIONS

In order for the present invention to be more readily understood, certain terms are first defined below. Additional definitions for the following terms and other terms are set forth  
25 throughout the specification.

*A or An:* The articles “a” and “an” are used herein to refer to one or to more than one (*i.e.*, to at least one) of the grammatical object of the article. By way of example, “an element” means one element or more than one element.

*Approximately or about:* As used herein, the term “approximately” or “about,” as  
30 applied to one or more values of interest, refers to a value that is similar to a stated reference

value. In certain embodiments, the term “approximately” or “about” refers to a range of values that fall within 25%, 20%, 19%, 18%, 17%, 16%, 15%, 14%, 13%, 12%, 11%, 10%, 9%, 8%, 7%, 6%, 5%, 4%, 3%, 2%, 1%, or less in either direction (greater than or less than) of the stated reference value unless otherwise stated or otherwise evident from the context  
5 (except where such number would exceed 100% of a possible value).

*Associated with:* Two events or entities are “associated” with one another, as that term is used herein, if the presence, level and/or form of one is correlated with that of the other. For example, a particular entity (e.g., polypeptide) is considered to be associated with a particular disease, disorder, or condition, if its presence, level and/or form correlates with  
10 incidence of and/or susceptibility to the disease, disorder, or condition (e.g., across a relevant population). In some embodiments, two or more entities are physically “associated” with one another if they interact, directly or indirectly, so that they are and remain in physical proximity with one another. In some embodiments, two or more entities that are physically associated with one another are covalently linked to one another; in some embodiments, two  
15 or more entities that are physically associated with one another are not covalently linked to one another but are non-covalently associated, for example by means of hydrogen bonds, van der Waals interaction, hydrophobic interactions, magnetism, and combinations thereof.

*Base Editor:* By “base editor (BE),” or “nucleobase editor (NBE)” is meant an agent that binds a polynucleotide and has nucleobase modifying activity. In various embodiments,  
20 the base editor comprises a nucleobase modifying polypeptide (e.g., a deaminase) and a polynucleotide programmable nucleotide binding domain in conjunction with a guide polynucleotide (e.g., guide RNA). In various embodiments, the agent is a biomolecular complex comprising a protein domain having base editing activity, *i.e.*, a domain capable of modifying a base (e.g., A, T, C, G, or U) within a nucleic acid molecule (e.g., DNA). In  
25 some embodiments, the polynucleotide programmable DNA binding domain is fused or linked to a deaminase domain. In one embodiment, the agent is a fusion protein comprising one or more domains having base editing activity. In another embodiment, the protein domains having base editing activity are linked to the guide RNA (e.g., via an RNA binding motif on the guide RNA and an RNA binding domain fused to the deaminase). In some  
30 embodiments, the domains having base editing activity are capable of deaminating a base within a nucleic acid molecule. In some embodiments, the base editor is capable of deaminating one or more bases within a DNA molecule. In some embodiments, the base editor is capable of deaminating a cytosine (C) or an adenosine (A) within DNA. In some



embodiments, the base editor is capable of deaminating a cytosine (C) and an adenosine (A) within DNA. In some embodiments, the base editor is a cytidine base editor (CBE). In some embodiments, the base editor is an adenosine base editor (ABE). In some embodiments, the base editor is an adenosine base editor (ABE) and a cytidine base editor (CBE). In some  
5 embodiments, the base editor is a nuclease-inactive Cas9 (dCas9) fused to an adenosine deaminase. In some embodiments, the base editor is fused to an inhibitor of base excision repair, for example, a UGI domain, or a dISN domain. In some embodiments, the fusion protein comprises a Cas9 nickase fused to a deaminase and an inhibitor of base excision repair, such as a UGI or dISN domain. In other embodiments the base editor is an abasic base  
10 editor. Details of base editors are described in International PCT Application Nos. PCT/2017/045381 (WO2018/027078) and PCT/US2016/058344 (WO2017/070632), each of which is incorporated herein by reference for its entirety. Also see Komor, A.C., *et al.*, “Programmable editing of a target base in genomic DNA without double-stranded DNA cleavage” *Nature* 533, 420-424 (2016); Gaudelli, N.M., *et al.*, “Programmable base editing of  
15 A•T to G•C in genomic DNA without DNA cleavage” *Nature* 551, 464-471 (2017); Komor, A.C., *et al.*, “Improved base excision repair inhibition and bacteriophage Mu Gam protein yields C:G-to-T:A base editors with higher efficiency and product purity” *Science Advances* 3:eao4774 (2017), and Rees, H.A., *et al.*, “Base editing: precision chemistry on the genome and transcriptome of living cells.” *Nat Rev Genet.* 2018 Dec;19(12):770-788. doi:  
20 10.1038/s41576-018-0059-1, the entire contents of which are hereby incorporated by reference.

*Base Editing Activity:* By “base editing activity” is meant acting to chemically alter a base within a polynucleotide. In one embodiment, a first base is converted to a second base. In one embodiment, the base editing activity is cytidine deaminase activity, *e.g.*, converting  
25 target C•G to T•A. In another embodiment, the base editing activity is adenosine or adenine deaminase activity, *e.g.*, converting A•T to G•C. In another embodiment, the base editing activity is cytosine or cytidine deaminase activity, *e.g.*, converting target C•G to T•A and adenosine or adenine deaminase activity, *e.g.*, converting A•T to G•C.

*Base Editor System:* The term “base editor system” refers to a system for editing a  
30 nucleobase of a target nucleotide sequence. In various embodiments, the base editor (BE) system comprises (1) a polynucleotide programmable nucleotide binding domain (*e.g.*, Cas9), a deaminase domain and a cytidine deaminase domain for deaminating nucleobases in the target nucleotide sequence; and (2) one or more guide polynucleotides (*e.g.*, guide RNA) in

conjunction with the polynucleotide programmable nucleotide binding domain. In various embodiments, the base editor (BE) system comprises a nucleobase editor domains selected from an adenosine deaminase or a cytidine deaminase, and a domain having nucleic acid sequence specific binding activity. In some embodiments, the base editor system comprises

5 (1) a base editor (BE) comprising a polynucleotide programmable DNA binding domain and a deaminase domain for deaminating one or more nucleobases in a target nucleotide sequence; and (2) one or more guide RNAs in conjunction with the polynucleotide programmable DNA binding domain. In some embodiments, the polynucleotide programmable nucleotide binding domain is a polynucleotide programmable DNA binding

10 domain. In some embodiments, the base editor is a cytidine base editor (CBE). In some embodiments, the base editor is an adenine or adenosine base editor (ABE). In some embodiments, the base editor is an adenine or adenosine base editor (ABE) or a cytidine base editor (CBE).

In some embodiments, a polynucleotide programmable nucleotide binding domain

15 can target a deaminase domain to a target nucleotide sequence by non-covalently interacting with or associating with the deaminase domain. For example, in some embodiments, the nucleobase editing component, *e.g.*, the deaminase component can comprise an additional heterologous portion or domain that is capable of interacting with, associating with, or capable of forming a complex with an additional heterologous portion or domain that is part

20 of a polynucleotide programmable nucleotide binding domain. In some embodiments, the additional heterologous portion may be capable of binding to, interacting with, associating with, or forming a complex with a polypeptide. In some embodiments, the additional heterologous portion may be capable of binding to, interacting with, associating with, or forming a complex with a polynucleotide. In some embodiments, the additional heterologous

25 portion may be capable of binding to a guide polynucleotide. In some embodiments, the additional heterologous portion may be capable of binding to a polypeptide linker. In some embodiments, the additional heterologous portion may be capable of binding to a polynucleotide linker. The additional heterologous portion may be a protein domain. In some embodiments, the additional heterologous portion may be a K Homology (KH) domain,

30 a MS2 coat protein domain, a PP7 coat protein domain, a SfMu Com coat protein domain, a steril alpha motif, a telomerase Ku binding motif and Ku protein, a telomerase Sm7 binding motif and Sm7 protein, or an RNA recognition motif.

*Biologically active:* As used herein, the phrase “biologically active” refers to a characteristic of any agent that has activity in a biological system, and particularly in an organism. For instance, an agent that, when administered to an organism, has a biological effect on that organism, is considered to be biologically active. In particular embodiments, where a peptide is biologically active, a portion of that peptide that shares at least one biological activity of the peptide is typically referred to as a “biologically active” portion.

*Cleavage:* As used herein, cleavage refers to a break in a target nucleic acid created by a nuclease of a CRISPR system described herein. In some embodiments, the cleavage event is a double-stranded DNA break. In some embodiments, the cleavage event is a single-stranded DNA break. In some embodiments, the cleavage event is a single-stranded RNA break. In some embodiments, the cleavage event is a double-stranded RNA break.

*Complementary:* As used herein, complementary refers to a nucleic acid strand that forms Watson-Crick base pairing, such that A base pairs with T, and C base pairs with G, or non-traditional base pairing with bases on a second nucleic acid strand. In other words, it refers to nucleic acids that hybridize with each other under appropriate conditions.

*Clustered Interspaced Short Palindromic Repeat (CRISPR)-associated (Cas) system:* As used herein, CRISPR-Cas9 system refers to nucleic acids and/or proteins involved in the expression of, or directing the activity of, CRISPR-effectors, including sequences encoding CRISPR effectors, RNA guides, and other sequences and transcripts from a CRISPR locus. In some embodiments, the CRISPR system is an engineered, non-naturally occurring CRISPR system. In some embodiments, the components of a CRISPR system may include a nucleic acid(s) (e.g., a vector) encoding one or more components of the system, a component(s) in protein form, or a combination thereof.

*CRISPR Array:* The term “CRISPR array”, as used herein, refers to the nucleic acid (e.g., DNA) segment that includes CRISPR repeats and spacers, starting with the first nucleotide of the first CRISPR repeat and ending with the last nucleotide of the last (terminal) CRISPR repeat. Typically, each spacer in a CRISPR array is located between two repeats. The terms “CRISPR repeat” or “CRISPR direct repeat,” or “direct repeat,” as used herein, refer to multiple short direct repeating sequences, which show very little or no sequence variation within a CRISPR array.

*CRISPR-associated protein (Cas):* The term “CRISPR-associated protein,” “CRISPR effector,” “effector,” or “CRISPR enzyme” as used herein refers to a protein that carries out

an enzymatic activity or that binds to a target site on a nucleic acid specified by a RNA guide. In different embodiments, a CRISPR effector has endonuclease activity, nickase activity, exonuclease activity, transposase activity, and/or excision activity. In some embodiments, the Cas is a high-accuracy Cas. In some embodiments, the Cas is a high-fidelity Cas. In some  
5 embodiments, the Cas is a SuperFi-Cas. In some embodiments, the high-accuracy, high-fidelity and SuperFi-Cas are as described in Bravo, J. *et al.* Structural basis for mismatch surveillance by CRISPR-Cas9 *Nature*, 603, March 2022.

*crRNA*: The term "CRISPR RNA" or "crRNA," as used herein, refers to a RNA molecule including a guide sequence used by a CRISPR effector to target a specific nucleic  
10 acid sequence. Typically, crRNAs contains a sequence that mediates target recognition and a sequence that forms a duplex with a tracrRNA. In some embodiments, the crRNA: tracrRNA duplex binds to a CRISPR effector.

*Ex Vivo*: As used herein, the term "ex vivo" refers to events that occur in cells or tissues, grown outside rather than within a multi-cellular organism.

*Functional equivalent or analog*: As used herein, the term "functional equivalent" or  
15 "functional analog" denotes, in the context of a functional derivative of an amino acid sequence, a molecule that retains a biological activity (either function or structural) that is substantially similar to that of the original sequence. A functional derivative or equivalent may be a natural derivative or is prepared synthetically. Exemplary functional derivatives  
20 include amino acid sequences having substitutions, deletions, or additions of one or more amino acids, provided that the biological activity of the protein is conserved. The substituting amino acid desirably has chemico-physical properties which are similar to that of the substituted amino acid. Desirable similar chemico-physical properties include, similarities in charge, bulkiness, hydrophobicity, hydrophilicity, and the like.

*Half-Life*: As used herein, the term "half-life" is the time required for a quantity such  
25 as protein concentration or activity to fall to half of its value as measured at the beginning of a time period.

*Improve, increase, or reduce*: As used herein, the terms "improve," "increase" or  
30 "reduce," or grammatical equivalents, indicate values that are relative to a baseline measurement, such as a measurement in the same individual prior to initiation of the treatment described herein, or a measurement in a control subject (or multiple control subject) in the absence of the treatment described herein. A "control subject" is a subject afflicted

with the same form of disease as the subject being treated, who is about the same age as the subject being treated.

*Inhibition:* As used herein, the terms “inhibition,” “inhibit” and “inhibiting” refer to processes or methods of decreasing or reducing activity and/or expression of a protein or a gene of interest. Typically, inhibiting a protein or a gene refers to reducing expression or a relevant activity of the protein or gene by at least 10% or more, for example, 20%, 30%, 40%, or 50%, 60%, 70%, 80%, 90% or more, or a decrease in expression or the relevant activity of greater than 1-fold, 2-fold, 3-fold, 4-fold, 5-fold, 10-fold, 50-fold, 100-fold or more as measured by one or more methods described herein or recognized in the art.

*Hybridization:* As used herein, the term “hybridization” refers to a reaction in which two or more nucleic acids bind with each other via hydrogen bonding by Watson-Crick pairing, Hoogsteen binding or other sequence-specific binding between the bases of the two nucleic acids. A sequence capable of hybridizing with another sequence is termed the “complement” of the sequence, and is said to be “complementary” or show “complementarity”.

*Indel:* As used herein, the term “indel” refers to insertion or deletion of bases in a nucleic acid sequence. It commonly results in mutations and is a common form of genetic variation.

*In Vitro:* As used herein, the term “*in vitro*” refers to events that occur in an artificial environment, *e.g.*, in a test tube or reaction vessel, in cell culture, *etc.*, rather than within a multi-cellular organism.

*In Vivo:* As used herein, the term “*in vivo*” refers to events that occur within a multi-cellular organism, such as a human and a non-human animal. In the context of cell-based systems, the term may be used to refer to events that occur within a living cell (as opposed to, for example, *in vitro* systems).

*Linker:* The term “linker” refers to any means, entity or moiety used to join two or more entities. In some embodiments, the linker is a covalent linker. In some embodiments, the linker is a non-covalent linker. Examples of covalent linkers include covalent bonds or a linker moiety covalently attached to one or more of the proteins or domains to be linked. In some embodiments, the linker is a non-covalent bond, *e.g.*, an organometallic bond through a metal center such as platinum atom. The joining can be permanent or reversible. For covalent linkages, various functionalities can be used, such as amide groups, including

carbonic acid derivatives, ethers, esters, including organic and inorganic esters, amino, urethane, urea and the like. To provide for linking, the domains can be modified by oxidation, hydroxylation, substitution, reduction etc. to provide a site for coupling. Methods for conjugation are well known by persons skilled in the art and are encompassed for use in the present invention. Linker moieties include, but are not limited to, chemical linker moieties, or for example a peptide linker moiety (a linker sequence). It will be appreciated that modification which do not significantly decrease the function of the RNA-binding domain and effector domain are preferred.

*Mutation:* As used herein, the term “mutation” has the ordinary meaning in the art, and includes, for example, point mutations, substitutions, insertions, deletions, inversions, and deletions.

*Oligonucleotide:* As used herein, the term “oligonucleotide” generally refers to polynucleotides of between about 5 and about 100 nucleotides of single- or double-stranded DNA. Oligonucleotides are also known as "oligomers" or "oligos" and may be isolated from genes, or chemically synthesized.

*PAM:* The term “PAM” or “Protospacer Adjacent Motif” refers to a short nucleic acid sequence (usually 2-6 base pairs in length) that follows the nucleic acid region targeted for cleavage by the CRISPR system, such as CRISPR-Cas9. The PAM is required for a Cas nuclease to cut and is generally found 3-4 nucleotides downstream from the cut site.

*Polypeptide:* The term “polypeptide” as used herein refers to a sequential chain of amino acids linked together via peptide bonds. The term is used to refer to an amino acid chain of any length, but one of ordinary skill in the art will understand that the term is not limited to lengthy chains and can refer to a minimal chain comprising two amino acids linked together via a peptide bond. As is known to those skilled in the art, polypeptides may be processed and/or modified. As used herein, the terms “polypeptide” and “peptide” are used inter-changeably.

*Prevent:* As used herein, the term “prevent” or “prevention”, when used in connection with the occurrence of a disease, disorder, and/or condition, refers to reducing the risk of developing the disease, disorder and/or condition.

*Protein:* The term “protein” as used herein refers to one or more polypeptides that function as a discrete unit. If a single polypeptide is the discrete functioning unit and does not require permanent or temporary physical association with other polypeptides in order to

form the discrete functioning unit, the terms “polypeptide” and “protein” may be used interchangeably. If the discrete functional unit is comprised of more than one polypeptide that physically associate with one another, the term “protein” refers to the multiple polypeptides that are physically coupled and function together as the discrete unit.

5           *Reference*: A “reference” entity, system, amount, set of conditions, etc., is one against which a test entity, system, amount, set of conditions, etc. is compared as described herein. For example, in some embodiments, a “reference” antibody is a control antibody that is not engineered as described herein.

10           *RNA guide*: The term RNA guide refers to an RNA molecule that facilitates the targeting of a protein described herein to a target nucleic acid. Exemplary “RNA guides” or “guide RNAs” include, but are not limited to, crRNAs or crRNAs in combination with cognate tracrRNAs. The latter may be independent RNAs or fused as a single RNA using a linker (sgRNAs). In some embodiments, the RNA guide is engineered to include a chemical or biochemical modification, in some embodiments, an RNA guide may include one or more  
15 nucleotides.

*Subject*: The term “subject”, as used herein, means any subject for whom diagnosis, prognosis, or therapy is desired. For example, a subject can be a mammal, *e.g.*, a human or non-human primate (such as an ape, monkey, orangutan, or chimpanzee), a dog, cat, guinea pig, rabbit, rat, mouse, horse, cattle, or cow.

20           *sgRNA*: The term “sgRNA” or “single guide RNA” refers to a single guide RNA containing (i) a guide sequence (crRNA sequence) and (ii) a Cas9 nuclease-recruiting sequence (tracrRNA).

*Substantial identity*: The phrase “substantial identity” is used herein to refer to a comparison between amino acid or nucleic acid sequences. As will be appreciated by those  
25 of ordinary skill in the art, two sequences are generally considered to be “substantially identical” if they contain identical residues in corresponding positions. As is well known in this art, amino acid or nucleic acid sequences may be compared using any of a variety of algorithms, including those available in commercial computer programs such as BLASTN for nucleotide sequences and BLASTP, gapped BLAST, and PSI-BLAST for amino acid  
30 sequences. Exemplary such programs are described in Altschul, et al., Basic local alignment search tool, *J. Mol. Biol.*, 215(3): 403-410, 1990; Altschul, et al., *Methods in Enzymology*; Altschul et al., *Nucleic Acids Res.* 25:3389-3402, 1997; Baxevanis et al., *Bioinformatics* : A

*Practical Guide to the Analysis of Genes and Proteins*, Wiley, 1998; and Misener, et al., (eds.), *Bioinformatics Methods and Protocols* (Methods in Molecular Biology, Vol. 132), Humana Press, 1999. In addition to identifying identical sequences, the programs mentioned above typically provide an indication of the degree of identity. In some embodiments, two  
5 sequences are considered to be substantially identical if at least 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more of their corresponding residues are identical over a relevant stretch of residues. In some  
embodiments, the relevant stretch is a complete sequence. In some embodiments, the relevant stretch is at least 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95,  
10 100, 125, 150, 175, 200, 225, 250, 275, 300, 325, 350, 375, 400, 425, 450, 475, 500 or more residues.

*Target Nucleic Acid*: The term “target nucleic acid” as used herein refers to nucleotides of any length (oligonucleotides or polynucleotides) to which the CRISPR-Cas9 system binds, either deoxyribonucleotides, ribonucleotides, or analogs thereof. Target nucleic  
15 acids may have three-dimensional structure, may including coding or non-coding regions, may include exons, introns, mRNA, tRNA, rRNA, siRNA, shRNA, miRNA, ribozymes, cDNA, plasmids, vectors, exogenous sequences, endogenous sequences. A target nucleic acid can comprise modified nucleotides, include methylated nucleotides, or nucleotide analogs. A target nucleic acid may be interspersed with non-nucleic acid components. A target nucleic  
20 acid is not limited to, single-, double-, or multi-stranded DNA or RNA, genomic DNA, cDNA, DNA-RNA hybrids, or a polymer comprising purine and pyrimidine bases or other natural, chemically or biochemically modified, non-natural, or derivatized nucleotide bases.

*Therapeutically effective amount*: As used herein, the term “therapeutically effective amount” refers to an amount of a therapeutic molecule (e.g., an engineered antibody  
25 described herein) which confers a therapeutic effect on a treated subject, at a reasonable benefit/risk ratio applicable to any medical treatment. The therapeutic effect may be objective (i.e., measurable by some test or marker) or subjective (i.e., subject gives an indication of or feels an effect). In particular, the “therapeutically effective amount” refers to an amount of a therapeutic molecule or composition effective to treat, ameliorate, or prevent  
30 a particular disease or condition, or to exhibit a detectable therapeutic or preventative effect, such as by ameliorating symptoms associated with the disease, preventing or delaying the onset of the disease, and/or also lessening the severity or frequency of symptoms of the disease. A therapeutically effective amount can be administered in a dosing regimen that may



comprise multiple unit doses. For any particular therapeutic molecule, a therapeutically effective amount (and/or an appropriate unit dose within an effective dosing regimen) may vary, for example, depending on route of administration, on combination with other pharmaceutical agents. Also, the specific therapeutically effective amount (and/or unit dose) for any particular subject may depend upon a variety of factors including the disorder being treated and the severity of the disorder; the activity of the specific pharmaceutical agent employed; the specific composition employed; the age, body weight, general health, sex and diet of the subject; the time of administration, route of administration, and/or rate of excretion or metabolism of the specific therapeutic molecule employed; the duration of the treatment; and like factors as is well known in the medical arts.

*tracrRNA*: The term "tracrRNA" or "trans-activating crRNA" as used herein refers to an RNA including a sequence that forms a structure required for a CRISPR-associated protein to bind to a specified target nucleic acid.

*Treatment*: As used herein, the term "treatment" (also "treat" or "treating") refers to any administration of a therapeutic molecule (e.g., a CRISPR-Cas therapeutic protein or system described herein) that partially or completely alleviates, ameliorates, relieves, inhibits, delays onset of, reduces severity of and/or reduces incidence of one or more symptoms or features of a particular disease, disorder, and/or condition. Such treatment may be of a subject who does not exhibit signs of the relevant disease, disorder and/or condition and/or of a subject who exhibits only early signs of the disease, disorder, and/or condition. Alternatively or additionally, such treatment may be of a subject who exhibits one or more established signs of the relevant disease, disorder and/or condition.

### BRIEF DESCRIPTION OF THE DRAWING

Drawings are for illustration purposes only; not for limitation.

FIG. 1A is a graph that shows a consensus PAM motif recognized by human codon-optimized *Streptococcus constellatus* Cas9. FIG. 1B is a graph that shows a consensus PAM motif recognized by human codon-optimized *Sharpea spp. isolate RUG017* Cas9. FIG. 1C is a graph that shows a consensus PAM motif recognized by human codon-optimized *Veillonella parvula* Cas9. FIG. 1D is a graph that shows a consensus PAM motif recognized by human codon-optimized *Ezakiella peruensis*. FIG. 1E is a graph that shows a consensus PAM motif recognized by human codon-optimized *Lactobacillus fermentum strain AF15-*

40LB. FIG. 1F is a graph that shows a consensus PAM motif recognized by human codon-optimized *Peptoniphilus sp. Marseille-P3761*.

FIG. 2A is a schematic that shows predicted RNA folding structure of sgRNA for human codon-optimized *Streptococcus constellatus* ScoCas9 using Geneious software. FIG. 2A depicts sgRNA comprising SEQ ID NO: 3. FIG. 2B is a schematic that shows predicted RNA folding structure of sgRNA for human codon-optimized *Sharpea spp. isolate RUG017* SirCas9 using Geneious software. FIG. 2B depicts sgRNA comprising SEQ ID NO: 7. FIG. 2C is a schematic that shows predicted RNA folding structure of sgRNA for human codon-optimized *Veillonella parvula* VapCas9 using Geneious software. FIG. 2C depicts sgRNA comprising SEQ ID NO: 13. FIG. 2D is a schematic that shows predicted RNA folding structure of sgRNA for human codon-optimized *Ezakiella peruensis* EpeCas9 using Geneious software. FIG. 2D depicts sgRNA comprising SEQ ID NO: 19. FIG. 2E is a schematic that shows predicted RNA folding structure of sgRNA for human codon-optimized *Lactobacillus fermentum strain AF15-40LB* LfeCas9 using Geneious software. FIG. 2E depicts sgRNA comprising SEQ ID NO: 95. FIG. 2F is a schematic that shows predicted RNA folding structure of sgRNA for human codon-optimized *Peptoniphilus sp. Marseille-P3761* PmaCas9 using Geneious software. FIG. 2F depicts sgRNA comprising SEQ ID NO: 96.

FIG. 3 is a graph that shows exemplary results of *ex vivo* cleavage activity of human codon-optimized ScoCas9 in HEK293T cells. The y-axis of the graph shows indel frequency obtained using various guide RNAs that targeted A-rich genomic test sites adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1A).

FIG. 4A is a schematic showing constructs of ScoCas9 D10A mutant fused at the N-terminal to an adenine base editor (ABE) or a cytosine base editor (CBE). FIG. 4B is a graph that shows results of indel frequency and adenine to guanine base (A-to-G) conversion percentage achieved with a base editor comprising an ABE fused to the N-terminus of a ScoCas9 D10A mutant. The A-to-G conversion percentage (y-axis) is plotted for various guide RNAs targeting A-rich genomic test sites (x-axis; Table 8) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1A). FIG. 4C is a graph that shows results of indel frequency and cytosine to thymine base (C-to-T) conversion percentage achieved with a base editor comprising an ABE fused to the N-terminus of a ScoCas9 D10A mutant. The C-to-T conversion percentage (y-axis) is plotted for various guide RNAs targeting C-rich genomic test sites (x-axis; Table 8) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1A).

FIG. 5A is a schematic showing constructs of WT SirCas9 and a SirCas9 D14A mutant fused at the N-terminus to an adenine base editor (ABE). FIG. 5B is a graph that shows results of the indel frequency and A-to-G conversion achieved with a base editor comprising an ABE fused to the N-terminus of a SirCas9 D14A mutant. The A-to-G conversion percentage (y-axis) is plotted for various guide RNAs targeting A-rich genomic test sites (x-axis; Table 9) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1B).

FIG. 6A is a schematic of constructs showing WT VapCas9 and VapCas9 D38A mutant fused at the N-terminus to an adenine base editor (ABE) or a cytosine base editor (CBE). FIG. 6B is a graph that shows results of the indel frequency, A-to-G conversion achieved with a base editor comprising an ABE fused to the N-terminus of a VapCas9 D38A mutant and C-to-T conversion achieved with a base editor comprising a CBE fused to the N-terminus of a VapCas9 D38A. The A-to-G conversion percentage (y-axis) is plotted for various guide RNAs targeting A-rich genomic test sites (x-axis; Table 10) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1C). The C-to-T conversion percentage (y-axis) is plotted for various guide RNAs targeting C-rich genomic test sites (x-axis; Table 10) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1C).

FIG. 7A is a schematic of constructs showing ABE fused to the N-terminus of VapCas9 or to the C-terminus of VapCas9. FIG. 7B is a graph that shows a comparison of A-to-G conversion achieved with a base editor comprising an ABE fused to the N-terminus and an ABE fused to the C-terminus of VapCas9. The A-to-G conversion percentage (y-axis) is plotted for various guide RNAs targeting A-rich genomic test sites (x-axis; Table 11) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1C)

FIG. 8A is a schematic of constructs showing WT EpeCas9 and EpeCas9 D38A mutant fused at the N-terminus to an ABE and a CBE. FIG. 8B is a graph that shows results of the indel frequency, A-to-G conversion achieved with a base editor comprising an ABE fused to the N-terminus of an EpeCas9 D38A mutant and C-to-T conversion achieved with a base editor comprising a CBE fused to the N-terminus of a EpeCas9 D38A. The A-to-G conversion percentage (y-axis) is plotted for various guide RNAs targeting A-rich genomic test sites (x-axis; Table 12) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1D). The C-to-T conversion percentage (y-axis) is plotted for various guide

RNAs targeting C-rich genomic test sites (x-axis; Table 12) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1D).

FIG. 9A is a schematic that shows WT LfeCas9 and LfeCas9 D9A mutant fused at the N-terminus to an ABE and a CBE. FIG. 9B is a graph that shows results of the indel  
5 frequency with LfeCas9. FIG. 9C is a graph that shows results of A-to-G conversion achieved with a base editor comprising an ABE fused to the N-terminus of an LfeCas9 D9A mutant. The A-to-G conversion percentage (y-axis) is plotted for various guide RNAs targeting A-rich genomic test sites (x-axis; Table 13) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1E). FIG. 9D is a graph that shows results of C-to-T conversion  
10 achieved with a base editor comprising a CBE fused to the N-terminus of an LfeCas9 D9A mutant. The C-to-T conversion percentage (y-axis) is plotted for various guide RNAs targeting C-rich genomic test sites (x-axis; Table 13) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1E).

FIG. 10A is a schematic that shows WT PmaCas9 and PmaCas9 D12A mutant fused  
15 at the N-terminus and C-terminus to an ABE and a CBE. FIG. 10B is a graph that shows results of A-to-G or C-to-T conversion achieved with a base editor comprising an ABE or a CBE fused to the N-terminus or C-terminus of an PmaCas9 D12A mutant. The A-to-G conversion percentage (y-axis) is plotted for various guide RNAs targeting A-rich genomic test sites (x-axis; Table 14) adjacent to a sequence corresponding to the PAM consensus  
20 motif (see FIG. 1F). The C-to-T conversion percentage (y-axis) is plotted for various guide RNAs targeting C-rich genomic test sites (x-axis; Table 14) adjacent to a sequence corresponding to the PAM consensus motif (see FIG. 1F).

FIG. 11A is a graph that shows exemplary results of indel frequency (y-axis; % indel frequency) measured by transfecting cells with two ScoCas9-NGC variants, ScoCas9-NGC-  
25 v1 and ScoCas9-NGC-v2 (x-axis). An untransfected cell control is also shown.

FIG. 11B is a graph that shows exemplary A-to-G conversion (y-axis; % A to G conversion) in HEK293T cells transfected with A-to-G base editors (ABE) comprising ScoCas9-NGC variants, ScoCas9-NGC-v1 and ScoCas9-NGC-v2 (x-axis) engineered to recognize an NGC PAM motif. The ScoCas9-NGG variant which does not recognize NGC  
30 showed no A-to-G conversion. A SpyCas9-NGC control vector showed A-to-G editing. An untransfected cell control is also shown.

## DETAILED DESCRIPTION

Clustered regularly interspaced short palindromic repeats (CRISPR) was first discovered as an adaptive immune system in bacteria and archaea, and then engineered to generate targeted DNA breaks in living cells and organisms. During the cellular DNA repair  
5 process, various DNA changes can be introduced. The diverse and expanding CRISPR toolbox allows programmable genome editing, epigenome editing and transcriptome regulation.

CRISPR-Cas systems comprise three main types (I, II, and III) based on their Cas gene organization, and the sequence and structure of component proteins. Each of the three  
10 CRISPR systems is characterized by a unique Cas gene: Cas3, a target-degrading nuclease/helicase in Type I; Cas9, an RNA-binding and target-degrading nuclease in type II; Cas10, a large protein for multiple functions in type III. The three CRISPR types also differ in their associated effector complexes. Type I Cas systems associate with Cascade effector complexes, type II effector complexes consist of a single Cas9 and one or more RNA  
15 molecules, and type III interference complexes are further divided into type III-A (Csm complex targeting DNA) and type III-B (Cmr complex targeting RNA). Cas proteins are important components of effector complexes in all CRISPR-Cas systems.

Current genome editing technologies have focused on Class II CRISPR-Cas systems, which contain single-protein effector nucleases for DNA cleavage, specifically, Cas9, a dual-  
20 RNA-guided nuclease which requires both CRISPR RNA (crRNA) and tracrRNA and contains both HNH and RuvC nuclease domains, and Cas12a, a single-RNA-guided nuclease which only requires crRNA and contains a single RuvC domain.

Various aspects of the invention are described in detail in the following sections. The use of sections is not meant to limit the invention. Each section can apply to any aspect of  
25 the invention. In this application, the use of “or” means “and/or” unless stated otherwise.

### **Engineered, Non-Naturally Occuring Cas9 Protein**

Described herein are engineered, non-naturally occurring Cas9 proteins modified from WT Cas9 obtained from *Streptococcus constellatus* (ScoCas9), *Sharpea spp. isolate RUG017* (SirCas9), *Veillonella parvula* (VapCas9 or VpaCas9, used interchangeably  
30 herein), *Ezakiella peruensis* (EpeCas9), *Lactobacillus fermentum* (LfeCas9) and *Peptoniphilus sp. Marseille-P3761* (PmaCas9) bacteria.

In some embodiments, the engineered non-naturally occurring Cas9 protein described herein comprises an amino acid sequence at least 60% (e.g., 60%, 65%, 70%, 75%, 80%,

81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more) identical to SEQ ID NO: 1, 4, 8, 14, 84 or 86. In some embodiments, the Cas9 protein has is 80% identical to SEQ ID NO: 1, 4, 8, 14, 84 or 86. In some embodiments, the amino acid sequence of the Cas9 protein is identical to SEQ ID NO: 1, 4, 8, 14, 84 or 86. Exemplary Cas9 amino acid sequences are provided in Table 1 below.

**Table 1. Exemplary Cas9 Amino Acid Sequences**

<b>Wild Type <i>Streptococcus constellatus</i> Cas9</b>
MGKPYSIGLDIGTNSVGVAVVTDDYKVPAAKMKVLGNTDKQSIKKNLLGALLFDSGETAEA TRLKRTARRRYTRRKNRLRYLQEI FTGEMNKVDENFFQRLDDSFVDEDKRGEHHP I FGNI AAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQA LFKDFVEEYDKTIEESHLS EITVDALSILTEKVS KSSRLENLIAHYPT EKKNTLFGNLI AL SLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEGVGEYADLFASAKNLYDAILLSGI LTVDDNSTKAPLSASMVKRYEEHQDLKKLKD FIKVNAPDQYNAI FKDKNKKGYASYIESG VKQDEFYKYLKGI LLKINGS GDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQG EHYPFLKENQDKIEKILTFRI PYYVGPLARKGSRFAWA EYKADEKITPWNFDDILDKEKSA EKFITRMTLNDLYLP EEKVLPKHSPLYEAF TVYNELTKVKYVNEQGEAKFFDTNMKQEIFD HVFKENRKVTKDKLLNYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSF LDD KANEKTIEDIIQTLTLFEDREMIRQLQKYSDI FTKAQLKKLERRHYTGWGRLSYKLINGI RNKENKKTILDYLDIDGYANRNFMLINDDALS FKEEIARAQIIDDVDDIANVVHDLPGSP AIKKGILQSVKIVDELVKVMGHN PANII IEMARENQTTDKGRRNSQQRLKLLQDSLKNLDN PVNIKNVENQQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHII PQAFIKDNSLDNR VLTRSDKNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGG LTEEDKAGF IKRQLVETRQITKHVAQILDERFNTEFDGNKRRI RNVKIITLKS NLVSNFRKEFELYKVRE INDYHHAHDAYLNAVVG NALLLKY PQLEPEFVYGEYPKYNSYRSRKSATEKFLFYSN I LRF FKKEDIQTNE DGEIAWNKEKH I KILRKVLSYPQVNIVKKTEEQTGGFSKESILPKGESDKL IPRKTKNSYWDPKKYGGFDS PVVAYSILVFADVEKGKSKKLRKVQDMVGITIMEKKRFEKN PVDFLEQRGYRNVRL EKI I KLPKYSLFELENKRRRL LASAKELQKGNELVIPQRFTTLLYH SYRIEKDYEP E HREYVEKHKDEFKELLEYISVFSRKYVLADNNLT KIEMLFSKNKDAEVSS LAKSFISLLTFTAFGAPAAFNFFGENIDRKRYTSVTECLNATLIHQ SITGLYETRIDL SKL GED (SEQ ID NO: 1).
<b><i>Streptococcus constellatus</i> Cas9 with Nuclear Localization Signal (NLS) and Linker</b>

**MPKKKRKV**GGPKPYSIGLDIGTNSVGVAVVTDDYKVPAKMKVLGNTDKQSIKKNLLGALLF  
 DSGETAEATRLKRTARRRYTRRKNRLRYLQEI FTGEMNKVDENFFQRLDDSFLVDEDKRGE  
 HHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLK  
 AENTDVQALFKDFVEEYDKTIEESHLSSEITVDALSILTEKVS KSSRLENLIAHYPTKKNNT  
 LFGNLIALSLDLHPNFKTNFQ LSEDAKLQFSKDTYEEDLEGFLGEGVDEYADLFASAKNLY  
 DAILLSGILTVDNSTKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDKNKKG  
 YASYIESGVKQDEFYKYLK GILLKINGS GDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEM  
 HAILRRQGEHY PFLKENQDKIEKILTFRIPIYYVGPLARKGSRFAWAEYKADEKITPWNFDD  
 ILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDT  
 NMKQEIFDHVFKENRKYTKDKLLNLYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKI  
 LDKSFLDDKANEKTI EDIIQTLTLFEDREMIRQLQKYSDI FTKAQLKKLERRHYTGWGR  
 SYKLINGIRNKENKKTILDYLI DDGYANRNF MQLINDDALSFKEEIARAQIIDDVDDIANV  
 VHDLPGPSPAIKKGILQSVKIVDELVKVMGHN PANIIEMARENQTTDKGRNSQQRLKLLQ  
 DSLKNLDNPVNIKNVENQQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHII PQAFI  
 KDNSLDNRVLT RSDKNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGL  
 TEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKII TLKSNLVS NFRKE  
 FELYKVREINDYHHAHDAYLNAVVG NALLKY PQLEPEFVYGEY PKYNSYRSRKSATEKFL  
 FYSNILRFFKKEDIQTNE DGEIAWNKEKH IKILRKVLSYPQVNI VKKTEEQTGGFSKESIL  
 PKGESDKLI PRKTKNSYWDPKKYGGFDS PVVAYSILVFADVEKGKSKKLRKVQDMVGITIM  
 EKRRFEKNPVDFLEQRGYRNVRL EKI IKLPKYSLFELENKRRRLLASAKELQKGNELVIPQ  
 RFTTLLYHSYRIEKDYEP EHYVEKHKDEFKELLEYSVFSRKYVLADNNLTKIEMLSK  
 NKDAEVSSLAKSFISLLTFTA FGAPAAFNFFGENIDRKRYTSVTECLNATLIHQSI TGLYE  
 TRIDL SKLGEDG**KRPAATKKAGQAKKKK**GSYPYDVPDYAYPYDVPDYAYPYDVPDYA  
 (SEQ ID NO: 2).

**Wild Type *Sharpea* Cas9**

MAKNKDIRYSIGLDIGTNSVGVAVMDEHYELLKKNHMMWGSRLFDAAEPAATRRASRSIR  
 RRYNKRREIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQKNLGNDYKDNYNLFIEK  
 DFNDKTYDYKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQSFAKVYEDI  
 EEKLDNTLKKFMSLNDLNDL FVDNDINSMITVLSKIYQRSKKADDLLKIMNPTKEERAAK  
 EFTKALVGLKFNVSKMILAQEVKKDDKDIELDFSNDYDSTVDGLQAE LGEYIEFIEMLHS  
 INSWVELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPKDYNEVFREDNPKLHNY  
 LGYIKYPKNTPV EEFY EYIKRLLAKVDTGEAREILERIDLEKFMLKQNSRTNGSIPYQMOK  
 DEMIQIIDNQSVYYYPQLKENREKLISILEFRIPY YFGPLNTHSEFAWIKKFEDKQKERILP

WNYDQIVDIDATAEGFIERMQNTGTYFPDKPVMKNSLTVSKFEVLNELNKIRINGKLI PV  
 ETKKELLSDLFMKNKTITDKKLDWLVTHQYYDTNEELKIEGYQKDLQFSTSLAPWIDFTK  
 IFGEINASNYQLIEKIIYDISIFEDKKILKRRLKVKYQLDDLLVDKILKLNVTGWSRLSEK  
 LLTGIKSKNSKETILSILENSNMNLMEIINDESLGFKQII EESNKKDIEGPF RYDEVK KLA  
 GSPAIKRGIWQALLVVQEITKFMKHEPSHIYIEFAREEQEKVRTESRIAKLQKIYKDLNLQ  
 TKEDQLVYESLKKEDAKKKIDTDALYLYYLQMGKSMYSGKPLDIDKLSTYHIDHILPRSLI  
 KDDSLDNRVVLVLPKENEWKLDSETVPPFEIRNKMMGFWQKLHENGLMSNKKFFSLIRTD FNE  
 KDKKRFINRQLVETRQIIKNVAVIINDHYTNTNVVTVRAELSHQFRERYKIYKNRDLNDLH  
 HAHDAYIACILGQFIHQNFGNMDVNMIYGQYKKNYKKDVQEHNNYGFILNSMNIHFNDN  
 SVIWDPSYIGKIKSCFCYKDVVYVTKKLEQNDAKLFDLTILPSDKNSENGVTKAKIPV NKYR  
 KDVNKYGGFSGDAPIMLAI EADKGKKHVRQVIAFPPLRLKNYNDEERIKFIEKEKNLKNVKI  
 LTEVKKNLILINHQYFFITGTNELVNATQLKLSAKNTKNLFNLVDANKHKNKLESIDDANF  
 NEVIQELICKLQEPYISRYNSIGKEFEDSYEKINAVTKQDKLYIIEYLIAIMSAKATQGYI  
 KPELAREIGTNGKNKGRIKSFTIDLNKTTFISTSVTGLFSK KYKL (SEQ ID NO: 4).

***Sharpea Cas9 with Nuclear Localization Signal (NLS) and Linker***

**MPKKKRKV**GAKNKDIRYSIGLDIGTNSVGVAVMDEHYELLKKGNNHMMWGSRLFDAEPAAT  
 RRASRSIRRRYNKRREIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQKNLGNDYKD  
 NYNLFIEKDFNDKTYDYKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQS  
 FAKVYEDIEEKLDNTLKKFMSLNDLNDLFDVNDINSMITVLSKIYQRSKKADDLLKIMNPT  
 KEERAAYKEFTKALVGLKFNVSKMILAQEVKDDKDIELDFSNVDYDSTVDGLQAE LGEYI  
 EFIEMLSINSWVELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPKDYNEVFRE  
 DNPKLHNYLGYIKYPKNTPVEEFYEYIKRLLAKVDTGEAREILERIDLEKFMKQNSRTNG  
 SIPYQMOKDEMIQIIDNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFED  
 KQKERILPWNVDQIVDIDATAEGFIERMQNTGTYFPDKPVMKNSLTVSKFEVLNELNKIR  
 INGKLI PVETKKELLSDLFMKNKTITDKKLDWLVTHQYYDTNEELKIEGYQKDLQFSTSL  
 APWIDFTKI FGEINASNYQLIEKIIYDISIFEDKKILKRRLKVKYQLDDLLVDKILKLNVT  
 GWSRLSEKLLTGIKSKNSKETILSILENSNMNLMEIINDESLGFKQII EESNKKDIEGPF R  
 YDEVK KLAGSPAIKRGIWQALLVVQEITKFMKHEPSHIYIEFAREEQEKVRTESRIAKLQK  
 IYKDLNLQTKEDQLVYESLKKEDAKKKIDTDALYLYYLQMGKSMYSGKPLDIDKLSTYHID  
 HILPRSLIKDDSLDNRVVLVLPKENEWKLDSETVPPFEIRNKMMGFWQKLHENGLMSNKKFFS  
 LIRTD FNEKDKKRFINRQLVETRQIIKNVAVIINDHYTNTNVVTVRAELSHQFRERYKIYK  
 NRDLNDLHHAHDAYIACILGQFIHQNFGNMDVNMIYGQYKKNYKKDVQEHNNYGFILNSM  
 NIHFNDNNSVIWDPSYIGKIKSCFCYKDVVYVTKKLEQNDAKLFDLTILPSDKNSENGVTKA



KIPV NKYRKDV NKYGGFSGDAPIMLAIEADKGGKHVRQVIAFPLRLKNYNDEERIKFIEKE  
 KNLKNVKILTEVKKNQLILINHQYFFITGTNELVNATQLKLSAKNTKNLFNLVDANKHNKL  
 ESIDDANFNEVIQELICKLQEPISRYNSIGKEFEDSYEKINAVTKQDKLYIIEYLIAMS  
 AKATQGYIKPELAREIGTNGKNKGRIKSF TIDLNKTTFISTSVTGLFSKKYKLGKRPAATK  
KAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 5).

**Wild Type *Veillonella parvula* Cas9**

MSIINFQRRGLMETQASNQLISSHLKGYPKDYFVGLDIGTSSVWAVTNKAYELLKFRSH  
 KMWGSRLFDEGESAVARRGFRSMRRLERRKRLRLKLEELFADAMAQVDPTFFMRLRESKY  
 HYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKFLAVHHILKY  
 RGNFLYEGATFDSNASTLDDVIKQALENITFNCFCDCNSAIISSIGQILMEAGTKSDKAKAI  
 EHLVDTYIATD TVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEEDLKK  
 LQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPGLTISESKVKAFNKHKD  
 DLAILKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETS CNREDFYKYTKKIVEGLS  
 DSKDKEYILSQIELQILLPLQRIKDNQVPIPYQLHLEELKAILAKCGPKFPFLNEVADGFSV  
 AEKLIKMLEFRIPIYYVGPLNTHHNVDNNGGFAVAVRKASGRVTPWNFDDKIDREKSAAAFIK  
 NLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNVRIDGKPLEKVVKEHLIEAVFKQDHKKM  
 TKNRIEQFLKDNQYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDIT  
 IFGESKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGT PET  
 I I ILMRNFSYNLMELLDGKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQA  
 LRIVDEVAHIKKALPARI FVEVTRSNKNEKKKDSRQKRLSDLYAAIKKDDVLLNGLNNEI  
 FGELKSSSLAKYDDAALRSKKLYLYTQMGRCAYTGEIIELSLLNTDNYDIDHIYPRSLTKD  
 DSFDNLVLCRKTANAQKSDAYPISEEIQKTQKPFWTF LKQQGLISERKYERLTRITPLTAD  
 DLSGFIARQLVETNQSVKAATTLRRLYPGVVVFVKAENVTD FRHDNNFIKVRSLNHHHH  
 AKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKD GKAWDVKTS  
 MDTVKKMMSNDVRVTKRLLEQT GALADATIYKATVAGKAKDGAYIGMKT KSSVFADVSKY  
 GGMTKIKNAYSIIVQYTGKKGEVIKEIVPLPIYL TNRNTTDQDLINYVASIIPQAKDISII  
 YGKLCINQLVKVNGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPYLVLEKFNMRKDNKDL  
 KANVVSTRALDNKHTIEVRIVEEKNI EFFDYLVSKLKMPIYQKMKGNKAAELSEKGYGLFK  
 KMSLEEQSIHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYS  
 NEVTIV (SEQ ID NO: 8).

***Veillonella parvula* Cas9 with Nuclear Localization Signal (NLS) and Linker**

**MPKKKRKV**GSIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLDIGTSSVGVAVTNKAY  
 ELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRLKLEELFADAMAQVDPTFF  
 MRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMSGTDDIRKFL  
 AVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAISSIGQILMEAGKT  
 KSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVE  
 ELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLISIKEPGLTISESKV  
 KAFNKHKDDLAILKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYT  
 KKIVEGLSDSKDKEYILSQIELQILLPLQRIKDNQVPIPYQLHLEELKAILAKCGPKFPFLN  
 EVADGFSVAEKLIKMLEFRIPYYVGPLNTHHNVDNGGFAWAVRKASGRVTPWNFDDKI DRE  
 KSAAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRI DGKPLEKVKEHLIEAV  
 FKQDHKMTKNRIEQFLKDNQYISETHKHEITGLDGEIKNDLAS YRDMVRILGDGFDRSMA  
 EEIITDITIFGESKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDK  
 AGDGTPETIIILMRNFSYNLMELLDGKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPA  
 VKRAVWQALRIVDEVAHIKKALPARI FVEVTRSNNKNEKKKDSRQKRLSDLYAAIKDDVL  
 LNGLNNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGRCAYTGEI IELSLNNTDNYDIDHI  
 YPRSLTKDDSDFNLVLCRRTANAQKSDAYPISEEIQKTQKPFWTFLKQQGLISERKYERLT  
 RITPLTADDLSGFIARQLVETNQS VKAATTLRRLYPGVVVFVKAENVTD FRHDNNFIKV  
 RSLNHHHAKDAYLNI VVGNVYHERFTRNFRAFFKNGANRTYNLAKMFNYDVNCTNAKDG  
 KAWDVKTSMDTVKKMMSNDVVRVTKRLLEQT GALADATIYKATVAGKAKDGAYIGMKT KSS  
 VFADVSKYGGMTKIKNAYSII VQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIP  
 QAKDISIIYGKLCINQLVKVNGFY YLGGKTNSKFCIDNAIQVIVSNEWIPYLVLEKFNN  
 MRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNI EFFDYLVSKLKMPIYQMKGNKAAELS  
 EKG YGLFKKMSLEEQS IHLIELLNLLTNQKTTFEVKPLGITASRSTVSGKISNQDEFKVIN  
 ESITGLYSNEVTIVKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA  
 (SEQ ID NO: 9).

**Wild Type *Ezakiella peruensis* Cas9**

MTKVKDYYIGLDIGTSSVGVAVTDEAYNVLFNSKKMWGVRLFDDAKTAEERRGQRGARRR  
 LDRKKERLSLLQDFFAEVAKVDPNFFLRDLNSDLYMEDKDQKLKSKYTLFNDKDFKDNF  
 HKKYPTIHHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIFEGQKFDTKSSFENSLNELK  
 VHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQKLV  
 DLFENPEDFDDSAIKSVDFSTTSFDDKYS DYELALGDKIALVNI LKEIYDSSILENLLKEA  
 DKSKDGNKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNNDNYVSYTKSSI  
 SNNKRVKANKFTDQEA FYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKI

KSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRIPYYVGPLNP  
 NSKYAWIKKQKDSEITPWNFKDVVDLSSREEFIDSLIGRCTYLKDEKVLPAKASLLYNEYM  
 VLNELNNLKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGD  
 FKQGLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGKYFTDSE  
 IKKMAGLNYKDWGRLSKKLLTGLEGANKITGERGSI IHFMREYNLNLMELEMSASFTFTEEI  
 QKLNVPVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIE MARGKEEV  
 KARKE SRKNQLLKFYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQLGRC  
 MYSLEPIDISELSSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPIPDEILNK  
 NCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTICK  
 NSEIVYSKAENASRFRQEFDIVKCRVNDLHHMHDAYINIIVGNVYNTKFTKDPMNPFVKKQ  
 EKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFRMNYIESG  
 ALFNATLQRKNKGSRPLKDKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLMFVERE  
 IYAKQKNDKKSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITGGSDKTLEL  
 RSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKS F  
 LNLFQDDKIENIDFKFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGRLLSSKTN  
 NIGNYQEFKIINQSITGLFENEVDLLKL (SEQ ID NO: 14).

***Ezakiella peruensis* Cas9 with Nuclear Localization Signal (NLS) and Linker**

**MPKKRKV**GTGVKDYIIGLDIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAEEER  
 GQRGARRRLDRKKERLSLLQDFFAEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFND  
 KDFKDNFHKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFI FEGQKFDTKSSF  
 ENSLNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIM  
 IGSSQKLVDLFENPEDFDDSAIKSVDFSTTSFDDKYSYELALGDKIALVNILKEIYDSSI  
 LENLLKEADKSKDGNKYISNAFVKYKXKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNNDNY  
 VSYTKSSISNNKRVKANKFTDQEA FYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDME  
 FKNFMPKIKSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRIP  
 YYVGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLSSREEFIDSLIGRCTYLKDEKVLPAK  
 SLLYNEYMVLNELNNLKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGE I  
 LLSGTDGDFKQGLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGY  
 GKYFTDSEIKKMAGLNYKDWGRLSKKLLTGLEGANKITGERGSI IHFMREYNLNLMELEMSA  
 SFTFTEEIQKLNVPVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIE  
 MARGKEEVKARKE SRKNQLLKFYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYL  
 YYTQLGRCMYSLEPIDISELSSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSY P  
 IPDEILNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETA

NLLKTI CKNSEIVVYSKAENASRFRQEFDIVKCRAVNDLHHMHDAYINI IVGNVYNTKFTKD  
 PMNFVKKQEKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFT  
 RMNYIESGALFNATLQRKNKGSRPLKDKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIER  
 KLMPVEREIYAKQKNDKCLSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITG  
 GSDKTLELRSALQLILPKKNEWAIKQIDKSSSENDYLTIERIQDLTEELVYNTFDIIVNKFK  
 TSVFKKSFNLNFQDDKIENIDFKFKSMDFKKCKTLLMLVKAIRASGVRQDLKSIDLKSDY  
 GRLSSKTNIGNYQEFKIINQSI TGLFENEVDLLKLGKRPAATKKAGQAKKKKGSYPYDVP  
 DYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 15).

**Wild Type *Lactobacillus fermentum* strain AF15-40LB Cas9**

MKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAERRTFRTRRRRLKR  
 RKWRLHYLDEIFAPHLQEVNENFLRRLKQSNIHPEPDAKNQAFIGKLLFPDLLKKNERGYP  
 TLIKMRDELVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAHVHIVKYRGHFLNNASV  
 DKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLDROKAVAK  
 LLEVKVADKEETKRNKQIATAMSKLVLYKADFATVAMANGNEWKIDLSSETSEDEIEKFR  
 EELSDAQNDILTEITSLSFSQIMLNEIVPNGMSISESMMDRYWOTHERQLAEVKEYLATQPAS  
 ARKEFDQVYNKYIGQAPKEKGFDELEKGLKKILSKKENWKEIDELKAGDFLPKQRTSANGV  
 IPHQMHOQELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPYYVGPLVTP  
 EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKDITYLLNEDVLPANS  
 LLYQKYNVLNELNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMAKTRGVNKPS  
 VEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVE  
 WLTDEQRSALVKKRYKGGWRLSKLLTGIVDENGQRIIDLWNTDQNFMQIVNQPVFKEQI  
 DQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSSISIEF  
 ARNEGNKGEITRSRRTQLQKLFEDQAHELVKDTSLTEELEKAPDLSDRYYFYFTQGGKDMY  
 TGDPINFDEISTKYDIDHILPQS FVKDDSLDNRVLVSRAENNKSDRVPKLYAAKMKPYW  
 NQLLKQGLITQRKFENLTMDVDQTIKYRSLGFVKRQLVETRQVIKLTANILGSMYQEAGTD  
 I IETRAGLTQQLREEFDLPKVREVNDYHHAVDAYLTTFAGQYLNRRYPKLRSEFFVYGEYMK  
 FKHGSDLKLRNFNFHHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLISNE  
 TYEETGKYFDASHESSESLYLKNQNKSKLVVPLKNKLQPEYYGAYTGITQGYMVILKLLDK  
 KGGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKSEVVVPRVLKGTFLVDGE  
 EKFILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLVEMQTKKLIETIDNIRQVDKYY  
 KLYDINKFRAKLHDGRSKFVELDDFGQDASKEKVI I KILRGLHFGSDLQNLKEIGFGTTP  
 GQFQVSEAGIRLSNTAFIIFKSPTGLFNRKLYLKNL (SEQ ID NO: 84).

***Lactobacillus fermentum* strain AF15-40LB Cas9 with Nuclear Localization Signal (NLS) and Linker**

**MPKKRKRKV**GKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTA AERRTFR  
TTRRRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEPAKNQAFIGKLLFPDLL  
KKNERGYPTLIKMRDELPVEQRAHYPVTNIYKLRAMINEDRQFDLREVYLA VHHIVKYRG  
HFLNNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKL  
DRQKAVAKLLEVKVADKEETKRNKQIATAMSKLVLGKADFATVAMANGNEWKIDLSSETS  
EDEIEKFREELSDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWTH ERQLAEVKE  
YLATQPASARKEFDQVYNKYIGQAPKEKGFLEKGLKKILSKKENWKEIDELLKAGDFLPK  
QRTSANGVI PHQMHQEELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPY  
YVGPLVTPEVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKD TYLLN  
EDVLPANSLLYQKYNVLNELNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVM AK  
TRGVNKPSVEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIF  
ADKLTEVEWLTDEQRSALVKKRYKGWGRLSKLLTGIVDENGQRIIDL MWNTDQNFMQIVN  
QPVFKEQIDQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNA  
PKSISIEFARNEGKGEITRSRRTQLQKLFEDQAHELVKDTSLTEELEKAPDLSDRYYFYF  
TQGGKDMYTGDPIINFDEISTKYDIDHILPQS FVKDDSLDNRVLVSR AENNKSDRVPKLY  
AAKMKPYWNQLLKQGLITQRKFENLTMDVDQTIKYRSLGFVKRQLVETRQVIKLTANILGS  
MYQEAGTDI IETRAGLTKQLREEFDLPKRVENDYHHAVDAYLTTFAGQYLNRRYPKLR SF  
FVYGEYMKFKHGSDLKLRNFNFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINL  
KVMLISNETYEETGKYFDASHESSLYLKNQNKSKLVVPLKNKLQPEYYGAYTGITQGYM  
VILKLLDKKGGFGVYRIPRYAADI LNKCHDEVAYRNKIAEIISSDPRAPKSF EVVVPRLK  
GTFLVDGEEKFILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLVNEMQTKKLI EIYDNI  
LRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEKVI I KILRGLHFGSDLQNLKE  
IGFGTTPLGQFQVSEAGIRLSNTAFIIFKSP TGLFNRKLYLKNL

GKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 85).

**Wild Type *Peptoniphilus* sp. Marseille-P3761 Cas9**

MEKKTNYTIGLDIGTDSVGVAVVKDDLELVKKRMKVLGNTETNYIKKNLWGSLLFESGQTA  
KDRRLKRVARRRYERRRNRLTELQKIFAPAIDEV DENFFFRLNESFLVPEDKAFSKNPIFG  
TLGEDKTYKYTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIAI  
NENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDIGRANKQS  
PMAFITLIVGKKAKFKAAFNLEEEISLNLTDSDYDENLEILLNTIGSDFADLFDHAQRVY

NAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQ  
 KKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQ  
 VHLAEMEAILDRQENYYPWLKENREKIISLLTFRIPIYYVGPLADGQSEFAWLERKSDEKIK  
 PWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIAYLDERQK  
 RMNLS SVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELA  
 KVPGMKSMDDPDNEDLMEEIVKILTVEFDRKMRRKQLEKYKERLSPEQIKELAKKHGTGW  
 GRLSKLLVGIIRDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDDRLSFKKTTIAELQM  
 IDPSADLYAQVQEIAGSPAIKKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTTARGKA  
 LSKRREAKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQY  
 DEDHIIPQSFTVDNSLDNLVLTNS SQNRGNKKDDVPSLEVVRQLAYWRS LK DAGLMTQRK  
 FDNLT KAMRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNIVL  
 LKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAAIAINLLALYPYMA DDFVYGEFRYKKK  
 PQA EKATYEKLRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNVVKA EER  
 TGMLFKETINGKTNKGSRIPIKKDLDP SKYGGYIEEKMAYYAVISYEDKKKKPGKTIVGIS  
 IMDKKEFEYDSISYLGKLGFSNPVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKA  
 NQIAMEQDLVNFIIYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINR  
 LRRIYEEEEKKDPVDIEALVASFIELLKLTSAGAPADFI FMGEAISRRRYNSMTGLFDGQV  
 IYQSLTGLYETRMRFED (SEQ ID NO: 86).

***Peptoniphilus sp. Marseille-P3761 Cas9 with Nuclear Localization Signal (NLS) and Linker***

**MPKKKRKV**GEKKTNYTIGLDIGTDSVGVAVVKDDLELVKKRMKVLGNTETNYIKKNLWGS L  
 LFESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEV DENFFRNLNESFLVPEDKA  
 FSKNPIFGTLGEDKTYKYPTIYHLRQH LADSEEKADVRLIYLALAHMIKYRGHFLIEGK  
 LDTEHIAINENLEQFFESYNALFSEEP IELRKEELIAIENILREKNSRTVKEKRITSFLKD  
 IGRANKQSPMAFITLIVGKKAKFKAAFNLEEEISLNLTD DSYDENLEILLNTIGSDFADL  
 FDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMT  
 FVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSF  
 YNGVIPNQVH LAEMEAILDRQENYYPWLKENREKIISLLTFRIPIYYVGPLADGQSEFAWLE  
 RKSDEKIKPWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKI  
 AYLDERQKRMNLS SVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNAD  
 YSTYVELAKVPGMKSMDDPDNEDLMEEIVKILTVEFDRKMRRKQLEKYKERLSPEQIKEL  
 AKKHGTGWGRLSKLLVGIIRDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDDRLSFK  
 KTTIAELQMIDPSADLYAQVQEIAGSPAIKKGILLGLKIVDEIIRVMGEKPENIVIEMAREN

QTTARGKALSKRREAKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDPEL  
DFDRLSQYDEDHIIPQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEVVNRQLAYWRS LKD  
AGLMTQRKFDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNK  
IRETNIVLLKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAAIAINLLALYPYMA DDFVY  
GEFRYKKKPQAEKATYEKLRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQVN  
VVKKAERTGMLFKETINGKTNKGSRIPIKKDLDP SKYGGYIEEKMAYYAVISYEDK KKKP  
GKTIVGISIMDKKEFEYDSISYLGKLGFSNPV VQIILKNYSLIAYPDGRRRYITGATKTTK  
GKVELQKANQIAMEQDLVNFYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFL  
DAETNINRLRRIYEEEEKKKDPVDIEALVASFIEL LKLT SAGAPADFI FMGEAISRRRYNSM  
TGLFDGQVIYQSLTGLYETRMRFEDGKRPAATKKAGQAKKKKGS*YPYDVPDYAYPYDVPDY*  
*AYPYDVPDYA* (SEQ ID NO: 87).

NLS (bold), can be substituted with different NLSs

Linker (underlined), can be removed or extended

3xHA tag (italics), can be substituted with different tags

In some embodiments, the Cas9 protein comprises one or more mutations in reference  
5 to SEQ ID NO: 1, 4, 8,14, 84 or 86. For example, the amino acid sequence of the Cas9  
protein comprises at least one, at least two, at least three, at least four, at least five, at least  
six, at least seven, at least eight, at least nine, at least 10 mutations in SEQ ID NO: 1, 4, 8, 14,  
84 or 86. Various mutations are known in the art, and include for example, amino acid  
substitutions.

10 In some embodiments, two or more catalytic domains of Cas9 (RuvC1, RuvCII,  
RuvCIII) are mutated to produce an inactive, or “dead” Cas9 (dCas9) that lacks nucleic acid  
cleavage activity. In some embodiments, the one or more mutations are in the PAM  
Interacting, HNH, and or the RuvC domains. In some embodiments, Cas9 is mutated to  
reduce DNA cleavage activity to less than about 25%, 15%, 10%, 5%, 1%, 0.1%, 0.01% or  
15 lower with respect to its non-mutated form.

In some embodiments a nickase-mutant version of Cas9 is provided. In some  
embodiments, the nickase mutant has one or more amino acid substitutions in the RuvC  
and/or the HNH domains. Various nickase mutations are known with respect to SpCas9  
(Streptococcus pyogenes) and include for example mutations at one or more of amino acid  
20 positions 10, 12, 17, 762, 840, 854, 863, 982, 983, 984, 986, 987 of wild type SpCas9. For  
example, an aspartic acid-to-alanine substitution that corresponds to D10A in SpCas9 results

in the creation of a nickase. In some embodiments, the Cas9 described herein has one or more mutations that result in the creation of a nickase. In some embodiments, the Cas9 described herein has one or more mutations at an amino acid position that corresponds to one or more of amino acids 10, 12, 17, 762, 840, 854, 863, 982, 983, 984, 986, 987 of SpCas9.

5 In some embodiments, the mutation is an aspartic acid-to-alanine substitution (D10A) in the RuvC domain of ScoCas9. In some embodiments, the mutation is an aspartic acid-to-alanine substitution (D14A) in the RuvC domain of SirCas9. In some embodiments, the mutation is an aspartic acid-to-alanine substitution (D38A) in the RuvC domain of VapCas9 (e.g., corresponding to D10A in SpCas9). In some embodiments, the mutation is an  
10 aspartic acid-to-alanine substitution (D12A) in the RuvC domain of EpeCas9. In some embodiments, the mutation is an aspartic acid-to-alanine substitution (D9A) in the RuvC domain of LfeCas9. In some embodiments, the mutation is an aspartic acid-to-alanine substitution (D12A) in the RuvC domain of PmaCas9.

In some embodiments, the mutation is an aspartic acid-to-glycine substitution (D10G)  
15 in the RuvC domain of ScoCas9. In some embodiments, the mutation is an aspartic acid-to-glycine substitution (D14G) in the RuvC domain of SirCas9. In some embodiments, the mutation is an aspartic acid-to-glycine substitution (D38G) in the RuvC domain of VapCas9 (e.g., corresponding to D10G in SpCas9). In some embodiments, the mutation is an aspartic acid-to-glycine substitution (D12G) in the RuvC domain of EpeCas9. In some embodiments,  
20 the mutation is an aspartic acid-to-glycine substitution (D9A) in the RuvC domain of LfeCas9. In some embodiments, the mutation is an aspartic acid-to-glycine substitution (D12G) in the RuvC domain of PmaCas9.

In some embodiments, such one or more mutations described herein converts Cas9 to an inactive, or “dead” version of Cas9 (dCas9). Accordingly, in some embodiments, the  
25 Cas9 protein comprises one or more mutations that inhibits the ability of Cas9 to cleave both strands of a DNA duplex.

In some embodiments, when coexpressed with a guide RNA, dead Cas9 generates a DNA recognition complex that can specifically interfere with transcriptional elongation, RNA polymerase binding, or transcription factor binding. In some embodiments, dead Cas9  
30 is used to specifically target effector proteins of various functions to specific nucleic acid target sites.



In some embodiments, a high-fidelity Cas9 variant comprises enhanced specificity, which minimizes off-target cleavage. In some embodiments, engineered variants, for example, ‘hyper-accurate Cas9’ (N692A, M694A, Q695A and/or H698A mutations corresponding to SpyCas9) and/or ‘high-fidelity Cas9’ (N467A, R661A, Q695A and/or Q926A mutations corresponding to SpyCas9) are used which comprise mutations mainly within the REC3 domain and achieve higher specificity and fidelity. High-fidelity variants reduce the capacity of Cas9 to stabilize mismatches and reduce off-target DNA cleavage. In some embodiments, the increase in specificity is accompanied by a loss in efficiency of on-target cleavage by about 100 fold. In some embodiments, a SuperFi-Cas9 is used, which is a high-fidelity variant that maintains on-target cleavage rates comparable to wild-type Cas9. In some embodiments, the SuperFi-Cas9 comprises mutations in the RuvC loop. In some embodiments, the mutations inhibit formation of a kinked conformation that facilitates subsequent cleavage of gRNA-TS duplex. In some embodiments, the Y1016, R1019, Y1010, Y1013, K1031, Q1027 and/or V1018 residues corresponding to SpyCas9 are mutated, for example, to aspartic acid. (Bravo, J. *et al.* Structural basis for mismatch surveillance by CRISPR-Cas9 *Nature*, 603, March 2022).

The engineered, non-naturally occurring Cas9 is has an amino acid sequence at least 80% (e.g., 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more) identical to a Cas9 amino sequence at SEQ ID NOs. 2, 5, 9, 15, 85, 87, 95, or 96.

In some embodiments, the engineered non-naturally occurring Cas9 is encoded in a nucleic acid molecule codon-optimized for human cells (e.g., codon optimized for expression, stability, etc.).

Exemplary Cas9 sequences with Nuclear Localization Signal (NLS) and a linker is provided in Table 2 below.

**Table 2. Exemplary Cas9 Sequence with NLS and Linker**

<b>Sequence of <i>Sco</i>Cas9 with Nuclear Localization Signal (NLS) and Linker</b>
MPKKRRKVGGKPYSIGLDIGTNSVGVAVVTDDYKVPAAKMKVLGNTDKQSIKKNLLGALLF DSGETAEATRRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDNFLVDEDKRGE HHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLK AENTDVQALFKDFVEEYDKTIEESHLSSEITVDALSILTEKVS KSSRLENLIAHYPTTEKKNT

LFGNLIALS LD LHPNF KTNFQLSEDAKLQFSKDTYEEDLEGFLGVEGDEYADLFASAKNLY  
 DAILLSGILTVDDNSTKAPLSASMVKRYEEHQKDLKCLKDFIKVNAPDQYNAI FKDKNKKG  
 YASYIESGVKQDEFYKYLK GILLKINGS GDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEM  
 HAILRRQGEHY PFLKENQDKIEKILTFRI PYYVGPLARKGSRFAWAEYKADEKITPWNFDD  
 I LDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFVYNELTKVKYVNEQGEAKFFDT  
 NMKQEIFDHVFKENRKVTKDKLLNYLNKEFEFRIVNLTGLDKENKAFNSSLGTYHDLRKI  
 LDKSFLDDKANEKTI EDIIQTTLTFEDREMIRQRLQKYSDI FTKAQLKKLERRHYTGWRL  
 SYKLINGIRNKENKKTILDYLI DDGYANRNF MQLINDDALS FKEE IARAQI ID DVDDIANV  
 VHDLPGSPA I KKGILQSVKIVDELVKVMGHN PANI I IEMARENQT TDKGRNSQORLKLLO  
 DSLKNLNDNPVNIKNVENQQQLQNDRLF LYI QNGKDMYTGETLDINNLSQYDIDHIIPQAFI  
 KDNSLDNRVLT RSDKNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGL  
 TEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRI RNVKIITLKS NLVSNFRKE  
 FELYKVREINDYHHAH DAYLNAVVG NALLLKY PQLEPEFVYGEY PKYNSYRSRKSATEKFL  
 FYSNILRFFKKEDIQT NEDGEI AWNKEKH I KILRKVLSYPQVNI VKKTEEQTGGFSKESIL  
 PKGESDKLI PRKTKNSYWDPKKYGGFDS PVVAYSILVFADVEK GKSKKLRKVQDMVGITIM  
 EKRF EKNPVDFLEQRGYRNVRL EKI I KLPKYSLFELENKRRRL LASAKELQKGNELVIPQ  
 RFTTLLYHSYRIEKDYEP EHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMLF SK  
 NKDAEVSSLAKSFISLLTFTA FGAPAAFNFFGENIDRKRYTSVTECLNATLIHQ SITGLYE  
 TRIDLSKLGEDG KRPAATKKAGQAKKKKGS YPYDVPDYAYPYDVPDYAYPYDVPDYA  
 (SEQ ID NO: 2).

**Sequence of *Sharpea Cas9* with Nuclear Localization Signal (NLS) and Linker**

**MPKKRRKV** GAKNKDIRYSIGLDIGTNSVGVAVMDEHYELLKKG NHHMWGSRLFDAEPAAT  
 RRASRSIRRRYNKR RERIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQK NLGNDYKD  
 NYNLFIEKDFNDKTY YDKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQS  
 FAKVYEDIEEKLDNTLKKFMSLNDLDNLFVDNDINSMITVLSKIYQRSKKADDLLKIMNPT  
 KEERAAYKEFTKALVGLKFNVSKMILAQEVKDDKDIELDFSNVDYDSTVDGLQAE LGEYI  
 EFIEMLSIN SWVELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPKYNEVFRE  
 DNPKLHNYLGYIKYPKNTPVEEFY EYIKRLLAKVDTGEAREILERIDLEKFM LKQNSRTNG  
 SIPYQMOKDEMIQIIDNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFED  
 KQKERILPWN YDQIVDIDATAEGFIERMQNTGTYPDPKPVMAKNSLTVSKFEVLNENLKNIR  
 INGKLIPVETK KELLSDLFMKNKTI TDKCLKDWLVTHQYYDTNEELKIEGYQKDLQFSTSL  
 APWIDFTKI FGEINASNYQLIEKI IYDISIFEDKKILKRRLK KVVYQLDDLLVDKILKLN YT  
 GWSRLSEKLLTG IKS KNSKETILSILENSNMNLMEI INDESLGFKQIIEESNKKDIEGPF R

YDEVKLAGSPAIKRGIWQALLVVQEITKFMKHEPSHIYIEFAREEQEKVRTESRIAKLQK  
 IYKDLNLQTKEDQLVYESLKKEDAKKKIDTDALYLYYLQMGKSMYSGKPLDIDKLSTYHID  
 HILPRSLIKDSDLDNRVVLVLPKENEWKLDSETVFPFEIRNKMMGFWQKLHENGLMSNKKFFS  
 LIRTFNEKDKKRFINRQLVETRQIIKNVAVIINDHYTNTNVVTVRAELSHQFRERYKIYK  
 NRDLNDLHHAHDAYIACILGQFIHQNFGNMDVNMIYGQYKKNYKKDVQEHNNYGFI LNSMN  
 HIFNDNSVIWDPSYIGKIKSCFCYKDVYVTKKLEQNDAKLFDLTILPSDKNSENGVTKA  
 KIPVNKYRKDVNKYGGFSGDAPIMLAIEADKGGKHVRQVIAFPLRLKNYNDEERIKFIEKE  
 KNLKNVKILTEVKKNQILILINHQYFFITGTNELVNATQLKLSAKNTKNLFNLVDANKHNKL  
 ESIDDANFNEVIQELICKLQEPISRYNSIGKEFEDSYEKINAVTKQDKLYIEYLIAIMS  
 AKATQGYIKPELAREIGTNGKNKGRIKSFIDLNKTTFISTSVTGLFSKKYKLGKRPAATK  
KAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 5).

**Sequence of *Veillonella parvula* Cas9 with Nuclear Localization Signal (NLS) and Linker**

**MPKKKRKV**GSIIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLDIGTSSVGWAVTNKAY  
 ELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRLKLEELFADAMAQVDPPTFF  
 MRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELKSGTDDIRKFLFL  
 AVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAISSIGQILMEAGKT  
 KSDKAKAIEHLVDTYIATDVTDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVE  
 ELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPGLTISESKV  
 KAFNKHKDDLAILKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYT  
 KKIVEGLSDSKDKEYILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLN  
 EVADGFSVAEKLIKMLEFRIPIYVGPLNTHHNVDNNGGFAWAVRKASGRVTPWNFDDKIDRE  
 KSAAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNVRIDGKPLEKVVKEHLIEAV  
 FKQDHKKMTKNRIEQFLKDNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMA  
 EEIITDITIFGESKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDK  
 AGDGTPETIIILMRNFSYNLMELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALS PA  
 VKRAVWQALRIVDEVAHIKKALPARIFVEVTRSNNKNEKKKKDSRQKRLSDLYAAIKKDDVL  
 LNGLNNEIFGELKSSLAKYDDAALRSKKLYLYTQMGRCAYTGEIIELSLLNTDNYDIDHI  
 YPRSLTKDDSDNLVLCRTANAQKSDAYPISEEIQKTQKPFWTFLKQQGLISERKYERLT  
 RITPLTADDLSGFIARQLVETNQSVAATTLRRLYPGVDVVFVKAENVTD<sup>R</sup>FRHDNNFIKV  
 RSLNHHHAKDAYLNIIVGNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKDG  
 KAWDVKTSMDTVKKMMSDNDVRVTKRLLLEQTGALADATIYKATVAGKAKDGAYIGMKTSS

VFADVSKYGGMTKIKNAYSIIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIP  
 QAKDISIIYGKLCINQLVKVNGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPYLVLEKFNN  
 MRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLVSCLKMPIYQMKMGNKAAELS  
 EKGYGLFKKMSLEEQS IHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVIN  
 ESITGLYSNEVTIVGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA  
 (SEQ ID NO: 9).

**Sequence of *Ezakiella peruensis* Cas9 with Nuclear Localization Signal (NLS) and Linker**

**MPKKKRKV**GTKVKDYYIGLDIGTSSVGWAVTDEAYNVLFNSKMKMWGVRLFDDAKTAEERR  
 GQRGARRRLDRKKERLSLLQDFFAEVAKVDPNFFLRDLSLDLYMEDKDQKLKSKYTLFND  
 KDFKDNFHKKYPTIHHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFI FEGQKFDTKSSF  
 ENSLNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIM  
 IGSSQKLVDLFENPEDFDDSAIKSVDFSTTSFDDKYSYELALGDKIALVNILKEIYDSSI  
 LENLLKEADKSKDGNKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNNDNY  
 VSYTKSSISNNKRVKANKFTDQEAIFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDME  
 FKNFMPKIKSSDNGVIPPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRIP  
 YYVGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLDSREEFIDSLIGRCTYLKDEKVLPA  
 SLLYNEYMVLNELNKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEI  
 LLSGTDGDFKQGLNSYNDFKAIIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGY  
 GKYFTDSEIKKMAGLNYKDWGRLSKLLTGLEGANKITGERGSI IHFMREYNLNLMELEMSA  
 SFTFTEEIQKLNVPDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIE  
 MARGKEEVKARKESRKNQLLKFYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYL  
 YYTQLGRCMYSLEPIDISELSSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYP  
 IPDEILNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETA  
 NLLKTICKNSEIVYSKAENASRFRQEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKD  
 PMNFVKKQEKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFT  
 RMNYIESGALFNATLQRKNKGSRPLKDKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIER  
 KLMPVEREIIYAKQKNDKLSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITG  
 GSDKTLELRSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFK  
 TSVFKKSFNLNFQDDKIENIDFKFKSMDFKCKTLLMLVKAIRASGVRQDLKSIDLKS DY  
 GRLSSKTNNIGNYQEFKIINQSITGLFENEVDLLKLG**KRPAATKKAGQAKKKKGS**YPYDVP  
 DYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 15).

**Sequence of *Lactobacillus fermentum* strain AF15-40LB Cas9 with Nuclear Localization Signal (NLS) and Linker**

**MPKKKRKV**GKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTA AERRTFR  
TTRRRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEDEPAKNQAFIGKLLFPDLL  
KKNERGYPTLIKMRDELPVEQRAHYPVTNIYKLCREAMINEDRQFDLREVYLAVHHIVKYRG  
HFLNNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKL  
DRQKAVAKLLEVKVADKEETKRNKQIATAMSKLVLGKADFATVAMANGNEWKIDLSSETS  
EDEIEKFREELESDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWOTHERQLAEVKE  
YLATQPASARKEFDQVYNKYIGQAPKEKGFDEKGLKKILSKKENWKEIDELKAGDFLPK  
QRTSANGVI PHQMHQOELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSRIPY  
YVGPLVTPVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAFAFIKRM TVKDTYLLN  
EDVLPANSLLYQKYNVLNELNNVRVNGRRLSVGIKQDIYTELFKKKTKVAGDVASLVMK  
TRGVNKPSVEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIF  
ADKLTEVEWLTDEQRSALVKKRYKGGWRLSKLLTGIVDENGQRIIDLWNTDQNFMQIVN  
QPVFKEQIDQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNA  
PKSISIEFARNEGKGEITRSRRTQLQKLFEDQAHELKDTSLTEELEKAPDLSDRYYFYF  
TQGGKDMYTGDPIINFDEISTKYDIDHILPQS FVKDDSLDNRVLVSR AENNKSDRVPKLY  
AAKMKPYWNQLLKQGLITQRKFENLTMDVDQTIKYRSLGFVKRQLVETRQVIKLTANILGS  
MYQEAGTDI IETRAGLTKQLREEFDLPKRVENDYHHAVDAYLTTFAGQYLNRRYPKLRSF  
FVYGEYMKFKHGS DLKLRNFNFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINL  
KVMLISNETYEETGKYFDASHESSLYLKNQNKSKLVVPLKNKLQPEYYGAYTGITQGYM  
VILKLLDKKGGFGVYRIPRYAADI LNKCHDEVAYRNKIAEIISSDPRAPKSEVVVPRVLK  
GTFLVDGEEKFILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLVNEMQTKKLIEIYDNI  
LRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEKVI I KILRGLHFGSDLQNLKE  
IGFGTTPLGQFQVSEAGIRLSNTAFIIFKSP TGLFNRKLYLKNL

GKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 85).

**Sequence of *Peptoniphilus* sp. Marseille-P3761 Cas9 with Nuclear Localization Signal (NLS) and Linker**

**MPKKKRKV**GEKKTNYTIGLDIGTDSVGVAVVKDDLELVKKRMKVLGNTETNYIKKNLWGS  
LFESGQTAKDRRLKRVARRRYERRRNLTELQKIFAPAIDEVDENFFRLNESFLVPEDKA  
FSKNPIFGTLGEDKTYKYTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGK  
LDTEHIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKD

IGRANKQSPMMAFITLIVGKKAKFKAAFNLEEEI SLNLTDDSYDENLEILLNTIGSDFADL  
 FDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMT  
 FVAPKDAQKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSF  
 YNGVIPNQVHLAEMEAILDRQENYYPWLKENREKIIISLLTFRIPYVGPLADGQSEFAWLE  
 RKSDEKIKPWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKSLEYQKYMVFNELTKI  
 AYLDERQKRMNLSVEKKEIFETLFFKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNAD  
 YSTYVELAKVPGMKSMDDPDNEDLMEEIVKILTVFEDRKMRRKQLEKYKERLSPEQIKEL  
 AKKHYTGWGRLSKKLLVGIIRDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDDRLSFK  
 KTIAELQ MIDPSADLYAQVQEIAGSPAIIKKGILLGLKIVDEIIRVMGEKPENIVIEMAREN  
 QTTARGKALSKRREAKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPL  
 DFDRLSQYDEDHII PQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEVVRQLAYWRS LKD  
 AGLMTQRKFDNLT KAMRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNK  
 IRETNI VLLKSAMASEFRKMFRLYK VRELNDYHHAHDAYLNAAIAINLLALYPYMADDFVY  
 GEFYK KKPQAEKATYEKLRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQVN  
 VVKKAEERTGMLFKETINGKTNKGSRIPIK KDLDP SKYGGYIEEKMAYYAVISYEDK KKKP  
 GKTIVGISIMDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIAYPDGRRRYITGATKTKK  
 GKVELQKANQIAMEQDLVNFYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFL  
 DAETNINRLRIYEEKKKDPVDIEALVASFIELLKLTSAGAPADFI FMGEAISRRRYNSM  
 TGLFDGQVIYQSLTGLYETRMRFEDGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDY  
 AYPYDVPDYA (SEQ ID NO: 87).

**Sequence of *ScoCas9* variant with Nuclear Localization Signal (NLS) and Linker  
 (*ScoCas9*-NGC-v1)**

**MPKKKRKVG**MGKPYSIGLDIGTNSVGWAVVTDDYKVPAKMKVLGNTDKQS I KKNLLGALL  
 FDSGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSFVDEDKRG  
 EHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDL  
 KAENTDVQALFKDFVEEYDKTIEESHLS EITVDALSILTEKVS KSSRLENLIAHYPT EKKN  
 TLFGNLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGEYADLFASAKNL  
 YDAILLSGILTVDDNSTKAPLSASMVKRYEEHQDLK KLDKDFIKVNAPDQYNAIFKDKNKK  
 GYASYIESGVKQDEFYKYLK GILLKINGS GDFLDKIDREDFLRKQRTFDNGSIPHQIHLQE  
 MHAILRRQGEHY PFLKENQDKIEKILTFRIPYVGPLARKGSRFAWAEYKADEKITPWNFD  
 DILDKEKSAEKFITRMTLNDLYLP EEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFD  
 TNMKQEIFDHVFKENRKVTKDKLLNYLNKEFEFRIVNL TGLDKENKAFNSSLGTYHDLRK  
 ILDKSFLDDKANEKTI EDIIQTLTLFEDREMIRQLRQKYS DIFTKAQLKKLERRHYTGWGR

LSYKLINGIRNKENKKTILDYLIIDGYANRNFQMLINDDALSFKEEIARAQIIDDVDDIAN  
 VVHDLPGSPAIAKKGILQSVKIVDELVKVMGHNPANIIEMARENQTTDKGRRNSQQRLKLL  
 QDSLKNLNDNPNVNIKNVENQQQLQNDRLFLLYYIQNGKDMYTGETLDINNLSQYDIDHIIIPQAF  
 IKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGG  
 LTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIIRNVKIITLKSNLVSNFRK  
 EFELYKVREINDYHHAHDAYLNAVVGNALLLKYPQLEPEFVYGEYPKYNSYRSRKSATEKF  
 LFYSNILRFFKKEDIQTNEEDGEIAWNKEKHKILRKVLSYPQVNIKKTEEQTGGFSKESI  
 LPKGESDKLIPRKTKNYSWDPKKYGGFMQPVVAYSILVFADVEKGGKSKLRKVQDMVGITII  
 MEKKRFEKNPVDFLEQRGYRNVRLKIIKLPKYSLFELENKRRRLASAKFLQKGNELVIP  
 QRFITLLYHSYRIEKDYEPHEHREYVEKHKDEFKELLEYSVFSRKYVLADNNLTKIEMLFS  
 KNKDAEVSSLAKSFISLLTFTAAGAPRAFNFGENIARKEYRSVTECLNATLIHQSIITGLY  
 ETRIDL SKLGEDGE**EGADKRTADGSEFESPKKKRKV** (SEQ ID NO: 95)

**Sequence of *ScoCas9* with Nuclear Localization Signal (NLS) and Linker (*ScoCas9*-*NGC-v2*)**

**MPKKKRKVG**MGKPYSIGLDIGTNSVGWAVVTDDYKVPKMKVGLGNTDKQSIKKNLLGALL  
 FDSGETAEATRLKRTARRRYTRRKNRLRYLQEI FTGEMNKVDENFFQRLDDSFVDEDEKRG  
 EHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDL  
 KAENTDVQALFKDFVEEYDKTIEESHLSIEITVDALSILTEKVS KSSRLENLIAHYPTKKN  
 TLFGNLIALSLDLHPNFKTNFQLS EDAKLQFSKDTYEEDLEGFLGEVGEYADLFASAKNL  
 YDAILLSGILTVDDNSTKAPLSASMVKRYEEHQDKLKKLKDIFIKVNAPDQYNAIFKDKNKK  
 GYASYIESGVKQDEFYKYLKGI LLKINGS GDFLDKIDREDFLRKQRTFDNGIIPHQIHLQE  
 MHAILRRQGEHY PFLKENQDKIEKILTFRIPIYVYVGPLARKGSRFAWAEYKADEKITPWNFD  
 DILDKEKSAEKFITRMTLNDLYLP EEKVLPKHSPLYEAF TVYNELTKVKYVNEQGEAKFFD  
 TNMKQEIFDHVFKENRKVTKDKLLNYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRK  
 ILDKSFLDDKANEKTI EDIIQTTLTFEDREMIRQLQKYSDI FTKAQLKKLERLHYTGWGR  
 LSYKLINGIRNKENKKTILDYLIIDGYANRNFQMLINDDALSFKEEIARAQIIDDVDDIAN  
 VVHDLPGSPAIAKKGILQSVKIVDELVKVMGHNPANIIEMARENQTTDKGRRNSQQRLKLL  
 QDSLKNLNDNPNVNIKNVENQQQLQNDRLFLLYYIQNGKDMYTGETLDINNLSQYDIDHIIIPQAF  
 IKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGG  
 LTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIIRNVKIITLKSNLVSNFRK  
 EFELYKVREINDYHHAHDAYLNAVVGNALLLKYPQLEPEFVYGEYPKYNSYRSRKSATEKF

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LFYSNILRFFKKEDIQTNEEDGEIAWNKEKHKILRKVLSYPQVNI VKKTEEQTGGFSKESI
LPKGESDKLI PRKTKNSYWDPKKYGGFMQPVVAYSILVFADVEKGKSKKLRKVQDMVGITI
MEKKRFEKNPVDFLEQRGYRNVRLKIIKLPKYSLFELENKRRRLASAKFLQKGNELVIP
QRFTTLLYHSYRIEKDYEPHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMLF
KNKDAEVSSLAKSFISLLTFTAAGAPRAFNFGENIARKEYRSVTECLNATLIHQSI TGLY
ETRIDLSKLGEDGEGADKRTADGSEFESPKKKRKV (SEQ ID NO: 96)

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NLS (bold), can be substituted with different NLSs

Linker (underlined), can be removed or extended

In some embodiments, the engineered non-naturally occurring Cas9 comprises a tag.  
 5 A variety of tags may be fused to the Cas9 variant (e.g., 3xHA tag), depending on purpose, as will be apparent to a skilled person.

Various species exhibit codon bias (i.e. differences in codon usage by organisms) which correlates with the efficiency of translation of messenger RNA (mRNA) by utilizing codons in mRNA that correspond with the abundance of tRNA species for that codon in a  
 10 particular organism. Various methods in the art can be used for computer optimization, including for example through use of software. In some embodiments, codon optimization refers to modification of nucleic acid sequences for enhanced expression in the host cells of interest by replacing at least one codon (e.g. 1, 2, 3, 4, 5, 10, 15, 20, 25, 50 or more codons)  
 15 the genes of the host cell while maintaining the native amino acid sequence.

In some embodiments, the Cas9 protein described herein is codon optimized. This type of optimization is known in the art and entails the mutation of foreign-derived DNA to mimic the codon preferences of the intended host organism or cell while encoding the same protein. Thus, the codons are changed, but the encoded protein remains unchanged. Codon  
 20 optimization improves soluble protein levels and increases activity and editing efficiency in a given species. Codon optimization also results in increased translation and protein expression.



In some embodiments, the Cas9 protein is codon optimized for expression in eukaryotic cells. In some embodiments, the Cas9 protein is codon optimized for expression in human cells.

### **Protospacer Adjacent Motif (PAM)**

5 Each Cas endonuclease binds to its target sequence only in the presence of a specific sequence, known as a protospacer adjacent motif (PAM), on the non-targeted i.e. complementary DNA strand. Cas nucleases isolated from different bacterial species recognize different PAM sequences. For example, the SpCas9 nuclease (from *Staphylococcus pyogenes*) cuts upstream of the PAM sequence 5'-NGG-3' (where "N" can be  
10 any nucleotide base), SaCas9 (from *Staphylococcus aureus*) recognizes the PAM sequence 5'-NNGRR (N)-3' in the target. Thus, the locations in the genome that can be targeted by different Cas proteins are limited by the locations of unique PAM sequences.

Disclosed herein Cas9 proteins engineered from *Streptococcus constellatus* and *Ezakiella peruensis* and *Peptoniphilus sp. Marseille-P3761* species recognize the consensus  
15 PAM sequence 5'-NGG-3'. Disclosed herein Cas9 proteins engineered from *Streptococcus constellatus* and *Ezakiella peruensis* and *Peptoniphilus sp. Marseille-P3761* species recognize the consensus PAM sequence 5'-NGG-3'. In some embodiments, Cas9 proteins disclosed herein are engineered to recognize the consensus PAM sequence 5'-NGC-3'. Exemplary embodiments are described below and should be nonlimiting. In some  
20 embodiments, Cas9 proteins from *Streptococcus constellatus* are engineered to recognize the consensus PAM sequence 5'-NGC-3'. In some embodiments, the NGC PAM variant includes one or more amino acid substitutions selected from or corresponding to D1117M, S118Q, E1201F, A1299R, D1309A, R1312E, and T1314R (collectively termed "MQFRAER") with reference to ScoCas9 (SEQ ID NO: 1). In some embodiments, the NGC PAM variant  
25 includes one or more amino acid substitutions selected from or corresponding to D1135M, S1136Q, G1218K, E1219F, A1322R, D1332A, R1335E, and T1337R (collectively termed "MQKFRAER") with reference to a naturally occurring SpyCas9 (SEQ ID NO: 173). In some embodiments, similar or corresponding amino acid substitutions can be made to SirCas9, VapCas9, EpeCas9, LfeCas9, or PmaCas9.

30 *Streptococcus pyogenes* Cas9 (*SpyCas9*; GenBank: QSG91308.1)

MDKKYSIGLDIGTNSVGVAVITDDYKVPSKKFKVLGNTDRHSIKKNLIGALLFDSGETAEAT  
 RLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDSFHRLEESFLVEEDKKHERHPIFGNIVD  
 EVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDLNPDNSDVKLFI  
 QLVQTYNQLFEEFNINASGVDAKAILSARLSKSRLENLIAQLPGEKKNGLFGNLIALSGLL  
 5 TPNFKSNFDLAEDAQLSKDQYDDDLNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNT  
 EITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEF  
 YKFIKPILEKMDGTEELLVKNLREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFY PFLK  
 DNREKIEKILTFRIPYYVGPLARGNSRFAMTRKSEETITPWNFEEVVDKGASAQSFIERMT  
 NFDKNLPNEKVLPKHSLLEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTRNK  
 10 VTVKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLKIIKDKDFLDNEENEDILEDIV  
 LTLTLFEDREMIEERLKYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSGKTILDF  
 LKSDGFANRFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIKKILQTVKVV  
 DELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEGIKELGSQILKEHPVENTQL  
 QNEKLYLYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDDSIDNKVLRSDKNRGKSD  
 15 NVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGLSELDKAGFIKRQLVETRQITKH  
 VAQILDSRMNTKYDENDKLIREVKVITLKSCLVSDFRKDFQFYKVVREINNYHHAHDAYLNAV  
 VGTALIKKYPKLESEFVYGDYKVYDVRKMIKSEQEIGKATAKYFFYSNIMNFFKTEITLAN  
 GEIRKRPLIETNGETGEIVWDKGRDFATVRKVLSPQVNIIVKKTQVQTTGGFSKESILPKRNS  
 DKLIARKKDWDPKKGFFDSPTVAYSVLVAKVEKGSKLLKSVKELLGITIMERSSEKPNP  
 20 IDFLEAKGYKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLAS  
 HYEKLGKSPEDNEQKQLFVEQHKHYLDEIEQISEFSKRVILADANLDKVL SAYNKHRDKPI  
 REQAENIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEVL DATLIHQSI TGLYETRIDL SQ  
 LGGD(SEQ ID NO:173).

25 In some embodiments, the Cas9 protein described herein does not bind or exhibit activity with any other PAM sequences.

### **RNA Guides**

An RNA guide comprises a polynucleotide sequence with complementarity to a target  
 sequence. The RNA guide hybridizes with the target nucleic acid sequence and directs  
 30 sequence-specific binding of a CRISPR complex to the target nucleic acid. In some  
 embodiments, an RNA guide has 50%, 60%, 70%, 75%, 80%, 85%, 90%, 95%, 96%, 97%,  
 98%, 99% or 100% complementarity to a target nucleic acid sequence.

In some embodiments, the RNA guides are about 5, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 35, 40, 45, 50, 75 or more nucleotides in length. In some embodiments, the RNA guides are about 18-24 nucleotides in length. In some embodiments, the RNA guide is complementary to about 18-24 nucleotides in the target  
5 nucleic acid sequence. For example, the RNA guide is complementary to about 18, 19, 20, 21, 22, 23, or 24 nucleotides in the target nucleic acid sequence. In some embodiments, the RNA guide is complementary to about 18-22 nucleotides. In some embodiments, the RNA guide is complementary to about 18-21 nucleotides. In some embodiments, the RNA guide is complementary to about 18-20 nucleotides. In some embodiments, the RNA guide is  
10 complementary to 20 nucleotides in the target nucleic acid sequence.

An RNA guide can be designed to target any target sequence. Optimal alignment is determined using any algorithm for aligning sequences, including the Needleman-Wunsch algorithm, Smith-Waterman algorithm, Burrows-Wheeler algorithm, ClustlW, ClustlX, BLAST, Novoalign, SOAP, Maq, and ELAND.

15 In some embodiments, an RNA guide is targeted to a unique target sequence within the genome of a cell. In some embodiments, an RNA guide is designed to lack a PAM sequence. In some embodiments, an RNA guide sequence is designed to have optimal secondary structure using a folding algorithm including mFold or Geneious. In some  
20 embodiments, expression of RNA guides may be under an inducible promoter, e.g. hormone inducible, tetracycline or doxycycline inducible, arabinose inducible, or light inducible.

In some embodiments, the CRISPR system includes one or more RNA guides e.g. crRNA, tracrRNA, and/or sgRNA. Accordingly, in some embodiments the RNA guide comprises a crRNA. In some embodiments, the RNA guide comprises a tracrRNA. In some  
25 embodiments, the RNA guide comprises a sgRNA. In some embodiments, the CRISPR system includes multiple RNA guides, comprising 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15 or more RNA guides.

In some embodiments, the RNA guide includes a crRNA. In some embodiments, the CRISPR system includes multiple crRNAs comprising 2-15 crRNAs. In some embodiments, the crRNA is a precursor crRNA (pre-crRNA), which includes a direct repeat sequence, a  
30 spacer sequence and a direct repeat sequence. In some embodiments, the crRNA is a processed or mature crRNA which includes a truncated direct repeat sequence.

In some embodiments, a CRISPR associated protein cleaves the pre-crRNA to form processed or mature crRNA.

In some embodiments, a CRISPR associated protein forms a complex with the mature crRNA and the spacer sequence targets the complex to a complementary sequence in the target nucleic acid. In some embodiments, an RNA guide comprises a direct repeat sequence  
5 and a spacer sequence capable of hybridizing under appropriate conditions to a target nucleic acid.

In some embodiments, the spacer length of crRNAs can range from about 15 to 50 nucleotides. In some embodiments, the spacer length of an RNA guide is at least 16  
10 nucleotides, at least 17 nucleotides, at least 18 nucleotides, at least 19 nucleotides, at least 20 nucleotides, at least 21 nucleotides, or at least 22 nucleotides. In some embodiments, the spacer length is from 15 to 17 nucleotides (e.g., 15, 16, or 17 nucleotides), from 17 to 20 nucleotides (e.g., 17, 18, 19, or 20 nucleotides), from 20 to 24 nucleotides (e.g., 20, 21, 22, 23, or 24 nucleotides), from 23 to 25 nucleotides (e.g., 23, 24, or 25 nucleotides), from 24 to  
15 27 nucleotides, from 27 to 30 nucleotides, from 30 to 45 nucleotides (e.g., 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, or 45 nucleotides), from 30 or 35 to 40 nucleotides, from 41 to 45 nucleotides, from 45 to 50 nucleotides (e.g., 45, 46, 47, 48, 49, or 50 nucleotides), or longer.

In some embodiments, the RNA guide comprises a direct repeat (DR) sequence of  
20 between about 16 and 26 nucleotides long. For example, in some embodiments, the DR is about 16 nucleotides long. In some embodiments, the DR is about 17 nucleotides long. In some embodiments, the DR is about 18 nucleotides long. In some embodiments, the DR is about 19 nucleotides long. In some embodiments, the DR is about 20 nucleotides long. In some embodiments, the DR is about 21 nucleotides long. In some embodiments, the DR is  
25 about 22 nucleotides long. In some embodiments, the DR is about 23 nucleotides long. In some embodiments, the DR is about 24 nucleotides long. In some embodiments, the DR is about 25 nucleotides long. In some embodiments, the DR is about 26 nucleotides long.

In some embodiments, the crRNA comprises a nucleotide guide sequence and a DR sequence. The nucleotide guide sequence can be between about 18 and 24 nucleotides long.  
30 Accordingly, in some embodiments, the nucleotide guide sequence is about 18 nucleotides long. In some embodiments, the nucleotide guide sequence is about 19 nucleotides long. In some embodiments, the nucleotide guide sequence is about 20 nucleotides long. In some

embodiments, the nucleotide guide sequence is about 21 nucleotides long. In some embodiments, the nucleotide guide sequence is about 22 nucleotides long. In some embodiments, the crRNA comprises a nucleotide guide sequence of about 22 nucleotides long and a direct repeat of about 22 nucleotides long.

5           In some embodiments, the crRNA sequences can be modified to "dead crRNAs," "dead guides," or "dead guide sequences" that can form a complex with a CRISPR-associated protein and bind specific targets without any substantial nuclease activity.

          In some embodiments, the crRNA may be chemically modified in the sugar phosphate backbone or base. In some embodiments, the crRNA may be modified using 2'-O-methyl, 2'-F  
10 or locked nucleic acids to improve nuclease resistance or base pairing. In some embodiments, the crRNA may contain modified bases such as 2-thiouridine or N6-methyladenosine.

          In some embodiments, the crRNA is conjugated with other oligonucleotides, peptides, proteins, tags, dyes, or polyethylene glycol.

          In some embodiments, the crRNA may include aptamer or riboswitch sequences that  
15 can bind specific target molecules due to their three-dimensional structure.

          In some embodiments, a trans-activating RNA (tracrRNA) is associated with crRNA to facilitate formation of a complex with Cas9 protein. In some embodiments, the tracrRNA sequence is about or more than about 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20,  
20 25, 30, 40, 50, 60, 70, 80, 90, 100 or more nucleotides in length. In some embodiments, the tracrRNA is about 70 nucleotides in length.

          In some embodiments, the tracrRNA and crRNA are contained in a single transcript called single guide RNA (sgRNA). In some embodiments, the sgRNA includes a loop between the tracrRNA and sgRNA.

          In some embodiments, the loop forming sequences are 3, 4, 5 or more nucleotides in  
25 length. In some embodiments, the loop has the sequence GAAA, AAAG, CAAA, AAAC, UUUU, UUAUUAU, UUA, UUU and/or AAUCA. In some embodiments, the loop has the sequence GAAA. In some embodiments, the loop has the sequence AAAG. In some embodiments, the loop has the sequence CAAA. In some embodiments, the loop has the sequence AAAC. In some embodiments, the loop has the sequence AAUCA. In some  
30 embodiments, the loop has the sequence UUUU. In some embodiments, the loop has the sequence UUAUUAU. In some embodiments, the loop has the sequence UUA. In some embodiments, the loop has the sequence UUU. In some embodiments, the loop has the sequence AAUCA.

In some embodiments, the tracrRNA and crRNA form a hairpin loop. In some embodiments, sgRNA has at least two or more hairpins. In some embodiments, sgRNA has two, three, four or five hairpins.

In some embodiments, sgRNA includes a transcription termination sequence, which  
5 includes a polyT sequences comprising six nucleotides.

In some embodiments, the sgRNA comprises a sequence having at least 80% identity to 5'-

GUUUUAGAGCUGUGCUGUUUAAACAACACAGCAAGUUAAAAUAAGGCUUUGU  
CCGUACUC (SEQ ID NO: 3) for ScoCas9,

10 5'-

GUUUUAGAGUUGUGUUUUAUUGAAAAUAACACAACGAGUUAAAAUAAAGCUUA  
UGCUUAAAUGCCAGCUUUGCUGGUGUCAUUUAGAUGACUUUACUAAGGUUGC  
UUCGGCAACCUUUUU-3' (SEQ ID NO: 7) for SirCas9,

15 5'-

GUUUGAGAGUAGUGUGAAAACAUAACGAGUUCAAAUACAAAUAAUUUACAA  
UGCCUUCGGGCUGCCCGACGUAGGGCACCUACUCUCAAUUCUUCGGAAUUGAG  
UU-3' (SEQ ID NO: 13) for VapCas9,

20 5'-

GUUUGAGAGUUAUGUAAUUGAAAAUUACAUGACGAGUUCAAAUAAAAUUU  
AUUCAAAACCGCCUAUUUAUAGGCCGCAGAUGUUCUGCAUUAUGCUUGCUAUU  
GCAAGCUU-3' (SEQ ID NO: 19) for EpeCas9,

25 5'-

GUCUUGGAUGAGUGUGAAAACACUCAUAGUCAAGAUCAAACGAGUGGUUUUC  
CACGAGUUAUUACUUUUGAGGUCUUAUAUGGCCCAUACAUAAAAGGAGUCG  
GAAUUUCCGGCUCUUUUUCUU-3' (SEQ ID NO: 95) for LfeCas9, and

5'-

GUUUUAGAGCCAUGUAGAAUACAUUGCAAGUUAAAAUAAGGCUUUGUCCGU  
AAUCAACUUGAAAAAGUGGGCGCUGUUUCGGCGCUUU-3' (SEQ ID NO: 96) for  
PmaCas9.

The guide RNA is added to the 5' end of the Cas9. In some embodiments, the sgRNA comprises a sequence having 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more identity to SEQ ID NO: 3, 7, 13, 19, 95 or 96. In some embodiments, the sgRNA comprises a sequence identical to SEQ ID NO: 3, 7, 13, 19, 95 or 96.

In some embodiments, the tracrRNA is a separate transcript, not contained with crRNA sequence in the same transcript.

### **Cas9 Fusion Proteins**

In some embodiments, the Cas9 enzyme is fused to one or more heterologous protein domains. In some embodiments, the Cas9 enzyme is fused to more than about 1, 2, 3, 4, 5, 6, 7, 8, 9, 10 or more protein domains. In some embodiments, the heterologous protein domain is fused to the C-terminus of the Cas9 enzyme. In some embodiments, the heterologous protein domain is fused to the N-terminus of the Cas9 enzyme. In some embodiments, the heterologous protein domain is fused internally, between the C-terminus and the N-terminus of the Cas9 enzyme. In some embodiments, the internal fusion is made within the Cas9 RuvCI, RuvC II, RuvCIII, HNH, REC I, or PAM interacting domain.

A Cas9 protein may be directly or indirectly linked to another protein domain. In some embodiments, a suitable CRISPR system contains a linker or spacer that joins a Cas9 protein and a heterologous protein. An amino acid linker or spacer is generally designed to be flexible or to interpose a structure, such as an alpha-helix, between the two protein moieties. A linker or spacer can be relatively short, or can be longer. Typically, a linker or spacer contains for example 1-100 (e.g., 1-100, 5-100, 10-100, 20-100, 30-100, 40-100, 50-100, 60-100, 70-100, 80-100, 90-100, 5-55, 10-50, 10-45, 10-40, 10-35, 10-30, 10-25, 10-20) amino acids in length. In some embodiments, a linker or spacer is equal to or longer than 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, or 100 amino acids in length. Typically, a longer linker may decrease steric hindrance. In some embodiments, a linker will comprise a mixture of glycine and serine residues. In some embodiments, the linker may additionally comprise threonine, proline and/or alanine residues.

In some embodiments, a Cas9 protein is fused to cellular localization signals, epitope tags, reporter genes, and protein domains with enzymatic activity, epigenetic modifying activity, RNA cleavage activity, nucleic acid binding activity, transcription modulation

activity. In some embodiments, the Cas9 protein is fused to a nuclear localization sequence (NLS), a FLAG tag, a HIS tag, and/or a HA tag.

Suitable fusion partners include, but are not limited to, a polypeptide that provides for methyltransferase activity, demethylase activity, acetyltransferase activity, deacetylase  
5 activity, kinase activity, phosphatase activity, ubiquitin ligase activity, deubiquitinating activity, adenylation activity, deadenylation activity, SUMOylating activity, deSUMOylating activity, ribosylation activity, deribosylation activity, myristoylation activity, demyristoylation activity, integrase activity, transposase activity, recombinase activity, polymerase activity, ligase activity, helicase activity, or nuclease activity, any of which can  
10 modify DNA or a DNA-associated polypeptide (e.g., a histone or DNA binding protein). In some embodiments, the Cas9 protein is fused to a histone demethylase, a transcriptional activator or a deaminase.

Further suitable fusion partners include, but are not limited to boundary elements (e.g., CTCF), proteins and fragments thereof that provide periphery recruitment (e.g., Lamin  
15 A, Lamin B, etc.), and protein docking elements (e.g., FKBP/FRB, Pill/Abyl, etc.).

In particular embodiments, a Cas9 is fused to a cytidine or adenosine deaminase domain, e.g., for use in base editing. In some embodiments, Cas9 is fused to an adenine and cytosine base editor (ACBE or CBE), wherein ACBE or CBE is generated by fusing a heterodimer of TadA and an activation-induced cytidine deaminase (AID) to the N- and C-  
20 terminals of Cas9 nickase (nCas9). In some embodiments, the ACBE or CBE simultaneously induces C-to-T and A-to-G base editing at the same target site. Xie, J *et al.* ACBE, a new base editor for simultaneous C-to-T and A-to-G substitutions in mammalian systems. *BMC Biology* (18: 131), 2020)

In some embodiments, the terms “cytidine deaminase” and “cytosine deaminase” can  
25 be used interchangeably. In certain embodiments, the cytidine deaminase domain may have sequence identity of 70%, 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more to any cytidine deaminase described herein. In some embodiments, the cytidine deaminase domain has cytidine deaminase activity, (e.g., converting C to U). In certain embodiments, the adenosine deaminase domain may have sequence identity of 70%,  
30 75%, 80%, 85%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% or more to any adenosine deaminase described herein. In some embodiments, the adenosine deaminase



domain has adenosine deaminase activity, (*e.g.*, converting A to I). In some embodiments, the terms “adenosine deaminase” and “adenine deaminase” can be used interchangeably.

In some embodiments, a cytidine deaminase can comprise all or a portion of an apolipoprotein B mRNA editing complex (APOBEC) family deaminase. APOBEC is a  
5 family of evolutionarily conserved cytidine deaminases. Members of this family are C-to-U editing enzymes. The N-terminal domain of APOBEC like proteins is the catalytic domain, while the C-terminal domain is a pseudocatalytic domain. More specifically, the catalytic domain is a zinc dependent cytidine deaminase domain and is important for cytidine deamination. APOBEC family members include APOBEC1, APOBEC2, APOBEC3A,  
10 APOBEC3B, APOBEC3C, APOBEC3D ("APOBEC3E" now refers to this), APOBEC3F, APOBEC3G, APOBEC3H, APOBEC4, and Activation-induced (cytidine or cytosine) deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of an APOBEC1 deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of APOBEC2 deaminase. In  
15 some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of is an APOBEC3 deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of an APOBEC3A deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of APOBEC3B deaminase. In some embodiments, a deaminase incorporated into a fusion  
20 protein comprises all or a portion of APOBEC3C deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of APOBEC3D deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of APOBEC3E deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of APOBEC3F deaminase. In some  
25 embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of APOBEC3G deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of APOBEC3H deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of APOBEC4 deaminase. In some embodiments, a deaminase incorporated into a fusion protein comprises  
30 all or a portion of activation-induced deaminase (AID). In some embodiments a deaminase incorporated into a fusion protein comprises all or a portion of cytidine deaminase 1 (CDA1). It should be appreciated that a fusion protein can comprise a deaminase from any suitable organism (*e.g.*, a human or a rat). In some embodiments, a deaminase domain of a fusion

protein is from a human, chimpanzee, gorilla, monkey, cow, dog, rat, or mouse. In some embodiments, the deaminase domain of the fusion protein is derived from rat (*e.g.*, rat APOBEC1). In some embodiments, the deaminase domain is human APOBEC1. In some embodiments, the deaminase domain is pmCDA1. Sequences of exemplary cytidine

5 deaminases are provided below.

pmCDA1 (*Petromyzon marinus*)

MTDAEYVRIHEKLDIYTFKKQFFNNKKS VSHRCYVLFELKRRGERRACFWGYAVNK  
PQSGTERGIHAEIFSIRKVEEYLRDNPQGFTINWYSSWSPCADCAEKILEWYNQELRG  
NGHTLKIWACKLYYEKNARNQIGLWNLRDNGVGLNVMVSEHYQCCRKIFIQSSHNQ  
10 LNENRWLEKTLKRAEKRRSELSIMI QVKILHTTKSPAV (SEQ ID NO: 22)

Human AID:

MDSLLMNRKFLYQFKNVRWAKGRRETYLCYVVKRRDSATSFSLDFGYLRNKNGC  
HVELLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGNPNLSLRIFTAR  
LYFCEDRKAPEPEGLRRLHRAGVQIAIMTFKAPV (SEQ ID NO: 23)

15 Human AID:

MDSLLMNRKFLYQFKNVRWAKGRRETYLCYVVKRRDSATSFSLDFGYLRNKNGC  
HVELLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGNPNLSLRIFTAR  
LYFCEDRKAPEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHEN  
SVRLSRQLRRILLPLYEVDDLRDAFRTLGL (underline: nuclear localization sequence;  
20 double underline: nuclear export signal) (SEQ ID NO: 24)

Mouse AID:

MDSLLMKQKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSCSLDFGHLRNKSGC  
HVELLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVAEFLRWPNLSLRIFTAR  
LYFCEDRKAPEPEGLRRLHRAGVQIGIMTFKDYFYCWNTFVENRERTFKAWEGLHEN  
25 SVRLTRQLRRILLPLYEVDDLRDAFRMLGF (underline: nuclear localization sequence;  
double underline: nuclear export signal) (SEQ ID NO: 25)

Canine AID:

MDSLLMKQRKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSFSLDFGHLRNKSGC  
HVELLFLRYISDWDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSLRIFAAR  
30 LYFCEDRKAPEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENREKTFKAWEGLHEN  
SVRLSRQLRRILLPLYEVDDLRDAFRTLGL (underline: nuclear localization sequence;  
double underline: nuclear export signal) (SEQ ID NO: 26)

Bovine AID:

MDSLLKKQRQFLYQFKNVRWAKGRHETYLCYVVKRRDSPTSFSLDFGHLRNKAGC  
 HVELLFLRYISDWLDPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSLRIFTAR  
 LYFCDKERKAEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHE  
 NSVRLSRQLRRILLPLYEVDDLRLDAFRTLGL (underline: nuclear localization sequence;  
 5 double underline: nuclear export signal) (SEQ ID NO: 27)

Rat AID:

MAVGSKPKAALVGPHWERERIWCFLCSTGLGTQQTGQTSRWLRPAATQDPVSPPRS  
 LLMKQRKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSFSLDFGYLRNKSGCHVE  
 LLFLRYISDWLDPGRCYRVTWFTSWSPCYDCARHVADFLRGNPNLSLRIFTARLTG  
 10 WGALPAGLMSPARPSDYFYCWNTFVENHERTFKAWEGLHENSVRLSRRLRRILLPL  
 YEVDLRLDAFRTLGL (SEQ ID NO: 28)

(underline: nuclear localization sequence; double underline: nuclear export signal)

clAID (*Canis lupus familiaris*):

MDSLLMKQRKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSFSLDFGHLRNKSGC  
 15 HVELLFLRYISDWLDPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSLRIFAAR  
 LYFCEDRKAPEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENREKTFKAWEGLHEN  
 SVRLSRQLRRILLPLYEVDDLRLDAFRTLGL (SEQ ID NO: 29)

btAID (*Bos taurus*):

MDSLLKKQRQFLYQFKNVRWAKGRHETYLCYVVKRRDSPTSFSLDFGHLRNKAGC  
 20 HVELLFLRYISDWLDPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSLRIFTAR  
 LYFCDKERKAEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHE  
 NSVRLSRQLRRILLPLYEVDDLRLDAFRTLGL (SEQ ID NO: 30)

mAID (*Mus musculus*):

MDSLLMNRKFLYQFKNVRWAKGRRETYLCYVVKRRDSATSFSLDFGYLRNKNGC  
 25 HVELLFLRYISDWLDPGRCYRVTWFTSWSPCYDCARHVADFLRGNPNLSLRIFTAR  
 LYFCEDRKAPEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHEN  
 SVRLSRQLRRILLPLYEVDDLRLDAFRTLGL (SEQ ID NO: 31)

rAPOBEC-1 (*Rattus norvegicus*):

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKTKETCLLYEINWGGRHSIWRHTSQNT  
 30 NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR  
 LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNEAHWPYPHILW  
 VRLYVLELYCIIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLK (SEQ  
 ID NO: 32)

maAPOBEC-1 (*Mesocricetus auratus*):

MSSETGPVVVDPTLRRRIEPHEFDAFFDQGELRKETCLLYEIRWGGRHNIWRHTGQN  
 TSRHVEINFIEKFTSERYFYFPSTRCSIVWFLSWSPCGECSKAITEFLSGHPNVTLFYAA  
 RLYHHTDQRNRQGLRDLISRGVTIRIMTEQEYCYCWRNFVNYPPSNEVYWPYPNL  
 WMRLYALELYCIHLGLPPCLKIKRRHQYPLTFFRLNLQSCHYQRIPPHILWATGFI

5 (SEQ ID NO: 33)

ppAPOBEC-1 (*Pongo pygmaeus*):

MTSEKGPSTGDPTLRRRIESWEFDVFYDPRELKRETCCLLYEIKWGMSRKIWRSSGKN  
 TTNHVEVNFIEKFTSERRFHSSISCSITWFLSWSPCWECQAIREFLSQHPGVTLVIVV  
 ARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYP  
 10 PLWMLLYALELHCILSLPPCLKISRRWQNHAFRLHLQNCYQTIPPHILLATGLIH  
 PSVTWR (SEQ ID NO: 34)

ocAPOBEC1 (*Oryctolagus cuniculus*):

MASEKGPSNKDYTLRRRIEPWEFEVFFDPQELRKEACCLLYEIKWGASSKTWRSSGKN  
 TTNHVEVNFLEKLTSEGRLLGPSTCCSITWFLSWSPCWECMAIREFLSQHPGVTLIIFV  
 15 ARLFQHMDRRNRQGLKDLVTSGVTVRVMSVSEYCYCWENFVNYPPGKAAQWPRY  
 PPRWMLMYALELYCIILGLPPCLKISRRHQKQLTFFSLTPQYCHYKMIPPYILLATGLL  
 QPSVPWR (SEQ ID NO: 35)

mdAPOBEC-1 (*Monodelphis domestica*):

MNSKTGPSVGDATLRRRIKPWEFVAFNPQELRRETCCLLYEIKWGNQNIWRHSNQN  
 20 TSQHAEIFMEKFTAERHFNSVRCSTWFLSWSPCWECASKAIRKFLDHYPNVTLAIFI  
 SRLYWHMDQQHRQGLKELVHSGVTIQIMSSEYHYHCWRNFVDYPQGEEDYWPKYV  
 YLWIMLYVLELHCILGLPPCLKISGSHSNQLALFSLDLQDCHYQKIPYNVLVATGLV  
 QPFVTWR (SEQ ID NO: 36)

ppAPOBEC-2 (*Pongo pygmaeus*):

MAQKEEAAAATEAASQNGEDLENLDDPEKLEKELIELPPFEIVTGERLPANFFKFQFRN  
 VEYSSGRNKTFLCYVVEAQKGGQVQASRGYLEDEHAAAHAEAEAFFNTILPAFDPA  
 LRYNVTWYVSSSPCAACADRIIKTLTKNLRLLLILVGRLFMWEELEIQDALKKLE  
 AGCKLRIMKPQDFEYVWQNFVEQEEGESKAFQPWEDIQENFLYYEEKLADILK (SEQ  
 25 ID NO: 37)

30 btAPOBEC-2 (*Bos taurus*):

MAQKEEAAAAAEPASQNGEEVENLEDPEKLEKELIELPPFEIVTGERLPAHYFKFQFRN  
 VEYSSGRNKTFLCYVVEAQSKGGQVQASRGYLEDEHATNHAEAEAFFNSIMPTFDPA  
 LRYMVTWYVSSSPCAACADRIVKTLNKTNLRLLLILVGRLFMWEEPEIQAALRKLKE

AGCRLRIMKPQDFEYIWQNFVEQEEGESKAFEPWEDIQENFLYYEEKLADILK (SEQ ID NO: 38)

mAPOBEC-3-(1) (*Mus musculus*):

MQPQRLGPRAGMGPFC LGCSHRKCYSPIRNLISQETFKFHFKNLGYAKGRKDTFLCY  
 5 EVTRKDCDSPVSLHHGVFKNKDNIHAEICFLYWFHDKVLKVLSPREEFKITWYMSW  
 SPCFECAEQIVRFLATHHNLSLDIFSSRLYNVQDPETQQNLCRLVQEGAQVAAMDLY  
 EFKKCWKKFVDNGGRRFRPWKRLLTNFRYQDSKLQEILRPCYISVPSSSSSTLSNICL  
 TKGLPETRFWVEGRRMDPLSEEEFY SQFYNQRVKHLCYYHRMKPYLCYQLEQFNG  
 QAPLKGCLLSEK GKQHAEILFLDKIRSMELSQVTITCYLTWSPCPNCAWQLAAFKRD  
 10 RPDILHIYTSRLYFHWKRPFQKGLCSLWQSGILVDVMDLPQFTDCWTN FVNPKRPF  
 WPWKGLEIISRRTQRRLRIKESWGLQDLVNDFGNLQLGPPMS (SEQ ID NO: 39)

Mouse APOBEC-3-(2):

MGPFC LGCSHRKCYSPIRNLISQETFKFHFKNLGYAKGRKDTFLCYEVTRKDCDSPV  
 SLHHGVFKNKDNIHAEICFLYWFHDKVLKVLSPREEFKITWYMSWSPCFECAEQIVRFL  
 15 ATHHNLSLDIFSSRLYNVQDPETQQNLCRLVQEGAQVAAMDLYEFKKCWKKFVDN  
 GGRRFRPWKRLLTNFRYQDSKLQEILRPCYIPVSSSSSTLSNICLTKGLPETRF CVEG  
 RRMDPLSEEEFY SQFYNQRVKHLCYYHRMKPYLCYQLEQFNGQAPLKGCLLSEK GK  
 QHAEILFLDKIRSMELSQVTITCYLTWSPCPNCAWQLAAFKRDRPDILHIYTSRLYFHW  
 KRPFQKGLCSLWQSGILVDVMDLPQFTDCWTN FVNPKRPFWPWKGLEIISRRTQRRL  
 20 RRIKESWGLQDLVNDFGNLQLGPPMS (italic: nucleic acid editing domain) (SEQ ID  
 NO: 40)

Rat APOBEC-3:

MGPFC LGCSHRKCYSPIRNLISQETFKFHFKNRLRYAIDRKDTFLCYEVTRKDCDSPV  
 SLHHGVFKNKDNIHAEICFLYWFHDKVLKVLSPREEFKITWYMSWSPCFECAEQVLRFL  
 25 ATHHNLSLDIFSSRLYNIRDPENQQNLCRLVQEGAQVAAMDLYEFKKCWKKFVDNG  
 GRRFRPWKLLTNFRYQDSKLQEILRPCYIPVSSSSSTLSNICLTKGLPETRF CVERR  
 RVHLLSEEEFY SQFYNQRVKHLCYYHGVPYLCYQLEQFNGQAPLKGCLLSEK GKQ  
 HAEILFLDKIRSMELSQVIITCYLTWSPCPNCAWQLAAFKRDRPDILHIYTSRLYFHWK  
 RPFQKGLCSLWQSGILVDVMDLPQFTDCWTN FVNPKRPFWPWKGLEIISRRTQRRLH  
 30 RIKESWGLQDLVNDFGNLQLGPPMS (italic: nucleic acid editing domain) (SEQ ID NO:  
 41)

hAPOBEC-3A (*Homo sapiens*):

MEASPASGPRHLMDPHIFTSN FNNGIGRHKTYLCYEVERLDNGT SVKMDQHRGFLH  
 NQAKNLLCGFYGRHAELRFLDLVPSLQLDPAQIYRVTWFISWSPCF SWGCAGEVRAF

LQENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIMTYDEFKHCWDTFVDHQ  
GCPFQPWDGLDEHSQALSGRLRAILQNQGN (SEQ ID NO: 42)

hAPOBEC-3F (*Homo sapiens*):

MKPHFRNTVERMYRDTFSYNFYNRPILSRRNTVWLCYEVKTKGSPRPLDAKIFRGQ  
5 VYSQPEHHAEMCFLSWFCGNQLPAYKCFQITWVFSWTPCPDCVAKLAEFLAEHPNV  
TLTISAARLYYYWERDYRRALCRLSQAGARVKIMDDEEFAYCWENFVYSEGQPFMP  
WYKFDDNYAFLHRTLKEILRNPMEAMYPHIFYHFKNLRKAYGRNESWLCFTMEV  
VKHHSVSWKRGVFRNQVDPETHCHAERCFLSWFCDDILSPNTNYEVTWYTSWSPC  
PECAGEVAEFLARHSNVNLTIFTARLYYFWDDTYQEGLRSLSQEGASVEIMGYKDFK  
10 YCWENFVYNDDEPFKPKWGLKYNFLFLDSKLQEILE (SEQ ID NO: 43)

Rhesus macaque APOBEC-3G:

MVEPMDPRTFVSNFNRPILSGLNTVWLCCEVKTCDPSGPPLDAKIFQGKVVSKAKY  
HPEMRFLRWFHKWRQLHHDQEYKVTWYVSWSPCTRCANSVATFLAKDPKVTLTIF  
VARLYYFWKPDYQQALRILCQKRGGPHATMKIMNYNEFQDCWKNFVDGRGKPFK  
15 RNNLPKHYTLLQATLGELLRHLMDPGTFTSNFNKPKWVSGQHETYLCYKVERLHND  
TWVPLNQHRGFLRNQAPNIHGFPKGRHAELCFLDLIPFWKLDGQQYRVTCFTSWSPC  
FSCAQEMAKFISNNEHVSLCIFAARIYDDQGRYQEGLRALHRDGAKIAMMNYSEFEY  
CWDTFVDRQGRPFQPWDGLDEHSQALSGRLRAI (italic: nucleic acid editing domain;  
underline: cytoplasmic localization signal) (SEQ ID NO: 44)

20 Chimpanzee APOBEC-3G:

MKPHFRNPVERMYQDTFSDNFYNRPILSHRNTVWLCYEVKTKGSPRPLDAKIFRGQ  
VYSKLYHPEMRFFHWFSSKWRKLHRDQEYEVTWYISWSPCTKCTRTRDVATFLAEDPKV  
TLTIFVARLYYFWDPDYQEALRSLCQKRDGPRATMKIMNYDEFQHCWSKFVYSQRE  
LFEPWNNLPKYIILLHIMLGEILRHSMDPPTFTSNFNELWVRGRHETYLCYEVEERL  
25 HNDTWVLLNQRRGFLCNQAPHKHGFLEGRHAELCFLDVIPFWKLDLHQDYRVTCFTS  
WSPCFSCAQEMAKFISNNKHVSLCIFAARIYDDQGRCQEGLRTLAKAGAKISIMTYSE  
FKHCWDTFVDHQGCPFQPWDGLEEHSQALSGRLRAILQNQGN (SEQ ID NO: 45)  
(italic: nucleic acid editing domain; underline: cytoplasmic localization signal)

Green monkey APOBEC-3G:

30 MNPQIRNMVEQMEPDIFVYFNNRPILSGRNTVWLCYEVKTKDPSGPPLDANIFQGK  
LYPEAKDHPEMKFLHWFRRKWRQLHRDQEYEVTWYVSWSPCTRCANSVATFLAEDPKV  
TLTIFVARLYYFWKPDYQQALRILCQERGGPHATMKIMNYNEFQHCWNEFVDGQG  
KPFKPRKNLPKHYTLLHATLGELLRHVMDPGTFTSNFNKPKWVSGQRETYLCYKVE  
RSHNDTWVLLNQHRGFLRNQAPDRHGFPKGRHAELCFLDLIPFWKLDLDDQQYRVTCFT

*SWSPCFSCAQKMAKFISNNKHVSLCIFAARIYDDQGRCQEGLRTLHRDGAKIAVMNY  
SEFEYCWDTFVDRQGRPFQPWDGLDEHSQALSGRLRAI* (SEQ ID NO: 46)

(italic: nucleic acid editing domain; underline: cytoplasmic localization signal)

Human APOBEC-3G:

5 MKPHFRNTVERMYRDTFSYNFYNRPILSRNNTVWLCYEVKTKGPSRPPLDAKIFRGQ  
VYSELKYHPEMRFHWFHWSKWRKLHRDQEYEV TWYISWSPCTKCTRDMATFLAEDPKV  
TLTIFVARLYYFWDPDYQEALRSLCQKRDGPRATMKIMNYDEFQHCWSKFVYSQRE  
LFEPWNNLPKYIILLHIMLGEILRHSMDPPTFTFNFNNEPWVRGRHETYLCYEVERM  
HNDTWVLLNQRRGFLCNQAPHKHGFLEGRHAELCFLDVIPFWKLDLDQDYRVTCTFS  
10 *WSPCFSCAQEMAKFISK NKHVSLCIFTARIYDDQGRCQEGLRTLAEAGAKISIMTYSE*  
*FKHCWDTFVDHQGCPFPWDGLDEHSQDLSGRLRAILQNQEN* (SEQ ID NO: 47)

(italic: nucleic acid editing domain; underline: cytoplasmic localization signal)

Human APOBEC-3F:

MKPHFRNTVERMYRDTFSYNFYNRPILSRNNTVWLCYEVKTKGPSRPRLDAKIFRGQ  
15 VYSQPEHHAEMCFLSWFCGNQLPAYKCFQITWVFSWTPCPDCVAKLAEFLAEHPNVTL  
TISAARLYYYWERDYRRALCRLS QAGARVKIMDDEEFAYCWENFVYSEGQPFMPW  
YKFDDNYAFLHRTLKEILRNPMEAMYPHIFYHFKNLRKAYGRNESWLCFTMEVVK  
HHSPVSWKRGVFRNQVDPETHCHAERCFLSWFCDDILSPNTNYEVTWYTSWSPCPECA  
GEVAEFLARHSNVNLTIFTARLYYFWDTDYQEGLRSLSQEGASVEIMGYKDFKYCW  
20 ENFVYNDDEPFKPWKGLKYNFLFLDSKLQEILE

(italic: nucleic acid editing domain)

Human APOBEC-3B:

MNPQIRNPMERMYRDTFYDNFENEPILYGRSYTWLCYEVKIKRGRSNLLWDTGVFR  
GQVYFKPQYHAEMCFLSWFCGNQLPAYKCFQITWVFSWTPCPDCVAKLAEFLSEHPN  
25 VTLTISAARLYYYWERDYRRALCRLS QAGARVTIMDYEEFAYCWENFVYNQGF  
MPWYKFDENY AFLHRTLKEILRYLMDPDTFTFNFNNDPLVLRRRQTYLCYEVERLD  
NGTWVLMQHMGLFCNEAKNLLCGFYGRHAELRFLDLVPSLQLDPAQIYRVTWFISWS  
PCFSWGCAGEVRAFLQENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIMTY  
DEFEYCWDTFVYRQGC PFQPWDGLEEHSQALSGRLRAILQNQGN

(italic: nucleic acid editing domain)

Rat APOBEC-3B:

MQPQGLGPNAGMGPVCLGCSHRRPYSPIRNPLKLYQQTFYHFKNVRYAWGRKN  
NFLCYEVNGMDCALPVPLRQGVFRKQGHIAELCFIYWFHDKVLRVLSPMEEFKVT  
WYMSWSPCSKCAEQVARFLAAHRNLSLAIFSSRLYYYLRNPNYQQKLCRLIQEGVH

VAAMDLPFKKWCWNKFVDNDGQPFPRPWRMLRINFSFYDCKLQEIFSRMNLLREDVF  
 YLQFNNSHRVKPVQNRYYRRKSYLCYQLERANGQEPLKGYLLYKKGEQHVEILFLE  
 KMRSMELSQVRITCYLTWSPCPNCARQLAAFKKDHDPDLILRIYTSRLYFWRKKFKQG  
 LCTLWRSGIHVDVMDLPQFADCWTNFNPNQRPFRPWNELEKNSWRIQRRLRRIKES  
 5 WGL (SEQ ID NO: 50)

Bovine APOBEC-3B:

MDGWEVAFRSGTVLKAGVLGVSMTEGWAGSGHPGQACVWTPGTRNTMNLREV  
 LFKQQFGNQPRVPAPYYRRKTYLCYQLKQRNDLTLDRGCFRNKKQRHAERFIDKIN  
 SLDLNPQSQYKIICYITWSPCPNCANELVNFITRNNHLKLEIFASRLYFHWIKSFKMGL  
 10 QDLQNAGISVAVMTHTEFEDCWEQFVDNQSRPFQPWDKLEQYSASIRRLQRILTAP  
 I (SEQ ID NO: 51)

Chimpanzee APOBEC-3B:

MNPQIRNPMEMYQRTFYNFENEPILYGRSYTWLCYEVKIRRGHSNLLWDTGVFR  
 GQMYSQPEHHAEMCFLSWFCGNQLSAYKCFQITWVFSWTPCPDCVAKLAKFLAEH  
 15 PNVTLTISAARLYYYWERDYRRALCRLSQAGARVKIMDDEEFAYCWENFVYNEGQP  
 FMPWYKFDDNYAFLHRTLKEIIRHLMDDPTFTFNFNNDPLVLRRHQTYLCYEVERLD  
 NGTWVLMQHMGLFCNEAKNLLCGFYGRHAELRFLDLVPSLQLDPAQIYRVTWFIS  
 WSPCFSWGACAGQVRAFLQENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIM  
 TYDEFEYCWDTFVYRQGCPFPWDGLEEHSQALSGRRLRAILQVRASSLCMVPHRPP  
 20 PPQSPGPCLPLCSEPPLGSLPTGRPAPSLPFLTASFSPPPASLPPLPSLSLSPGHLPVP  
 SFHSLTSCSIQPPCSSRIRETEGWASVSKEGRDLG (SEQ ID NO: 52)

Human APOBEC-3C:

MNPQIRNPMKAMYPGTFYFQFKNLWEANDRNETWLCFTVEGIKRRSVVSWKTGVF  
 RNQVDSETH*CHAERCFLSWFCDDILSPNTKYQVTWYTSWSPCPDC*AGEVAEFLARHSN  
 25 VNLTIFTARLYYFQYPCYQEGRLSLSQEGVAVEIMDYEDFKYCWENFVYNDNEPFKP  
 WKGLKTNFRLLKRRLRESLQ (SEQ ID NO: 53)

(italic: nucleic acid editing domain)

Gorilla APOBEC-3C

MNPQIRNPMKAMYPGTFYFQFKNLWEANDRNETWLCFTVEGIKRRSVVSWKTGVF  
 30 RNQVDSETH*CHAERCFLSWECDDILSPNTNYQVTWYTSWSPCPEC*AGEVAEFLARHSN  
 VNLTIFTARLYYFQDQDYQEGRLSLSQEGVAVKIMDYKDFKYCWENFVYNDDEPFK  
 PWKGLKYNFRFLKRRLQEILE (SEQ ID NO: 54)

(italic: nucleic acid editing domain)

Human APOBEC-3A:



MEASPASGPRHLMDPHIFTSNFNNGIGRHKTYLCYEVERLDNGT SVKMDQHRGFLH  
NQAKNLLCGFYGR*HAELRFLDLVPSLQLDPAQIYRVTWFISWSPCF*SWGCAGEVRAFLQ  
ENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIMTYDEFKHCWDTFVDHQGC  
PFQPWDGLDEHSQALSGRLRAILQNQGN (SEQ ID NO: 55)

5 (italic: nucleic acid editing domain)

Rhesus macaque APOBEC-3A:

MDGSPASRPRHLMDPNTFTFNFNNDLSVRGRHQTYLCYEVERLDNGTWVPMDERR  
GFLCNKAKNVPCGDYGC*HVELRFLCEVPSWQLDPAQTYRVTWFISWSPCF*FRRGCAGQ  
VRVFLQENKHVRLRIFAARIYDYDPLYQEALRTL RDAGAQVSIMTYEEFKHCWDTF  
10 VDRQGRPFQPWDGLDEHSQALSGRLRAILQNQGN (SEQ ID NO: 56)

(italic: nucleic acid editing domain)

Bovine APOBEC-3A:

MDEYTFTENFNNQGWPSKTYLCYEMERLDGDATIPLDEYKGFVRNKGLDQPEKPC*H*  
*AELYFLGKIHSWNLDNRNQHYRLTCFISWSPCYDCAQKLTFLKENHHISLHILASRIYTH*  
15 NRFGCHQSGLC ELQAAGARITIMTFEDFKHCWETFVDHKGKPFQWEG LNVKSQAL  
CTELQAILKTQQN (SEQ ID NO: 57)

(italic: nucleic acid editing domain)

Human APOBEC-3H:

MALLTAETFRLQFNKRRLRRPYYPRKALLCYQLTPQNGSTPTRGYFENKKK*CHAEI*  
20 *CFINEIKSMGLDETQCYQVTCYLTWSPCSSCAWELVDFIKAHDHLNLGIFASRLYYHWC*  
KPQQKGLRLLCGSQVPVEVMGFPKFADCWENFVDHEKPLSFNPYKMLEELDKNSRA  
IKRRLERIKIPGVRAQGRYMDILCDAEV (SEQ ID NO: 58)

(italic: nucleic acid editing domain)

Rhesus macaque APOBEC-3H:

MALLTAKTFSLQFNKRRLVKNPYYPRKALLCYQLTPQNGSTPTRGHLKNNKKDHA*E*  
25 IRFINKIKSMGLDETQCYQVTCYLTWSPCPCAGELVDFIKAHRHLNLRIFASRLYYH  
WRPNYQEGLLLLCGSQVPVEVMGLPEFTDCWENFVDHKEPPSFNPSEKLEELDKNS  
QAIKRRLERIKSRSDVLENGLRSLQLGPVTPSSSIRNSR (SEQ ID NO: 59)

Human APOBEC-3D:

MNPQIRNPMERMYRDTFYDNFENEPILYGRSYTWLCYEVKIKRGRSNLLWDTGVFR  
30 GPVLPKRQSNHRQEVYFRFEN*HAEMCFLSWFCGNRLPANRRFQITWFWVSWNPCLPCVV*  
KVTKFLAEHPNVTLTISAARLYYYRDRDWRWVLLRLHKAGARVKIMDYEDFAYCW  
ENFVCNEGQPFMPWYKFDDNYASLHRTLKEILRNPMEAMYPHIFYHFKNLLKACG  
RNESWLCFTMEVTKHHSVFRKRGVFRNQVDPETH*CHAERCFLSWFCDDILSPNTNY*

*EVTWYTSWSPCPE*CAGEVAEFLARHSNVNLTIFTARLCYFWDTDYQEGLCSLSQEGAS  
VKIMGYKDFVSCWKNFVYSDDEPFKPKWGLQTNFRLKRRRLREILQ (SEQ ID NO:  
60)

(italic: nucleic acid editing domain)

5 Human APOBEC-1:

MTSEKGPSTGDPTLRRRIEPWEFDVFYDPRELKKEACLLYEIKWGMSRKIWRSSGKN  
TTNHVEVNFIIKFTSERDFHPSMSCSITWFLSWSPCWECQAIREFLSRHPGVTLVIYV  
ARLFWHMDQQNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQY  
PPLWMMMLYALELHCIIISLPPCLKISRWRQNHLTFFRLHLQNCHYQTIPPHILLATGLI

10 HPSVAWR (SEQ ID NO: 61)

Mouse APOBEC-1:

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKKEACLLYEINWGGRHSVWRHTSQN  
TSNHVEVNFLEKFTTERYFRPNTRCSITWFLSWSPCGECSRAITEFLSRHPYVTLFIYIA  
RLYHHTDQRNRQGLRDLISSGVTIQIMTEQEYCYCWRNFVNYPPSNEAYWPRYPHL  
15 WVKLYVLELYCIIILGLPPCLKILRRKQPQLTFFTITLQTCHYQRIPPHLLWATGLK

(SEQ ID NO: 62)

Rat APOBEC-1:

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKKEACLLYEINWGGRHSIWRHTSQNT  
NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR  
20 LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSPSNEAHWPRYPHLW  
VRLYVLELYCIIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLK (SEQ

ID NO: 63)

Human APOBEC-2:

MAQKEEA AVATEAASQNGEDLENLDDPEKLELIELPPFEIVTGERLPANFFKFQFRN  
25 VEYSSGRNKTFLCYVVEAQGKGGQVQASRGYLEDEHAAAHAEAEAFFNTILPAFDPA  
LRYNVTWYVSSSPCAACADRIIKTLTKNLRLLLILVGRLFMWEEPEIQ AALKKLKE  
AGCKLRIMKPQDFEYVWQNFVEQEEGESKAFQPWEDIQENFLYYEEKLADILK (SEQ

ID NO: 64)

Mouse APOBEC-2:

30 MAQKEEA AEAAAPASQNGDDLLENLEDPEKLELIDLPPFEIVTGVRLPVNFFKFQFR  
NVEYSSGRNKTFLCYVVEVQSKGGQAQATQGYLEDEHAGAHAEAEAFFNTILPAFDP  
ALKYNVTWYVSSSPCAACADRILKTLTKNLRLLLILVSRLFMWEEPEVQAALKKLK  
EAGCKLRIMKPQDFEYIWQNFVEQEEGESKAFEPWEDIQENFLYYEEKLADILK (SEQ

ID NO: 65)

Rat APOBEC-2:

MAQKEEAAEAAAPASQNGDDLENLEDPEKCLKELIDLPPFEIVTGVRLPVNFFKFQFR  
 NVEYSSGRNKTFLCYVVEAQSKGGQVQATQGYLEDEHAGAHAEAEFFNTILPAFDPA  
 ALKYNVTWYVSSSPCAACADRILKTLKTKNLRLLLILVSRLFMWEEPEVQAALKKLLK  
 5 EAGCKLRIMKPQDFEYLWQNFVEQEEGESKAFEPWEDIQENFLYYEEKLADILK  
 (SEQ ID NO: 66)

Bovine APOBEC-2:

MAQKEEAAAAAEPASQNGEEVENLEDPEKCLKELIELPPFEIVTGERLPAHYFKFQFRN  
 VEYSSGRNKTFLCYVVEAQSKGGQVQASRGYLEDEHATNHAEAEFFNSIMPTFDPA  
 10 LRYMVTWYVSSSPCAACADRIVKTLNKTKNLRLLLILVGRLFMWEEPEIQAALRKLKE  
 AGCRLRIMKPQDFEYIWQNFVEQEEGESKAFEPWEDIQENFLYYEEKLADILK (SEQ  
 ID NO: 67)

*Petromyzon marinus* CDA1 (pmCDA1):

MTDAEYVRIHEKLDIYTFKKQFFNNKKSVMHRCYVLFELKRRGERRACFWGYAVNK  
 15 PQSGTERGIHAEIFSIRKVEEYLRDNPQGFTINWYSSWSPCADCAEKILEWYNQELRG  
 NGHTLKIWACKLYYEKNARNQIGLWNLRDNGVGLNVMVSEHYQCCRKIFIQSSHNQ  
 LNENRWLEKTLKRAEKRRSELSFMIQVKILHTTKSPAV (SEQ ID NO: 68)

Human APOBEC3G D316R D317R:

MKPHFRNTVERMYRDTFSYNFYNRPILSRRNTVWLCYEVKTKGPSRPPLDAKIFRGQ  
 20 VYSELKYHPEMRFFHWFSKWRKLHRDQEYEV TWYISWSPCTKCTRDMATFLAEDP  
 KVTLTIFVARLYYFWDPDYQEALRSLCQKRDGPRATMKFN YDEFQHCWSKFVYSQ  
 RELFEPWNNLPKYIILLHFMLGEILRHSMDPPTFTFNFNNEPWVRGRHETYLCYEVE  
 RMHNDTWVLLNQRRGFLCNQAPHKHGFLEGRHAELCFLDVIPFWKLDLDQDYRVT  
 CFTSWSPCFSCAQEMAKFISKKHVSLCIFTARIYRRQGRCQEGLRTLAEAGAKISFTY  
 25 SEFKHCWDTFVDHQGCPFQPWDGLDEHSQDLSGRLRAILQNQEN (SEQ ID NO: 69)

Human APOBEC3G chain A:

MDPPTFTFNFNNEPWVRGRHETYLCYEVE RMHNDTWVLLNQRRGFLCNQAPHKHGF  
 LEGRHAELCFLDVIPFWKLDLDQDYRVTCFTSWSPCFSCAQEMAKFISKKNKHVSLCIF  
 TARIYDDQGRCQEGLRTLAEAGAKISFTYSEFKHCWDTFVDHQGCPFQPWDGLD  
 30 EHSQDLSGRLRAILQ (SEQ ID NO: 70)

Human APOBEC3G chain A D120R D121R:

MDPPTFTFNFNNEPWVRGRHETYLCYEVE RMHNDTWVLLNQRRGFLCNQAPHKHGF  
 FLEGRHAELCFLDVIPFWKLDLDQDYRVTCFTSWSPCFSCAQEMAKFISKKNKHVSLCI

FTARIYRRQGRQCQEGLRTLAEAGAKISFMTYSEFKHCWDTFVDHQGCPFQPWDGLD  
EHSQDLSGRLRAILQ (SEQ ID NO: 71)

hAPOBEC-4 (*Homo sapiens*):

MEPIYEEYLANHGTIVKPYYWLSFSLDCSNCPYHIRTGEEARVSLTEFCQIFGFPYGT  
5 FPQTKHLTFYELKTSSGSLVQKGHASSCTGNYIHPESMLFEMNGYLDSAIYNND  
SIRH  
IILYSNNSPCNEANHCISKMYNFLITYPGITLSIYFSQLYHTEMDFPASAWNREALRS  
LASLWPRVVLSPISGGIWHSVLHSFISGVSGSHVFQPILTGRALADRHNAYEINAITGV  
KPYFTDVLQTKRNPNTKAQEALESYPLNNAFPGQFFQMPSGQLQPNLPPDLRAPVV  
FVLVPLRDLPPMHMGQNPKNPRNIVRHLNMPQMSFQETKDLGRLPTGRSVEIVEITE  
10 QFASSEADEKGGGGGGK (SEQ ID NO: 72)

mAPOBEC-4 (*Mus musculus*):

MDSLLMKQKKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSCSLDFGHLRNKSGC  
HVELLFLRYISDWLDPGRCYRVTWFTSWSPCYDCARHVAEFLRWPNLSLRIFTAR  
LYFCEDRKAPEGLRRLHRAGVQIGIMTFKDYFYCWNTFVENRERTFKAWEGLHEN  
15 SVRLTRQLRRILLPLYEVDDLDAFRMLGF (SEQ ID NO: 73)

rAPOBEC-4 (*Rattus norvegicus*):

MEPLYEEYLTHSGTIVKPYYWLSVSLNCTNCPYHIRTGEEARVPYTEFHQTFGFPWST  
YPQTKHLTFYELRSSSGNLIQKGLASNCTGSHTHPESMLFERDGYLDSLIFHDSNIRHI  
IILYSNNSPCDEANHCISKMYNFMNYPEVTLVFFSPLYHTENQFPTSAWNREALR  
20 GLASLWPQVTLAISGGIWQSILETFVSGISEGLTAVRPFTAGRTLTDRYNAYEINCIT  
EVKPYFTDALHSWQKENQDQKVAASENQPLHNTTPAQWQPDMSQDCRTPAVFM  
LVPYRDLPPIHVNPSQKPRTVVRHLNLTQLSASKVKALRKSPSGRPVKKEEARKGS  
TRSQEANETNKSQKQTLFIKSNICHLLEREQKKIGILSSWSV (SEQ ID NO: 74)

mfAPOBEC-4 (*Macaca fascicularis*):

MEPTYEEYLANHGTIVKPYYWLSFSLDCSNCPYHIRTGEEARVSLTEFCQIFGFPYGT  
25 TYPQTKHLTFYELKTSSGSLVQKGHASSCTGNYIHPESMLFEMNGYLDSAIYNND  
SIRH  
HIILYCNNSPCNEANHCISKVYNFLITYPGITLSIYFSQLYHTEMDFPASAWNREALR  
SLASLWPRVVLSPISGGIWHSVLHSFVSGVSGSHVFQPILTGRALTDRYNAYEINAITG  
VKPFFTDVLLHTKRNPNNTKAQMALESYPLNNAFPGQSFQMTSGIPDLRAPVVFVLL  
30 PLRDLPPMHMGQDPNKPRNIIRHLNMPQMSFQETKDLERLPTRRSVETVEITERFAS  
KQAEKTKGGGGGGK (SEQ ID NO: 75)

pmCDA-1 (*Petromyzon marinus*):

MAGYECVRVSEKLDFTFEFQFENLHYATERHRTYVIFDVKPKQSAGGRSRRLWGYII  
NNPNVCHAELILMSMIDRHLESNPGVYAMTWYMSWSPCANCSSKLNPLWLNLEE

QGHTLTMHFSRIYDRDREGDHRGLRGLKHVSNSFRMGVVGRAEVKECLA EYVEAS  
 RRTL TWLDTTESMAAKMRRKLCILVRCAGMRESGIPLHLFTLQTPLLSGRV VWR  
 V (SEQ ID NO: 76)

pmCDA-2 (*Petromyzon marinus*):

5 MELREVVDCALASCVRHEPLSRV AFLRCFAAPSQKPRGTVILFYVEGAGRGTGGH  
 AVNYNKQGTSIHAEVLLLSAVRAALLRRRRCEDEGEEATR GCTLHCYSTYSPCRDCVE  
 YIQEFGASTGVRVVIHCCRLYELDVNRRRSEAEGLRSL SRLGRDFRLMGPRDAIAL  
 LLGGRLANTADGESGASNAWVTETNVVEPLVDMTGFGDEDLHAQVQRNKQIREA  
 YANYASAVSLMLGELHVDPKFPFLAEFLAQTSVEPSGTPRETRGRPRGASSRGP EIG  
 10 RQRPADFERALGAYGLFLHPRIVSREADREEIKRDLIVVMRKHNYQGP (SEQ ID NO:  
 77)

pmCDA-5 (*Petromyzon marinus*):

MAGDENVRVSEKLDFTFEFQFENLHY ATERHRTYVIFDVK PQSAGGRSRRLWGYII  
 NNPVCHAELILMSMIDRHLESNPGVYAMTWYMSWSPCANCSSKLN PWLKNLLEE  
 15 QGHTLMMHFSRIYDRDREGDHRGLRGLKHVSNSFRMGVVGRAEVKECLA EYVEAS  
 RRTL TWLDTTESMAAKMRRKLCILVRCAGMRESGMPLHLFT (SEQ ID NO: 78)

yCD (*Saccharomyces cerevisiae*):

MVTGGMASKWDQKGMDIAYEEAALGYKEGGVPIGGCLINNKDGSVLGRGHNMRF  
 QKGSATLHGEISTLENCGRLEGKVYKDTTLYTTLSPCDMCTGAIIMYGIPRCV VGEN  
 20 VNFKSKGEKYLQTRGHEVVVDDERCKKIMKQFIDERPQDWFEDIGE (SEQ ID NO:  
 79)

rAPOBEC-1 (delta 177-186):

MSSETGPVAVDPTLRRRIEPHEFEVFFDPREL RKETCLLYEINWGGRHSIWRHTSQNT  
 NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR  
 25 LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNEAHWP RYPHLW  
 VRGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLK (SEQ ID NO: 80)

rAPOBEC-1 (delta 202-213):

MSSETGPVAVDPTLRRRIEPHEFEVFFDPREL RKETCLLYEINWGGRHSIWRHTSQNT  
 NKHVEVNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR  
 30 LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNEAHWP RYPHLW  
 VRLYVLELYCIILGLPPCLNILRRKQPQHYQRLPPHILWATGLK (SEQ ID NO: 81)

Mouse APOBEC-3:

MGPFCLGC SHRKCYSPIRNLISQETFKFHFKNLGYAKGRKDTFLCYEVTRKDCDSPV  
 SLHHGVFKNKDNIHAEICFLYWFHDKVLKVLSPREEFKITWYMSWSPCFECAEQIVRFL

ATHHNLSLDIFSSRLYNVQDPETQQNLCLRVQEGAQVAAMDLYEFKKCWKKFVDN  
 GGRRFRPWKRLLTNFRYQDSKLQEILRPCYIPVSSSSSTLSNICLTKGLPETRFCVEG  
 RRMDPLSEEFYSQFYNQRVKHLCCYYHRMKPYLCYQLEQFNGQAPLKGCLLSEKGG  
 QHAEILFLDKIRSMELSQVTITCYLTWSPCPNCAWQLAAFKRDRPDLILHIYTSRLYFHW  
 5 KRPFQKGLCSLWQSGILVDVMDLPQFTDCWTNFPKRPFWPWKGLEIISRRTQRRR  
 RRIKESWGLQDLVNDVDFGNLQLGPPMS (SEQ ID NO: 82)

(italic: nucleic acid editing domain)

In some embodiments, an adenosine deaminase can comprise all or a portion of an  
 adenosine deaminase ADAR (e.g., ADAR1 or ADAR2). In another embodiment, an  
 10 adenosine deaminase can comprise all or a portion of an adenosine deaminase ADAT. In  
 some embodiments, an adenosine deaminase can comprise all or a portion of an ADAT from  
 Escherichia coli (EcTadA) comprising one or more of the following mutations: D108N,  
 A106V, D147Y, E155V, L84F, H123Y, I157F, or a corresponding mutation in another  
 adenosine deaminase. The adenosine deaminase can be derived from any suitable organism  
 15 (e.g., *E. coli*). In some embodiments, the adenosine deaminase is from *Escherichia coli*,  
*Staphylococcus aureus*, *Salmonella typhi*, *Shewanella putrefaciens*, *Haemophilus influenzae*,  
*Caulobacter crescentus*, or *Bacillus subtilis*. In some embodiments, the adenosine deaminase  
 is from *E. coli*. In some embodiments, the adenine deaminase is a naturally-occurring  
 adenosine deaminase that includes one or more mutations corresponding to any of the  
 20 mutations provided herein (e.g., mutations in ecTadA). The corresponding residue in any  
 homologous protein can be identified by e.g., sequence alignment and determination of  
 homologous residues. The mutations in any naturally-occurring adenosine deaminase (e.g.,  
 having homology to ecTadA) that corresponds to any of the mutations described herein (e.g.,  
 any of the mutations identified in ecTadA) can be generated accordingly. In particular  
 25 embodiments, the TadA is any one of the TadA described in PCT/US2017/045381 (WO  
 2018/027078), which is incorporated herein by reference in its entirety. Mutations were  
 identified through rounds of evolution and selection (e.g., TadA\*7.10 = variant 10 from  
 seventh round of evolution) having desirable adenosine deaminase activity on single stranded  
 DNA as shown in Table 3.

30 Table 3. Genotypes of TadA Variants

TadA	23	26	36	37	48	49	51	72	84	87	105	108	123	125	142	145	147	152	155	156	157	16
0.1	W	R	H	N	P		R	N	L	S	A	D	H	G	A	S	D	R	E	I	K	K
0.2	W	R	H	N	P		R	N	L	S	A	D	H	G	A	S	D	R	E	I	K	K

TadA	23	26	36	37	48	49	51	72	84	87	105	108	123	125	142	145	147	152	155	156	157	16
1.1	W	R	H	N	P		R	N	L	S	A	N	H	G	A	S	D	R	E	I	K	K
1.2	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	D	R	E	I	K	K
2.1	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.2	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.3	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.4	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.5	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.6	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.7	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.8	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.9	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.10	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.11	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
2.12	W	R	H	N	P		R	N	L	S	V	N	H	G	A	S	Y	R	V	I	K	K
3.1	W	R	H	N	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	K
3.2	W	R	H	N	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	K
3.3	W	R	H	N	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	K
3.4	W	R	H	N	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	K
3.5	W	R	H	N	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	K
3.6	W	R	H	N	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	K
3.7	W	R	H	N	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	K
3.8	W	R	H	N	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	K
4.1	W	R	H	N	P		R	N	L	S	V	N	H	G	N	S	Y	R	V	I	K	K
4.2	W	G	H	N	P		R	N	L	S	V	N	H	G	N	S	Y	R	V	I	K	K
4.3	W	R	H	N	P		R	N	F	S	V	N	Y	G	N	S	Y	R	V	F	K	K
5.1	W	R	L	N	P		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
5.2	W	R	H	S	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	T
5.3	W	R	L	N	P		L	N	I	S	V	N	Y	G	A	C	Y	R	V	I	N	K
5.4	W	R	H	S	P		R	N	F	S	V	N	Y	G	A	S	Y	R	V	F	K	T
5.5	W	R	L	N	P		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
5.6	W	R	L	N	P		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
5.7	W	R	L	N	P		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
5.8	W	R	L	N	P		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
5.9	W	R	L	N	P		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K

TadA	23	26	36	37	48	49	51	72	84	87	105	108	123	125	142	145	147	152	155	156	157	16
5.10	W	R	L	N	P		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
5.11	W	R	L	N	P		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
5.12	W	R	L	N	P		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
5.13	W	R	H	N	P		L	D	F	S	V	N	Y	A	A	S	Y	R	V	F	K	K
5.14	W	R	H	N	S		L	N	F	C	V	N	Y	G	A	S	Y	R	V	F	K	K
6.1	W	R	H	N	S		L	N	F	S	V	N	Y	G	N	S	Y	R	V	F	K	K
6.2	W	R	H	N	T	V	L	N	F	S	V	N	Y	G	N	S	Y	R	V	F	N	K
6.3	W	R	L	N	S		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
6.4	W	R	L	N	S		L	N	F	S	V	N	Y	G	N	C	Y	R	V	F	N	K
6.5	W	R	L	N	I	V	L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
6.6	W	R	L	N	T	V	L	N	F	S	V	N	Y	G	N	C	Y	R	V	F	N	K
7.1	W	R	L	N	A		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
7.2	W	R	L	N	A		L	N	F	S	V	N	Y	G	N	C	Y	R	V	F	N	K
7.3	I	R	L	N	A		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
7.4	R	R	L	N	A		L	N	F	S	V	N	Y	G	A	C	Y	R	V	F	N	K
7.5	W	R	L	N	A		L	N	F	S	V	N	Y	G	A	C	Y	H	V	F	N	K
7.6	W	R	L	N	A		L	N	I	S	V	N	Y	G	A	C	Y	P	V	I	N	K
7.7	L	R	L	N	A		L	N	F	S	V	N	Y	G	A	C	Y	P	V	F	N	K
7.8	I	R	L	N	A		L	N	F	S	V	N	Y	G	N	C	Y	R	V	F	N	K
7.9	L	R	L	N	A		L	N	F	S	V	N	Y	G	N	C	Y	P	V	F	N	K
7.10	R	R	L	N	A		L	N	F	S	V	N	Y	G	A	C	Y	P	V	F	N	K

In some embodiments, the TadA is provided as a monomer or dimer (e.g., a heterodimer of wild-type *E. coli* TadA and an engineered TadA variant). In some embodiments, the adenosine deaminase is an eighth generation TadA\*8 variant as shown in Table 4 below.

5 Table 4: TadA8\* Adenosine Deaminase Variants

<b>Adenosine Deaminase</b>	<b>Adenosine Deaminase Description</b>
TadA*8.1	Monomer_TadA*7.10 + Y147T
TadA*8.2	Monomer_TadA*7.10 + Y147R
TadA*8.3	Monomer_TadA*7.10 + Q154S
TadA*8.4	Monomer_TadA*7.10 + Y123H



<b>Adenosine Deaminase</b>	<b>Adenosine Deaminase Description</b>
TadA*8.5	Monomer_TadA*7.10 + V82S
TadA*8.6	Monomer_TadA*7.10 + T166R
TadA*8.7	Monomer_TadA*7.10 + Q154R
TadA*8.8	Monomer_TadA*7.10 + Y147R_Q154R_Y123H
TadA*8.9	Monomer_TadA*7.10 + Y147R_Q154R_I76Y
TadA*8.10	Monomer_TadA*7.10 + Y147R_Q154R_T166R
TadA*8.11	Monomer_TadA*7.10 + Y147T_Q154R
TadA*8.12	Monomer_TadA*7.10 + Y147T_Q154S
TadA*8.13	Monomer_TadA*7.10 + H123H_Y147R_Q154R_I76Y
TadA*8.14	Heterodimer_(WT) + (TadA*7.10 + Y147T)
TadA*8.15	Heterodimer_(WT) + (TadA*7.10 + Y147R)
TadA*8.16	Heterodimer_(WT) + (TadA*7.10 + Q154S)
TadA*8.17	Heterodimer_(WT) + (TadA*7.10 + Y123H)
TadA*8.18	Heterodimer_(WT) + (TadA*7.10 + V82S)
TadA*8.19	Heterodimer_(WT) + (TadA*7.10 + T166R)
TadA*8.20	Heterodimer_(WT) + (TadA*7.10 + Q154R)
TadA*8.21	Heterodimer_(WT) + (TadA*7.10 + Y147R_Q154R_Y123H)
TadA*8.22	Heterodimer_(WT) + (TadA*7.10 + Y147R_Q154R_I76Y)
TadA*8.23	Heterodimer_(WT) + (TadA*7.10 + Y147R_Q154R_T166R)
TadA*8.24	Heterodimer_(WT) + (TadA*7.10 + Y147T_Q154R)
TadA*8.25	Heterodimer_(WT) + (TadA*7.10 + Y147T_Q154S)
TadA*8.26	Heterodimer_(WT) + (TadA*7.10 + H123H_Y147T_Q154R_I76Y)

In some embodiments, the adenosine deaminase is a ninth generation TadA\*9 variant containing an alteration at an amino acid position selected from the following: 21, 23, 25, 38,

51, 54, 70, 71, 72, 72, 94, 124, 133, 138, 139, 146, and 158 of a TadA variant as shown in the reference sequence below:

```

      10          20          30          40          50
MSEVEFSHEY WMRHALTLAK RARDEREVPV GAVLVLNNRV IGEGWNRAIG
5          60          70          80          90          100
LHDPTAHA EI MALRQGGLVM QNYRLIDATL YVTFEPCVMC AGAMIHSRIG
      110          120          130          140          150
RVVFGVRNAK TGAAGSLMDV LHYPGMNHRV EITEGILADE CAALLCYFFR
      160
10 MPRQVFNAAQK KAQSSTD (SEQ ID NO: 83)

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In one embodiment, the adenosine deaminase variant contains alterations at two or more amino acid positions selected from the following: 21, 23, 25, 38, 51, 54, 70, 71, 72, 94, 124, 133, 138, 139, 146, and 158 of the TadA reference sequence above. In another embodiment, the adenosine deaminase variant contains one or more (e.g., 2, 3, 4) alterations selected from the following: R21N, R23H, E25F, N38G, L51W, P54C, M70V, Q71M, N72K, Y73S, M94V, P124W, T133K, D139L, D139M, C146R, and A158K of SEQ ID NO.

1. In other embodiments, the adenosine deaminase variant further contains one or more of the following alterations: Y147T, Y147R, Q154S, Y123H, and Q154R. In still other embodiments, the adenosine deaminase variant contains a combination of alterations relative to the above TadA reference sequence selected from the following: E25F + V82S + Y123H, T133K + Y147R + Q154R; E25F + V82S + Y123H + Y147R + Q154R; L51W + V82S + Y123H + C146R + Y147R + Q154R; Y73S + V82S + Y123H + Y147R + Q154R; P54C + V82S + Y123H + Y147R + Q154R; N38G + V82T + Y123H + Y147R + Q154R; N72K + V82S + Y123H + D139L + Y147R + Q154R; E25F + V82S + Y123H + D139M + Y147R + Q154R; Q71M + V82S + Y123H + Y147R + Q154R; E25F + V82S + Y123H + T133K + Y147R + Q154R; E25F + V82S + Y123H + Y147R + Q154R; V82S + Y123H + P124W + Y147R + Q154R; L51W + V82S + Y123H + C146R + Y147R + Q154R; P54C + V82S + Y123H + Y147R + Q154R; Y73S + V82S + Y123H + Y147R + Q154R; N38G + V82T + Y123H + Y147R + Q154R; R23H + V82S + Y123H + Y147R + Q154R; R21N + V82S + Y123H + Y147R + Q154R; V82S + Y123H + Y147R + Q154R + A158K; N72K + V82S + Y123H + D139L + Y147R + Q154R; E25F + V82S + Y123H + D139M + Y147R + Q154R; M70V + V82S + M94V + Y123H + Y147R + Q154R; Q71M + V82S + Y123H + Y147R + Q154R; E25F + I76Y + V82S + Y123H + Y147R + Q154R; I76Y + V82T + Y123H +

Y147R + Q154R; N38G + I76Y + V82S + Y123H + Y147R + Q154R; R23H + I76Y + V82S  
 + Y123H + Y147R + Q154R; P54C + I76Y + V82S + Y123H + Y147R + Q154R; R21N +  
 I76Y + V82S + Y123H + Y147R + Q154R; I76Y + V82S + Y123H + D138M + Y147R +  
 Q154R; Y72S + I76Y + V82S + Y123H + Y147R + Q154R; E25F + I76Y + V82S + Y123H  
 5 + Y147R + Q154R; I76Y + V82T + Y123H + Y147R + Q154R; N38G + I76Y + V82S +  
 Y123H + Y147R + Q154R; R23H + I76Y + V82S + Y123H + Y147R + Q154R; P54C +  
 I76Y + V82S + Y123H + Y147R + Q154R; R21N + I76Y + V82S + Y123H + Y147R +  
 Q154R; I76Y + V82S + Y123H + D138M + Y147R + Q154R; Y72S + I76Y + V82S +  
 Y123H + Y147R + Q154R; and V82S + Q154R; N72K\_V82S + Y123H + Y147R + Q154R;  
 10 Q71M\_V82S + Y123H + Y147R + Q154R; V82S + Y123H + T133K + Y147R + Q154R;  
 V82S + Y123H + T133K + Y147R + Q154R + A158K; M70V + Q71M + N72K + V82S +  
 Y123H + Y147R + Q154R; N72K\_V82S + Y123H + Y147R + Q154R; Q71M\_V82S +  
 Y123H + Y147R + Q154R; M70V + V82S + M94V + Y123H + Y147R + Q154R; V82S +  
 Y123H + T133K + Y147R + Q154R; V82S + Y123H + T133K + Y147R + Q154R +  
 15 A158K; and M70V + Q71M + N72K + V82S + Y123H + Y147R + Q154R. In some  
 embodiments, the deaminase or other polypeptide sequence lacks a methionine, for example  
 when included as a component of a fusion protein. This can alter the numbering of positions.  
 However, the skilled person will understand that such corresponding mutations refer to the  
 same mutation, e.g., Y73S and Y72S and D139M and D138M.

20 In some embodiments, Cas9 is fused to nuclear localization sequences, including an  
 NLS of the SV40 large T antigen, nucleoplasmin, c-myc, hRNPA1 M9, IBB domain from  
 importin-alpha, NLS of myoma T protein, human p53, c-abl IV, influenza virus NS1,  
 hepatitis virus delta antigen, mouse Mx1, human poly(ADP-ribose) polymerase, steroid  
 hormone receptor (human) glucocorticoid.

25 In some embodiments, a Cas9 protein is fused to epitope tags including, but not  
 limited to hemagglutinin (HA) tags, histidine (His) tags, FLAG tags, Myc tags, V5 tags,  
 VSV-G tags, SNAP tags, thioredoxin (Trx) tags.

30 In some embodiments, Cas9 is fused to reporter genes including, but not limited to  
 glutathione-S-transferase (GST), horseradish peroxidase (HRP), chloramphenicol transferase  
 (CAT), HcRed, DsRed, cyan fluorescent protein, yellow fluorescent protein and blue  
 fluorescent protein, green fluorescent protein (GFP), including enhanced versions or  
 superfolded GFP, as well as other modified versions of reporter genes.

In some embodiments, serum half-life of an engineered Cas9 protein is increased by fusion with heterologous proteins such as a human serum albumin protein, transferrin protein, human IgG and/or sialylated peptide, such as the carboxy-terminal peptide (CTP, of chorionic gonadotropin  $\beta$  chain).

5 In some embodiments, serum half-life of an engineered Cas9 protein is decreased by fusion with destabilizing domains, including but not limited to geminin, ubiquitin, FKBP12-L106P, and/or dihydrofolate reductase.

Suitable fusion partners that provide for increased or decreased stability include, but are not limited to degen sequences. Degrons are readily understood by one of ordinary skill  
10 in the art to be amino acid sequences that control the stability of the protein of which they are part. For example, the stability of a protein comprising a degen sequence is controlled at least in part by the degen sequence. In some cases, a suitable degen is constitutive such that the degen exerts its influence on protein stability independent of experimental control (i.e., the degen is not drug inducible, temperature inducible, etc.) In some cases, the degen  
15 provides the variant Cas9 polypeptide with controllable stability such that the variant Cas9 polypeptide can be turned "on" (i.e., stable) or "off" (i.e., unstable, degraded) depending on the desired conditions. For example, if the degen is a temperature sensitive degen, the variant Cas9 polypeptide may be functional (i.e., "on", stable) below a threshold temperature (e.g., 42°C, 41°C, 40°C, 39°C, 38°C, 37°C, 36°C, 35°C, 34°C, 33°C, 32°C, 31°C, 30°C, etc.) but  
20 non-functional (i.e., "off, degraded) above the threshold temperature. As another example, if the degen is a drug inducible degen, the presence or absence of drug can switch the protein from an "off (i.e., unstable) state to an "on" (i.e., stable) state or vice versa. An exemplary drug inducible degen is derived from the FKBP12 protein. The stability of the degen is controlled by the presence or absence of a small molecule that binds to the degen.

25 Examples of suitable degrons include, but are not limited to those degrons controlled by Shield-1, DHFR, auxins, and/or temperature. Non-limiting examples of suitable degrons are known in the art (e.g., Dohmen et al., Science, 1994, 263(5151): p. 1273-1276: Heat-inducible degen: a method for constructing temperature-sensitive mutants; Schoeber et al., Am J Physiol Renal Physiol. 2009 Jan;296(1):F204-11 : Conditional fast expression and  
30 function of multimeric TRPV5 channels using Shield-1 ; Chu et al., Bioorg Med Chem Lett. 2008 Nov 15;18(22):5941-4: Recent progress with FKBP-derived destabilizing domains ; Kanemaki, Pflugers Arch. 2012 Dec 28: Frontiers of protein expression control with conditional degrons; Yang et al., Mol Cell. 2012 Nov 30;48(4):487-8: Titivated for

destruction: the methyl degnon; Barbour et al., Biosci Rep. 2013 Jan 18;33(1).: Characterization of the bipartite degnon that regulates ubiquitin-independent degradation of thymidylate synthase; and Greussing et al., J Vis Exp. 2012 Nov 10;(69): Monitoring of ubiquitin-proteasome activity in living cells using a Degron (dgn)-destabilized green fluorescent protein (GFP)-based reporter protein; all of which are hereby incorporated in their entirety by reference).

Exemplary degnon sequences have been well-characterized and tested in both cells and animals. Thus, fusing dead Cas9 to a degnon sequence produces a "tunable" and "inducible" dead Cas9 polypeptide.

Any of the fusion partners described herein can be used in any desirable combination. As one non-limiting example to illustrate this point, a Cas9 fusion protein can comprise a YFP sequence for detection, a degnon sequence for stability, and transcription activator sequence to increase transcription of the target DNA. Furthermore, the number of fusion partners that can be used in a dCas9 fusion protein is unlimited. In some cases, a Cas9 fusion protein comprises one or more (e.g. two or more, three or more, four or more, or five or more) heterologous sequences.

### **Target Nucleic Acids**

A target nucleic acid is a DNA molecule, RNA molecule, which is single-, double-, or multi-stranded DNA or RNA, genomic DNA, cDNA, DNA-RNA hybrids, or a polymer comprising purine and pyrimidine bases or other natural, chemically or biochemically modified, non-natural, or derivatized nucleotide bases either deoxyribonucleotides, ribonucleotides, or analogs thereof. Target nucleic acids may have three-dimensional structure, may include coding or non-coding regions, may include exons, introns, mRNA, tRNA, rRNA, siRNA, shRNA, miRNA, ribozymes, cDNA, plasmids, vectors, exogenous sequences, endogenous sequences. A target nucleic acid can comprise modified nucleotides, include methylated nucleotides, or nucleotide analogs. In some embodiments, a target nucleic acid may be interspersed with non-nucleic acid components.

A target nucleic acid is recognized by CRISPR-Cas9 system and binds Cas9. In some embodiments, it is modified or cleaved or has altered expression due to the binding of Cas9. A target nucleic acid contains a specific recognizable PAM motif, for example, 5'-NGG-3', 5'-NGC-3', 5'-NAGHC-3', 5'-NRHRRH-3' or 5'-NNAAA-3' (H=A, C or T; R=A or G).

### **Recombinant Gene Technology**

In accordance with the present disclosure, there may be employed conventional molecular biology, microbiology, and recombinant DNA techniques within the skill of the art. Such techniques are described in the literature (see, *e.g.*, Sambrook, Fritsch & Maniatis, *Molecular Cloning: A Laboratory Manual*, Second Edition (1989) Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.; *DNA Cloning: A Practical Approach*, Volumes I and II (D. N. Glover ed. 1985); *Oligonucleotide Synthesis* (M. J. Gait ed. 1984); *Nucleic Acid Hybridization* (B. D. Hames & S. J. Higgins eds. (1985)); *Transcription And Translation* (B. D. Hames & S. J. Higgins, eds. (1984)); *Animal Cell Culture* (R. I. Freshney, ed. (1986)); *Immobilized Cells and Enzymes* (IRL Press, (1986)); B. Perbal, *A Practical Guide To Molecular Cloning* (1984); F. M. Ausubel *et al.* (eds.), *Current Protocols in Molecular Biology*, John Wiley & Sons, Inc. (1994).

Recombinant expression of a gene, such as a nucleic acid encoding a polypeptide, such as an engineered Cas9 enzyme described herein, can include construction of an expression vector containing a nucleic acid that encodes the polypeptide. Once a polynucleotide has been obtained, a vector for the production of the polypeptide can be produced by recombinant DNA technology using techniques known in the art. Known methods can be used to construct expression vectors containing polypeptide coding sequences and appropriate transcriptional and translational control signals. These methods include, for example, *in vitro* recombinant DNA techniques, synthetic techniques, and *in vivo* genetic recombination.

An expression vector can be transferred to a host cell by conventional techniques, and the transfected cells can then be cultured by conventional techniques to produce polypeptides.

In some embodiments, a nucleotide sequence encoding a DNA-targeting RNA and/or Cas9 protein is operably linked to a control element, *e.g.*, a transcriptional control element, such as a promoter. The transcriptional control element may be functional in either a eukaryotic cell, *e.g.*, a mammalian cell; or a prokaryotic cell (*e.g.*, bacterial or archaeal cell). In some embodiments, the eukaryotic cell is a human cell. In some embodiments, a nucleotide sequence encoding a DNA-targeting RNA and/or a novel Cas9 protein is operably linked to multiple control elements that allow expression of the encoded nucleotide sequence in both prokaryotic and eukaryotic cells.

A promoter can be a constitutively active promoter (*i.e.*, a promoter that is constitutively in an active/"ON" state), it may be an inducible promoter (*i.e.*, a promoter

whose state, active/"ON" or inactive/"OFF", is controlled by an external stimulus, e.g., the presence of a particular temperature, compound, or protein.), it may be a spatially restricted promoter (i.e., transcriptional control element, enhancer, etc.)(e.g., tissue specific promoter, cell type specific promoter, etc.), and it may be a temporally restricted promoter (i.e., the promoter is in the "ON" state or "OFF" state during specific stages of embryonic development or during specific stages of a biological process, e.g., hair follicle cycle in mice).

Suitable promoters can be derived from viruses and can therefore be referred to as viral promoters, or they can be derived from any organism, including prokaryotic or eukaryotic organisms. Suitable promoters can be used to drive expression by any RNA polymerase (e.g., pol I, pol II, pol III). Exemplary promoters include, but are not limited to the SV40 early promoter, mouse mammary tumor virus long terminal repeat (LTR) promoter; adenovirus major late promoter (Ad MLP); a herpes simplex virus (HSV) promoter, a cytomegalovirus (CMV) promoter such as the CMV immediate early promoter region (CMVIE), a rous sarcoma virus (RSV) promoter, a human U6 small nuclear promoter (U6) (Miyagishi et al. , Nature Biotechnology 20, 497 - 500 (2002)), an enhanced U6 promoter (e.g., Xia et al., Nucleic Acids Res. 2003 Sep 1;31(17)), and/or a human HI promoter (HI).

Examples of inducible promoters include, but are not limited to T7 RNA polymerase promoter, T3 RNA polymerase promoter, Isopropyl-beta-D-thiogalactopyranoside (IPTG) - regulated promoter, lactose induced promoter, heat shock promoter, Tetracycline-regulated promoter (e.g., Tet-ON, Tet-OFF, etc.), Steroid-regulated promoter, Metal-regulated promoter, estrogen receptor-regulated promoter, etc. Inducible promoters can therefore be regulated by molecules including, but not limited to, doxycycline, RNA polymerase, e.g., T7 RNA polymerase, an estrogen receptor and/or an estrogen receptor fusion.

In some embodiments, the promoter is a spatially restricted promoter (i.e., cell type specific promoter, tissue specific promoter, etc.) such that in a multi-cellular organism, the promoter is active (i.e., "ON") in a subset of specific cells. Spatially restricted promoters may also be referred to as enhancers, transcriptional control elements, control sequences, etc. Any convenient spatially restricted promoter may be used and the choice of suitable promoter (e.g., a brain specific promoter, a promoter that drives expression in a subset of neurons, a promoter that drives expression in the germline, a promoter that drives expression in the lungs, a promoter that drives expression in muscles, a promoter that drives expression in islet cells of the pancreas, etc.) will depend on the organism. Thus, a spatially restricted promoter

can be used to regulate the expression of a nucleic acid encoding a subject site-directed polypeptide in a wide variety of different tissues and cell types, depending on the organism. Some spatially restricted promoters are also temporally restricted such that the promoter is in the "ON" state or "OFF" state during specific stages of embryonic development or during  
5 specific stages of a biological process (e.g., hair follicle cycle).

For illustration purposes, examples of spatially restricted promoters include, but are not limited to, neuron-specific promoters, adipocyte-specific promoters, cardiomyocyte-specific promoters, smooth muscle-specific promoters, photoreceptor-specific promoters, etc. Neuron-specific spatially restricted promoters include, but are not limited to, a neuron-  
10 specific enolase (NSE) promoter, an aromatic amino acid decarboxylase (AADC) promoter, a neurofilament promoter, a synapsin promoter, a thy-1 promoter, a serotonin receptor promoter, a tyrosine hydroxylase promoter (TH), a GnRH promoter, an L7 promoter, a DNMT promoter, an enkephalin promoter, a myelin basic protein (MBP) promoter, a Ca<sup>2+</sup>-calmodulin- dependent protein kinase II-alpha (CamKIIa) promoter and/or a CMV  
15 enhancer/platelet-derived growth factor-β promoter.

Adipocyte-specific spatially restricted promoters include, but are not limited to aP2 gene promoter/enhancer, e.g., a region from -5.4 kb to +21 bp of a human aP2 gene, a glucose transporter-4 (GLUT4) promoter, a fatty acid translocase (FAT/CD36) promoter, a stearoyl-CoA desaturase-1 (SCD1) promoter, a leptin promoter, and an adiponectin promoter,  
20 an adipsin promoter and/or a resistin promoter.

Cardiomyocyte-specific spatially restricted promoters include, but are not limited to control sequences derived from the following genes: myosin light chain-2, a-myosin heavy chain, AE3, cardiac troponin C, and/or cardiac actin.

Smooth muscle-specific spatially restricted promoters include, but are not limited to  
25 an SM22a promoter, a smoothelin promoter, and/or an a-smooth muscle actin promoter.

Photoreceptor-specific spatially restricted promoters include, but are not limited to, a rhodopsin promoter, a rhodopsin kinase promoter, a beta phosphodiesterase gene promoter, a retinitis pigmentosa gene promoter, an interphotoreceptor retinoid-binding protein (IRBP) gene enhancer, and/or an IRBP gene promoter.

30 **Gene Editing Uses of CRISPR-Cas9**



The CRISPR-Cas9 system described herein can be used for gene editing, which can result in a gene silencing event, or an alteration of the expression (e.g., an increase or a decrease) in the expression of a desired target gene. Accordingly, in some embodiments, the CRISPR-Cas9 system described herein is used in a method of altering the expression of a target nucleic acid. In some embodiments the CRISPR-Cas9 system described herein is used in a method of modifying a target nucleic acid in a desired target cell. In some embodiments, the invention provides methods for site-specific modification of a target nucleic acid in eukaryotic cells to effectuate a desired modification in gene expression.

In some embodiments, the invention provides an engineered, non-naturally occurring CRISPR-Cas system comprising: an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to a target nucleic acid; and a codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to SEQ ID NO: 1, 4, 8, 14, 84 or 86, and wherein the Cas protein is capable of binding to the RNA guide and of causing a break in the target nucleic acid sequence complementary to the RNA guide.

In some embodiments, the invention provides engineered, non-naturally occurring CRISPR-Cas system comprising: an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to a target nucleic acid; and a codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to SEQ ID NO: 1, 4, 8, 14, 84 or 86; wherein the Cas protein is fused to a deaminase, and wherein the Cas protein fusion is capable of binding to the RNA guide and of editing the target nucleic acid sequence complementary to the RNA guide.

In some embodiments, provided herein is an engineered, non-naturally occurring CRISPR-Cas system comprising a codon-optimized CRISPR-associated (Cas) protein, further comprising a nuclear localization sequence (NLS) and/or a FLAG, HIS or HA tag.

In some embodiments, provided herein is an engineered, non-naturally occurring Cas9 fusion protein further comprising a nuclear localization sequence (NLS) and/or a FLAG, HIS or HA tag.

In some embodiments, provided herein is an engineered, non-naturally occurring Cas9 fusion protein having at least 80% identity to SEQ ID NOs: 2, 5, 9, 15, 85, 87, 95 or 96.

In some embodiments, the invention provides a method of altering expression of a target nucleic acid in a eukaryotic cell comprising: contacting the cell with a Cas9 described herein, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide  
5 comprises a direct repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and wherein the Cas9 protein is capable of binding to the RNA guide and of causing a break in the target nucleic acid sequence complementary to the RNA guide.

In some embodiments, the invention provides a method of altering expression of a target nucleic acid in a eukaryotic cell comprising: contacting the cell with a Cas9 described  
10 herein, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and wherein the Cas9 protein is capable of binding to the RNA guide and editing the target nucleic acid sequence complementary to the RNA guide.

In some embodiments, the invention provides a method of modifying a target nucleic  
15 acid in a eukaryotic cell comprising: contacting the cell with a Cas9 described herein, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and wherein the Cas9 protein is capable of binding to the RNA guide and editing the target nucleic acid sequence complementary to the RNA guide.

20 Accordingly, in some embodiments, the Cas protein has about 80%, 81%, 82%, 83%, 84%, 85%, 86%, 87%, 88%, 89%, 90%, 91%, 92%, 93%, 94%, 95%, 96%, 97%, 98%, 99% identity to SEQ ID NO: 1, 4, 8, 14, 84 or 86. In some embodiments, the Cas protein is identical to SEQ ID NO: 1, 4, 8, 14, 84 or 86.

Suitable guide RNA, Cas9 mutations and fusion proteins for use in the CRISPR-Cas9  
25 system and method are as described throughout this disclosure.

In one aspect, the method comprises binding of the CRISPR-Cas9 to a target nucleic acid and effecting cleavage of a target nucleic acids. In some embodiments, the CRISPR-Cas9 system cleaves target DNA or RNA duplexes by introducing double-stranded breaks. In some embodiments, the CRISPR-Cas9 system cleaves target DNA or RNA by introducing  
30 single-stranded breaks or nicks.

In some embodiments, the CRISPR-Cas9 method or system comprises a fusion protein with an effector that modifies target DNA in a site-specific manner, where the

modifying activity includes methyltransferase activity, demethylase activity, acetyltransferase activity, deacetylase activity, kinase activity, phosphatase activity, ubiquitin ligase activity, deubiquitinating activity, adenylation activity, deadenylation activity, SUMOylating activity, deSUMOylating activity, ribosylation activity, deribosylation activity, myristoylation  
5 activity, demyristoylation activity, integrase activity, transposase activity, recombinase activity, polymerase activity, ligase activity, helicase activity, or nuclease activity, any of which can modify DNA or a DNA-associated polypeptide (e.g., a histone or DNA binding protein).

In some embodiments, the CRISPR-Cas9 method or system comprises a fusion  
10 protein with enzymes that can edit DNA sequences by chemically modifying nucleotide bases, including deaminase enzymes that can modify adenosine or cytosine bases and function as site-specific base editors. For example, APOBEC1 cytidine deaminase, which usually uses RNA as a substrate, can be targeted to single-stranded and double-stranded DNA when it is fused to Cas9, converting cytidine to uridine directly, and ADAR enzymes  
15 deaminate adenosine to inosine. Thus, 'base editing' using deaminases enables programmable conversion of one target DNA base into another. Various base editors are known in the art and can be used in the method and systems described herein. Exemplary base editors are described in, for example, Rees and Liu *Nature Review Genetics*, 2018, 19(12): 770-788, the contents of which are incorporated herein. Accordingly, in some embodiments, the Cas9  
20 enzymes (ScoCas9, SirCas9, VapCas9, EpeCas9, LfeCas9, PmaCas9) described herein is a component of a nucleobase editor. In some embodiments, the base editor is the adenine deaminase TadA8 or TadA9.

In some embodiments, base editing results in the introduction of stop codons to  
25 silence genes. In some embodiments, base editing results in altered protein function by altering amino acid sequences.

In some embodiments, the CRISPR-Cas9 method or system comprises epigenetic  
modification of target DNA by fusion with a histone. In some embodiments, the CRISPR-  
Cas9 system comprises epigenetic modification of target DNA by fusion with an epigenetic  
modifying enzyme such as a reader, writer or eraser protein. In some embodiments, the  
30 CRISPR-Cas9 system comprises fusion with a histone modifying enzyme to alter the histone modification pattern in a selected region of target DNA. Histone modifications can occur in many different ways including methylation, acetylation, ubiquitination, phosphorylation, and

in many different combinations, leading to structural changes in DNA. In some embodiments, histone modification leads to transcriptional repression or activation.

In some embodiments, the CRISPR-Cas9 method or system modulates transcription of target DNA by increasing or decreasing transcription through fusion with transcriptional  
5 activator proteins or transcriptional repressor proteins, small molecule/drug-responsive transcriptional regulators, inducible transcription regulators. In some embodiments, the CRISPR-Cas9 system is used to control the expression of a target coding mRNA (i.e. a protein encoding gene) where binding results in increased or decreased gene expression.

In some embodiments, the CRISPR-Cas9 method or system is used to control gene  
10 regulation by editing genetic regulatory elements such as promoters or enhancers.

In some embodiments, the CRISPR-Cas9 method or system is used to control the expression of a target non-coding RNA, including tRNA, rRNA, snoRNA, siRNA, miRNA, and long ncRNA.

In some embodiments, the CRISPR-Cas9 method or system is used for targeted  
15 engineering of chromatin loop structures. Targeted engineering of chromatin loops between regulatory genomic regions provides a means to manipulate endogenous chromatin structures and enable the formation of new enhancer-promoter connections to overcome genetic deficiencies or inhibit aberrant enhancer-promoter connections.

In some embodiments, CRISPR-Cas9 is used for live cell imaging. Fluorescently  
20 labelled Cas9 is targeted to repetitive genomic regions such as centromeres and telomeres to track native chromatin loci throughout the cell cycle and determine differential positioning of transcriptionally active and inactive regions in the 3D nuclear space.

In some embodiments, the CRISPR-Cas9 method or system is used for correction of pathogenic mutations by insertion of beneficial clinical variants or suppressor mutations.

## 25 **Nucleobase Editors**

Disclosed herein, are novel base editors or nucleobase editors for editing, modifying or altering a target nucleotide sequence of a polynucleotide comprising a Cas9. Described herein is a nucleobase editor or a base editor comprising a polynucleotide programmable nucleotide binding domain (e.g., Cas9) and a nucleobase editing domain (e.g., adenosine  
30 deaminase). A polynucleotide programmable nucleotide binding domain (e.g., Cas9), when in conjunction with a bound guide polynucleotide (e.g., gRNA), can specifically bind to a

target polynucleotide sequence (*i.e.*, via complementary base pairing between bases of the bound guide nucleic acid and bases of the target polynucleotide sequence) and thereby localize the base editor to the target nucleic acid sequence desired to be edited. In some embodiments, the target polynucleotide sequence comprises single-stranded DNA or double-stranded DNA. In some embodiments, the target polynucleotide sequence comprises RNA. In some embodiments, the target polynucleotide sequence comprises a DNA-RNA hybrid. As most of the known genetic variations associated with human disease are point mutations, methods that can more efficiently and cleanly make precise point mutations are needed. Base editing systems as provided herein provide a new way to provide genome editing without generating double-strand DNA breaks, without requiring a donor DNA template, and without inducing an excess of stochastic insertions and deletions.

The base editors provided herein are capable of modifying a specific nucleotide base without generating a significant proportion of indels. The term “indel(s)”, as used herein, refers to the insertion or deletion of a nucleotide base within a nucleic acid. Such insertions or deletions can lead to frame shift mutations within a coding region of a gene. In some embodiments, it is desirable to generate base editors that efficiently modify (*e.g.*, mutate or deaminate) a specific nucleotide within a nucleic acid, without generating a large number of insertions or deletions (*i.e.*, indels) in the target nucleotide sequence. In certain embodiments, any of the base editors provided herein are capable of generating a greater proportion of intended modifications (*e.g.*, point mutations or deaminations) versus indels.

In some embodiments, any of base editor systems provided herein result in less than 50%, less than 40%, less than 30%, less than 20%, less than 19%, less than 18%, less than 17%, less than 16%, less than 15%, less than 14%, less than 13%, less than 12%, less than 11%, less than 10%, less than 9%, less than 8%, less than 7%, less than 6%, less than 5%, less than 4%, less than 3%, less than 2%, less than 1%, less than 0.9%, less than 0.8%, less than 0.7%, less than 0.6%, less than 0.5%, less than 0.4%, less than 0.3%, less than 0.2%, less than 0.1%, less than 0.09%, less than 0.08%, less than 0.07%, less than 0.06%, less than 0.05%, less than 0.04%, less than 0.03%, less than 0.02%, or less than 0.01% indel formation in the target polynucleotide sequence.

Some aspects of the disclosure are based on the recognition that any of the base editors provided herein are capable of efficiently generating an intended mutation, such as a point mutation, in a nucleic acid (*e.g.*, a nucleic acid within a genome of a subject) without generating a significant number of unintended mutations, such as unintended point mutations.

In some embodiments, any of the base editors provided herein are capable of generating at least 0.01% of intended mutations (*i.e.* at least 0.01% base editing efficiency). In some embodiments, any of the base editors provided herein are capable of generating at least 0.01%, 1%, 2%, 3%, 4%, 5%, 10%, 15%, 20%, 25%, 30%, 40%, 45%, 50%, 60%, 70%,  
5 80%, 90%, 95%, or 99% of intended mutations.

In some embodiments, the base editors provided herein are capable of generating a ratio of intended point mutations to indels that is greater than 1:1. In some embodiments, the base editors provided herein are capable of generating a ratio of intended point mutations to indels that is at least 1.5:1, at least 2:1, at least 2.5:1, at least 3:1, at least 3.5:1, at least 4:1, at  
10 least 4.5:1, at least 5:1, at least 5.5:1, at least 6:1, at least 6.5:1, at least 7:1, at least 7.5:1, at least 8:1, at least 8.5:1, at least 9:1, at least 10:1, at least 11:1, at least 12:1, at least 13:1, at least 14:1, at least 15:1, at least 20:1, at least 25:1, at least 30:1, at least 40:1, at least 50:1, at least 100:1, at least 200:1, at least 300:1, at least 400:1, at least 500:1, at least 600:1, at least 700:1, at least 800:1, at least 900:1, or at least 1000:1, or more.

15 The number of intended mutations and indels can be determined using any suitable method, for example, as described in International PCT Application Nos. PCT/2017/045381 (WO2018/027078) and PCT/US2016/058344 (WO2017/070632); Komor, A.C., *et al.*, “Programmable editing of a target base in genomic DNA without double-stranded DNA cleavage” *Nature* 533, 420-424 (2016); Gaudelli, N.M., *et al.*, “Programmable base editing of  
20 A•T to G•C in genomic DNA without DNA cleavage” *Nature* 551, 464-471 (2017); and Komor, A.C., *et al.*, “Improved base excision repair inhibition and bacteriophage Mu Gam protein yields C:G-to-T:A base editors with higher efficiency and product purity” *Science Advances* 3:eaa04774 (2017); the entire contents of which are hereby incorporated by reference.

25 In some embodiments, to calculate indel frequencies, sequencing reads are scanned for exact matches to two 10-bp sequences that flank both sides of a window in which indels can occur. If no exact matches are located, the read is excluded from analysis. If the length of this indel window exactly matches the reference sequence the read is classified as not containing an indel. If the indel window is two or more bases longer or shorter than the  
30 reference sequence, then the sequencing read is classified as an insertion or deletion, respectively. In some embodiments, the base editors provided herein can limit formation of indels in a region of a nucleic acid. In some embodiments, the region is at a nucleotide

targeted by a base editor or a region within 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides of a nucleotide targeted by a base editor.

The number of indels formed at a target nucleotide region can depend on the amount of time a nucleic acid (*e.g.*, a nucleic acid within the genome of a cell) is exposed to a base editor. In some embodiments, the number or proportion of indels is determined after at least 5 1 hour, at least 2 hours, at least 6 hours, at least 12 hours, at least 24 hours, at least 36 hours, at least 48 hours, at least 3 days, at least 4 days, at least 5 days, at least 7 days, at least 10 days, or at least 14 days of exposing the target nucleotide sequence (*e.g.*, a nucleic acid within the genome of a cell) to a base editor. It should be appreciated that the characteristics 10 of the base editors as described herein can be applied to any of the fusion proteins, or methods of using the fusion proteins provided herein.

### **Therapeutic Applications**

The CRISPR-Cas9 methods or systems described herein can have various therapeutic applications. Accordingly, in some embodiments, a method of treating a disorder or a disease 15 in a subject in need thereof is provided, the method comprising administering to the subject a CRISPR-Cas9 system comprising a Cas9 as described herein, wherein the guide RNA is complementary to at least 10 nucleotides of a target nucleic acid associated with the condition or disease; wherein the Cas protein associates with the guide RNA; wherein the guide RNA binds to the target nucleic acid; wherein the Cas protein causes a break in the target nucleic 20 acid, optionally wherein the Cas9 is an inactive Cas9 (dCas9) fused to a deaminase and results in one or more base edits in the target nucleic acid, thereby treating the disorder or disease.

In some embodiments, the CRISPR-Cas9 methods or systems can be used to treat various diseases and disorders, *e.g.*, genetic disorders (*e.g.*, monogenetic diseases), diseases 25 that can be treated by nuclease activity, and various cancers, etc.

In some embodiments, the CRISPR methods or systems described herein can be used to edit a target nucleic acid to modify the target nucleic acid (*e.g.*, by inserting, deleting, or mutating one or more nucleic acid residues). For example, in some embodiments the CRISPR systems described herein comprise an exogenous donor template nucleic acid (*e.g.*, a DNA 30 molecule or a RNA molecule), which comprises a desirable nucleic acid sequence. Upon resolution of a cleavage event induced with the CRISPR system described herein, the molecular machinery of the cell will utilize the exogenous donor template nucleic acid in

repairing and/or resolving the cleavage event. Alternatively, the molecular machinery of the cell can utilize an endogenous template in repairing and/or resolving the cleavage event. In some embodiments, the CRISPR systems described herein may be used to alter a target nucleic acid resulting in an insertion, a deletion, and/or a point mutation). In some  
5 embodiments, the insertion is a scarless insertion (i.e., the insertion of an intended nucleic acid sequence into a target nucleic acid resulting in no additional unintended nucleic acid sequence upon resolution of the cleavage event). Donor template nucleic acids may be double stranded or single stranded nucleic acid molecules (e.g., DNA or RNA). In some  
10 embodiments, the CRISPR methods or systems described herein comprise a nucleobase editor. For example, in some embodiments, the Cas9 proteins described herein are fused to a polypeptide having nucleobase editing activity.

In one aspect, the CRISPR methods or systems described herein can be used for treating a disease caused by overexpression of RNAs, toxic RNAs, and/or mutated RNAs (e.g., splicing defects or truncations).

15 In some embodiments, the CRISPR methods or systems described herein can also target trans-acting mutations affecting RNA- dependent functions that cause various diseases.

In some embodiments, the CRISPR methods or systems described herein can also be used to target mutations disrupting the cis-acting splicing codes that can cause splicing defects and diseases.

20 The CRISPR methods or systems described herein can further be used for antiviral activity, in particular against RNA viruses. The CRISPR-associated proteins can target the viral RNAs using suitable RNA guides selected to target viral RNA sequences.

The CRISPR methods or systems described herein can also be used to treat a cancer in a subject (e.g., a human subject). For example, the CRISPR-associated proteins described  
25 herein can be programmed with crRNA targeting a RNA molecule that is aberrant (e.g., comprises a point mutation or are alternatively-spliced) and found in cancer cells to induce cell death in the cancer cells (e.g., via apoptosis).

Further, the CRISPR methods or systems described herein can also be used to treat an infectious disease in a subject. For example, the CRISPR-associated proteins described herein  
30 can be programmed with crRNA targeting a RNA molecule expressed by an infectious agent (e.g., a bacteria, a virus, a parasite or a protozoan) in order to target and induce cell death in the infectious agent cell. The CRISPR systems may also be used to treat diseases where an



intracellular infectious agent infects the cells of a host subject. By programming the CRISPR-associated protein to target a RNA molecule encoded by an infectious agent gene, cells infected with the infectious agent can be targeted and cell death induced.

Furthermore, in vitro RNA sensing assays can be used to detect specific RNA  
5 substrates. The CRISPR-associated proteins can be used for RNA-based sensing in living cells. Examples of applications are diagnostics by sensing of, for examples, disease-specific RNAs.

In applications in which it is desirable to insert a polynucleotide sequence into a target DNA sequence, a polynucleotide comprising a donor sequence to be inserted is also  
10 provided to the cell. By a "donor sequence" or "donor polynucleotide" it is meant a nucleic acid sequence to be inserted at the cleavage site induced by a site-directed modifying polypeptide. The donor polynucleotide will contain sufficient homology to a genomic sequence at the cleavage site, e.g. 70%, 80%, 85%, 90%, 95%, or 100% homology with the nucleotide sequences flanking the cleavage site, e.g. within about 50 bases or less of the  
15 cleavage site, e.g. within about 30 bases, within about 15 bases, within about 10 bases, within about 5 bases, or immediately flanking the cleavage site, to support homology-directed repair between it and the genomic sequence to which it bears homology. Approximately 25, 50, 100, or 200 nucleotides, or more than 200 nucleotides, of sequence homology between a donor and a genomic sequence (or any integral value between 10 and 200 nucleotides, or  
20 more) will support homology-directed repair. Donor sequences can be of any length, e.g. 10 nucleotides or more, 50 nucleotides or more, 100 nucleotides or more, 250 nucleotides or more, 500 nucleotides or more, 1000 nucleotides or more, 5000 nucleotides or more, etc.

The donor sequence is typically not identical to the genomic sequence that it replaces. Rather, the donor sequence may contain at least one or more single base changes, insertions,  
25 deletions, inversions or rearrangements with respect to the genomic sequence, so long as sufficient homology is present to support homology-directed repair. In some embodiments, the donor sequence comprises a non-homologous sequence flanked by two regions of homology, such that homology-directed repair between the target DNA region and the two flanking sequences results in insertion of the non-homologous sequence at the target region.  
30 Donor sequences may also comprise a vector backbone containing sequences that are not homologous to the DNA region of interest and that are not intended for insertion into the DNA region of interest. Generally, the homologous region(s) of a donor sequence will have at least 50% sequence identity to a genomic sequence with which recombination is desired. In

certain embodiments, 60%, 70%, 80%, 90%, 95%, 98%, 99%, or 99.9% sequence identity is present. Any value between 1% and 100% sequence identity can be present, depending upon the length of the donor polynucleotide.

5 The donor sequence may comprise certain sequence differences as compared to the genomic sequence, e.g. restriction sites, nucleotide polymorphisms, selectable markers (e.g., drug resistance genes, fluorescent proteins, enzymes etc.), etc., which may be used to assess for successful insertion of the donor sequence at the cleavage site or in some cases may be used for other purposes (e.g., to signify expression at the targeted genomic locus). In some cases, if located in a coding region, such nucleotide sequence differences will not change the amino acid sequence, or will make silent amino acid changes (i.e., changes which do not  
10 affect the structure or function of the protein). Alternatively, these sequences differences may include flanking recombination sequences such as FLPs, loxP sequences, or the like, that can be activated at a later time for removal of the marker sequence.

The donor sequence may be provided to the cell as single-stranded DNA, single-  
15 stranded RNA, double-stranded DNA, or double-stranded RNA. It may be introduced into a cell in linear or circular form. If introduced in linear form, the ends of the donor sequence may be protected (e.g., from exonucleolytic degradation) by methods known to those of skill in the art. For example, one or more dideoxynucleotide residues are added to the 3' terminus of a linear molecule and/or self-complementary oligonucleotides are ligated to one or both  
20 ends. Additional methods for protecting exogenous polynucleotides from degradation include, but are not limited to, addition of terminal amino group(s) and the use of modified internucleotide linkages such as, for example, phosphorothioates, phosphor amidates, and O-methyl ribose or deoxyribose residues. As an alternative to protecting the termini of a linear donor sequence, additional lengths of sequence may be included outside of the regions of  
25 homology that can be degraded without impacting recombination. A donor sequence can be introduced into a cell as part of a vector molecule having additional sequences such as, for example, replication origins, promoters and genes encoding antibiotic resistance. Moreover, donor sequences can be introduced as naked nucleic acid, as nucleic acid complexed with an agent such as a liposome or poloxamer, or can be delivered by viruses (e.g., adenovirus,  
30 AAV), as described above for nucleic acids encoding a DNA -targeting RNA and/or site -directed modifying polypeptide and/or donor polynucleotide.

Following the methods described above, a DNA region of interest may be cleaved and modified, i.e. "genetically modified", ex vivo. In some embodiments, as when a selectable

marker has been inserted into the DNA region of interest, the population of cells may be enriched for those comprising the genetic modification by separating the genetically modified cells from the remaining population. Prior to enriching, the "genetically modified" cells may make up only about 1% or more (e.g., 2% or more, 3% or more, 4% or more, 5% or more, 5  
6% or more, 7% or more, 8% or more, 9% or more, 10% or more, 15% or more, or 20% or more) of the cellular population. Separation of "genetically modified" cells may be achieved by any convenient separation technique appropriate for the selectable marker used. For example, if a fluorescent marker has been inserted, cells may be separated by fluorescence activated cell sorting, whereas if a cell surface marker has been inserted, cells may be  
10 separated from the heterogeneous population by affinity separation techniques, e.g. magnetic separation, affinity chromatography, "panning" with an affinity reagent attached to a solid matrix, or other convenient technique. Techniques providing accurate separation include fluorescence activated cell sorters, which can have varying degrees of sophistication, such as multiple color channels, low angle and obtuse light scattering detecting channels, impedance  
15 channels, etc. The cells may be selected against dead cells by employing dyes associated with dead cells (e.g. propidium iodide). Any technique may be employed which is not unduly detrimental to the viability of the genetically modified cells. Cell compositions that are highly enriched for cells comprising modified DNA are achieved in this manner. By "highly enriched", it is meant that the genetically modified cells will be 70% or more, 75% or more,  
20 80% or more, 85% or more, 90% or more of the cell composition, for example, about 95% or more, or 98% or more of the cell composition. In other words, the composition may be a substantially pure composition of genetically modified cells.

Genetically modified cells produced by the methods described herein may be used immediately. Alternatively, the cells may be frozen at liquid nitrogen temperatures and stored  
25 for long periods of time, being thawed and capable of being reused. In such cases, the cells will usually be frozen in 10% dimethylsulfoxide (DMSO), 50% serum, 40% buffered medium, or some other such solution as is commonly used in the art to preserve cells at such freezing temperatures, and thawed in a manner as commonly known in the art for thawing frozen cultured cells.

30 The genetically modified cells may be cultured *in vitro* under various culture conditions. The cells may be expanded in culture, i.e. grown under conditions that promote their proliferation. Culture medium may be liquid or semi-solid, e.g. containing agar, methylcellulose, etc. The cell population may be suspended in an appropriate nutrient

medium, such as Iscove's modified DMEM or RPMI 1640, normally supplemented with fetal calf serum (about 5-10%),

L-glutamine, a thiol, particularly 2-mercaptoethanol, and antibiotics, e.g. penicillin and streptomycin. The culture may contain growth factors to which the regulatory T cells are responsive. Growth factors, as defined herein, are molecules capable of promoting survival, growth and/or differentiation of cells, either in culture or in the intact tissue, through specific effects on a transmembrane receptor. Growth factors include polypeptides and non-polypeptide factors.

Cells that have been genetically modified in this way may be transplanted to a subject for purposes such as gene therapy, e.g. to treat a disease or as an antiviral, antipathogenic, or anticancer therapeutic, for the production of genetically modified organisms in agriculture, or for biological research. The subject may be a neonate, a juvenile, or an adult. Of particular interest are mammalian subjects. Mammalian species that may be treated with the present methods include canines and felines; equines; bovines; ovines; etc. and primates, particularly humans. Animal models, particularly small mammals (e.g. mouse, rat, guinea pig, hamster, lagomorpha (e.g., rabbit), etc.) may be used for experimental investigations.

Cells may be provided to the subject alone or with a suitable substrate or matrix, e.g. to support their growth and/or organization in the tissue to which they are being transplanted. Usually, at least  $1 \times 10^3$  cells will be administered, for example  $5 \times 10^3$  cells,  $1 \times 10^4$  cells,  $5 \times 10^4$  cells,  $1 \times 10^5$  cells,  $1 \times 10^6$  cells or more. The cells may be introduced to the subject via any of the following routes: parenteral, subcutaneous, intravenous, intracranial, intraspinal, intraocular, or into spinal fluid. The cells may be introduced by injection, catheter, or the like. Cells may also be introduced into an embryo (e.g., a blastocyst) for the purpose of generating a transgenic animal (e.g., a transgenic mouse).

The number of administrations of treatment to a subject may vary. Introducing the genetically modified cells into the subject may be a one-time event; but in certain situations, such treatment may elicit improvement for a limited period of time and require an on-going series of repeated treatments. In other situations, multiple administrations of the genetically modified cells may be required before an effect is observed. The exact protocols depend upon the disease or condition, the stage of the disease and parameters of the individual subject being treated.

In other aspects of the invention, the DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotide are employed to modify cellular DNA in vivo, again for purposes such as gene therapy, e.g. to treat a disease or as an antiviral, antipathogenic, or anticancer therapeutic, for the production of genetically modified organisms in agriculture, or for biological research. In these in vivo embodiments, a DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotide are administered directly to the individual. A DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotide may be administered by any of a number of well-known methods in the art for the administration of peptides, small molecules and nucleic acids to a subject. A DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotide can be incorporated into a variety of formulations. More particularly, a DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotide of the present invention can be formulated into pharmaceutical compositions by combination with appropriate pharmaceutically acceptable carriers or diluents.

Pharmaceutical preparations are compositions that include one or more a DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotide present in a pharmaceutically acceptable vehicle. "Pharmaceutically acceptable vehicles" may be vehicles approved by a regulatory agency of the Federal or a state government or listed in the U.S.

Pharmacopeia or other generally recognized pharmacopeia for use in mammals, such as humans. The term "vehicle" refers to a diluent, adjuvant, excipient, or carrier with which a compound of the invention is formulated for administration to a mammal. Such pharmaceutical vehicles can be lipids, e.g. liposomes, e.g. liposome dendrimers; liquids, such as water and oils, including those of petroleum, animal, vegetable or synthetic origin, such as peanut oil, soybean oil, mineral oil, sesame oil and the like, saline; gum acacia, gelatin, starch paste, talc, keratin, colloidal silica, urea, and the like. In addition, auxiliary, stabilizing, thickening, lubricating and coloring agents may be used. Pharmaceutical compositions may be formulated into preparations in solid, semisolid, liquid or gaseous forms, such as tablets, capsules, powders, granules, ointments, solutions, suppositories, injections, inhalants, gels, microspheres, and aerosols. As such, administration of the a DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotide can be achieved in various ways, including oral, buccal, rectal, parenteral, intraperitoneal, intradermal, transdermal, intratracheal, intraocular, etc., administration. The active agent may be systemic after

administration or may be localized by the use of regional administration, intramural administration, or use of an implant that acts to retain the active dose at the site of implantation. The active agent may be formulated for immediate activity or it may be formulated for sustained release.

5           For some conditions, particularly central nervous system conditions, it may be necessary to formulate agents to cross the blood-brain barrier (BBB). One strategy for drug delivery through the blood-brain barrier (BBB) entails disruption of the BBB, either by osmotic means such as mannitol or leukotrienes, or biochemically by the use of vasoactive substances such as bradykinin. The potential for using BBB opening to target specific agents  
10 to brain tumors is also an option. A BBB disrupting agent can be co-administered with the therapeutic compositions of the invention when the compositions are administered by intravascular injection. Other strategies to go through the BBB may entail the use of endogenous transport systems, including Caveolin-1 mediated transcytosis, carrier-mediated transporters such as glucose and amino acid carriers, receptor-mediated transcytosis for  
15 insulin or transferrin, and active efflux transporters such as p- glycoprotein. Active transport moieties may also be conjugated to the therapeutic compounds for use in the invention to facilitate transport across the endothelial wall of the blood vessel.

          Alternatively, drug delivery of therapeutics agents behind the BBB may be by local delivery, for example by intrathecal delivery.

20           Typically, an effective amount of a DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotide are provided. As discussed above with regard to ex vivo methods, an effective amount or effective dose of a DNA-targeting RNA and/or site- directed modifying polypeptide and/or donor polynucleotide in vivo is the amount to induce a 2 fold increase or more in the amount of recombination observed between  
25 two homologous sequences relative to a negative control, e.g. a cell contacted with an empty vector or irrelevant polypeptide. The amount of recombination may be measured by any convenient method, e.g. as described above and known in the art. The calculation of the effective amount or effective dose of a DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotide to be administered is within the skill of one of  
30 ordinary skill in the art, and will be routine to those persons skilled in the art. The final amount to be administered will be dependent upon the route of administration and upon the nature of the disorder or condition that is to be treated.

The effective amount given to a particular patient will depend on a variety of factors, several of which will differ from patient to patient. A competent clinician will be able to determine an effective amount of a therapeutic agent to administer to a patient to halt or reverse the progression the disease condition as required. Utilizing LD50 animal data, and  
5 other information available for the agent, a clinician can determine the maximum safe dose for an individual, depending on the route of administration. For instance, an intravenously administered dose may be more than an intrathecally administered dose, given the greater body of fluid into which the therapeutic composition is being administered. Similarly, compositions which are rapidly cleared from the body may be administered at higher doses,  
10 or in repeated doses, in order to maintain a therapeutic concentration. Utilizing ordinary skill, the competent clinician will be able to optimize the dosage of a particular therapeutic in the course of routine clinical trials.

For inclusion in a medicament, a DNA-targeting RNA and/or site -directed modifying polypeptide and/or donor polynucleotide may be obtained from a suitable commercial source.  
15 As a general proposition, the total pharmaceutically effective amount of the a DNA-targeting RNA and/or site -directed modifying polypeptide and/or donor polynucleotide administered parenterally per dose will be in a range that can be measured by a dose response curve.

Therapies based on a DNA-targeting RNA and/or site-directed modifying polypeptide and/or donor polynucleotides, i.e. preparations of a DNA-targeting RNA and/or site-directed  
20 modifying polypeptide and/or donor polynucleotide to be used for therapeutic administration, must be sterile. Sterility is readily accomplished by filtration through sterile filtration membranes (e.g., 0.2  $\mu\text{m}$  membranes). Therapeutic compositions generally are placed into a container having a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle. The therapies based on a DNA-  
25 targeting RNA and/or site- directed modifying polypeptide and/or donor polynucleotide may be stored in unit or multi-dose containers, for example, sealed ampules or vials, as an aqueous solution or as a lyophilized formulation for reconstitution. As an example of a lyophilized formulation, 10-mL vials are filled with 5 ml of sterile-filtered 1 % (w/v) aqueous solution of compound, and the resulting mixture is lyophilized. The infusion solution is prepared by  
30 reconstituting the lyophilized compound using bacteriostatic Water-for-Injection.

Pharmaceutical compositions can include, depending on the formulation desired, pharmaceutically-acceptable, non-toxic carriers of diluents, which are defined as vehicles commonly used to formulate pharmaceutical compositions for animal or human

administration. The diluent is selected so as not to affect the biological activity of the combination. Examples of such diluents are distilled water, buffered water, physiological saline, PBS, Ringer's solution, dextrose solution, and Hank's solution. In addition, the pharmaceutical composition or formulation can include other carriers, adjuvants, or non-  
5 toxic, nontherapeutic, nonimmunogenic stabilizers, excipients and the like. The compositions can also include additional substances to approximate physiological conditions, such as pH adjusting and buffering agents, toxicity adjusting agents, wetting agents and detergents.

The composition can also include any of a variety of stabilizing agents, such as an antioxidant for example. When the pharmaceutical composition includes a polypeptide, the  
10 polypeptide can be complexed with various well-known compounds that enhance the *in vivo* stability of the polypeptide, or otherwise enhance its pharmacological properties (e.g., increase the half-life of the polypeptide, reduce its toxicity, and enhance solubility or uptake). Examples of such modifications or complexing agents include sulfate, gluconate, citrate and phosphate. The nucleic acids or polypeptides of a composition can also be complexed with  
15 molecules that enhance their *in vivo* attributes. Such molecules include, for example, carbohydrates, polyamines, amino acids, other peptides, ions (e.g., sodium, potassium, calcium, magnesium, manganese), and lipids.

The pharmaceutical compositions can be administered for prophylactic and/or therapeutic treatments. Toxicity and therapeutic efficacy of the active ingredient can be  
20 determined according to standard pharmaceutical procedures in cell cultures and/or experimental animals, including, for example, determining the LD50 (the dose lethal to 50% of the population) and the ED50 (the dose therapeutically effective in 50% of the population). The dose ratio between toxic and therapeutic effects is the therapeutic index and it can be expressed as the ratio LD50/ED50. Therapies that exhibit large therapeutic indices are  
25 preferred.

The data obtained from cell culture and/or animal studies can be used in formulating a range of dosages for humans. The dosage of the active ingredient typically lies within a range of circulating concentrations that include the ED50 with low toxicity. The dosage can vary within this range depending upon the dosage form employed and the route of  
30 administration utilized.

The components used to formulate the pharmaceutical compositions are preferably of high purity and are substantially free of potentially harmful contaminants (e.g., at least



National Food (NF) grade, generally at least analytical grade, and more typically at least pharmaceutical grade). Moreover, compositions intended for *in vivo* use are usually sterile. To the extent that a given compound must be synthesized prior to use, the resulting product is typically substantially free of any potentially toxic agents, particularly any endotoxins, which  
5 may be present during the synthesis or purification process. Compositions for parental administration are also sterile, substantially isotonic and made under GMP conditions.

### **Delivery Systems**

The CRISPR systems described herein, or components thereof, nucleic acid molecules thereof, and/or nucleic acid molecules encoding or providing components thereof, CRISPR-  
10 associated proteins, or RNA guides, can be delivered by various delivery systems such as vectors, e.g., plasmids and delivery vectors. Exemplary embodiments are described below. The CRISPR systems (e.g., including the Cas9 comprising nucleobase editor described herein) can be encoded on a nucleic acid that is contained in a viral vector. Viral vectors can include lentivirus, Adenovirus, Retrovirus, and Adeno-associated viruses (AAVs). Viral  
15 vectors can be selected based on the application. For example, AAVs are commonly used for gene delivery *in vivo* due to their mild immunogenicity. Adenoviruses are commonly used as vaccines because of the strong immunogenic response they induce. Packaging capacity of the viral vectors can limit the size of the base editor that can be packaged into the vector. For example, the packaging capacity of the AAVs is ~4.5 kb including two 145 base inverted  
20 terminal repeats (ITRs).

AAV is a small, single-stranded DNA dependent virus belonging to the parvovirus family. The 4.7 kb wild-type (wt) AAV genome is made up of two genes that encode four replication proteins and three capsid proteins, respectively, and is flanked on either side by 145-bp inverted terminal repeats (ITRs). The virion is composed of three capsid proteins,  
25 Vp1, Vp2, and Vp3, produced in a 1:1:10 ratio from the same open reading frame but from differential splicing (Vp1) and alternative translational start sites (Vp2 and Vp3, respectively). Vp3 is the most abundant subunit in the virion and participates in receptor recognition at the cell surface defining the tropism of the virus. A phospholipase domain, which functions in viral infectivity, has been identified in the unique N terminus of Vp1.

30 Similar to wt AAV, recombinant AAV (rAAV) utilizes the *cis*-acting 145-bp ITRs to flank vector transgene cassettes, providing up to 4.5 kb for packaging of foreign DNA. Subsequent to infection, rAAV can express a fusion protein of the invention and persist

without integration into the host genome by existing episomally in circular head-to-tail concatemers. Although there are numerous examples of rAAV success using this system, *in vitro* and *in vivo*, the limited packaging capacity has limited the use of AAV-mediated gene delivery when the length of the coding sequence of the gene is equal or greater in size than  
5 the wt AAV genome.

The small packaging capacity of AAV vectors makes the delivery of a number of genes that exceed this size and/or the use of large physiological regulatory elements challenging. These challenges can be addressed, for example, by dividing the protein(s) to be delivered into two or more fragments, wherein the N-terminal fragment is fused to a split  
10 intein-N and the C-terminal fragment is fused to a split intein-C. These fragments are then packaged into two or more AAV vectors. As used herein, "intein" refers to a self-splicing protein intron (*e.g.*, peptide) that ligates flanking N-terminal and C-terminal exteins (*e.g.*, fragments to be joined). The use of certain inteins for joining heterologous protein fragments is described, for example, in Wood *et al.*, J. Biol. Chem. 289(21); 14512-9 (2014). For  
15 example, when fused to separate protein fragments, the inteins IntN and IntC recognize each other, splice themselves out and simultaneously ligate the flanking N- and C-terminal exteins of the protein fragments to which they were fused, thereby reconstituting a full-length protein from the two protein fragments. Other suitable inteins will be apparent to a person of skill in the art.

20 In some embodiments, the CRISPR system of the invention can vary in length. In some embodiments, a protein fragment ranges from 2 amino acids to about 1000 amino acids in length. In some embodiments, a protein fragment ranges from about 5 amino acids to about 500 amino acids in length. In some embodiments, a protein fragment ranges from about 20 amino acids to about 200 amino acids in length. In some embodiments, a protein fragment  
25 ranges from about 10 amino acids to about 100 amino acids in length. Suitable protein fragments of other lengths will be apparent to a person of skill in the art.

In some embodiments, a portion or fragment of a nuclease (*e.g.*, Cas9) is fused to an intein. The nuclease can be fused to the N-terminus or the C-terminus of the intein. In some  
30 embodiments, a portion or fragment of a fusion protein is fused to an intein and fused to an AAV capsid protein. The intein, nuclease and capsid protein can be fused together in any arrangement (*e.g.*, nuclease-intein-capsid, intein-nuclease-capsid, capsid-intein-nuclease, etc.). In some embodiments, the N-terminus of an intein is fused to the C-terminus of a fusion protein and the C-terminus of the intein is fused to the N-terminus of an AAV capsid protein.

In one embodiment, dual AAV vectors are generated by splitting a large transgene expression cassette in two separate halves (5' and 3' ends, or head and tail), where each half of the cassette is packaged in a single AAV vector (of <5 kb). The re-assembly of the full-length transgene expression cassette is then achieved upon co-infection of the same cell by  
5 both dual AAV vectors followed by: (1) homologous recombination (HR) between 5' and 3' genomes (dual AAV overlapping vectors); (2) ITR-mediated tail-to-head concatemerization of 5' and 3' genomes (dual AAV *trans*-splicing vectors); or (3) a combination of these two mechanisms (dual AAV hybrid vectors). The use of dual AAV vectors *in vivo* results in the expression of full-length proteins. The use of the dual AAV vector platform represents an  
10 efficient and viable gene transfer strategy for transgenes of >4.7 kb in size.

The disclosed strategies for designing CRISPR systems including the Cas9 described herein can be useful for generating CRISPR systems capable of being packaged into a viral vector. The use of RNA or DNA viral based systems for the delivery of a base editor takes advantage of highly evolved processes for targeting a virus to specific cells in culture or in  
15 the host and trafficking the viral payload to the nucleus or host cell genome. Viral vectors can be administered directly to cells in culture, patients (*in vivo*), or they can be used to treat cells *in vitro*, and the modified cells can optionally be administered to patients (*ex vivo*). Conventional viral based systems could include retroviral, lentivirus, adenoviral, adeno-associated and herpes simplex virus vectors for gene transfer. Integration in the host genome  
20 is possible with the retrovirus, lentivirus, and adeno-associated virus gene transfer methods, often resulting in long term expression of the inserted transgene. Additionally, high transduction efficiencies have been observed in many different cell types and target tissues.

The tropism of a retrovirus can be altered by incorporating foreign envelope proteins, expanding the potential target population of target cells. Lentiviral vectors are retroviral  
25 vectors that are able to transduce or infect non-dividing cells and typically produce high viral titers. Selection of a retroviral gene transfer system would therefore depend on the target tissue. Retroviral vectors are comprised of cis-acting long terminal repeats with packaging capacity for up to 6-10 kb of foreign sequence. The minimum cis-acting LTRs are sufficient for replication and packaging of the vectors, which are then used to integrate the therapeutic  
30 gene into the target cell to provide permanent transgene expression. Widely used retroviral vectors include those based upon murine leukemia virus (MuLV), gibbon ape leukemia virus (GaLV), Simian Immuno deficiency virus (SIV), human immuno deficiency virus (HIV), and combinations thereof (*See, e.g., Buchscher et al., J. Virol. 66:2731-2739 (1992); Johann et*

*al.*, J. Virol. 66:1635-1640 (1992); Somnerfelt *et al.*, Virol. 176:58-59 (1990); Wilson *et al.*, J. Virol. 63:2374-2378 (1989); Miller *et al.*, J. Virol. 65:2220-2224 (1991); PCT/US94/05700).

5 Retroviral vectors, especially lentiviral vectors, can require polynucleotide sequences smaller than a given length for efficient integration into a target cell. For example, retroviral vectors of length greater than 9 kb can result in low viral titers compared with those of smaller size. In some aspects, a CRISPR system (e.g., including the Cas9 disclosed herein) of the present disclosure is of sufficient size so as to enable efficient packaging and delivery into a target cell via a retroviral vector. In some cases, a Cas9 is of a size so as to allow  
10 efficient packing and delivery even when expressed together with a guide nucleic acid and/or other components of a targetable nuclease system.

In applications where transient expression is preferred, adenoviral based systems can be used. Adenoviral based vectors are capable of very high transduction efficiency in many cell types and do not require cell division. With such vectors, high titer and levels of  
15 expression have been obtained. This vector can be produced in large quantities in a relatively simple system. Adeno-associated virus (“AAV”) vectors can also be used to transduce cells with target nucleic acids, e.g., in the *in vitro* production of nucleic acids and peptides, and for *in vivo* and *ex vivo* gene therapy procedures (*See, e.g.*, West *et al.*, Virology 160:38-47 (1987); U.S. Patent No. 4,797,368; WO 93/24641; Kotin, Human Gene Therapy 5:793-801  
20 (1994); Muzyczka, J. Clin. Invest. 94:1351 (1994). The construction of recombinant AAV vectors is described in a number of publications, including U.S. Patent No. 5,173,414; Tratschin *et al.*, Mol. Cell. Biol. 5:3251-3260 (1985); Tratschin, *et al.*, Mol. Cell. Biol. 4:2072-2081 (1984); Hermonat & Muzyczka, PNAS 81:6466-6470 (1984); and Samulski *et al.*, J. Virol. 63:03822-3828 (1989).

25 A CRISPR system (e.g., including the Cas9 disclosed herein) described herein can therefore be delivered with viral vectors. One or more components of the base editor system can be encoded on one or more viral vectors. For example, a base editor and guide nucleic acid can be encoded on a single viral vector. In other cases, the base editor and guide nucleic acid are encoded on different viral vectors. In either case, the base editor and guide nucleic  
30 acid can each be operably linked to a promoter and terminator.

The combination of components encoded on a viral vector can be determined by the cargo size constraints of the chosen viral vector.

***Non-Viral Delivery of Base Editors***

Non-viral delivery approaches for CRISPR are also available. One important category of non-viral nucleic acid vectors are nanoparticles, which can be organic or inorganic. Nanoparticles are well known in the art. Any suitable nanoparticle design can be used to deliver genome editing system components or nucleic acids encoding such components. For instance, organic (*e.g.* lipid and/or polymer) nanoparticles can be suitable for use as delivery vehicles in certain embodiments of this disclosure. Exemplary lipids for use in nanoparticle formulations, and/or gene transfer are shown in Table 5 (below).

**Table 5**

Lipids Used for Gene Transfer		
Lipid	Abbreviation	Feature
1,2-Dioleoyl-sn-glycero-3-phosphatidylcholine	DOPC	Helper
1,2-Dioleoyl-sn-glycero-3-phosphatidylethanolamine	DOPE	Helper
Cholesterol		Helper
N-[1-(2,3-Dioleoyloxy)propyl]N,N,N-trimethylammonium chloride	DOTMA	Cationic
1,2-Dioleoyloxy-3-trimethylammonium-propane	DOTAP	Cationic
Diocetadecylamidoglycylspermine	DOGS	Cationic
N-(3-Aminopropyl)-N,N-dimethyl-2,3-bis(dodecyloxy)-1-propanaminium bromide	GAP-DLRIE	Cationic
Cetyltrimethylammonium bromide	CTAB	Cationic
6-Lauroxyhexyl ornithinate	LHON	Cationic
1-(2,3-Dioleoyloxypropyl)-2,4,6-trimethylpyridinium	2Oc	Cationic
2,3-Dioleoyloxy-N-[2(sperminecarboxamido-ethyl)-N,N-dimethyl-1-propanaminium trifluoroacetate	DOSPA	Cationic
1,2-Dioleoyl-3-trimethylammonium-propane	DOPA	Cationic
N-(2-Hydroxyethyl)-N,N-dimethyl-2,3-bis(tetradecyloxy)-1-propanaminium bromide	MDRIE	Cationic
Dimyristooxypropyl dimethyl hydroxyethyl ammonium bromide	DMRI	Cationic
3 $\beta$ -[N-(N',N'-Dimethylaminoethane)-carbamoyl]cholesterol	DC-Chol	Cationic
Bis-guanidium-tren-cholesterol	BGTC	Cationic

Lipids Used for Gene Transfer		
Lipid	Abbreviation	Feature
1,3-Diideoxy-2-(6-carboxy-spermyl)-propylamide	DOSPER	Cationic
Dimethyloctadecylammonium bromide	DDAB	Cationic
Diocadecylamidoglycylspermidin	DSL	Cationic
rac-[(2,3-Dioctadecyloxypropyl)(2-hydroxyethyl)]-dimethylammonium chloride	CLIP-1	Cationic
rac-[2(2,3-Dihexadecyloxypropyl-oxymethyloxy)ethyl]trimethylammonium bromide	CLIP-6	Cationic
Ethyl dimyristoyl phosphatidylcholine	EDMPC	Cationic
1,2-Distearoyloxy-N,N-dimethyl-3-aminopropane	DSDMA	Cationic
1,2-Dimyristoyl-trimethylammonium propane	DMTAP	Cationic
O,O'-Dimyristyl-N-lysyl aspartate	DMKE	Cationic
1,2-Distearoyl-sn-glycero-3-ethylphosphocholine	DSEPC	Cationic
N-Palmitoyl D-erythro-sphingosyl carbamoyl-spermine	CCS	Cationic
N-t-Butyl-N0-tetradecyl-3-tetradecylaminopropionamide	diC14-amidine	Cationic
Octadecenolyoxy[ethyl-2-heptadecenyl-3 hydroxyethyl]imidazolium chloride	DOTIM	Cationic
N1 -Cholesteryloxy carbonyl-3,7-diazanonane-1,9-diamine	CDAN	Cationic
2-(3-[Bis(3-amino-propyl)-amino]propylamino)-N-ditetradecylcarbamoylme-ethyl-acetamide	RPR209120	Cationic
1,2-dilinoleoyloxy-3-dimethylaminopropane	DLinDMA	Cationic
2,2-dilinoleyl-4-dimethylaminoethyl-[1,3]-dioxolane	DLin-KC2-DMA	Cationic
dilinoleyl-methyl-4-dimethylaminobutyrate	DLin-MC3-DMA	Cationic

Table 6 lists exemplary polymers for use in gene transfer and/or nanoparticle formulations.

**Table 6**

Polymers Used for Gene Transfer	
Polymer	Abbreviation
Poly(ethylene)glycol	PEG
Polyethylenimine	PEI
Dithiobis (succinimidylpropionate)	DSP
Dimethyl-3,3'-dithiobispropionimidate	DTBP
Poly(ethylene imine)biscarbamate	PEIC
Poly(L-lysine)	PLL
Histidine modified PLL	
Poly(N-vinylpyrrolidone)	PVP
Poly(propylenimine)	PPI
Poly(amidoamine)	PAMAM
Poly(amidoethylenimine)	SS-PAEI
Triethylenetetramine	TETA
Poly( $\beta$ -aminoester)	
Poly(4-hydroxy-L-proline ester)	PHP
Poly(allylamine)	
Poly( $\alpha$ -[4-aminobutyl]-L-glycolic acid)	PAGA
Poly(D,L-lactic-co-glycolic acid)	PLGA
Poly(N-ethyl-4-vinylpyridinium bromide)	
Poly(phosphazene)s	PPZ
Poly(phosphoester)s	PPE
Poly(phosphoramidate)s	PPA
Poly(N-2-hydroxypropylmethacrylamide)	pHPMA
Poly (2-(dimethylamino)ethyl methacrylate)	pDMAEMA
Poly(2-aminoethyl propylene phosphate)	PPE-EA
Chitosan	
Galactosylated chitosan	
N-Dodacylated chitosan	
Histone	
Collagen	

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 Polymers Used for Gene Transfer
 

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Polymer	Abbreviation
Dextran-spermine	D-SPM

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Table 7 summarizes delivery methods for a polynucleotide encoding a Cas9 described herein.

**Table 7**

Delivery	Vector/Mode	Delivery into Non-Dividing Cells	Duration of Expression	Genome Integration	Type of Molecule Delivered
Physical	( <i>e.g.</i> , electroporation, particle gun, Calcium Phosphate transfection)	YES	Transient	NO	Nucleic Acids and Proteins
Viral	Retrovirus	NO	Stable	YES	RNA
	Lentivirus	YES	Stable	YES/NO with modification	RNA
	Adenovirus	YES	Transient	NO	DNA
	Adeno-Associated Virus (AAV)	YES	Stable	NO	DNA
	Vaccinia Virus	YES	Very Transient	NO	DNA
	Herpes Simplex Virus	YES	Stable	NO	DNA
Non-Viral	Cationic Liposomes	YES	Transient	Depends on what is delivered	Nucleic Acids and Proteins
	Polymeric Nanoparticles	YES	Transient	Depends on what is delivered	Nucleic Acids and Proteins
Biological Non-Viral	Attenuated Bacteria	YES	Transient	NO	Nucleic Acids



Delivery	Vector/Mode	Delivery into Non-Dividing Cells	Duration of Expression	Genome Integration	Type of Molecule Delivered
Delivery Vehicles	Engineered Bacteriophages	YES	Transient	NO	Nucleic Acids
	Mammalian Virus-like Particles	YES	Transient	NO	Nucleic Acids
	Biological liposomes: Erythrocyte Ghosts and Exosomes	YES	Transient	NO	Nucleic Acids

In another aspect, the delivery of genome editing system components or nucleic acids encoding such components, for example, a nucleic acid binding protein such as, for example, Cas9 or variants thereof, optionally fused to a polypeptide having biological activity (e.g., a nucleobase editor), and a gRNA targeting a genomic nucleic acid sequence of interest, may be accomplished by delivering a ribonucleoprotein (RNP) to cells. The RNP comprises the nucleic acid binding protein, e.g., Cas9, in complex with the targeting gRNA. RNPs may be delivered to cells using known methods, such as electroporation, nucleofection, or cationic lipid-mediated methods, for example, as reported by Zuris, J.A. et al., 2015, *Nat. Biotechnology*, 33(1):73-80. RNPs are advantageous for use in CRISPR base editing systems, particularly for cells that are difficult to transfect, such as primary cells. In addition, RNPs can also alleviate difficulties that may occur with protein expression in cells, especially when eukaryotic promoters, e.g., CMV or EF1A, which may be used in CRISPR plasmids, are not well-expressed. Advantageously, the use of RNPs does not require the delivery of foreign DNA into cells. Moreover, because an RNP comprising a nucleic acid binding protein and gRNA complex is degraded over time, the use of RNPs has the potential to limit off-target effects. In a manner similar to that for plasmid based techniques, RNPs can be used to deliver binding protein (e.g., Cas9 variants) and to direct homology directed repair (HDR).

A promoter used to drive the CRISPR system (e.g., including the Cas9 described herein) can include AAV ITR. This can be advantageous for eliminating the need for an additional promoter element, which can take up space in the vector. The additional space

freed up can be used to drive the expression of additional elements, such as a guide nucleic acid or a selectable marker. ITR activity is relatively weak, so it can be used to reduce potential toxicity due to over expression of the chosen nuclease.

Any suitable promoter can be used to drive expression of the Cas9 and, where  
5 appropriate, the guide nucleic acid. For ubiquitous expression, promoters that can be used include CMV, CAG, CBh, PGK, SV40, Ferritin heavy or light chains, etc. For brain or other CNS cell expression, suitable promoters can include: SynapsinI for all neurons, CaMKIIalpha for excitatory neurons, GAD67 or GAD65 or VGAT for GABAergic neurons, etc. For liver cell expression, suitable promoters include the Albumin promoter. For lung cell expression,  
10 suitable promoters can include SP-B. For endothelial cells, suitable promoters can include ICAM. For hematopoietic cells suitable promoters can include IFNbeta or CD45. For Osteoblasts suitable promoters can include OG-2.

In some cases, a Cas9 of the present disclosure is of small enough size to allow separate promoters to drive expression of the base editor and a compatible guide nucleic acid  
15 within the same nucleic acid molecule. For instance, a vector or viral vector can comprise a first promoter operably linked to a nucleic acid encoding the base editor and a second promoter operably linked to the guide nucleic acid.

The promoter used to drive expression of a guide nucleic acid can include: Pol III promoters such as U6 or H1 Use of Pol II promoter and intronic cassettes to express gRNA  
20 Adeno Associated Virus (AAV).

A Cas9 described herein with or without one or more guide nucleic can be delivered using adeno associated virus (AAV), lentivirus, adenovirus or other plasmid or viral vector types, in particular, using formulations and doses from, for example, U.S. Patent No. 8,454,972 (formulations, doses for adenovirus), U.S. Patent No. 8,404,658 (formulations,  
25 doses for AAV) and U.S. Patent No. 5,846,946 (formulations, doses for DNA plasmids) and from clinical trials and publications regarding the clinical trials involving lentivirus, AAV and adenovirus. For example, for AAV, the route of administration, formulation and dose can be as in U.S. Patent No. 8,454,972 and as in clinical trials involving AAV. For Adenovirus, the route of administration, formulation and dose can be as in U.S. Patent No.  
30 8,404,658 and as in clinical trials involving adenovirus. For plasmid delivery, the route of administration, formulation and dose can be as in U.S. Patent No. 5,846,946 and as in clinical studies involving plasmids. Doses can be based on or extrapolated to an average 70 kg

individual (*e.g.* a male adult human), and can be adjusted for patients, subjects, mammals of different weight and species. Frequency of administration is within the ambit of the medical or veterinary practitioner (*e.g.*, physician, veterinarian), depending on usual factors including the age, sex, general health, other conditions of the patient or subject and the particular  
5 condition or symptoms being addressed. The viral vectors can be injected into the tissue of interest. For cell-type specific base editing, the expression of the base editor and optional guide nucleic acid can be driven by a cell-type specific promoter.

For *in vivo* delivery, AAV can be advantageous over other viral vectors. In some cases, AAV allows low toxicity, which can be due to the purification method not requiring  
10 ultra-centrifugation of cell particles that can activate the immune response. In some cases, AAV allows low probability of causing insertional mutagenesis because it doesn't integrate into the host genome.

AAV has a packaging limit of 4.5 or 4.75 Kb. Constructs larger than 4.5 or 4.75 Kb can lead to significantly reduced virus production. For example, SpCas9 is quite large, the  
15 gene itself is over 4.1 Kb, which makes it difficult for packing into AAV. Therefore, embodiments of the present disclosure include utilizing a disclosed Cas9 which is shorter in length than conventional Cas9.

An AAV can be AAV1, AAV2, AAV5 or any combination thereof. One can select the type of AAV with regard to the cells to be targeted; *e.g.*, one can select AAV serotypes 1,  
20 2, 5 or a hybrid capsid AAV1, AAV2, AAV5 or any combination thereof for targeting brain or neuronal cells; and one can select AAV4 for targeting cardiac tissue. AAV8 is useful for delivery to the liver. A tabulation of certain AAV serotypes as to these cells can be found in Grimm, D. et al, J. Virol. 82: 5887-5911 (2008)).

Lentiviruses are complex retroviruses that have the ability to infect and express their  
25 genes in both mitotic and post-mitotic cells. The most commonly known lentivirus is the human immunodeficiency virus (HIV), which uses the envelope glycoproteins of other viruses to target a broad range of cell types.

Lentiviruses can be prepared as follows. After cloning pCasES10 (which contains a lentiviral transfer plasmid backbone), HEK293FT at low passage (p=5) were seeded in a T-75  
30 flask to 50% confluence the day before transfection in DMEM with 10% fetal bovine serum and without antibiotics. After 20 hours, media is changed to OptiMEM (serum-free) media and transfection was done 4 hours later. Cells are transfected with 10 µg of lentiviral transfer

plasmid (pCasES10) and the following packaging plasmids: 5 µg of pMD2.G (VSV-g pseudotype), and 7.5 µg of psPAX2 (gag/pol/rev/tat). Transfection can be done in 4 mL OptiMEM with a cationic lipid delivery agent (50 µl Lipofectamine 2000 and 100 ul Plus reagent). After 6 hours, the media is changed to antibiotic-free DMEM with 10% fetal  
5 bovine serum. These methods use serum during cell culture, but serum-free methods are preferred.

Lentivirus can be purified as follows. Viral supernatants are harvested after 48 hours. Supernatants are first cleared of debris and filtered through a 0.45 µm low protein binding (PVDF) filter. They are then spun in an ultracentrifuge for 2 hours at 24,000 rpm. Viral  
10 pellets are resuspended in 50 µl of DMEM overnight at 4° C. They are then aliquoted and immediately frozen at -80° C.

In another embodiment, minimal non-primate lentiviral vectors based on the equine infectious anemia virus (EIAV) are also contemplated. In another embodiment, RetinoStat®, an equine infectious anemia virus-based lentiviral gene therapy vector that expresses  
15 angiostatic proteins endostatin and angiostatin that is contemplated to be delivered via a subretinal injection. In another embodiment, use of self-inactivating lentiviral vectors is contemplated.

Any RNA of the systems, for example a guide RNA or a Cas9-encoding mRNA, can be delivered in the form of RNA. Cas9 encoding mRNA can be generated using *in vitro*  
20 transcription. For example, Cas9 mRNA can be synthesized using a PCR cassette containing the following elements: T7 promoter, optional kozak sequence (GCCACC), nuclease sequence, and 3' UTR such as a 3' UTR from beta globin-polyA tail. The cassette can be used for transcription by T7 polymerase. Guide polynucleotides (*e.g.*, gRNA) can also be transcribed using *in vitro* transcription from a cassette containing a T7 promoter, followed by  
25 the sequence "GG", and guide polynucleotide sequence.

To enhance expression and reduce possible toxicity, the Cas9 sequence and/or the guide nucleic acid can be modified to include one or more modified nucleoside *e.g.* using pseudo-U or 5-Methyl-C.

The disclosure in some embodiments comprehends a method of modifying a cell or  
30 organism. The cell can be a prokaryotic cell or a eukaryotic cell. The cell can be a mammalian cell. The mammalian cell may be a non-human primate, bovine, porcine, rodent or mouse cell. The modification introduced to the cell by the base editors,

compositions and methods of the present disclosure can be such that the cell and progeny of the cell are altered for improved production of biologic products such as an antibody, starch, alcohol or other desired cellular output. The modification introduced to the cell by the methods of the present disclosure can be such that the cell and progeny of the cell include an alteration that changes the biologic product produced.

The system can comprise one or more different vectors. In an aspect, the Cas9 is codon optimized for expression the desired cell type, preferentially a eukaryotic cell, preferably a mammalian cell or a human cell.

In general, codon optimization refers to a process of modifying a nucleic acid sequence for enhanced expression in the host cells of interest by replacing at least one codon (*e.g.* about or more than about 1, 2, 3, 4, 5, 10, 15, 20, 25, 50, or more codons) of the native sequence with codons that are more frequently or most frequently used in the genes of that host cell while maintaining the native amino acid sequence. Various species exhibit particular bias for certain codons of a particular amino acid. Codon bias (differences in codon usage between organisms) often correlates with the efficiency of translation of messenger RNA (mRNA), which is in turn believed to be dependent on, among other things, the properties of the codons being translated and the availability of particular transfer RNA (tRNA) molecules. The predominance of selected tRNAs in a cell is generally a reflection of the codons used most frequently in peptide synthesis. Accordingly, genes can be tailored for optimal gene expression in a given organism based on codon optimization. Codon usage tables are readily available, for example, at the "Codon Usage Database" available at [www.kazusa.or.jp/codon/](http://www.kazusa.or.jp/codon/) (visited Jul. 9, 2002), and these tables can be adapted in a number of ways. See, Nakamura, Y., *et al.* "Codon usage tabulated from the international DNA sequence databases: status for the year 2000" *Nucl. Acids Res.* 28:292 (2000). Computer algorithms for codon optimizing a particular sequence for expression in a particular host cell are also available, such as Gene Forge (Aptagen; Jacobus, Pa.), are also available. In some embodiments, one or more codons (*e.g.* 1, 2, 3, 4, 5, 10, 15, 20, 25, 50, or more, or all codons) in a sequence encoding an engineered nuclease correspond to the most frequently used codon for a particular amino acid.

Packaging cells are typically used to form virus particles that are capable of infecting a host cell. Such cells include 293 cells, which package adenovirus, and psi.2 cells or PA317 cells, which package retrovirus. Viral vectors used in gene therapy are usually generated by producing a cell line that packages a nucleic acid vector into a viral particle. The vectors

typically contain the minimal viral sequences required for packaging and subsequent integration into a host, other viral sequences being replaced by an expression cassette for the polynucleotide(s) to be expressed. The missing viral functions are typically supplied in trans by the packaging cell line. For example, AAV vectors used in gene therapy typically only  
5 possess ITR sequences from the AAV genome which are required for packaging and integration into the host genome. Viral DNA can be packaged in a cell line, which contains a helper plasmid encoding the other AAV genes, namely rep and cap, but lacking ITR sequences. The cell line can also be infected with adenovirus as a helper. The helper virus can promote replication of the AAV vector and expression of AAV genes from the helper  
10 plasmid. The helper plasmid in some cases is not packaged in significant amounts due to a lack of ITR sequences. Contamination with adenovirus can be reduced by, *e.g.*, heat treatment to which adenovirus is more sensitive than AAV.

### **Pharmaceutical Compositions**

Other aspects of the present disclosure relate to pharmaceutical compositions  
15 comprising CRISPR system (*e.g.*, including Cas9 disclosed herein). The term “pharmaceutical composition”, as used herein, refers to a composition formulated for pharmaceutical use. In some embodiments, the pharmaceutical composition further comprises a pharmaceutically acceptable carrier. In some embodiments, the pharmaceutical composition comprises additional agents (*e.g.*, for specific delivery, increasing half-life, or  
20 other therapeutic compounds).

As used here, the term “pharmaceutically-acceptable carrier” means a pharmaceutically-acceptable material, composition or vehicle, such as a liquid or solid filler, diluent, excipient, manufacturing aid (*e.g.*, lubricant, talc magnesium, calcium or zinc stearate, or steric acid), or solvent encapsulating material, involved in carrying or transporting  
25 the compound from one site (*e.g.*, the delivery site) of the body, to another site (*e.g.*, organ, tissue or portion of the body). A pharmaceutically acceptable carrier is “acceptable” in the sense of being compatible with the other ingredients of the formulation and not injurious to the tissue of the subject (*e.g.*, physiologically compatible, sterile, physiologic pH, etc.).

Some nonlimiting examples of materials which can serve as pharmaceutically-  
30 acceptable carriers include: (1) sugars, such as lactose, glucose and sucrose; (2) starches, such as corn starch and potato starch; (3) cellulose, and its derivatives, such as sodium carboxymethyl cellulose, methylcellulose, ethyl cellulose, microcrystalline cellulose and

cellulose acetate; (4) powdered tragacanth; (5) malt; (6) gelatin; (7) lubricating agents, such as magnesium stearate, sodium lauryl sulfate and talc; (8) excipients, such as cocoa butter and suppository waxes; (9) oils, such as peanut oil, cottonseed oil, safflower oil, sesame oil, olive oil, corn oil and soybean oil; (10) glycols, such as propylene glycol; (11) polyols, such as glycerin, sorbitol, mannitol and polyethylene glycol (PEG); (12) esters, such as ethyl oleate and ethyl laurate; (13) agar; (14) buffering agents, such as magnesium hydroxide and aluminum hydroxide; (15) alginic acid; (16) pyrogen-free water; (17) isotonic saline; (18) Ringer's solution; (19) ethyl alcohol; (20) pH buffered solutions; (21) polyesters, polycarbonates and/or polyanhydrides; (22) bulking agents, such as polypeptides and amino acids (23) serum alcohols, such as ethanol; and (23) other non-toxic compatible substances employed in pharmaceutical formulations. Wetting agents, coloring agents, release agents, coating agents, sweetening agents, flavoring agents, perfuming agents, preservative and antioxidants can also be present in the formulation. The terms such as "excipient," "carrier," "pharmaceutically acceptable carrier," "vehicle," or the like are used interchangeably herein.

Pharmaceutical compositions can comprise one or more pH buffering compounds to maintain the pH of the formulation at a predetermined level that reflects physiological pH, such as in the range of about 5.0 to about 8.0. The pH buffering compound used in the aqueous liquid formulation can be an amino acid or mixture of amino acids, such as histidine or a mixture of amino acids such as histidine and glycine. Alternatively, the pH buffering compound is preferably an agent which maintains the pH of the formulation at a predetermined level, such as in the range of about 5.0 to about 8.0, and which does not chelate calcium ions. Illustrative examples of such pH buffering compounds include, but are not limited to, imidazole and acetate ions. The pH buffering compound may be present in any amount suitable to maintain the pH of the formulation at a predetermined level.

Pharmaceutical compositions can also contain one or more osmotic modulating agents, *i.e.*, a compound that modulates the osmotic properties (*e.g.*, tonicity, osmolality, and/or osmotic pressure) of the formulation to a level that is acceptable to the blood stream and blood cells of recipient individuals. The osmotic modulating agent can be an agent that does not chelate calcium ions. The osmotic modulating agent can be any compound known or available to those skilled in the art that modulates the osmotic properties of the formulation. One skilled in the art may empirically determine the suitability of a given osmotic modulating agent for use in the inventive formulation. Illustrative examples of suitable types of osmotic modulating agents include, but are not limited to: salts, such as

sodium chloride and sodium acetate; sugars, such as sucrose, dextrose, and mannitol; amino acids, such as glycine; and mixtures of one or more of these agents and/or types of agents. The osmotic modulating agent(s) may be present in any concentration sufficient to modulate the osmotic properties of the formulation.

5           In some embodiments, the pharmaceutical composition is formulated for delivery to a subject, *e.g.*, for gene editing. Suitable routes of administering the pharmaceutical composition described herein include, without limitation: topical, subcutaneous, transdermal, intradermal, intralesional, intraarticular, intraperitoneal, intravesical, transmucosal, gingival, intradental, intracochlear, transtympanic, intraorgan, epidural, intrathecal, intramuscular,  
10 intravenous, intravascular, intraosseous, periocular, intratumoral, intracerebral, and intracerebroventricular administration.

          In some embodiments, the pharmaceutical composition described herein is administered locally to a diseased site. In some embodiments, the pharmaceutical composition described herein is administered to a subject by injection, by means of a  
15 catheter, by means of a suppository, or by means of an implant, the implant being of a porous, non-porous, or gelatinous material, including a membrane, such as a sialastic membrane, or a fiber.

          In other embodiments, the pharmaceutical composition described herein is delivered in a controlled release system. In one embodiment, a pump can be used (*See, e.g.*, Langer, 20 1990, *Science* 249: 1527-1533; Sefton, 1989, *CRC Crit. Ref. Biomed. Eng.* 14:201; Buchwald *et al.*, 1980, *Surgery* 88:507; Saudek *et al.*, 1989, *N. Engl. J. Med.* 321:574). In another embodiment, polymeric materials can be used. (*See, e.g.*, *Medical Applications of Controlled Release* (Langer and Wise eds., CRC Press, Boca Raton, Fla., 1974); *Controlled Drug Bioavailability, Drug Product Design and Performance* (Smolen and Ball eds., Wiley, 25 New York, 1984); Ranger and Peppas, 1983, *Macromol. Sci. Rev. Macromol. Chem.* 23:61. See also Levy *et al.*, 1985, *Science* 228: 190; During *et al.*, 1989, *Ann. Neurol.* 25:351; Howard *et al.*, 1989, *J. Neurosurg.* 71: 105.) Other controlled release systems are discussed, for example, in Langer, *supra*.

          In some embodiments, the pharmaceutical composition is formulated in accordance  
30 with routine procedures as a composition adapted for intravenous or subcutaneous administration to a subject, *e.g.*, a human. In some embodiments, pharmaceutical composition for administration by injection are solutions in sterile isotonic use as solubilizing



agent and a local anesthetic such as lignocaine to ease pain at the site of the injection. Generally, the ingredients are supplied either separately or mixed together in unit dosage form, for example, as a dry lyophilized powder or water free concentrate in a hermetically sealed container such as an ampoule or sachette indicating the quantity of active agent.

5 Where the pharmaceutical is to be administered by infusion, it can be dispensed with an infusion bottle containing sterile pharmaceutical grade water or saline. Where the pharmaceutical composition is administered by injection, an ampoule of sterile water for injection or saline can be provided so that the ingredients can be mixed prior to administration.

10 A pharmaceutical composition for systemic administration can be a liquid, *e.g.*, sterile saline, lactated Ringer's or Hank's solution. In addition, the pharmaceutical composition can be in solid forms and re-dissolved or suspended immediately prior to use. Lyophilized forms are also contemplated. The pharmaceutical composition can be contained within a lipid particle or vesicle, such as a liposome or microcrystal, which is also suitable for parenteral  
15 administration. The particles can be of any suitable structure, such as unilamellar or plurilamellar, so long as compositions are contained therein. Compounds can be entrapped in "stabilized plasmid-lipid particles" (SPLP) containing the fusogenic lipid dioleoylphosphatidylethanolamine (DOPE), low levels (5-10 mol%) of cationic lipid, and stabilized by a polyethyleneglycol (PEG) coating (Zhang Y. P. et al, *Gene Ther.* 1999, 6:  
20 1438-47). Positively charged lipids such as N-[1-(2,3-dioleoyloxy)propyl]-N,N,N-trimethylammoniummethylsulfate, or "DOTAP," are particularly preferred for such particles and vesicles. The preparation of such lipid particles is well known. *See, e.g.*, U.S. Patent Nos. 4,880,635; 4,906,477; 4,911,928; 4,917,951; 4,920,016; and 4,921,757; each of which is incorporated herein by reference.

25 The pharmaceutical composition described herein can be administered or packaged as a unit dose, for example. The term "unit dose" when used in reference to a pharmaceutical composition of the present disclosure refers to physically discrete units suitable as unitary dosage for the subject, each unit containing a predetermined quantity of active material calculated to produce the desired therapeutic effect in association with the required diluent;  
30 *i.e.*, carrier, or vehicle.

Further, the pharmaceutical composition can be provided as a pharmaceutical kit comprising (a) a container containing a compound of the invention in lyophilized form and (b) a second container containing a pharmaceutically acceptable diluent (*e.g.*, sterile used for

reconstitution or dilution of the lyophilized compound of the invention. Optionally associated with such container(s) can be a notice in the form prescribed by a governmental agency regulating the manufacture, use or sale of pharmaceuticals or biological products, which notice reflects approval by the agency of manufacture, use or sale for human  
5 administration.

In another aspect, an article of manufacture containing materials useful for the treatment of the diseases described above is included. In some embodiments, the article of manufacture comprises a container and a label. Suitable containers include, for example, bottles, vials, syringes, and test tubes. The containers can be formed from a variety of  
10 materials such as glass or plastic. In some embodiments, the container holds a composition that is effective for treating a disease described herein and can have a sterile access port. For example, the container can be an intravenous solution bag or a vial having a stopper pierceable by a hypodermic injection needle. The active agent in the composition is a compound of the invention. In some embodiments, the label on or associated with the  
15 container indicates that the composition is used for treating the disease of choice. The article of manufacture can further comprise a second container comprising a pharmaceutically-acceptable buffer, such as phosphate-buffered saline, Ringer's solution, or dextrose solution. It can further include other materials desirable from a commercial and user standpoint, including other buffers, diluents, filters, needles, syringes, and package inserts with  
20 instructions for use.

In some embodiments, the CRISPR system (e.g., including the Cas9 described herein) are provided as part of a pharmaceutical composition. In some embodiments, the pharmaceutical composition comprises any of the fusion proteins provided herein (e.g., including the nucleobase editor described herein comprising LubCas9). In some  
25 embodiments, the pharmaceutical composition comprises any of the complexes provided herein. In some embodiments, the pharmaceutical composition comprises a ribonucleoprotein complex comprising an RNA-guided nuclease (e.g., Cas9) that forms a complex with a gRNA and a cationic lipid. In some embodiments pharmaceutical composition comprises a gRNA, a nucleic acid programmable DNA binding protein, a  
30 cationic lipid, and a pharmaceutically acceptable excipient. Pharmaceutical compositions can optionally comprise one or more additional therapeutically active substances.

### **Kits**

In one aspect, the invention provides kits containing any one or more of the elements disclosed in the above methods and compositions. In some embodiments, the kit comprises a vector system and instructions for using the kit. In some embodiments, the vector system comprises one or more insertion sites for inserting a guide sequence, wherein when  
5 expressed, the guide sequence directs sequence-specific binding of a CRISPR complex to a target sequence in a eukaryotic cell, wherein the CRISPR complex comprises a CRISPR enzyme complexed with (1) the guide sequence that is hybridized to the target sequence, and (2) a sequence that is hybridized to the tracr sequence; and/or (b) a second regulatory element operably linked to an enzyme-coding sequence encoding said CRISPR enzyme comprising a  
10 nuclear localization sequence. Elements may be provide individually or in combinations, and may be provided in any suitable container, such as a vial, a bottle, or a tube. In some embodiments, the kit includes instructions in one or more languages, for example in more than one language.

In some embodiments, the kit comprises a nucleobase editor. For example, in some  
15 embodiments, the kit includes a nucleobase editor comprising the Cas9 enzymes (ScaCas9, SirCas9, VapCas9, EpeCas9, LfeCas9, PmaCas9) described herein.

In some embodiments, a kit comprises one or more reagents for use in a process utilizing one or more of the elements described herein. Reagents may be provided in any suitable container. For example, a kit may provide one or more reaction or storage buffers.  
20 Reagents may be provided in a form that is usable in a particular assay, or in a form that requires addition of one or more other components before use (e.g. in concentrate or lyophilized form). A buffer can be any buffer, including but not limited to a sodium carbonate buffer, a sodium bicarbonate buffer, a borate buffer, a Tris buffer, a MOPS buffer, a HEPES buffer, and combinations thereof. In some embodiments, the buffer is alkaline. In  
25 some embodiments, the buffer has a pH from about 7 to about 10. In some embodiments, the kit comprises one or more oligonucleotides corresponding to a guide sequence for insertion into a vector so as to operably link the guide sequence and a regulatory element. In some embodiments, the kit comprises a homologous recombination template polynucleotide.

All publications, patent applications, patents, and other references mentioned herein  
30 are incorporated by reference in their entirety. In addition, the materials, methods, and examples are illustrative only and not intended to be limiting. Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although methods and

materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, suitable methods and materials are described herein.

## EXAMPLES

5           The following examples describe some of the preferred modes of making and practicing the present invention. However, it should be understood that these examples are for illustrative purposes only and are not meant to limit the scope of the invention.

Example 1. Screening for novel Cas9 enzymes, discovery and optimization of novel Cas9 enzymes

10           This example describes a screen for the discovery of novel Cas9 enzymes. As described herein, using this screen novel Cas9 enzymes from *Streptococcus constellatus*, *Sharpea spp. isolate RUG017*, *Veillonella parvula*, *Ezakiella peruensis*, *Lactobacillus fermentum strain AF15-40LB* and *Peptoniphilus sp. Marseille-P3761* bacteria were isolated and optimized.

15           In a search to discover new Cas9 enzymes which recognize novel PAM sequences, a bioinformatics screen was used to search for additional enzymes to expand CRISPR's targeting range. The screen utilized seed sequences of Cas9 from *S. pyogenes*, *S. aureus*, *S. thermophilus*, and *F. novicida*. Bioinformatics was carried out using the tblastn variant of BLAST with an e-value threshold of 1e-6 for considering BLAST hits. Briefly, loci selected  
20 for testing were loci that remained intact in the presence of Cas9 proteins from other species. Loci were selected that had greater than three spacers within the CRISPR array and greater than 1 kb endogenous sequence 5' of Cas9 and greater than 300 nt 3' of the CRISPR array. Using this approach, novel Cas9 enzymes were identified from different bacterial species and codon optimized for expression in human cells. The novel engineered Cas9 enzymes were  
25 then recombinantly produced and tested.

Example 2. Identifying 3' PAM consensus motif for novel Cas9 enzymes from *Streptococcus constellatus*, *Sharpea spp. isolate RUG017*, *Veillonella parvula*, *Ezakiella peruensis*, *Lactobacillus fermentum strain AF15-40LB* and *Peptoniphilus sp. Marseille-P3761* bacteria

30           This example illustrates the identification of the protospacer adjacent motif (PAM) sequence for human codon-optimized Cas9 originally isolated from *Streptococcus*

*constellatus*, *Sharpea* spp. isolate RUG017, *Veillonella parvula*, *Ezakiella peruensis*, ,  
*Lactobacillus fermentum* strain AF15-40LB and *Peptoniphilus* sp. Marseille-P3761 species.

The human, codon-optimized Cas9 was tested for its recognition of a PAM sequence using an *in vitro* PAM identification assay. A library of plasmids bearing randomized PAM sequences were incubated with Cas9 isolated from different bacteria. Uncleaved plasmid was purified and sequenced to identify specific PAM motifs that were cleaved. The consensus PAM sequence recognized by *Streptococcus constellatus* Cas9 was identified as 5'-NGG-3' (FIG. 1A). The consensus PAM sequence recognized by *Sharpea* spp. isolate RUG017 Cas9 was identified as 5'-NAGHC-3' (FIG. 1B). The consensus PAM sequence recognized by *Veillonella parvula* Cas9 was identified as 5'-NRHRRH-3' (H=A, C or T; R=A or G) (FIG. 1C). The consensus PAM sequence recognized by *Ezakiella peruensis* Cas9 was identified as 5'-NGG-3' (FIG. 1D). The consensus PAM sequence recognized by *Lactobacillus fermentum* strain AF15-40LB Cas9 was identified as 5'-NNAAA-3' (FIG. 1E). The consensus PAM sequence recognized by *Peptoniphilus* sp. Marseille-P3761 Cas9 was identified as 5'-NGG-3' (FIG. 1F).

Example 3. Predicting RNA folding structure of sgRNA for novel Cas9 enzymes from *Streptococcus constellatus*, *Sharpea* spp. isolate RUG017, *Veillonella parvula*, *Ezakiella peruensis*, *Lactobacillus fermentum* strain AF15-40LB and *Peptoniphilus* sp. Marseille-P3761 bacteria

This example demonstrates the predicted RNA folding structure of exemplary sgRNA comprising crRNA and tracrRNA for use with novel Cas9 enzymes.

Small RNA sequencing was carried out on RNA derived from an *E. coli* strain heterologously expressing Cas9 Crispr loci. Briefly, RNA was isolated from stationary phase bacteria by first resuspending the *E. coli* in Trizol, then homogenizing the bacteria with zirconia/silica beads in a homogenizer for three 1 min cycles. Total RNA was purified from homogenized samples, DNase treated and 3' dephosphorylated with T4 polynucleotide kinase and rRNA was removed. RNA libraries were prepared from rRNA-depleted RNA, and size selected for small RNA.

For RNA sequencing, transcripts were poly-A tailed with *E. coli* Poly (A) polymerase, ligated with 5' RNA adapters using T4 RNA ligase 1 and reverse transcribed, followed by PCR amplification of cDNA with barcoded primers, and sequencing on a MiSeq. Reads from each sample were identified on the basis of their associated barcode and aligned to a

reference sequence using BWA. Paired-end alignments were used to extract transcript sequences using Picard tools and the sequences were analyzed using Geneious software.

RNA folding was based on prediction from Geneious 11.1.2 software. The single sgRNA transcript fuses the crRNA to tracrRNA mimicking the dual RNA structure required to guide site-specific Cas9 activity. The predicted RNA folding structure for the chimeric sgRNA for use with ScoCas9 from *Streptococcus constellatus* is shown in FIG. 2A, sgRNA for use with SirCas9 from *Sharpea spp. isolate RUG017* is shown in FIG. 2B, sgRNA for use with VapCas9 from *Veillonella parvula* is shown in FIG. 2C, sgRNA for use with EpeCas9 from *Ezakiella peruensis* is shown in FIG. 2D, sgRNA for use with LfeCas9 from *Lactobacillus fermentum strain AF15-40LB* is shown in FIG. 2E and sgRNA for use with PmaCas9 from *Peptoniphilus sp. Marseille-P3761* is shown in FIG. 2F.

Example 4. Ex vivo cleavage activity by WT ScoCas9 in HEK293T cells

This example illustrates *ex vivo* nucleic acid cleavage activity by WT *ScoCas9* from *Streptococcus constellatus* in HEK293T cells.

HEK293T cells were plated in a 96-well plate. Cells were transfected with expression vectors containing Cas9 and guide RNAs (Table 10), 24 hours after plating. Cells were harvested 72 hours post-transfection and total DNA was extracted.

Deep sequencing was carried out to characterize indel patterns in the HEK293T cells. Briefly, exemplary targets (Table 8) were amplified using a two-round PCR to add Illumina adapters as well as unique barcodes to the target amplicons. PCR products were run on a 2% gel and gel extracted. Samples were pooled, quantified and cDNA libraries were prepared and sequenced on MiSeq. Indel frequency was determined by deep sequencing (FIG. 3).

**Table 8. Exemplary Guide RNA Sequences and PAM Sequences**

ID (Sco/Pma)	5'->3' guide sequence	3' PAM
guide 2	GAAACAATGATAACAAGACC (SEQ ID NO: 97)	TGG
guide 3	GTGGCCCCTGTGCCAGCCC (SEQ ID NO: 98)	TGG
guide 4	GTCCCAAATATGTAGCTGTT (SEQ ID NO: 99)	TGG
guide 6	GCTCCCATCACATCAACCGG (SEQ ID NO: 100)	TGG

guide 7	GATGTCACCTCCAATGACTA (SEQ ID NO: 101)	GGG
guide 9	GTTGAAGATGAAGCCCAGAG (SEQ ID NO: 102)	CGG
guide 10	GCCAACACCAACCAGAACTT (SEQ ID NO: 103)	GGG
guide 11	TGCTGCACACAGCAGGCCTT (SEQ ID NO: 104)	TGG

The data showed that that WT *ScoCas9* achieved between 2-32% indel frequency. Guide RNAs 2 and 9 resulted in greater than 30% indel mutations, while guide RNA 11 resulted in about 2% indel mutations.

Example 5. Base editing by *Cas9* enzyme with an N-terminal fusion of an adenine base editor (ABE) or a cytidine base editor (CBE)

This example illustrates base conversion efficiency of a *Cas9* enzyme fused to an adenine base editor (ABE), or to a cytidine base editor (CBE).

Briefly, 25,000 HEK293T cells were plated per 96-well. 100 ng of *Cas9* expression plasmid and 100 ng of guide expression plasmid were transfected 24h after plating. Cells were harvested 5 days after transfection and DNA was extracted.

Deep sequencing was carried out to characterize A-to-G conversion or C-to-T conversion in the HEK293T cells. Exemplary targets were amplified using a two-round PCR region to add Illumina adapters as well as unique barcodes to the target amplicons. PCR products were run on a 2% gel and gel extracted. Samples were pooled, quantified and cDNA libraries were prepared and sequenced on MiSeq. The percent A-to-G conversion was determined by deep sequencing for the N-terminal as well as the C-terminal Tada8 fusion constructs. The percent C-to-T conversion was determined by deep sequencing for the N-terminal as well as the C-terminal ppAPOBEC1 fusion constructs.

FIG. 4A shows a schematic diagram of constructs of *ScoCas9* fused to ABE or CBE at the N-terminal. Table 9 shows the guide RNA sequences used with *ScoCas9*. FIG. 4B shows a graph of indel mutations and targeted adenine to guanine conversion percentage achieved with an N-terminal fusion of *ScoCas9* to an adenine base editor (ABE) (FIG. 4B), which are directed to genomic sites in a human cell line (HEK293T). FIG. 4C shows a graph of indel mutations and targeted cytosine to thymine conversion percentage achieved with an

N-terminal fusion of *ScoCas9* to a cytidine base editor (FIG. 4C), which are directed to genomic sites in a human cell line (HEK293T).

**Table 9. Guide RNA Sequences and PAM Sequences used with *ScoCas9***

ID (Sco)	5'→3' guide sequence	3' PAM
guide 1	GAACACAAAGCATAGACTGC (SEQ ID NO: 105)	GGG
guide 2	GAAACAATGATAACAAGACC (SEQ ID NO: 106)	TGG
guide 3	GTGGCCCCTGTGCCAGCCC (SEQ ID NO: 107)	TGG
guide 4	GTCCCAAATATGTAGCTGTT (SEQ ID NO: 108)	TGG
guide 5	AGAGGGACACACAGATCTAT (SEQ ID NO: 109)	TGG
guide 6	GCTCCCATCACATCAACCGG (SEQ ID NO: 110)	TGG
guide 7	GATGTCACCTCCAATGACTA (SEQ ID NO: 111)	GGG
guide 8	GGGCAACCACAAACCCACGA (SEQ ID NO: 112)	GGG
guide 9	GTTGAAGATGAAGCCCAGAG (SEQ ID NO: 113)	CGG
guide 10	GCCAACACCAACCAGAACTT (SEQ ID NO: 114)	GGG
guide 11	TGCTGCACACAGCAGGCCTT (SEQ ID NO: 115)	TGG
guide 12	GTGCCAGAAACAGGGGTGAC (SEQ ID NO: 116)	GGG

FIG. 5A shows a schematic diagram of constructs of WT *SirCas9* as well as *SirCas9* (“D14A” mutant) fused to an ABE at the N-terminal. Table 10 shows the exemplary NAGMC guide RNA sequences used with *SirCas9*. FIG. 5B shows a graph of indel mutations and targeted adenine to guanine conversion percentage achieved with an N-terminal fusion of *SirCas9* to an adenine base editor (ABE) (FIG. 5B), which are directed to genomic sites in a human cell line (HEK293T).

**Table 10. Guide RNA Sequences and PAM Sequences used with *SirCas9***



ID (Sir)	5'->3' sequence	3' PAM
guide 1	CCTGCCTCAGCTGCTCACTT (SEQ ID NO: 117)	GAGCC
guide 2	AAACGGTCCCCAGAGGGTTC (SEQ ID NO: 118)	TAGAC
guide 3	GCCACCGGTTGATGTGATGG (SEQ ID NO: 119)	GAGCC
guide 4	AAGTGGTCCCAGGCCTCAGC (SEQ ID NO: 120)	CAGCC
guide 5	AGAGAAAATGAAACTTTCAA (SEQ ID NO: 121)	AAGCC
guide 6	CCAAACCCA ACTCCATCTAC (SEQ ID NO: 122)	CAGCC
guide 7	GGTCCTTGAATTGCAGTATC (SEQ ID NO: 123)	TAGCC
guide 8	GCATAGACTGCGGGGCGGGC (SEQ ID NO: 124)	CAGCC
guide 9	GGAAACTGGAACACAAAGCA (SEQ ID NO: 125)	TAGAC
guide 10	GACAGCATGTGGTAATTTTC (SEQ ID NO: 126)	CAGCC
guide 11	GCCCCGGA AACTCTGTCCA (SEQ ID NO: 127)	GAGAC
guide 12	TCGACCCCCACCAAGGTTCA (SEQ ID NO: 128)	CAGCC

FIG. 6A shows a schematic diagram of constructs showing WT VapCas9, as well as VapCas9 (“D38A” mutant) fused to an ABE or CBE at the N-terminal. Table 11 shows the exemplary NRHRRH [wherein H is adenine, cytosine or thymine, and R is adenine or guanine] guide RNA sequences used with *VapCas9*. FIG. 6B shows a graph of indel mutations and targeted adenine to guanine conversion percentage achieved with an N-terminal fusion of *VapCas9* to an adenine base editor (ABE) as well as targeted cytosine to thymine conversion percentage achieved with an N-terminal fusion of *VapCas9* to a cytidine base editor (CBE) (FIG. 6B), which are directed to genomic sites in a human cell line (HEK293T).

10 **Table 11. Guide RNA Sequences and PAM Sequences for use with VapCas9**

ID (Vap)	5'->3' sequence	3' PAM
guide 1	TGTTAACAGCTGACCCAATA (SEQ ID NO: 129)	AGTGGC
guide 2	GTTACTCGCCTGTCAAGTGG (SEQ ID NO: 130)	CGTGAC
guide 3	GGGCTCCCATCACATCAACC (SEQ ID NO: 131)	GGTGGC
guide 4	GCTTTGGGGAGGCCTGGAGT (SEQ ID NO: 132)	CATGGC
guide 5	TAGCTGCCAATGACTATAGC (SEQ ID NO: 133)	AATAGC
guide 6	TTAAAATAGGATCTACATCA (SEQ ID NO: 134)	CGTAAC
guide 7	GAATCCTGCCATACACTTTG (SEQ ID NO: 135)	AATAGC
guide 8	CTGCGGGGCGGGCCAGCCTG (SEQ ID NO: 136)	AATAGC
guide 9	ACATTGTCAGAGGGACACAC (SEQ ID NO: 137)	TGTGGC
guide 10	AGCAACTCCAGTCCCAAATA (SEQ ID NO: 138)	TGTAGC
guide 11	GTGGTGGCCGAGCGCCCCCT (SEQ ID NO: 139)	AGTGAC
guide 12	CATTCACCCAGCTTCCCTGT (SEQ ID NO: 140)	GGTGGC

FIG. 7A shows a schematic diagram of constructs showing an N-terminal fusion of ABE and a C-terminal fusion of ABE to VapCas9. FIG. 7B shows a graph of targeted adenine to guanine conversion percentage achieved with an N-terminal fusion and C-terminal fusion to an adenine base editor (ABE).

5 FIG. 8A shows a schematic diagram of constructs showing an N-terminal fusion of ABE and CBE to EpeCas9. Table 12 shows the exemplary guide RNA sequences used with *EpeCas9*. FIG. 8B shows a graph of indel mutations, a graph of targeted adenine to guanine conversion percentage achieved with an N-terminal fusion to an ABE and targeted cytosine to thymine conversion percentage achieved with an N-terminal fusion to a CBE.

10 **Table 12. Guide RNA Sequences and PAM Sequences for use with EpeCas9**

<u>ID (Epe)</u>	<u>Sequence</u>	<u>PAM</u>
guide 1	GAACACAAAGCATAGACTGC (SEQ ID NO: 141)	GGG
guide 2	GAAACAATGATAACAAGACC (SEQ ID NO: 142)	TGG
guide 3	GTGGCCCCTGTGCCAGCCC (SEQ ID NO: 143)	TGG
guide 4	GTCCCAAATATGTAGCTGTT (SEQ ID NO: 144)	TGG
guide 5	AGAGGGACACACAGATCTAT (SEQ ID NO: 145)	TGG
guide 6	GCTCCCATCACATCAACCGG (SEQ ID NO: 146)	TGG
guide 7	GATGTCACCTCCAATGACTA (SEQ ID NO: 147)	GGG
guide 8	GGGCAACCACAAACCCACGA (SEQ ID NO: 148)	GGG
guide 9	GTTGAAGATGAAGCCCAGAG (SEQ ID NO: 149)	CGG
guide 10	GCCAACACCAACCAGAACTT (SEQ ID NO: 150)	GGG
guide 11	TGCTGCACACAGCAGGCCTT (SEQ ID NO: 151)	TGG
guide 12	GTGCCAGAAACAGGGGTGAC (SEQ ID NO: 152)	GGG

FIG. 9A shows a schematic diagram of constructs showing WT LfeCas9 and LfeCas9 D9A mutant fused at the N-terminus to an ABE and a CBE. Table 13 shows the exemplary guide RNA sequences used with *LfeCas9*. FIG. 9B shows a graph that shows results of the indel mutation frequency achieved with LfeCas9. FIG. 9C shows a graph of targeted adenine to guanine conversion achieved with an N-terminal fusion of LfeCas9 to an adenine base editor. FIG. 9D shows a graph of targeted cytosine to thymine conversion achieved with a base editor comprising a CBE fused to the N-terminus of an LfeCas9 D9A mutant.

**Table 13. Guide RNA Sequences and PAM Sequences for use with LfeCas9**

<u>ID (Lfe)</u>	<u>Sequence</u>	<u>PAM</u>
-----------------	-----------------	------------

guide 1	TCACGGAGACTGAACACTCC (SEQ ID NO: 153)	TCAAA
guide 2	GTAACAGACATGGACCATCA (SEQ ID NO: 154)	GGAAA
guide 3	GGGAGGGAGGGGCACAGATG (SEQ ID NO: 155)	AGAAA
guide 4	TGTGGTTCCAGAACCGGAGG (SEQ ID NO: 156)	ACAAA
guide 5	AATGAGAGAAAATGAACTT (SEQ ID NO: 157)	TCAAA
guide 6	GGCCATCAAGGATGCCACG (SEQ ID NO: 158)	AGAAA
guide 7	AAATTGTCCAGCCCCATCTG (SEQ ID NO: 159)	TCAAA
guide 8	CCTGTAAAGGAACTGGAAC (SEQ ID NO: 160)	ACAAA
guide 9	TACATGAAGCAACTCCAGTC (SEQ ID NO: 161)	CCAAA
guide 10	AAACTCCCCCACCCCCTTT (SEQ ID NO: 162)	CCAAA
guide 11	GAGTTGGGTTTGGTGCTCAA (SEQ ID NO: 163)	TGAAA
guide 12	GCGGGCCAGCCTGAATAGCT (SEQ ID NO: 164)	GCAAA

FIG. 10A shows a schematic of constructs showing WT PmaCas9 and PmaCas9 D12A mutant fused at the N-terminus and C-terminus to an ABE and a CBE. FIG. 10B shows a graph that shows results of A-to-G or C-to-T conversion achieved with a base editor comprising an ABE or a CBE fused to the N-terminus or C-terminus of an PmaCas9 D12A mutant.

**Table 14. Guide RNA Sequences and PAM Sequences for use with PmaCas9**

<u>ID (Pma)</u>	<u>Sequence</u>	<u>PAM</u>
guide 2	GAAACAATGATAACAAGACC (SEQ ID NO: 165)	TGG
guide 3	GTGGCCCCTGTGCCAGCCC (SEQ ID NO: 166)	TGG
guide 4	GTCCCAAATATGTAGCTGTT (SEQ ID NO: 167)	TGG

guide 6	GCTCCCATCACATCAACCGG (SEQ ID NO: 168)	TGG
guide 7	GATGTCACCTCCAATGACTA (SEQ ID NO: 169)	GGG
guide 9	GTTGAAGATGAAGCCCAGAG (SEQ ID NO: 170)	CGG
guide 10	GCCAACACCAACCAGAACTT (SEQ ID NO: 171)	GGG
guide 11	TGCTGCACACAGCAGGCCTT (SEQ ID NO: 172)	TGG

Table 15 discloses sequences for exemplary Cas9 adenosine or adenine and cytosine or cytidine base editors for base editing functions.

**Table 15. Sequences of exemplary Cas9 adenosine or adenine and cytosine or cytidine base editors**

Sequence ID No. (description)	Components of DNA cleavage assay
	<p><b>Amino Acid Sequence of Adenine Deaminase, TadA8.13m-nickase fused to the N-terminal of nickase <i>ScoCas9</i> (ABE-n<i>ScoCas9</i>, D10A mutant)</b></p> <p><i>M</i><b>PAAKRVKLD</b><u><i>GSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDP</i></u>  <i>TAHAEIMALRQ</i><u><i>GGLVMQNYRLYDATLYVTTFPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLM</i></u>  <i>DVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESAT</i>  <u><i>P</i></u><b>ESSG</b><u><i>PKKKRKV</i></u><b>G</b><u><i>GKPYSIGLA</i></u><b>I</b><u><i>GTNSVGWAVVTDDYKVP</i></u><b>AK</b><u><i>KMKVLGNTDKQSIKKNLLGALL</i></u>  <i>FDSGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRGEHH</i>  <i>PIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTD</i>  <i>VQALFKDFVEEYDKTIEESHLS</i><u><i>EITVDALSILTEKVS</i></u><b>K</b><u><i>SSRLENLIAHYPT</i></u><b>E</b><u><i>KKNTLFGNLIAL</i></u>  <i>SLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAILLSGILTV</i>  <i>DDNSTKAPLSASMVKRYEEHQDLKCLKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEF</i>  <i>YKYLKGILLKINGS</i><u><i>GDFLDKIDREDFLRKQRTFDNGSI</i></u><b>P</b><u><i>HQIHLQEMHAILRRQGEHY</i></u><b>P</b><u><i>FLKEN</i></u>  <i>QDKIEKILTFRIPYYVGPLARKGS</i><u><i>RFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDL</i></u>  <i>YLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVT</i><u><i>KDKLL</i></u>  <i>NYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKI</i><u><i>LDKSFLDDKANEKTI</i></u><b>E</b><u><i>DI</i></u><b>I</b><u><i>QTLTLFE</i></u>  <i>DREMIRQRLQKYS</i><u><i>DI</i></u><b>F</b><u><i>TKAQLKKLERRHYTGWGRLSYKLINGIRNKENKKT</i></u><b>I</b><u><i>LDYLIDDGYANR</i></u>  <i>NFMQLINDDALS</i><u><i>FKEE</i></u><b>I</b><u><i>ARAQI</i></u><b>I</b><u><i>DDVDDIANVVHDLPGSPA</i></u><b>I</b><u><i>KKGILQSVKIVDELVKVMGHNP</i></u></p>

ANIIIEMARENQTTDKGRRNSQQRLKLLQDSLKNLNDNPVNIKNVENQQQLQNDRLFLLYYIQNGKD  
 MYTGETLDINNLSQYDIDHIIIPQAFIKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKSFWSK  
 LLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRR  
 IRNVKIITLKSNIIVSNFRKEFELYKVREINDYHHAHDAYLNAVVGNALLLKYPQLEPEFVYGEY  
 PKYNSYRSRKSATEKFLFYSNIIIRFFKKEDIQTNEDGEIAWNKEKHIIKILRKVLSYPQVNIIVKK  
 TEEQTGGFSKESILPKGESDKLIIPRKTKNZYWDPKKGFFSDSPVVAISILVFADVEKGGKSKLR  
 KVQDMVGITIMEKKRFEKNPVDFLEQRGYRNVRLKIIKLPKYSLELENKRRRLLASAKELQK  
 GNELVIPQRFITLLYHSYRIEKDYEPEHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIE  
 MLFSKNKDAEVSSLAKSFIISLLTFTAAGAPAAFNFFGENIDRKRYTSVTECLNATLIHQSIITGL  
 YETRIDLSKLGEDGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ  
 ID NO: 20)

**Amino Acid Sequence of Cytidine Deaminase, ppAPOBEC1 fused to the N-terminus of nickase nScoCas9 (CBE-nScoCas9, D10A mutant)**

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWRSSG  
KNTTNHVEVNFIIKKFTSERRFHSSISCSITWFLSWSPWECSQAIREFLSQHPGVTLVIYVARL  
FWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYPPLWMMLYALELH  
CIILSLPPCLKISRRWQNHIAFFRLHLQONCHYQTIPPHILLATGLIHPSVTWRLKSGSSGGSS  
GSETPGTSESATPESGGSSGGSPKKKRKVGGKPYSIGLAIGTNSVGVAVVTDDYKVPKMKV  
 LGNTDKQSIKKNLLGALLFDSGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFF  
 QRLDDNFLVDEDEKRGHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMI  
 KFRGHFLYEGDLKAENTDVQALFKDFVEEYDKTIEESHLSITVDALSILTEKVSKSSRLENLI  
 AHYPTEKKNLTFGNLIALSLDLHPNFKTNFQLSSEDAKLQFSKDTYEEDLEGFLGEVGDYADLF  
 ASAKNLYDAILLSGILTVDDNSTKAPLSASMVKRYEEHQKDLKLLKDFIKVNAPDQYNAIFKDK  
 NKKGYASYIESGVKQDEFYKYLKGIILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQE  
 MHAILRRQGEHYPFLENQDKIEKILTRIPYVVGPLARKGSRFAWAEYKADEKITPWNFDDIL  
 DKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQE  
 IFDHVFKENRKVTKDKLLNYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSFDD  
 KANEKTIEDIIQTLTLFEDREMIRQLQKYSDIPTKAQLKKLERRHYTGWGRLSYKLINGIRNK  
 ENKKTILDYLIDGYANRNFQMLINDDALSFKEEIARAQIIDVDDIANVVHDLPGSPAIIKGI  
 LQSVKIVDELVKVMGHN PANIIIEMARENQTTDKGRRNSQQRLKLLQDSLKNLNDNPVNIKNVEN  
 QQQLQNDRLFLLYYIQNGKDMYTGETLDINNLSQYDIDHIIIPQAFIKDNSLDNRVLTRSDKNRGK  
 SDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHV  
 AQILDERFNTEFDGNKRRIRNVKIITLKSNIIVSNFRKEFELYKVREINDYHHAHDAYLNAVVG

ALLLKYPQLEPEFVYGEYPKYNSYRSRKSATEKFLFYSNILRFFKKEDIQTNEDGEIAWNKEKH  
 IKILRKVLSYPQVNIIVKKTTEEQTGGFSKESILPKGESDKLIPRKTKNSYWDPKKYGGFDSPVVA  
 YSILVFADVEKGGKSKLRKVQDMVGITIMEKKRFEKNPVDFLEQRGYRNVRLKIIKLPKYSLF  
 ELENKRRRLASAKELQKGNELVIPQRFTTLLYHSYRIEKDYEPHREYVEKHKDEFKELLEYI  
 SVFSRKVVLADNNLTKIEMLFSSKNKDAEVSSLAKSFISLLTFTAAGAPAAFNFFGENIDRKRYT  
 SVTECLNATLIHQSI TGLYETRIDL SKLGEDGKRPAATKAGQAKKKKSGSSGGSSGGSS TNLS  
DIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWA  
LVIQDSNGENKIKMLSGSSGGSSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESD  
ILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAYPYDVPDYAYP  
 YDVPDYA (SEQ ID NO: 21)

**Amino Acid Sequence of Adenine Deaminase, TadA8.13m fused to the N-terminal of nickase SirCas9 (ABE-nSirCas9, D14A mutant).**

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDP  
TAHAEIMALRQGGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLM  
DVLHHPGMNRHVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESAT  
PESSGPKKKRKVGAKNKDIRYSIGLAIGTNSVGWAVMDEHYELLKGNHMHMWSRLFDAEPAA  
 TRRASRSIRRRYNKRREIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQKNLGNDYKDNY  
 NLFIEKDFNDKTYDYKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQSFQVY  
 EDIEEKLDNTLKKFMSLNDLDNLFVDNDINSMITVLSKIYQRSKKADDLLKIMNPTKEERAAYK  
 EFTKALVGLKFNVSKMILAQEVKKDDKDIELDFSNVDYDSTVDGLQAEUGEYIEFIEMLHSINS  
 WVLEQDILGNNSTISAAMVERYEEHKNDLRVLKVKVIREELPKYNEVFREDNPKLHNYLGYIKY  
 PKNTPVEEFYEYIKRLLAKVDTGEAREILERIDLEKFMKQNSRTNGSIPYQMOKDEMIQIIDN  
 QSVYYPQLKENREKLISILEFRIPYFGPLNTHSEFAWIKKFEDKQKERILPWNYDQIVDIDAT  
 AEGFIERMQNTGTYPDPKPVMAKNSLTVSKFEVLNELNKIRINGKLI PVETKKELLSDFMKNK  
 TITDKKLKDWLVTHQYYDTNEELKIEGYQKDLQFSTSLAPWIDFTKIFGEINASNYQLIEKIIY  
 DISIFEDKKILKRRLKKVYQLDDLLVDKILKLNVTGWSRLSEKLLTGIKSKNSKETILSILENS  
 NMNLMEIINDESLGFKQIIIEESNKKDIEGPFYRYDEVKKLAGSPA I KRGIWQALLVVQEITKFMK  
 HEP SHIYIEFAREEQEKV RTESRIAKLQKIYKDLNLQTKEDQLVYESLKKEDAKKKIDTDALYL  
 YYLQMGKSMYS GKPLDIDKLSTYHIDHILPRSLIKDDSLDNRVVLVLPKENEWKLDSETVPFEIR  
 NKMMGFQK L HENGLMSNKKFFSLIRTD FNEKDKKRFINRQLVETRQIIKNVAVIINDHYTNTN  
 VVTVRAELSHQFRERYKIYKNRDLNDLHHAHDAYIACILGQFIHQNFGNMDVNMIIYGQYKKNYK  
 KDVQEHNNYGFILNSMNHIHFNDDNSVIWDPSYIGKIKSCFCYKD VYVTKKLEQNDAKLFDLTI  
 LPSDKNSENGVTKAKIPVNKYRKDVNKYGGFSGDAPIMLAIEADKGGKHVRQVIAFPPLRLKNYN

DEERIKFIEKEKNLKNVKILTEVKKNQLILINHQYFFITGTNELVNATQLKLSAKNTKNLFNLV  
 DANKHNKLESIDDANFNEVIQELICKLQEPYISRYNSIGKEFEDSYEKINAVTKQDKLYIIIEYL  
 IAIMSAKATQGYIKPELAREIGTNGKNKGRIKSFTIDLNKTTFISTSVTGLFSKKYKLGKRPA  
TKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 6)

**Amino Acid Sequence of Adenine Deaminase, TadA8.13m fused to the N-terminal of nickase VapCas9 (ABE-nVapCas9, D38A mutant)**

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAFLVLNNRVIGEGWNRAIGLHDP  
TAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLM  
DVLHHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESAT  
PESSGPKKKRKVGSIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLAIGTSSVGWAVTNKA  
 YELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRLLKLEELFADAMAQVDPFFMR  
 LRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHHI  
 LKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFCNSAISSIGQILMEAGKTKSDKAKAI  
 EHLVDTYIATDVTVDTSKTKQKQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEEDLKKLQI  
 TGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPGLTISESKVKAFNKHKDDLAILK  
 SLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEYIL  
 SQIELQIILLPLQRIKDNQVIPIYQLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKMLEFRI  
 PYYVGPLNTHHNVNDNGGFAWAVRKASGRVTPWNFDDKI DREKSAAAFIKNLTNKCTYLLGEDVL  
 PKSSLLYSEFMLLNELNVRIDGKPLEKVVEHLIEAVFKQDHKKMTKNRIEQFLKDNQYISET  
 HKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGESKMLRETLRKKFASCL  
 DDEAIKKLTCLRDRDWRSLSQKLLNGIEGCDKAGDGPETIIILMRNFSYNLMELLGDKFSFME  
 RIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVDEVAHIKKALPARI FVEVTRS  
 NEKKKDSRQRLSDLYAAIKKDDVLLNGLNNEIFGELKSSLAKYDDAALRSKKLYLYYTQMG  
 CAYTGEIIELSLLNTDNYDIDHIYPRSLTKDSDFDNLVLCRRTANAQKSDAYPISEEIQKTQKP  
 FWTFLKQOGLISERKYERLTRITPLTADDLSGFARQLVETNQSVAATTLRRLYPGVDVVFV  
 KAENVTDVFRHDNFIKVRSLNHHHHAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAK  
 MFNYDVNCTNAKDGKAWDVKTSMDTVKKMMSNDVVRVTKRLLLEQTGALADATIYKATVAGKAKD  
 GAYIGMKTSSVFADVSKYGGMTKIKNAYSIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLI  
 NYVASIIPQAKDISIIYGKLCINQLVKVNGFYLLGGKTNKFCIDNAIQVIVSNEWIPLYKVL  
 EKFNMRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLVSKLKMPIYQKMKGNKAAE  
 LSEKGYGLFKMSLEEQSIHLIELLNLLTNQKTTFEVKPLGITASRSTVSGSKISNQDEFKVINE  
 SITGLYSNEVTIVGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ  
 ID NO: 10)



**Amino Acid Sequence of Adenine Deaminase, TadA8.13m fused to the C-terminal of nickase *VapCas9* (n*VapCas9*-ABE8, D38A mutant)**

**MPK**KKRKVSIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGL**A**IGTSSVGVAVTNKAYELLK  
FRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRKLLLEELFADAMAQVDPTFFMRLRESK  
YHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKFLAVHHILKYRG  
NFLYEGATFDSNASTLDDVIKQALENITFNCFCNSAIISSIGQILMEAGKTKSDKAKAIEHLVD  
TYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEEDLKKLQITGDTY  
DDKRDELAKAWSDEIYIIDCKSVYDAIILLISIKEPGLTISESKVKAFNKHKDDLAILKSLKLS  
DRSIYNTMFKVDEKGLHNYVHYIKQGRTEETS CNREDFYKYTKKIVEGLSDSKDKEYILSQIEL  
QILLPLQRIKDNQVI PYQLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKMLEFRIPYYVG  
PLNTHHNVNDGGFAVAVRKASGRVTPWNFDDKIDREKSAAAFIKNLTNKCTYLLGEDVLPKSSL  
LYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTKNRIEQFLKDNGYISETHKHEI  
TGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGESKKMLRETLRKKFASCLDDEAI  
KKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGTPETIIILMRNFSYNLMELLGDKFSFMERIQEI  
NAKLTGQIVNPHDIIDDLALSPAVKRAVWQALRIVDEVAHIKKALPARI FVEVTRSNKNEKKK  
KDSRQKRLSDLYAAIKKDDVLLNGLNNEIFGELKSSLAKYDDAALRSKKLYLYTQMGRCAYTG  
EIIELSLNNTDNYDIDHIYPRSLTKDDSFNVLVCKRTANAQKSDAYPISEEIQKTQKPFWTF  
KQQGLISERKYERLTRITPLTADDLSGFIARQLVETNQSVAATTLRRLYPGVDVVFVKAENV  
TDFRHDNFIKVRSLNHHHAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYD  
VNCTNAKD GKAWDVKTSM DTVKMMDSNDVRVTKRLL EQT GALADATIYKATVAGKAKDGAYIG  
MKTSSVFADVSKYGGMTKIKNAYSIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVAS  
IIPQAKDISIIYGKLCINQLVKVNGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNN  
MRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLVSKLKMPIYQKMKGNKAAEELSEKG  
YGLFKMSLEEQSIHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKI SNQDEFKVINESITGL  
YSNEVTIV**KRPAATKKAGQAKKKK**SGSETPGTSESATPESSG**SEVEFSHEYWMRHALTLAKRAR**  
**DEREVPVGAVLVLNNRVIGEGWNRAIGLHDP**TAHAE**IMALRQGGLVMQNYRLYDATLYVTFEPC**  
**VMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRM**  
**PRRVFNAQKKAQSSTDPAAKRVKLDGS**YPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO:  
11)

**Amino Acid Sequence of Cytidine Deaminase, ppAPOBEC1 fused to the N-terminal of nickase *VapCas9* (CBE-n*VapCas9*, D38A mutant)**

MPAAKRVKLD TSEKGPSTGDPTLRRRIESWEFDVIFYDPRELKRETCLLYEIKWGMSRKIWRSSG  
KNTTNHVEVNFIIKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIYVARL  
FWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMMLYALELH  
CIILSLPPCLKISRRWQNHLAFFRLHLQONCHYQTIPPHILLATGLIHPSVTWRLKSGSSGGSS  
GSETPGTSESATPESSSGSSGGSPKKKRKVGSIINFQRRGLMETQASNQLISSHLKGYPIKDYF  
VGLAIGTSSVGWAVTNKAYELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRKLL  
EELFADAMAQVDPTFFMRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSEL  
MKSGETDIRKFLFLAVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFCNSAIISSI  
GQILMEAGKTKSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLI  
DLFGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLISIKEPGLTIS  
ESKVKA FNKHKDDLAILKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKY  
TKKIVEGLSDSKDKEYILSQIELQILLPLQRIKDNQVI PYQLHLEELKAILAKCGPKFPFLNEV  
ADGFSVAEKLIKMLEFRIPYYVGPLNTHHNVNDNGGFAWAVRKASGRVT PWNFDDKIDREKSAAA  
FIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKM  
TKNRIEQFLKDNQYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFG  
ESKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGT PETIIILMR  
NFSYNLMELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVDEVAH  
IKKALPARI FVEVTRSNKNEKKKKDSRQKRLSDLYAAIKKDDVLLNGLNNEIFGELKSSLAKYD  
DAALRSKKLYLYYTQMGRCAYTGEIIELSLLNTDNYDIDHIYPRSLTKDDSFDNLVLCKRTANA  
QKSDAYPISEEQKTQKPFWTFLKQQGLISERKYERLTRITPLTADDLSGFARQLVETNQSVK  
AATLLRRLYPGVDVVFVKAENVTDFRHDNFIKVRSLNHHHAKDAYLNIVVGNVYHERFTRN  
FRAFFKKNGANRTYNLAKMFNYDVNCTNAKDGAWDVKTSMDTVKKMMSNDVVRVTKRLLLEQTG  
ALADATIYKATVAGKAKDGAYIGMKTSSVFADVSKYGGMTKIKNAYSIIVQYTGKKGEVIKEI  
VPLPIYL TNRNTTDQDLINYVASIIPQAKDISIIYGKLCINQLVKVNGFYYYLGGKTNSKFCID  
NAIQVIVSNEWIPYLKVLEKFNMRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLV  
SKLKMPIYQKMKGNAEELSEKGYGLFKKMSLEEQSIHLIELLNLLTNQKTTFEVKPLGITASR  
STVGSKISNQDEFKVINESITGLYSNEVTIVGKRPAATKKAGQAKKKKGSSGGSSGGSSGGSS TNLS  
DIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWA  
LVIQDSNGENKIKMLSGGSSGGSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESD  
ILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAYPYDVPDYAY  
(SEQ ID NO: 12)

**Amino Acid Sequence of Adenine Deaminase, TadA8.13m fused to the N-terminal of nickase EpeCas9 (ABE-nEpeCas9, D12A mutant)**

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDP  
TAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLM  
DVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESAT  
PESSGPKKKRKVGTKVKDYYIGLAIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAEER  
RGQRGARRRLDRKKERLSLLQDFFAAEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKD  
FKDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFI FEGQKFDTKSSFENSLN  
ELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQKLV  
DLFENPEDFDDSAIKSVDFSTTSFDDKYS DYELALGDKIALVNI LKEIYDSSILENLLKEADKS  
KDG NKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEK VNDNYVS YTKSSISNNKRV  
KANKFTDQEAFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNGVIP  
YQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRI PYVGPLNPNSKYAWIKKQKD  
SEITPWNFKDVVDLDS SREEFIDSLIGRCTYLKDEKVL PKASLLYNEYMVLNELLNKLNDLPI  
TEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGDFKQGLNSYNDFKAI VGDK  
VSDDYRDKIEEIIKLVLYGDDKSYLQKKIKAGYGYFTDSEIKKMAGLNYKDWGRLSKLLT  
GLEGANKITGERGSI IHFMREYNLNLME LMSASFTFTEEIQKLN PVDDRKLSYEMVDELYLSPS  
VKRMLWQSLRIVDEIKNIMGTDSKKI FIEMARGKEEVKARKESRKNQLLK FYKDGKKA FISEIG  
EERYSYLLSEIEGEEENKFRWDNLYLYTQLGRCMYSLEPIDISELSSKNIYDQDHIYPKSKIY  
DDSIENRVLVKKDLNSKKGNSYPI PDEILNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDE  
LVQFISRQIVETRQATKETANLLKTICKNSEIVYSKAENASRFRQEFDIVKCRAVNDLHHMHDA  
YINIIVGNVYNTKFTKDP MNFVKKQEKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIK  
RIRKELEGTNYRFRMNYIESGALFNATLQRKNKGSRPLKDKGPKSSIEKYGGYTNINKACFAV  
LDIKSKNKIERKLPVEREIIYAKQKNDKKLSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKID  
GSYYFITGGSDKTLELRSALQLILPKKNEWAIKQIDKSENDYLTIERIQDLTEELVYNTFDII  
VNFKTSVFKKSFLNLFQDDKIENIDFKFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKS  
DYGRLLSSKTNNIGNYQEFKIINQSITGLFENEVDLLKLGKRPAATKKAGQAKKKGSYPYDVPD  
YAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 16)

**Amino Acid Sequence of Adenine Deaminase, TadA8.13m fused to the C-terminal of nickase EpeCas9 (nEpeCas9-ABE8, D12A mutant)**

MPKKRKVTVKDYYIGLAIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAEERRGQRG  
ARRRLDRKKERLSLLQDFFAAEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKDFKDN  
FHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFI FEGQKFDTKSSFENSLNELKVH  
LNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQKLVDLFEN  
PEDFDDSAIKSVDFSTTSFDDKYS DYELALGDKIALVNI LKEIYDSSILENLLKEADKSKDG NK

YISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNNDNYVSYTKSSISNNKRVKANKF  
 TDQEAFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNGVIPYQLKL  
 MELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRIPIYYVGPLNPNISKYAWIKKQKDEITP  
 WNFKDVVDLDSREEFIDSLIGRCTYLKDEKVLPKASLLYNEYMVLNELNNLKLNDLPITEEMK  
 KKI F D Q L F K T R K K V T L K A V A N L L K K E F N I N G E I L L S G T D G D F K Q G L N S Y N D F K A I V G D K V D S D D  
 Y R D K I E E I I K L I V L Y G D D K S Y L Q K K I K A G Y G K Y F T D S E I K K M A G L N Y K D W G R L S K K L L T G L E G A  
 N K I T G E R G S I I H F M R E Y N L N M E L M S A S F T F T E E I Q K L N P V D D R K L S Y E M V D E L Y L S P S V K R M L  
 W Q S L R I V D E I K N I M G T D S K K I F I E M A R G K E E V K A R K E S R K N Q L L K F Y K D G K K A F I S E I G E E R Y S  
 Y L L S E I E G E E E N K F R W D N L Y L Y T Q L G R C M Y S L E P I D I S E L S S K N I Y D Q D H I Y P K S K I Y D D S I E  
 N R V L V K K D L N S K K G N S Y P I P D E I L N K N C Y A Y W K I L Y D K G L I G Q K K Y T R L T R R T G F T D D E L V Q F I  
 S R Q I V E T R Q A T K E T A N L L K T I C K N S E I V Y S K A E N A S R F R Q E F D I V K C R A V N D L H H M H D A Y I N I I  
 V G N V Y N T K F T K D P M N F V K K Q E K A R S Y N L E N M F K Y D V K R G G Y T A W I A D D E K G T V K N A S I K R I R K E  
 L E G T N Y R F T R M N Y I E S G A L F N A T L Q R K N K G S R P L K D K G P K S S I E K Y G G Y T N I N K A C F A V L D I K S  
 K N K I E R K L M P V E R E I Y A K Q K N D K K L S D E I F S K Y L K D R F G I E D Y R V V Y P V V K M R T L L K I D G S Y Y F  
 I T G G S D K T L E L R S A L Q L I L P K K N E W A I K Q I D K S S E N D Y L T I E R I Q D L T E E L V Y N T F D I I V N K F K  
 T S V F K K S F L N L F Q D D K I E N I D F K F K S M D F K E K C K T L L M L V K A I R A S G V R Q D L K S I D L K S D Y G R L  
 S S K T N N I G N Y Q E F K I I N Q S I T G L F E N E V D L L K L KRPAATKKAGQAKKKKSGSETPGTSESATPE  
SSGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDPTAHAEIMA  
LRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGM  
NHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPYDVPDYAYPY  
 DVPDYAYPYDVPDYA (SEQ ID NO: 17)

**Amino Acid Sequence of Adenine Deaminase, TadA8.13m fused to the C-terminal of nickase EpeCas9 (nEpeCas9-ABE8, D12A mutant)**

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWRSSG  
KNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWSPWECSQAIREFLSQHPGVTLVIYVARL  
FWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYPPLWMLYALELH  
CIILSLPPCLKISRWRQNHLAFFRLHLQNCYQTIPPHILLATGLIHPSVTWRLKSGGSSGGSS  
GSETPGTSESATPESSGGSSGGSPKKKRKVGTKVKDYIYIGLAIGTSSVGWAVTDEAYNVLKFNS  
 KKMWGVRLFDDAKTAEERRGQRGARRRLDRKKERLSLLQDFFAEVAVKVDPNFFLRLDNSDLYM  
 EDKDQKLKSKYTLFNDKDFKDNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHF  
 IFEGQKFDTKSSFENSLNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDT  
 KFLKAVSAIMIGSSQKLVDLFENPEDFDDSAIKSVDFSTTSFDDKYSYELALGDKIALVNILK  
 EIYDSSILENLLKEADKSKDGNKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKV

NDNYVSYTKSSISNNKRVKANKFTDQEAIFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDM  
 EFKNFMPKIKSSDNGVIPPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRIPYY  
 VGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLDS SREEFIDSLIGRCTYLKDEKVLPKASLLYN  
 EYMVLNELNKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTGDG  
 FKQGLNSYNDFKAIIVGDKVSDDDYRDKIEEIIKLVLYGDDKSYLQKKIKAGYGKYFTDSEIKK  
 MAGLNYKDWGRLSKLLTGLEGANKITGERGSI IHFMREYNLNLMELEMSASFTFTEEIQKLNPV  
 DDRKLSYEMVDELYLSPSVKRWLWQSLRIVDEIKNIMGTDSKKIFIE MARGKEEVKARKESRKN  
 QLLKFYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLVLYYTQLGRCMYSLEPIDISEL  
 SSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPIPDEILNKNCYAYWKILYDKGLI  
 GQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTI CKNSEIVYSKAENASRFRQE  
 FDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKDP MNFVKKQEKARSYNLENMFKYDVKRGGY  
 TAWIADDEKGTVKNASIKRIRKELEGTNYRFRMNYIESGALFNATLQRKNKGSRPLKDKGPKS  
 SIEKYGGYTNINKACFAVLDIKSKNKIERKLPVERE IYAKQKNDKLSDEIFSKYLKDRFGIE  
 DYRVVYPVVKMRTLLKIDGSYYFITGGSDKTLELRSALQLILPKKNEWAIKQIDKSSENDYLT I  
 ERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNFLQDDKIENIDFKFKSMDFKKCKTLLMLVK  
 AIRASGVRQDLKSIDLKS DYGRLS SKTNNIGNYQEFKIINQSITGLFENEVDLLKLGKRPAA TK  
 KAGQAKKKKGGSSGGSGGSGGS TNLSDIIEKETGKQLVIQESI LMLPEEVEEVIGNKPESDILVH  
TAYDESTDENVMLLTS DAPEYKPWALVIQDSNGENKIKMLS GSGSGSGSTNLSDIIEKETGKQ  
LVIQESI LMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTS DAPEYKPWALVIQDSNGEN  
KIKMLYPYDVDPDYAYPYDVDPDYAY (SEQ ID NO: 18)

**Amino Acid Sequence of Adenine Deaminase, TadA8.13m-nickase fused to the N-terminal of nickase *LfeCas9* (ABE-nLfeCas9, D9A mutant)**

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGA VLVLNNRVIGEGWNRAIGLHDPT  
AHAEIMALRQGGLVMQNYRLYDATLYVTTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDV  
LHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSS TDGSSGSETPGTSESATPES  
SGPKKKRKVGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTA AERTFRTR  
 RRLKRRKWRLHYLDEIFAPHLQEV DENFLRRLKQSNIHPE DPAKNQAFIGKLLFPDLLKKNERGY  
 PTLIKMRDELPVEQRAHYPVTNIYKLRAMINEDRQFDLRE VYLAVHHIVKYRGHFLNNASVDKF  
 KVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLD RQKAVAKLLEVKVA  
 DKEETKRNKQIATAMSKLVLYGKADFATVAMANGNEWKIDLSSETSEDEIEKFREELSDA QNDIL  
 TEITSLFSQIMLNEIVPNGMSISESMMDRYW THERQLAEVKEYLATQPASARKEFDQVY NKYIGQ  
 APKEKGF DLEKGLKKILSKKENWKEIDELLKAGDFLPKQRTSANGVI PHQM HQQELDR IIEKQAK  
 YYPWLATENPATGERDRHQAKYELDQLVSFRIPYVVGPLVTP EVQKATSGAKFAWAKRKEDGEIT

PWNLWDKIDRAESAEAFIKRMTVKDITYLLNEDVLPANSLLYQKYNVLNELNVRVNGRRLSVGIK  
 QDIYTELFKKKKTVKAGDVASLVMKTRGVNKPVSVEGLSDPKKFNSNLATYLDLKSIVGDKVDDN  
 RYQMDLENIIEWRSVFEDGEIFADKLTEVEWLTDEQRSALVKKRYKGGWRLSKKLLTGIVDENGQ  
 RIIDLWNTDQNFMQIVNQPVFKEQIDQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQV  
 VRVVEDIVKAVGNAPKSSISIEFARNEGKGEITRSRRTQLQKLFEDQAHELKVDTSLTEELEKAP  
 DLSDRYYFYFTQGGKDMYTGDPIINFDEISTKYDIDHILPQS FVKDDSLDNRVLSRAENNKSDR  
 VPAKLYAAKMKPYWNQLLKQGLITQRKFENLTMDVDQTIKYRSLGFVKRQLVETRQVIKLTANIL  
 GSMYQEAGTDIIEETRAGLTKQLREEFDLPKVREVNDYHHAVDAYLTTFAGQYLNRRYPKLSRFFV  
 YGEYMKFKHGS DLKLRNFNFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLIS  
 NETYEETGKYFDASHESLKLKNQNKSKLVPLKNKLQPEYYGAYTGITQGYMVILKLLDKKG  
 GFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAKSFVVVPRVLKGTFLVDGEEKFILS  
 SYRYKVNATQLILPVSDIKLIQDNFKALKKLNEMQTKKLEIYDNILRQVDKYKLYDINKFRA  
 KLHDGRSKFVELDDFGQDASKEKVIKILRGLHFSGDLQNLKEIGFGTTPLGQFQVSEAGIRLSN  
 TAFIIFKSP TGLFNRKLYLKNLGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVP  
 DYA (SEQ ID NO: 88)

**Amino Acid Sequence of Adenine Deaminase, TadA8.13m-nickase fused to the C-terminal of nickase *LfeCas9* (nLfeCas9-ABE, D9A mutant)**

MPKKKRKVGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTA AERRTFRTTR  
 RRLKRRKWRLHYLDEIFAPHLQEV DENFLRRLKQSNIHPE DPAKNQAFIGKLLFPDLLKKNERG  
 YPTLIKMRDEL PVEQRAHY PVTNIYKLREAMINEDRQFDLREVYLAVHHIVKYRGHFLNNASVD  
 KFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLD RQKAVAKLLEV  
 KVADKEETKRNKQIATAMSKLV LGYKADFATVAMANGNEWKIDLSSETSEDEIEKFREE LSDAQ  
 NDILTEITSLFSQIMLNEIVPNGMSISESMMDRYW THERQLAEVKEYLATQPASARKEFDQVYN  
 KYIGQAPKEKGF DLEKGLKKILSKKENWKEIDELKAGDFLPKQRTSANGVIPHQM HQE LDRI  
 IEKQAKYYPW LATENPATGERDRHQAKYELDQLVSFRIPYYVGPLVTP EVQKATSGAKFAWAKR  
 KEDGEITPWNLWDKIDRAESAEAFIKRMTVKDITYLLNEDVLPANSLLYQKYNVLNELNVRVNG  
 RRLSVGIKQDIYTELFKKKKTVKAGDVASLVMKTRGVNKPVSVEGLSDPKKFNSNLATYLDLKS  
 IVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVEWLTDEQRSALVKKRYKGGWRLSKKLL  
 LTGIVDENGQRIIDLWNTDQNFMQIVNQPVFKEQIDQLNQKAITNDGMTLRERVESVLDDAYT  
 SPQNKKAIWQVVRVVEDIVKAVGNAPKSSISIEFARNEGKGEITRSRRTQLQKLFEDQAHELK  
 DTSLTEELEKAPDLSDRYYFYFTQGGKDMYTGDPIINFDEISTKYDIDHILPQS FVKDDSLDNRV  
 LVSRAENNKSDRVP AKLYAAKMKPYWNQLLKQGLITQRKFENLTMDVDQTIKYRSLGFVKRQL

VETRQVIKLTANILGSMYQEAGTDIIETRAGLTKQLREEFDLPKVREVNDYHHAVDAYLTTFAG  
 QYLNRRYPKLRSEFFVYGEYMKFKHGSDLKLRNFNFHELMEGDKSQGKVVDDQQTGELITTRDEV  
 ADYFDWVINLKVMLISNETYEETGKYFDASHSSSLYLKNQNKSKLVVPLKNKLOPEYYGAYT  
 GITQGYMVIKLLDCKGGFGVYRIPRYAADI LNKCHDEVAYRNKIAEIISSDPRAPKSEFVVVP  
 RVLKGTFLVDGEEKFILSSYRYKVNATQLILPVS DIKLIQDNFKALKKLNVMQTKKLEIYDN  
 I LRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEKVI IKILRGLHFGSDLQNLKEIG  
 FGTTP LGQFQVSEAGIRLSNTAFIIFKSP TGLFNRKLYLKNL **KRPAATKKAGQAKKKKSGSETP**  
GTSESATPESSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLN RVIGEGWNRAIGLHD  
PTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSL  
MDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTD **PAAKRVKLDGSPY**  
 DVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 89)

**Amino Acid Sequence of Cytidine Deaminase, ppAPOBEC1 fused to the N-terminal of nickase LfeCas9 (CBE-nLfeCas9, D9A mutant)**

**MPAAKRVKLD** TSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWRSSG  
KNTTNHVEVNF IKKFTSERRFHSSISCSITWFLSWSPWECSQAIREFLSQHPGVTLVIYVARL  
FWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMMLYALELH  
CIILSLPPCLKISRRWQNH LAFFRLHLQ NCHYQTIPPHILLATGLIHPSVTWRLKSGSSGGSS  
GSETPGTSESATPESSEGGSSGGSPKKKRKV GKEYHIGLAIGTSSIGWAVTDSQFKLMRIKPKTA  
 IGVRLFEEGKTAAERRTFRTRRRRLKRRKWRLHYLDEIFAPHLQEV DENFLRRLKQSNIHPEDP  
 AKNQAFIGKLLFPDLLKKNRGYPTLIKMRDEL PVEQRAHYPVTNIYKLREAMINEDRQFDLRE  
 VYLAVHHIVKYRGHFLNNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQ  
 LLLDTKMRKLD RQKAVAKLLEVKVADKEETKRNKQIATAMSKLV LGYKADFATVAMANGNEWKI  
 DLSSETSEDEIEKFREELSDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWTHERQLA  
 EVKEYLATQPASARKEFDQVYNKYIGQAPKEKGF DLEKGLKKILSKKENWKEIDELLKAGDFLP  
 KQRTSANGVI PHQM HQE LDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRI PYYV  
 GPLVTP EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKD TYLLNEDVLP  
 ANSLLYQKYNVLNELNVRVNGRRLSVGIKQDIYTELFK KKT VKAGDVASLVM AKTRGVNKPS  
 VEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVEWLT  
 DEQRSALVKRYKGWGRLSK KLLTGIVDENGQRIIDL MWNTDQNF MQIVNQPVFKEQIDQLNQK  
 AITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSI SIEFARNEGKGE  
 ITRSRRTQLQKLFEDQAHEL VKDTSLTEELEKAPDLSDRYYFYFTQGGKDMYTGDPINFDEIST  
 KYDIDHILPQS FVKDDSLDNRVLVSRAENNKSDRVPAKLYAAKMPYWNQLLKQGLITQRKFE  
 NLTMDVDQTIKYRSLGFVQRQLVETRQVIKLTANILGSMYQEAGTDIIETRAGLTKQLREEFDL

PKVREVNDYHHAVDAYLTT FAGQYLNRRYPKLRSEFFVYGEYMKFKHGS DLKLRNFNFFHELMEG  
 DKSQGKVVDDQQTGELITTRDEVADYFDWVINLKVMLISNETYEETGKYFDASHESSSLYLKNQN  
 KKSCLVPLKNKLQPEYYGAYTGITQGYMVILKLLDKKGGFGVYRIPRYAADILNKCHDEVAYR  
 NKIAEIISSDPAPKSEFVVVPRVLKGTFLVDGEEKFILSSYRYKVNATQLILPVSDIKLIQDN  
 FKALKKLNVMQTKKLEIYDNILRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEK  
 VIKILRGLHFSDLQNLKEIGFGTTPGQFQVSEAGIRLSNTAFIIFKSP TGLFNRKLYLKNL  
GKRPAATKKAGQAKKKKGSSGGSGGSGGS **TNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNK**  
**PESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGSGGSGGS TNLSDI**  
**IEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALV**  
**IQDSNGENKIKML**YPYDVDPDYAYPYDVDPDYAY (SEQ ID NO: 90)

**Amino Acid Sequence of Adenine Deaminase, TadA8.13m-nickase fused to the N-terminal of nickase PmaCas9 (ABE-PmaCas9, D12A mutant)**

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLHDP  
TAHAEIMALRQGGLVMQNYRLYDATLYVTTFPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLM  
DVLHHPGMNRHVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESAT  
PESSGPKKKRKVGEKKTNYTIGLAIGTDSVGWAVVKDDLELVKKRMKVLGNTETNYIKKNLWGS  
 LLFESGQTAKDRRLKRVARRRYERRRNLTELQKIFAPAIDEVDFFRNLNESFLVPEDKAFS  
 KNPIFGTLGEDKTYKYTYPTIYHLRQHLADSEKADVRLIYLALAHMIKYRGHFLIEGKLDTEH  
 IAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDIGRANKQS  
 PMMAFITLIVGKAKFKAAFNLEEEISLNLTDSDYDENLEILLNTIGSDFADLFDHAQRVYNAV  
 ELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKDLKG  
 YAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVHLAEMEI  
 LDRQENYYPWLKENREKIIISLLTFRIPIYYVGPLADGQSEFAWLERKSDEKIKPWNFSDVVDLDR  
 SAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIAYLDERQKRMNLSVEKKEIFET  
 LFKRKRKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMDDPDNEDLM  
 EEIVKILTVFEDRKMRKQLEKYKERLSPEQIKELAKKHGTGWGRLSKLLVGI RDKETQKTIL  
 DYLVEDDNHSGGRQHLNRNLMQLINDDRLSFKKTAELQMI DPSADLYAQVQEIAGSPA KKG I  
 LLGLKIVDEIIRVMGEKPENIVIEMARENQTTARGKALSKRREAKIKEGLAALGSSLLKENLPG  
 NADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHIIPQSFTVDNSLDNLVLTNS SQNRGN  
 KKDDVPSLEVNRQLAYWRS LK DAGLMTQRKFDNLTKAMRGGLTDKDRERFIQRQLVETRQITK  
 NVAKLLDMRLNDKKDEAGNKIRETNIVLLKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAAI  
 AINLLALYPYMADDFVYGEFRYKPKPQAEKATYEKLRQWNLIKRFGEKQLFTPDHEDCWNKERD



IKTIKKVMGYRQVNVVKKAEERTGMLFKETINGKTNKGSRIPIKKDLDP SKYGGYIEEKMAYYA  
 VISYEDK K K K K P G K T I V G I S I M D K K E F E Y D S I S Y L G K L G F S N P V V Q I I L K N Y S L I A Y P D G R R R Y I  
 T G A T K T T K G K V E L Q K A N Q I A M E Q D L V N F I Y H L K N Y D E I S H P E S Y A F V Q S H T D Y F D R L F D S I E H Y  
 T R R F L D A E T N I N R L R R I Y E E E K K K D P V D I E A L V A S F I E L L K L T S A G A P A D F I F M G E A I S R R R Y N  
 S M T G L F D G Q V I Y Q S L T G L Y E T R M R F E D K R P A A T K K A G Q A K K K K G S Y P Y D V P D Y A Y P Y D V P D Y A  
Y P Y D V P D Y A (SEQ ID NO: 91)

**Amino Acid Sequence of Adenine Deaminase, TadA8.13m-nickase fused to the C-terminal of nickase *PmaCas9* (n*PmaCas9*-ABE, D12A mutant)**

M P K K K R K V E K K T N Y T I G L A I G T D S V G W A V V K D D L E L V K K R M K V L G N T E T N Y I K K N L W G S L L F E S  
 G Q T A K D R R L K R V A R R R Y E R R R N R L T E L Q K I F A P A I D E V D E N F F F R L N E S F L V P E D K A F S K N P I F  
 G T L G E D K T Y Y K T Y P T I Y H L R Q H L A D S E E K A D V R L I Y L A L A H M I K Y R G H F L I E G K L D T E H I A I N E  
 N L E Q F F E S Y N A L F S E E P I E L R K E E L I A I E N I L R E K N S R T V K E K R I T S F L K D I G R A N K Q S P M M A F  
 I T L I V G K K A K F K A A F N L E E E I S L N L T D D S Y D E N L E I L L N T I G S D F A D L F D H A Q R V Y N A V E L A G I  
 L S G D V K N T H A K L S A Q M V A M Y E R H K E Q L K E Y K S F I K A N L P D Q Y D M T F V A P K D A Q K K D L K G Y A G Y I  
 D G N M S Q D S F Y K F V K D Q L K E V P G S E K F L D S I E K E D F L R K Q R S F Y N G V I P N Q V H L A E M E A I L D R Q E  
 N Y Y P W L K E N R E K I I S L L T F R I P Y Y V G P L A D G Q S E F A W L E R K S D E K I K P W N F S D V V D L D R S A E K F  
 I E Q L I G R D T Y L P D E Y V L P K K S L I Y Q K Y M V F N E L T K I A Y L D E R Q K R M N L S S V E K K E I F E T L F K K R  
 S K V T E K Q L V K F F E N Y L Q I D N P T I F G I E D A F N A D Y S T Y V E L A K V P G M K S M M D D P D N E D L M E E I V K  
 I L T V F E D R K M R R K Q L E K Y K E R L S P E Q I K E L A K K H Y T G W G R L S K K L L V G I R D K E T Q K T I L D Y L V E  
 D D N H S G G R Q H L N R N L M Q L I N D D R L S F K K T I A E L Q M I D P S A D L Y A Q V Q E I A G S P A I K K G I L L G L K  
 I V D E I I R V M G E K P E N I V I E M A R E N Q T T A R G K A L S K R R E A K I K E G L A A L G S S L L K E N L P G N A D L S  
 Q R K I Y L Y Y T Q N G K D I Y L D E P L D F D R L S Q Y D E D H I I P Q S F T V D N S L D N L V L T N S S Q N R G N K K D D V  
 P S L E V V N R Q L A Y W R S L K D A G L M T Q R K F D N L T K A M R G G L T D K D R E R F I Q R Q L V E T R Q I T K N V A K L  
 L D M R L N D K K D E A G N K I R E T N I V L L K S A M A S E F R K M F R L Y K V R E L N D Y H H A H D A Y L N A A I A I N L L  
 A L Y P Y M A D D F V Y G E F R Y K K K P Q A E K A T Y E K L R Q W N L I K R F G E K Q L F T P D H E D C W N K E R D I K T I K  
 K V M G Y R Q V N V V K K A E E R T G M L F K E T I N G K T N K G S R I P I K K D L D P S K Y G G Y I E E K M A Y Y A V I S Y E  
 D K K K K P G K T I V G I S I M D K K E F E Y D S I S Y L G K L G F S N P V V Q I I L K N Y S L I A Y P D G R R R Y I T G A T K  
 T T K G K V E L Q K A N Q I A M E Q D L V N F I Y H L K N Y D E I S H P E S Y A F V Q S H T D Y F D R L F D S I E H Y T R R F L  
 D A E T N I N R L R R I Y E E E K K K D P V D I E A L V A S F I E L L K L T S A G A P A D F I F M G E A I S R R R Y N S M T G L  
 F D G Q V I Y Q S L T G L Y E T R M R F E D K R P A A T K K A G Q A K K K K G S G S E T P G T S E S A T P E S S G S E V E F S H E  
Y W M R H A L T L A K R A R D E R E V P V G A V L V L N N R V I G E G W N R A I G L H D P T A H A E I M A L R Q G G L V M Q N Y  
R L Y D A T L Y V T F E P C V M C A G A M I H S R I G R V V F G V R N A K T G A A G S L M D V L H H P G M N H R V E I T E G I L

ADECAALLCRFFRMPRRVFNAQKKAQSSTD**PAAKRVKLD**GS YPYDVPDYAYPYDVPDYAYPYDV  
PDYA (SEQ ID NO: 92)

**Amino Acid Sequence of Cytidine Deaminase, ppAPOBEC1 fused to the N-terminal of nickase PmaCas9 (CBE-nPmaCas9, D12A mutant)**

M**PAAKRVKLD**TSEKGPSTGDPTLRRRIESWEFDVFDPRELRKETCLLYEIKWGMSRKIWRSSG  
KNTTNHVEVNF~~IKKFTSERRFHSS~~ISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIYVARL  
FWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYPPLWMMLYALELH  
CIILSLPPCLKISR~~RWQNH~~LAF~~FR~~LHLQ~~NCHYQTIPPHILLATGLIHPSVTWRLKSGGSSGGSS~~  
GSETPGTSESATPES~~SGGSSGGSS~~**PKKKRKV**GEKKTNYTIGLAIGTDSVGWAVVKDDLELVKKRM  
KVLGNTETNYIKKNLWGSLLFESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEV  
DENFFFRLNESFLVPEDKAFSKNPIFGTLGEDKTYKYTYPTIYHLRQHLADSEEKADVRLIYLALAH  
MIKYRGHFLIEGKLDTEHIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVK  
EKRITSFLKDIGRANKQSPMMAFITLIVGKAKFKAAFNLEEEISLNLTDDSYDENLEILLNTI  
GSDFADLFDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQ  
YDMTFVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRS  
FYNGVIPNQVHLAEMEAILDROENYYPWLKENREKIIISLLTFRIPIYYVGPLADGQSEFAWLERK  
SDEKIKPWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIAYLDE  
RQKRMNLS~~SSVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELA  
KVPGMKSMDDPDNEDLMEEIVKILTVFEDRKMRRKQLEKYKERLSPEQIKELAKKHYTGWGR  
L~~SKLLVGIRDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDRSLFKKTI AELQ MIDPSAD  
LYAQVQEIAGSPAIKKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTTARGKALSKRREAKI  
KEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHII PQSFTV  
DNSLDNLVLTNSSQNRGNKKDDVPSLEV~~VNRQLAYWRS~~LKDAGLMTQRKFDNLTKAMRGGLTDK  
DRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNIVLLKSAMASEFRKMFRLYKV  
RELNDYHHAHDAYLNAAIAINLLALYPYMA~~DDFVYGEFRY~~KKKPKQAEKATY~~EKLRQWNLI~~KRFG  
EKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNVVKKAEERTGMLFKETINGKTNKGSRIPIK  
KDLDPKYGGYIEEKMAYYAVISYEDKKKKPGKTIVGISIMDKKEFEYDSISYLGKLGFSNPV  
VQIILKNYSLIAYPDGRRRYITGATKT~~TKG~~VELQKANQIAMEQDLVNFYHLKNYDEISHPESYAF  
VQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRIYEEKKKDPVDIEALVASFIELLKLTSAG  
APADFI~~FMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMR~~FED**GRPAATKKAGQAKKKKGS**  
SGGSSGGSSGGSS**TNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENV  
MLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSSGGSSGTNLSDIIEKETGKQLVIQESILMLP**

EEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKML YPYDVP  
DYAYPYDVPDYAY (SEQ ID NO: 93)

**Amino Acid Sequence of Cytidine Deaminase, ppAPOBEC1 fused to the C-terminal of nickase PmaCas9 (nPmaCas9-CBE, D12A mutant)**

MPAAKRVKLD TNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENV  
MLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSGGSGGSTNLSDIIEKETGKQLVIQESILMLP  
EEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSGG  
SGGS PKKKRKVEKKKNTNYTIGLAIGTDSVGVAVKDDLELVKKRMKVLGNTETNYIKKNLWGSLL  
FESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFRRLNESFLVPEDKAFSKN  
PIFGTLGEDKTYKYTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIA  
INENLEQFFESYNALFSEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDIGRANKQSPM  
MAFITLIVGKKAKFKAAFNLEEEISLNLTDSDYDENLEILLNTIGSDFADLFDHAQRVYNAVEL  
AGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKDLKGYA  
GYIDGNMSQDSFYKFKVDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVHLAEMEAILD  
RQENYYPWLKENREKIIISLLTFRIPIYVGPLADGQSEFAWLERKSDEKIKPWNFSDVVDLDRSA  
EKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIAYLDERQKRMNLS SVEKKEIFETLF  
KKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMDDPDNEDLMEE  
IVKILTVFEDRKMRRKQLEKYKERLSPEQIKELAKKHYTGWGRLSKLLLVGIRDKETQKTILDY  
LVEDDNHSGGRQHLNRNLMQLINDRSLFKKTI AELQ MIDPSADLYAQVQEIAGSPA I KKGILL  
GLKIVDEIIRVMGEKPENIVIEMARENQTTARGKALSKRREAKIKEGLAALGSSLLKENLPGNA  
DLSQRKIYLYYTQNGKDIYLDEPLDFDRLS QYDEDHII PQSFTVDNSLDNLVLTNSSQNRGNKK  
DDVPSLEVVRQLAYWRS LKDAGLMTQRKFDNLT KAMRGGLTDKDRERFIQRQLVETRQITKNV  
AKLLDMRLNDKKDEAGNKIRETNIVLLKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAAIAI  
NLLALYPYMADDFVYGEFRYKKKQAEKATYEKLRQWNLIKRFGEKQLFTPDHEDCWNKERDIK  
TIKKVMGYRQVNVVKKAEERTGMLFKETINGKTNKGSRIPIKKDLPSKYGGYIEEKMAYYAVI  
SYEDKKKKPGKTIVGISIMDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIAYPDGRRRYITG  
ATKTTKGKVELQKANQIAMEQDLVNFIIYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTR  
RFLDAETNINRLRRIYEEKKKDPVDIEALVASFIELLKLTSAGAPADFI FMGEAISRRRYNSM  
TGLFDGQVIYQSLTGLYETRMRFED KRPAATKKAGQAKKKKSGSSGGSSGGSSGSETPGTSESAT  
PESGGSSGGS TSEKGPSTGDPTLRRRIESWEFDFVYDPRELRKETCLLYEIKWGMSRKIWRSS  
GKNTTNHVEVNFIIKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIYVAR  
LFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYPPLWMMLYALEL

HCIIILSLPPCLKISRWRQNHLLAFFRLHLQNCYQTIPPHILLATGLIHPSVTWRYPYDVPDYAY  
PYDVPDYAYPYDVPDYA (SEQ ID NO: 94)

Linker (underlined, no italics or bolding)

TadA8 (ABE) or ppABOBEC1 (CBE) (italics and underlined)

Nickase mutation: D10A mutation in ScoCas9, D14A mutation in SirCas9, D38A in VapCas9, D12A in EpeCas9, D9A in LfeCas9, D12A in PmaCas9 (bold and italics)

5 2xUGI (bold, italics and underlined)

3xHA tag (italics), can be substituted with different tags

### Example 6. Engineered *Streptococcus constellatus* (ScoCas9) NGC PAM variants

10 This example illustrates the engineering of ScoCas9 variants that recognize NGC PAM variants.

Briefly, two variants were engineered, ScoCas9-NGC-v1, which contains amino acid substitutions for NGC PAM recognition and ScoCas9-NGC-v2, which contains amino acid substitutions for NGC PAM recognition and additional amino acid substitutions that enhance SpyCas9 activity. The amino acid residues were identified by structural comparison between  
 15 *S. pyogenes* SpyCas9 and *S. constellatus* ScoCas9. The amino acid sequence of ScoCas9-NGC-v1 (SEQ ID NO: 95) comprised the following mutations from wild type ScoCas9 sequence: D1117M, S118Q, E1201F, A1299R, D1309A, R1312E, T1314R. The amino acid sequence of ScoCas9-NGC-v2 (SEQ ID NO: 96) comprised the following mutations from wild type ScoCas9 sequence: S409I, R655L, D1117M, S118Q, E1201F, A1299R, D1309A,  
 20 R1312E, T1314R.

#### Amino acid sequence of *Streptococcus constellatus* (ScoCas9) variant (ScoCas9-NGC-v1)

**MPKKKRKVG**MGKPYSIGLDIGTNSVGWAVVTDDYKVPAAKMKVLGNTDKQSIKKNLLGALLFDS  
 GETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDSDFLVDEDKRGEHHPIF  
 GNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQA  
 LFKDFVEEYDKTIEESHLSIEITVDALSILTEKVS KSSRLENLIAHYPTKKNLTFGNLIALSLD  
 LHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDYADLFASAKNLYDAILLSGILTVDN  
 STKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEFYKY

LKGILLKINGS GDFLDKIDREDFLRKQRTFDNGSIIPHQIHLQEMHAILRRQGEHY PFLKENQDK  
 IEKILTFRI PYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLP  
 EEKVLPHKSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTKDKLLNYL  
 NKEFEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSFDDKANAKTIEDIIQTTLTFEDRE  
 MIRQRLQKYSDI FTKAQLKKLERRH YTGWRLSYKLINGIRNKENKKTILDYLI DDGYANRNF  
 QLINDALS FKEEIARAQIIDDVDDIANVVHDLPGSPA I KKGILQSVKIVDELVKVMGHNPANI  
 I IEMARENQT TDKGRNSQORLKL LQDSLKNLDPVNIKNVENQQ LQNDRLFLYYIQNGKDMYT  
 GETLDINNLSQYDIDHII PQA FIKDNSLDNRVLT RSDKNRGKSDDVPSIEVVHEMKS FWSKLLS  
 VKLITQRKFDNLT KAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRN  
 VKIITLKS NLVSNFRKEFELYKVREINDYHHAHDAYLNAVVG NALLKYPQLEPEFVYGEYPKY  
 NSYRSRKSATEKFLFY SNILRFFKEDIQTNE DGEIAWNKEKHIKILRKVLSYPQVNI VKKTEE  
 QTGGFSKESILPKGESDKLI PRKTKNSYWDPKKYGGFMQPVVAYSILVFADVEK GKSKKLRKVQ  
 DMVGITIMEKKRFEKNPVDFLEQRGYRNVRL EKI I KLPKYSLFELENKRRRL LASAKFLQKNE  
 LVIPQRFTTLLYHSYRIEKDYEP EPREYVEKHKDEFKELLEYISVFSRKYVLADNNLT KIEMLF  
 SKNKDAEVSS LAKSFISLLTFTA FGA PRAFNFFGENIARKEYRSVTECLNATLIHQSI TGLYET  
 RIDLSKLGEDGE **EGADKRTADGSEFE SPKKRKV** (SEQ ID NO: 95)

**Amino acid sequence of *Streptococcus constellatus* (ScoCas9) variant (ScoCas9-NGC-v2)**

**MPKKKRKV**GMGKPYSIGLDIGTNSVGWAVVTDDYKVP AKKMKVLGNTDKQSIKKNLLGALLFDS  
 GETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDSSFLVDEDKRGEHHPIF  
 GNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQA  
 LFKDFVEEYDKTIEESHLS EITVDALSILTEKVS KSSRLENLIAHYPT EKKNTLFGNLI ALSLD  
 LHPNFKTNFQ LSEDAKLQFSKDTYEEDLE GFLGEV GDEYADLFASAKNLYDAILLSGILT VDDN  
 STKAPLSASMVKRYEEHQKDLK KLDKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEFYKY  
 LKGILLKINGS GDFLDKIDREDFLRKQRTFDNGSIIPHQIHLQEMHAILRRQGEHY PFLKENQDK  
 IEKILTFRI PYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLP  
 EEKVLPHKSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTKDKLLNYL  
 NKEFEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSFDDKANAKTIEDIIQTTLTFEDRE  
 MIRQRLQKYSDI FTKAQLKKLERLHYTGWRLSYKLINGIRNKENKKTILDYLI DDGYANRNF  
 QLINDALS FKEEIARAQIIDDVDDIANVVHDLPGSPA I KKGILQSVKIVDELVKVMGHNPANI  
 I IEMARENQT TDKGRNSQORLKL LQDSLKNLDPVNIKNVENQQ LQNDRLFLYYIQNGKDMYT

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GETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKS FWSKLLS
VKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRN
VKIITLKSNLVSNFRKEFELYKVREINDYHHAHDAYLNAVVGNALLLKYPQLEPEFVYGEYPKY
NSYRSRKSATEKFLFYSNILRFFKKEDIQTNEDEGEIAWNKEKHIKILRKVLSYPQVNIKKTEE
QTGGFSKESILPKGESDKLIPRKTKN SYWDPKKYGGFMQPVVAYSILVFADVEK GKSKKLRKVQ
DMVGITIMEKKRFEKNPVDFLEQRGYRNVRLKIKL PKYSLFELENKRRRL LASAKFLQKGNE
LVIPQRFTTLLYHSYRIEKDYEP E HREYVEKHKDEFKELLEYISVFSRKYVLADNNLT KIEMLF
SKNKDAEVSSLAKSFISLLTFTA F GAPRAFNF F GENIARKEYRSVTECLNATLIHQSI TGLYET
RIDLSKLGEDGEGADKRTADGSEFE SPKKRKV (SEQ ID NO: 96)

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NLS (bold italics)

Linker (bold underlined)

ScoCas9-NGC variants were used to target a genomic locus that was randomly  
5 integrated into the genome of HEK293T cells by lentivirus mediated insertion and tested for  
nuclease and base editing activities.

Briefly, HEK293T cells were plated in a 96-well plate. Cells were transfected with  
expression vectors containing ScoCas9-NGC variants, and guide RNA sequence  
ATCGACAAGAAAGGGACTGA (SEQ ID NO: 97), 24 hours after plating. The ScoCas9  
10 variants recognized an exemplary NGC 3' PAM sequence, AGC. Cells were harvested 72  
hours post-transfection and total DNA was extracted.

Deep sequencing was carried out to characterize indel patterns in the HEK293T cells.  
Exemplary targets were amplified using a two-round PCR to add Illumina adapters as well as  
unique barcodes to the target amplicons. PCR products were run on a 2% gel and gel  
15 extracted. Samples were pooled, quantified and cDNA libraries were prepared and sequenced  
on MiSeq. Indel frequency was determined by deep sequencing 4 days after transfection.

The results showed nuclease activity of both ScoCas9-NGC variants. An indel  
frequency of between about 20-35% was achieved with ScoCas9-NGC-v1 and ScoCas9-  
NGC-v2 (FIG. 11A).

20 Fusions were constructed of ScoCas9-NGC variants with ABE base editors.

**Amino acid sequence of a ScoCas9 variant fused to an adenine base editor (ABE-nScoCas9-NGC-v1)**

MSEVEFSHEYWMRHALTLAKRARDEREVPGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMALR  
 QGGLVMQNYRLYDATLYSTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNH  
 RVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSDSGSSGGSSGSETPGTSESATPESS  
GGSSGGSGKPYSIGLAIGTNSVGWAVVTDDYKVPKMKVGLGNTDKQSIKKNLLGALLFDSGET  
 AEATRLKRTARRRYTRRNRLRYLQEIFTGEMNKVDENFFQRLDDSLVDEDKRGEHHPIFGNI  
 AAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFK  
 DFVEEYDKTIEESHLSEITVDALSILTEKVS KSSRLENLIAHYPTKKNLTFGNLIALSLDLHP  
 NFKTNFQLSEDAKLQFSKDTYEEDLEGFLGVEGDEYADLFASAKNLYDAILLSGILTVDNSTK  
 APLSASMVKRYEEHQDLKCLKDFIKVNA PDQYNAIFKDKNKKGYASYIESGVKQDEFYKYLKG  
 ILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHYPFLKENQDKIEK  
 IILTFRIPIYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLPEEK  
 VLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKYTKDKLLNYLNKE  
 FEEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSLDDKAN EKTIEDI IQTLTLFEDREMIR  
 QRLQKYSDFTKAQLKKLERRHYTGWRLSYKLINGIRNKENKKTILDYLI DDGYANRNFMQLI  
 NDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPA IKKGILQSVKIVDELVKVMGHN PANIIIE  
 MARENQTTDKGRRNSQQRLKLLQDSLKNLDNPVNIKNVENQQQLQNDRLFLYYIQNGKDMYTGET  
 LDINNLSQYDIDHII PQAFIKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKS FWSKLLSVKL  
 ITQRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKI  
 ITLKS NLVSNFRKEFELYKVREINDYHHAHDAYLNAVVG NALLLKYPQLEPEFVYGEYPKYNSY  
 RSRKSATEKFLFYSNILRFFKKEDIQTNEDEGEIAWNKEKH IKILRKVLSYPQVNIVKKTEEQTG  
 GFSKESILPKGESDKLIPRKTKN SYWDPKKGGMQPVVAYSILVFADVEKGKSKKLRKVQDMV  
 GITIMEKKRFEKNPVDFLEQRGYRNRLEKIIKLPKYSLFELENKRRRL LASAKFLQKGNELVI  
 PQRFTTLLYHSYRIEKDYEPHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMLFSKN  
 KDAEVSSLAKSFISLLTFTA FGA PRAFNF GENIARKEYRSVTECLNATLIHQ SITGLYETRID  
 LSKLGEDGEGADKRTADGSEFE SPKKRKY (SEQ ID NO: 98)

**Amino acid sequence of a ScoCas9 variant fused to an adenine base editor (ABE-nScoCas9-NGC-v2)**

MSEVEFSHEYWMRHALTLAKRARDEREVPGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMALR

*QGGLVMQNYRLYDATLYSTFFPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNH*  
*RVEITEGILADECAALLCRFFRMPRRVFNQAQKAQSS***TD****SGGSSGGSSGSETPGTSESATPESS**  
**GGSSGGS**GKPYSIGLAIGTNSVGWAVVTDDYKVPKMKVGLGNTDKQSIKKNLLGALLFDSGET  
 AEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRGEHHPIFGNI  
 AAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFK  
 DFVEEYDKTIEESHLSEITVDALSILTEKVS KSSRLENLIAHYPTKKNLTFGNLIALSLDLHP  
 NFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDYADLFASAKNLYDAILLSGILTVDDNSTK  
 APLSASMVKRYEEHQKDLKDLKDFIKVNA PDQYNAIFKDKNKKGYASYIESGVKQDEFYKYLKG  
 ILLKINGSGDFLDKIDREDFLRKQRTFDNGIIPHQIHLQEMHAILRRQGEHY PFLKENQDKIEK  
 ILTFRIPYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLPEEK  
 VLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTKDKLLNYLNKE  
 FEEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSLDDKAN EKTIEDIIQTTLTFEDREMIR  
 QRLQKYSDFTKAQLKKLERLHYTGWGRLSYKLINGIRNKENKKTILDYLI DDGYANRNFMQLI  
 NDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPA IKKGILQSVKIVDELVKVMGHNPANIIIE  
 MARENQTTDKGRNSQQRLKLLQDSLKNLNDPNVNIKNVENQQQLQNDRLFLYYIQNGKDMYTGET  
 LDINNLSQYDIDHII PQAFIKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKS FWSKLLSVKL  
 ITQRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKI  
 ITLKSNLVSNFRKEFELYKVREINDYHHAHDAYLNAVVG NALLLKYPQLEPEFVYGEYPKYNSY  
 RSRKSATEKFLFYSNILRFFKKEDIQTNE DGEIAWNKEKH IKILRKVLSYPQVNIVKKTEEQTG  
 GFSKESILPKGESDKLIPRKTKN SYWDPKKGGMQPVVAYSILVFADVEKGKSKKLRKVQDMV  
 GITIMEKKRFEKNPVDFLEQRGYRNRLEKIIKLPKYSLFELENKRRRL LASAKFLQKGNELVI  
 PQRFTTLLYHSYRIEKDYEPHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMLF SKN  
 KDAEVSSLAKSFISLLTFTAFGAPRAFNF GENIARKEYRSVTECLNATLIHQ SITGLYETRID  
 LSKLGEDG***EGADKRTADGSEFE SPKKRKY*** (SEQ ID NO: 99)

Linker (bold underlined)

TadA8 (ABE) (italics)

NLS (bold italics)

5 Deep sequencing was also carried out to characterize A-to-G conversion in the HEK293T cells (FIG. 11B). Adenine-to-Guanine (A-to-G) conversions were measured by NGS 4 days post transfection. The results showed base editing activity by both ABE-nScoCas9-NGC variants. Both variants showed between about 20-30% A-to-G conversion.



ScoCas9 that recognized NGG was used as a negative control and showed no base editing. SpyCas9 was used as a positive control and showed about 40% A-to-G conversion.

Overall, the results showed that ScoCas9 variants engineered to recognize NGC PAM sequences could carry out nuclease as well as base editing activities.

5

### **EQUIVALENTS AND SCOPE**

Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. The scope of the present invention is not intended to be limited to the  
10 above Description, but rather is as set forth in the following claims.

CLAIMS

1. An engineered, non-naturally occurring Cas9 protein modified from *Streptococcus constellatus* Cas9, *Sharpea* Cas9, *Veillonella parvula* Cas9, *Ezakiella peruensis* Cas9, *Lactobacillus fermentum* strain AF15-40LB Cas9, or *Peptoniphilus* sp. Marseille-P3761

5 Cas9.

2. The Cas9 protein of claim 1, wherein the *Streptococcus constellatus* Cas9 has at least 80% sequence identity to

MGKPYSIGLDIGTNSVGVAVVTDDYKVPAAKMKVLGNTDKQSIKKNLLGALLFDSGETAEAT  
 RLKRTARRRYTRRKNRLRYLQEI FTGEMNKVDENFFQRLDDSFLVDEDEKRGHEHPIFGNIAA  
 10 EVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFK  
 DFVEEYDKTIEESHLSSEITVDALSILTEKVS KSSRLENLIAHYPTKKNLTFGNLIALSLDL  
 HPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDYADLFASAKNLYDAILLSGILTVDD  
 NSTKAPLSASMVKRYEEHQKDLKLLKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEF  
 YKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHY PFLK  
 15 ENQDKIEKILTFRIPIYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMT  
 LNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKV  
 TKDKLLNYLNKEFEEFRIVNLTGLDKENKAFNSSLGT YHDLRKI LDKSFLDDKAN EKTIEDI  
 IQTLTLFEDREMIRQRLQKYSDFTKAQLKLERRH YTGWRLSYKLINGIRNKENKKTILD  
 YLIDDG YANRNFQMLINDDALS FKEE IARAQIIDDVDDIANVVHDLPGSPA I KKGILQSVKI  
 20 VDELVKVMGHNPANIIEMARENQTTDKGRNSQORLKL LQDSLKNLDNPNVNIKNVENQQQLQ  
 NDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSD  
 VPSIEVVHEMKS FWSKLLSVKLITQRKFDNLT KAERGGLTEEDKAGFIKRQLVETRQITKHV  
 AQILDERFNTEFDGNKRRIRNVKIITLKS NLVSNFRKEFELYKVREINDYHHAHDAYLNAV  
 GNALLLKYPQLEPEFVYGEYPKYNSYRSRKSATEKFLFYSNILRFFKKEDIQTNE DGEIAWN  
 25 KEKHIKILRKVLSYPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKT KNSYWDPKKYGGF  
 DSPVVAYSILVFADVEKGKSKKLRKVQDMVGITIMEKRF EKNPVDFLEQRGYRNVRLKII  
 KLPKYSLFELENKRRRLLASAKELQKGNELVIPQRFTLLYHSYRIEKDYEP EPREYVEKHK  
 DEFKELLEYISVFSRKYVLADNNLTKIEMLF SKNKDAEVSSLAKSFISLLTFTAFGAPAAFN  
 FFGENIDRKRYTSVTECLNATLIHQSI TGLYETRIDL SKLGED (SEQ ID NO: 1).

30 3. The Cas9 protein of claim 1, wherein the *Sharpea* Cas9 has at least 80% sequence identity to

MAKNKDIRYSIGLDIGTNSVGVAVMDEHYELLKKGNNHMMWGSRLFDAEPAATTRASRSIRR  
 RYNKRREIRILLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQKNLGN DYKDNYNLFIEKDF  
 NDKTYDYKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQSFAKVYEDI EEK  
 LDNTLKKFMSLNDLDNLFVDNDINSMITVLSKIYQRSKKADDLLKIMNPTKEERAAYKEFTK  
 5 ALVGLKFNVS KMILAQEVKDDKDIELD FSNVDYDSTVDGLQAE LGEYIEFIEMLHSINSW  
 ELQDILGNNSTISAAMVERYEEHKNDLRVLK KVI REELPKYNEVFREDNPKLHNYLGYIKY  
 PKNTPVEEFY EYIKRLLAKVDTGEAREILERIDLEK FMLKQNSRTNGSIPYQMOKDEMIQII  
 DNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFEDKQKERILPWN YDQIVD  
 IDATAEGFIERMONTGTYPDPKPVMAKNSLTVSKFEVLNELNKIRINGKLI PVETKKELLS  
 10 LFMKNKTITDKKLDWLVT HQYYDTNEELKIEGYQKDLQFSTSLAPWIDFTKIFGEINASNY  
 QLIEKIIYDISIFEDKKILKRRLK K VYQLDDLLVDKILKLN YTGWSRLSEKLLTG IKSNSK  
 ETILSILENSNMNLMEIINDESLGFKQIIEESNKKDIEGPF RYDEVK KLAGSPA I KRGIWQA  
 LLVVQEITKFMKHEP SHIYIEFAREEQEKV RTESRIAKLQKIYKDLNLQTKEDQLVYESLKK  
 EDAKKKIDTDALYLYLQMGKSMYSGKPLDIDKLSTYHIDHILPRSLIKDDSLDNRVLVLPK  
 15 ENEWKLDSETVPFEIRNKMMGFWQKLHENG LMSNKKFFSLIRTD FNEKDKKRFINRQLVETR  
 QIIKNVAVIINDHYTNTNVVTVRAELSHQFRERYKIYKNRDLNDLHHAHDAYIACILGQFIH  
 QNFGNMDVNMIYGQYKKNYK KDVQEHNNYGFILNSMNHIFND DNSVIWDPSYIGKIKSCFC  
 YKDVYVTKKLEQNDAKLFDLTI LPSDKNSENGVTKAKI PVNKYRKDVNKYGGFSGDAPIMLA  
 IEADKGGK KHRQVIAFPLRLKNYNDEERIKFIEKEKNLKNVKILTEVKKNQLILINHQYFFI  
 20 TGTNELVNATQLKLSAKNTKNL FNLDANKHNKLESID DANFNEVIQELICKLQEP IYSRYN  
 SIGKEFEDSYEKINAVTKQDKLYIIEYLIAIMSAKATQGYIKPELAREIGTNGKNKGRIKSF  
 TIDLNKTTFISTSVTGLFSK KYKL (SEQ ID NO: 4).

4. The Cas9 protein of claim 1, wherein the *Veillonella parvula* Cas9 has at least 80% sequence identity to

MSIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLDIGTSSVGVAVTNKAYELLKFRSHK  
 MWGSRLFDEGESAVARRGFRSMRRRLERRKLRLLKLEELFADAMAQVDPTFFMRLRESKYHY  
 EDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHHILKYRGN  
 FLYEGATFDSNASTLDDVIKQALENITFNCFCNSAIISSIGQILMEAGKTKSDKAKAIEHLV  
 DTYIATDVTDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEDLKKLQITG  
 25 DTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILL SIKEPGLTISESKVKA FNKHKDDLAILK  
 SLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEY  
 ILSQIELQILLPLQRIKDN GVI PYQLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKML  
 EFRIPIYYVGPLNTHHNVDNNGGFAVA VRKASGRVTPWNFDDKIDREKSAAAFIKNLTNKCTYL  
 30

LGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVEHLIEAVFKQDHKKMTKNRIEQFLK  
 DNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGESKMLRE  
 TLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGTPEIIILMRNFSYNL  
 MELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVDEVAHIKKA  
 5 LPARIFVEVTRSNNKNEKKKKDSRQRLSDLYAAIKKDDVLLNGLNNEIFGELKSSLAKYDDA  
 ALRSKKLYLYYTQMGRCAYTGEIIELSLLNTDNYDIDHIYPRSLTKDDSFDNLVLCKRTANA  
 QKSDAYPISEEIQKTQKPFWTFKQOGLISERKYERLTRITPLTADDLSGFIARQLVETNQS  
 VKAATTLRLRYPGVDVVFVKAENVTDNRHDNFIKVRSLNHHHHAKDAYLNIIVGVNHYHER  
 FTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKDGGKAWDVKTSMDTVKKMMSNDVVRVTKR  
 10 LLEQTGALADATIYKATVAGKAKDGAYIGMKTSSVFADVSKYGGMTKIKNAYSIIIVQYTGK  
 KGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISIIYGKLCINQLVKVNGFYYYLG  
 GKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNNMRKDNKDLKANVVSTRALDNKHTIEVRIV  
 EEKNIEFFDYLVSKLKMPIYQMKGNKAAELSEKGYGLFKKMSLEEQSIHLIELLNLLTNQK  
 TTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIV (SEQ ID NO: 8).

15 5. The Cas9 protein of claim 1, wherein the *Ezakiella peruensis* Cas9 has at least 80%  
 sequence identity to

MTKVKDYYIGLDIGTSSVGWAVTDEAYNVLFKNSKKMWGVRLFDDAKTAEERRGQRGARRRL  
 DRKKERLSLLQDFFAEVAKVDPNFFLRDLNSDLYMEDKDQKLKSKYTLFNDKDFKDKNFHK  
 KYPTIHHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIFEGQKFDTKSSFENSLNELKVHL  
 20 NDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQKLVDLFE  
 NPEDFDDSAIKSVDFSTTSFDDKYSYELALGDKIALVNILKEIYDSSILENLLKEADKSKD  
 GNKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNNDNYVSYTKSSISNNKRV  
 KANKFTDQEAIFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNGV  
 IPYQLKLMELNKILENQSKHHEFLNVSDYGSVCDKIASIMEFRIPYYVGPLNPNSKYAWIK  
 25 KQKDESEITPWNFKDVVDLDS SREEFIDSLIGRCTYLKDEKVLPKASLLYNEYMVLNELNLLK  
 LNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGDFKQGLNSYND  
 FKAIVGDKVDSDDYRDKIEEIIKLVLYGDDKSYLQKKIKAGYGKYFTDSEIKKMAGLNYKD  
 WGRLSKKLLTGLEGANKITGERGSIHFMREYNLNMELMSASFTEETEEIQKLNVPVDDRKLS  
 YEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIEMARGKEEVKARKE SRKNQLLK  
 30 FYKDGKKAIFISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQLGRCMYSLEPIDISELSS  
 KNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPIPDEILNKNCYAYWKILYDKGLI  
 GQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTIKCNSEIVYSKAENASRFR  
 QEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKDPMNFVKKQEKARSYNLENMFKYDVK

RGGYTAWIADDEKGTVKNAS IKRIRKELEGTNYRFTRMNYIESGALFNATLQRKNKGSRPLK  
 DKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLMVEREIYAKQKNDKKSDEIFSKY  
 LKDRFGIEDYRVVYPVVKMRTLKIDGSYYFITGGSDKTLELRSALQLILPKKNEWAIKQID  
 KSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLFQDDKIENIDFKFKSMDF  
 5 KEKCKTLLMLVKAIRASGVRQDLKSIDLKS DYGRLLSSKTNNIGNYQEFKIINQSITGLFENE  
 VDLLKL (SEQ ID NO: 14).

6. The Cas9 protein of claim 1, wherein the *Lactobacillus fermentum strain AF15-40LB* Cas9 has at least 80% sequence identity to

MKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGTAKIGVRLFEEGKTAERRTFRTRRRRLKRR  
 10 KWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEPDAKNQAFIGKLLFPDLLKKNERGYPTL  
 IKMRDELPEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAVHHIVKYRGHFLNNASVDKF  
 KVGRI DFDKS FNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLDKQAVAKLLEV  
 KVADKEETKRKQIATAMSKLVLYGKADFATVAMANGNEWKIDLSSETSEDEIEKFREE LSD  
 AQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWTHERQLAEVKEYLATQPASARKEFD  
 15 QVYNKYIGQAPKEKGFDFLEKGLKILSKKENWKEIDELLKAGDFLPKQRTSANGVI PHQMHO  
 QELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPIYYVGPLVTPVQKATSG  
 AKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKDTYLLNEDVLPANSLLYQKYNVL  
 NELNNVRVNGRRLSVGIKQDIYTELFKKKTKVAGDVASLVMAKTRGVNKP SVEGLSDPKKF  
 NSNLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVEWLTDEQRSALV  
 20 KKRYKGWGRLSKLLTGIVDENGQRIIDL MWNTDQNFMQIVNQPVFKEQIDQLNQKAITNDG  
 MTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSI SIEFARNEGKGEITRS  
 RRTQLQKLFEDQAHELKDTSLTEELEKAPDLSDRYFYFTQGGKDMYTGDPINFDEISTKY  
 DIDHILPQS FVKDDSLDNRVLVSRAENNKSDRVPKLYAAKMKPYWNQLLKQGLITQRKFE  
 NLTMDVDQTIKYRSLGFVKRQLVETRQVIKLTANILGSMYQEAGTDIIETRAGLTKQLREEF  
 25 DLPKVREVNDYHHAVDAYLTTFAGQYLNRRYPKLR SFFVYGEYMKFKHGS DLKLRNFNF FHE  
 LMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLISNETYEETGKYFDASHES SL  
 YLKNQNKKS KLVVPLKNKLQPEYYGAYTGITQGYMVILKLLDKKGGFGVYRIPRYAADILNK  
 CHDEVAYRNKIAEIISSDPAPKSFVVVPRVLKGTFLVDGEEKFILSSYRYKVNATQLILP  
 VSDIKLIQDNFKALKKLVEMQTKKLIETIDNLRQVDKYYKLYDINKFRAKLHDGRSKFVE  
 30 LDDFGQDASKEKVIKILRGLHFGSDLQNLKEIGFGTTP LGQFQVSEAGIRLSNTAFIIFKS  
 PTGLFNRKLYLKNL (SEQ ID NO: 84).

7. The Cas9 protein of claim 1, wherein the *Peptoniphilus sp. Marseille-P3761* Cas9 has at least 80% sequence identity to

MEKKTNYTIGLDIGTDSVGVAVVKDDLELVKKRMKVLGNTETNYIKKNLWGSLLFESGQTAK  
 DRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDFNFFRLNESFLVPEDKAFSKNPIFGTL  
 5 GEDKTYKYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIAINEN  
 LEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDIGRANKQSPMMA  
 FITLIVGKKAKFKAAFNLEEEISLNLTDSDYDENLEILLNTIGSDFADLFDHAQRVYNAVEL  
 AGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKKDLKG  
 YAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVHLAEME  
 10 AILDRQENYYPWLKENREKIIISLLTFRIPIYVGPLADGQSEFAWLERKSDEKIKPWNFSDVV  
 DLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIAYLDERQKRMNLS SVEK  
 KEIFETLFFKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMMD  
 DPNEDLMEEIVKILTVFEDRKMRRKQLEKYKERLSPQIKELAKKHHTGWGRLSKKLLVGI  
 RDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDRSLFKKTI AELQ MIDPSADLYAQVQ  
 15 EIAGSPAIKKGILLGLKIVDEIRVMGEKPENIVIEMARENQTTARGKALS KRREAKIKEGL  
 AALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHII PQSFTVDN  
 SLDNLVLTNSSQNRGNKKDDVPSLEVVRQLAYWRS LKDAGLMTQRKFDNLT KAMRGGLTDK  
 DRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNIVLLKSAMASEFRKMFRLY  
 KVRELNDYHHAHDAYLNAAI AINLLALYPYMA DDFVYGEFRYKKKPQAEKATYEKLRQWNLI  
 20 KRFGKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNVVKKAEERTGMLFKETINGKTNKGSR  
 IPIKKDLDP SKYGGYIEEKMAYYAVISYEDKKKPKGKTIVGISIMDKKEFEYDSISYLGKLG  
 FSNPVVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKANQIAMEQDLVNFYHLLKNYD  
 EISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIYEEKKKDPVDIEALVA  
 SFIELLKLTSAGAPADFI FMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMRFED (SEQ  
 25 ID NO: 86).

8. The Cas9 protein of any one of claims 2-7 comprising an amino acid sequence that is at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% identical to SEQ ID NOs: 1, 4, 8, 14, 84 or 86.

9. The Cas9 protein of any one of the preceding claims, further comprising a nuclear  
 30 localization sequence (NLS) and/or a FLAG, HIS or HA tag.

10. The Cas9 protein of claim 9, wherein the *Streptococcus constellatus* Cas9 has an amino acid sequence at least 80% identical to

MPKKKRKVGKPYISGLDIGTNSVGVAVVTDDYKVPAKKMKVLGNTDKQSIKKNLLGALLFD  
 SGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSFVDEDEKRGHH  
 5 PIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAEN  
 TDVQALFKDFVEEYDKTIEESHLSSEITVDALSILTEKVS KSSRLENLIAHYPTKKNLTFGN  
 LIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGEYADLFASAKNLYDAILL  
 SGILTVDNSTKAPLSASMVKRYEEHQDLKCLKDFIKVNAPDQYNAIFKDKNKKGYASYIE  
 SGVKQDEFYKYLKGI LLKINGS GDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQ  
 10 GEHY PFLKENQDKIEKILTRIPYVYVGPLARKGSRFAWA EYKADEKITPWNFDDILDKEKSA  
 EKFITRMTLNLDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDH  
 VFKENRKVTKDKLLNYLNKEFEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSFLDDKA  
 NEKTIEDIIQTLTLFEDREMIRQLQKYSDI FTKAQLKKLERRHYTGWGRLSYKLINGIRNK  
 ENKKTILDYLI DDGYANRNFQMLINDDALS FKEE IARAQIIDDVDDIANVVHDLPGSPA I KK  
 15 GILQSVKIVDELVKVMGHN PANI I IEMARENQTTDKGRRNSQQRLKLLQDSLKNLDNPNVNIK  
 NVENQQQLQNDRLFLYYIQNGKDMYTGETLDINNL SQYDIDHIIPQAFIKDNSLDNRVLRSD  
 KNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLVE  
 TRQITKHVAQILDERFNTEFDGNKRRI RNVKIITLKS NLVSNFRKEFELYK VREINDYHHAH  
 DAYLNAVVG NALLLKY PQLEPEFVYGEY PKYNSYRSRKSATEKFLFY SNILRFFKKEDIQTN  
 20 EDGEIAWNKEKHIKILRKVLSYPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKTKN SYW  
 DPKKYGGFDS PVVAYSILVFADVEK GKSKLKRKVQDMVGITIMEKKRFEKNPVDFLEQRGYR  
 NVRLEKIIKLPKYSLFELENKRRRLLASAKELQKGNELVIPQRFTTLLYHSYRIEKDYEP E H  
 REYVEKHKDEFKELLEYISVFSRKYVLADN NLTKIEMLF SKNKDAEVSSLAKSFISLLTFTA  
 FGAPAAFNFFGENIDRKRYTSVTECLNATLIHQSI TGLYETRIDL SKLGEDGKRPAATKKAG  
 25 QAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 2).

10b. The Cas9 protein of claim 9, wherein the *Streptococcus constellatus* Cas9 has an amino acid sequence at least 80% identical to

MPKKKRKVGGMKPYISGLDIGTNSVGVAVVTDDYKVPAKKMKVLGNTDKQSIKKNLLGALLF  
 DSGETAETRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSFVDEDEKRGHH  
 30 HPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAE  
 NTDVQALFKDFVEEYDKTIEESHLSSEITVDALSILTEKVS KSSRLENLIAHYPTKKNLTFG  
 NLIALS LLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGEYADLFASAKNLYDAILL

LSGILTVDDNSTKAPLSASMVKRYEEHQDLKCLKDFIKVNAPDQYNAI FKDKNKKGYASYI  
 ESGVKQDEFYKYLKGI LLKINGS GDFLDKI DREDFLRKQRTFDNGS I PHQIHLQEMHAILRR  
 QGEHY PFLKENQDKIEKILTFRIPYYVGPLARKGSRFAWA EYKADEKITPWNFDDI LDKEKS  
 AEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFD  
 5 HVFKENRKVTKDKLLNLYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSFLDDK  
 ANEKTIEDIIQTLTLFEDREMIRQLQKYS DI FTKAQLKKLERRHYTGWGRLSYKLLINGIRN  
 KENKKTILDYLI DDGYANRNFQMLINDDALS FKEEIARAQIIDDVDDIANVHDLPGSPA IK  
 KGILQSVKIVDELVKVMGHNPANII IEMARENQTTDKGRRNSQQRLKLLQDSLKNLNDPNVI  
 KNVENQQLQNDRLFLYYIQNGKDMYTGETLDINNSQYDIDHII PQAFIKDNSLDNRVLTRS  
 10 DKNRGKSDDVPSIEVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLV  
 ETRQITKHVAQILDERFNTEFDGNKRRI RNVKIITLKS NLVSNFRKEFELYK VREINDYHHA  
 HDAYLNAVVG NALLKY PQLEPEFVYGEY PKYNSYRSRKSATEKFLFY SNILRFFKKEDIQT  
 NEDGEIAWNKEKH IKILRKVLSYPQVNI VKKTEEQTGGFSKESILPKGESDKLI PRKTKNSY  
 WDPKKYGGFMQPVVAYSILVFADVEKGKSKKLRKVQDMVGITIMEKKRFEKNPVDFLEQRGY  
 15 RNVRL EKI IKLPKYS LFELENKRRLLASAKFLQGNELVIPQRFTLLYHSYRIEKDYEP E  
 HREYVEKHKDEFKELLEYISVFSRKYVLADNNLT KIEMLFSKNKDAEVSS LAKSFISLLTFT  
 AFGAPRAFNF FGENIARKEYRSVTECLNATLIHQSI TGLYETRIDLSKLGEDGEGADKRTAD  
 GSEFESP KKRKV (SEQ ID NO: 95).

10c. The Cas9 protein of claim 9, wherein the *Streptococcus constellatus* Cas9 has an amino  
 20 acid sequence at least 80% identical to

MPKKKRKVG MGKPYSIGLDIGTNSVGVAVVTDDYKVP AKKMKVLGNTDKQS I KKNLLGALLF  
 DSGETA EATRLKRTARRRYTRRKNRLRYLQE I FTGEMNKVDENFFQRLDDS FLVDEDKRGEH  
 HPIFGNIAAEVKYHDDFPTIYHLRRHLADT SKKADLRVYLALAHMIKFRGHFLYEGDLKAE  
 NTDVQALFKDFVEEYDKTIEESHLS EITVDALSILTEKVS KSSRLENLIAHYPTEKKNTLFG  
 25 NLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEV GDEYADLFASAKNLYDAIL  
 LSGILTVDDNSTKAPLSASMVKRYEEHQDLKCLKDFIKVNAPDQYNAI FKDKNKKGYASYI  
 ESGVKQDEFYKYLKGI LLKINGS GDFLDKI DREDFLRKQRTFDNGI I PHQIHLQEMHAILRR  
 QGEHY PFLKENQDKIEKILTFRIPYYVGPLARKGSRFAWA EYKADEKITPWNFDDI LDKEKS  
 AEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFD  
 30 HVFKENRKVTKDKLLNLYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSFLDDK  
 ANEKTIEDIIQTLTLFEDREMIRQLQKYS DI FTKAQLKKLERLHYTGWGRLSYKLLINGIRN  
 KENKKTILDYLI DDGYANRNFQMLINDDALS FKEEIARAQIIDDVDDIANVHDLPGSPA IK  
 KGILQSVKIVDELVKVMGHNPANII IEMARENQTTDKGRRNSQQRLKLLQDSLKNLNDPNVI



KNVENQQLQNDRLFLYYIQNGKDMYTGETLDINNLQYDIDHIIIPQAFIKDNSLDNRVLTRS  
 DKNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLV  
 ETRQITKHVAQILDERFNTEFDGNKRRI RNVKIITLKS NLVSNFRKEFELYK VREINDYHHA  
 HDAYLNAVVG NALLLKY P QLEPEFVYGEY PKYNSYRSRKSATEKFLFYSN I L R F F K K E D I Q T  
 5 NEDGEI A W N K E K H I K I L R K V L S Y P Q V N I V K K T E E Q T G G F S K E S I L P K G E S D K L I P R K T K N S Y  
 WDPKKYGGFMQPVVAYSILVFADVEK GKSKKLRKVQDMVGITIMEKKRFEKNPVDFLEQ RGY  
 RNVRL E K I I K L P K Y S L F E L E N K R R R L L A S A K F L Q K G N E L V I P Q R F T T L L Y H S Y R I E K D Y E P E  
 HREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMLF SKNKDAEVSS LAKS FISLLTFT  
 AFGAPRAFNF FGENIARKEYRSVTECLNATLIHQSI TGLYETRIDLSKLGEDGE GADKRTAD  
 10 GSEFESP K K R K V (SEQ ID NO: 96).

11. The Cas9 protein of claim 9, wherein the *Sharpea Cas9* has an amino acid sequence at least 80% identical to

MPKKKRKVGAKNKDIRYSIGLDIGTNSVGVAVMDEHYELLKKG NHHMWGSRLFDAAEPAATR  
 RASRSIRRRYNKRREIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQKNLGNDYKDNY  
 15 NLFIEKDFNDKTYDYKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQSFAK  
 VYEDIIEKLDNTLKKFMSLNDLNDL FVDNDINSMITVLSKIYQRSKADDLLKIMNPTKEER  
 AAYKEFTKALVGLKFNVS KMILAQE V K K D D K D I E L D F S N V D Y D S T V D G L Q A E L G E Y I E F I E M  
 LHSINSWVELQDILGNNSTISAAMVERYEEHKNDLRVLK KVI REELPKYNEVFREDNP KLH  
 NYLGYIKYPKNTPVEEFY EYIKRLLAKVDTGEAREILERIDLEK FMLKQNSRTNGSIPYQMQ  
 20 KDEMIQIIDNQSVYYPQLKENREKLISILEFRIPY YFGPLNTHSEFAWIKKFEDKQKERILP  
 WNYDQIVDIDATAEGFIERMQNTGTYPDPKPVMAKNSLTVSKFEVLNELNKIRINGKLI PVE  
 TKKELLSDFMKNKTITDKKLDWLVTHQY YDTNEELKIEGYQKDLQFSTSLAPWIDFTKIF  
 GEINASNYQLIEKIIYDISIFEDK KILKRR LK K V Y Q L D D L L V D K I L K L N Y T G W S R L S E K L L T  
 GIKSKNSKETILSILENSNMNLMEIINDESLGFKQIIEESNKKDIEGPF RYDEVKKLAGSPA  
 25 IKRGIWQALLVQEI TKFMKHEP SHIYIEFAREEQE KVRTESRIAKLQKIYKDLNLQTKEDQ  
 LVYESLKKEDAKKKIDTDALYLYYLQMGKSMYS GKPLDIDKLSTYHIDHILPRSLIKDDSLD  
 NRVLVLPKENEWKLDSETVPFEIRNKMMGFWQKLHENG LMSNKKFFSLIRTD FNEKDKKRFI  
 NRQLVETRQI IKNVAVIINDHYTNTNVVTVRAELSHQFRERYKIYKNRDLNDLHHAHDAYIA  
 CILGQFIHQNFGNMDVNMIYGQYKKNYK KDVQEHN NYGFILNSMNH IHFNDDNSVIWDPSYI  
 30 G K I K S C F C Y K D V Y V T K K L E Q N D A K L F D L T I L P S D K N S E N G V T K A K I P V N K Y R K D V N K Y G G F S  
 GDAPIMLAIEADKGGKHVRQVIAFPLRLKNYNDEERIKFIEKEKNLKNVKILTEVKK NQLIL  
 INHQYFFITGTNELVNATQLKLSAKNTKNLFNLVDANKHNKLESIDDANFNEVIQELICKLQ  
 EPIYSRYNSIGKEFEDSYEKINAVTKQDKLYIIEYLIAIMSAKATQGYIKPELAREIGTNGK

NRGRIKSFETIDLNKTTFISTSVTGLFSKYYKLGKRPAAATKKAGQAKKKKGSYPYDVPDYAYP  
YDVPDYAYPYDVPDYA (SEQ ID NO: 5).

12. The Cas9 protein of claim 9, wherein the *Veillonella parvula* Cas9 has an amino acid sequence at least 80% identical to

5 MPKKKRKVGSIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLDIGTSSVGWAVTNKAYE  
LLKFRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRLLKLEELFADAMAQVDPTFFMR  
LRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKFLAVH  
HILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGKTKSDK  
AKAIEHLVDTYIATDVTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEED  
10 LKKLQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPGLTISESKVKA FNKH  
KDDLAILKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGL  
SDSKDKEYILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLNEVADGFSV  
AEKLIKMLEFRIPIYVGPLNTHHNVDNNGGFAWAVRKASGRVTPWNFDDKIDREKSAAAFIKN  
LTNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTK  
15 NRIEQFLKDNNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFG  
ESKMLRETLRKKFASCLDDEAIKLTCLRDRWGRLSQKLLNGIEGCDKAGDGTPETIIIL  
MRNFSYNMELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVD  
EVAHIKKALPARI FVEVTRSNKNEKKKKDSRQKRLSDLYAAIKKDDVLLNGLNNEIFGELKS  
SLAKYDDAALRSKKLYLYYTQMGRCAYTGEIIELSLLNTDNYDIDHIYPRSLTKDDSDNLV  
20 LCKRTANAQKSDAYPISEEIQKTQKPFWTF LKQOGLISERKYERLTRITPLTADDLSGFIAR  
QLVETNQSVKAATTLRRLYPGVDVVFVKAENVTFDRHDNNFIKVRSLNHHHHAKDAYLNIV  
VGNVYHERFTRNFRAF FKKNGANRTYNLAKMFNYDVNCTNAKDGKAWDVKTSMDTVKKMMDS  
NDVRVTKRLL EQTGALADATIYKATVAGKAKDGAYIGMKTSSVFADVSKYGGMTKIKNAYS  
IIVQYTGKKGEVIKEIVPLPIYLTNRNTDQDLINIVASIIIPQAKDISIIYGKLCINQLVKV  
25 NGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNMRKDNKDLKANVVSTRALDNK  
HTIEVRIVEEKNIIEFFDYLVSKLKMPIYQKMKGNKAAELSEKGYGLFKKMSLEEQSIHLIEL  
LNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIVGKRPAATKK  
AGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 9).

13. The Cas9 protein of claim 9, wherein the *Ezakiella peruensis* Cas9 has an amino acid sequence at least 80% identical to

MPKKKRKVGTKVKDYIIGLDIGTSSVGWAVTDEAYNVLFNSKKMWGVRLFDDAKTAEERRG  
QRGARRRLDRKKERLSLLQDFFAEVAKVDPNFFLRDLNSDLYMEDKDQKLKSKYTLFNDKD

FKDKNFHKYPTIHHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFI FEGQKFDTKSSFENS  
 LNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSS  
 QKLVDFENPEDFDDSAIKSVDFSTTSFDDKYSYELALGDKIALVNILKEIYDSSILENLL  
 KEADKSKDGNKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDI FRSEKVNNDNYVSYTKS  
 5 SISNNKRVKANKFTDQEAIFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPK  
 IKSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDYGSVCDKIASIMEFRI PYYVGPLNP  
 NSKYAWIKKQKDSEITPWNFKDVVDLDS SREEFIDSLIGRCTYLKDEKVL PKASLLYNEYMV  
 LNELNNLKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGDFK  
 QGLNSYNDFKAIVGDKVSDDYRDKIEEIKLIVLYGDDKSYLQKKIKAGYGYFTDSEIKK  
 10 MAGLNYKDWGRLSKLLTGLEGANKITGERGSI IHFMREYNLNLMEELMSASFTFTEEIQKLN  
 PVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKI FIEMARGKEEVKARKE  
 SRKNQLLKIFYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQLGRCMYSLEP  
 IDISELSSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPI PDEILNKNCYAYWK  
 ILYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTICKNSEIVYSK  
 15 AENASRFRQEFDIVKRAVNDLHHMHDAYINIIVGNVYNTKFTKDPMNFVKKQEKARSYNLE  
 NMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRK  
 NKGSRPLKDKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLPVEREIYAKQKNDKKL  
 SDEIFSKYLKDRFGIEDYRVVYPVVKMRTLKIDGSYYFITGGS DKTLELRSALQLILPKKN  
 EWAIKQIDKSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLFQDDKIENID  
 20 FKFKSMDFEKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGRLLSKTNNIGNYQEFKINQS  
 ITGLFENEVDLLKLGKRPAAATKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA  
 (SEQ ID NO: 15) (D12A mutant in bold).

14. The Cas9 protein of claim 9, wherein the *Lactobacillus fermentum* strain AF15-40LB Cas9 has an amino acid sequence at least 80% identical to

25 MPKKKRKVGKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGTAI GVRLEFEEGKTAAERTFRT  
 TRRRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPE DPAKNQAFIGKLLFPDLLKK  
 NERGYPTLIKMRDEL PVEQRAHY PVTNIYKLRAMINEDRQFDLREVYLAVHHIVKYRGHFL  
 NNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLD RQK  
 AVAKLLEVKVADKEETKRNKQIATAMSKLVLGYKADFATVAMANGNEWKIDLSSETSEDEIE  
 30 KFREELSDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYW THERQLAEVKEYLATQP  
 ASARKEFDQVYNKYIGQAPKEKGFLEKGLKKILSKKENWKEIDELLKAGDFLPKQRTSANG  
 VIPHQMHQQELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVS FRI PYYVGPLVTP  
 EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKD TYLLNEDVLPANSL

LYQKYNVLNELNNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMKTRGVNKPVSVE  
 GLSDPKKFNENLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVEWLT  
 DEQRSALVKKRYKGGWGRSLSKKLLTGIVDENGQRIIDLWNTDQNFMQIVNQPVFKEQIDQLN  
 QKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSI SIEFARNEG  
 5 NKGEITRSRRTQLQKLFEDQAHELKDTSLTEELEKAPDLSDRYFYFTQGGKDMYTGDPIIN  
 FDEISTKYDIDHILPQS FVKDDSLDNRVLVSRAENNKSDRVPAKLYAAKMKPYWNQLLKQG  
 LITQRKFENLTMDVDQTIKYRSLGFVQRQLVETRQVIKLTANILGSMYQEAGTDIIETRAGL  
 TKQLREEFDLPKRVENDYHHAVDAYLTTFAGQYLNRRYPKLRSEFFVYGEYMKFKHGS DLKL  
 RNFNFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLISNETYEETGKYFD  
 10 ASHESSSLYLKNQNKSKLVVPLKNKLQPEYYGAYTGITQGYMVLKLLDKKGGFGVYRIPR  
 YAADILNKCHDEVAYRNKIAEIISSDPRAPKSEVVVPRVLKGTFLVDGEEKFILSSYRYKV  
 NATQLILPVSDIKLIQDNFKALKKLVEMQTKKLEIYDNILRQVDKYKLYDINKFRAKLH  
 DGRSKFVELDDFGQDASKEKVIKILRGLHFGSDLQNLKEIGFGTTPLGQFQVSEAGIRLSN  
 TAFIIFKSPTGLFNRKLYLKNLGRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPY  
 15 DVPDYA (SEQ ID NO: 85).

15. The Cas9 protein of claim 9, wherein the *Peptoniphilus sp. Marseille-P3761* Cas9 has an amino acid sequence at least 80% identical to

MPKKKRKVGEKKNYITIGLDIGTDSVGVAVVKKDDLELVKKRMKVLGNTETNYIKKNLWGSLL  
 FESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFRLNESFLVPEDKAFS  
 20 KNPIFGTLGEDKTYKYTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDT  
 EHIAINENLEQFFESYNALFSEPEIELRKEELIAIENILREKNSRTVKEKRITSFLKDIGRA  
 NKQSPMMAFITLIVGKKAKFKAAFNLEEEISLNLTDSDYDENLEILLNTIGSDFADLFDHAQ  
 RVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKD  
 AQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPN  
 25 QVHLAEMEAILDRQENYYPWLKENREKIIISLLTFRIPYYVGPLADGQSEFAWLERKSDEKIK  
 PWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIAYLDERQKR  
 MNLSSVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKV  
 PGMKSMDDPDNEDLMEEIVKILTVFEDRKMRRKQLEKYKERLSPEQIKELAKKHGTGWGR  
 SKKLLVGIRDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDRRLSFKKTI AELQMI DPS  
 30 ADLYAQVQEIAGSPAIKKGI LLGLKIVDEIIRVMGEKPENIVIEMARENQTTARGKALSRR  
 EAKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHII  
 PQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEVVRQLAYWRS LKDAGLMTQRKFDNLTKA  
 MRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNIVLLKSAMASE

FRKMFRLYKVRELNDYHHAHDAYLNAAIAINLLALYPYMADDFVYGEFRYKKKPKQAEKATYE  
 KLRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNVVKKAEERTGMLFKETIN  
 GKTNKGSRPIPKKDLDPSTKYGGYIEEKMAYYAVISYEDKPKKPGKTIVGISIMDKKEFEYDS  
 ISYLGKLGFSNPVVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKANQIAMEQDLVNF  
 5 IYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIYEEEEKKDP  
 VDIEALVASFIELLKLTSAGAPADFI FMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETMRM  
 FEDGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 87).

16. The Cas9 protein of any one of the preceding claims, wherein the amino acid  
 sequence of the Cas9 protein comprises at least one, at least two, at least three, at least four,  
 10 at least five, at least six, at least seven, at least eight, at least nine, or at least 10 mutations in  
 SEQ ID NO: 1, 4, 8, 14, 84 or 86.
17. The Cas9 protein of claim 16, wherein the mutation is an amino acid substitution.
18. The Cas9 protein of any one of the preceding claims, wherein the Cas9 protein has  
 nickase activity.
- 15 18b. The Cas9 protein of claim 18, wherein the nickase mutation at an amino acid positions  
 corresponds to one or more amino acids 10, 12, 17, 762, 840, 854, 863, 982, 983, 984, 986,  
 987 of wild type SpCas9.
19. The Cas9 protein of claim 16, wherein the at least one mutation results in an inactive  
 Cas9 (dCas9).
- 20 20. The Cas9 protein of any one of the preceding claims, wherein the Cas9 protein  
 comprises at least one amino acid mutation in PAM Interacting, HNH and/or RuvC domain.
- 20b. The Cas9 protein of claim 20, wherein the mutation at an amino acid position  
 corresponds to amino acid 14 in the RuvC domain of SirCas9.
- 20c. The Cas9 protein of claim 20, wherein the mutation at an amino acid position  
 25 corresponds to amino acid 12 in the RuvC domain of EpeCas9.
- 20d. The Cas9 protein of claim 20, wherein the mutation at an amino acid position  
 corresponds to amino acid 9 in the RuvC domain of LfeCas9.

20e. The Cas9 protein of claim 20, wherein the mutation at an amino acid position corresponds to amino acid 12 in the RuvC domain of PmaCas9.

20f. The Cas9 protein of claim 20, wherein the Cas9 protein is a hyper-accurate Cas9.

20g. The Cas9 protein of claim 20, wherein the Cas9 protein comprises mutations  
5 corresponding to N692A, M694A, Q695A and/or H698A with reference to SpyCas9 (SEQ ID NO: 173).

20h. The Cas9 protein of claim 20, wherein the Cas9 protein is a high-fidelity Cas9.

20i. The Cas9 protein of claim 20, wherein the Cas9 protein comprises mutations  
10 corresponding to N467A, R661A, Q695A and/or Q926A with reference to SpyCas9 (SEQ ID NO: 173).

20j. The Cas9 protein of claim 20, wherein the Cas9 protein is a SuperFi-Cas9.

20k. The Cas9 protein of claim 20, wherein Y1016, R1019, Y1010, Y1013, K1031, Q1027  
and/or V1018 residues corresponding to SpyCas9 are mutated to aspartic acid.

21. An engineered, non-naturally occurring Cas9 fusion protein comprising a Cas9  
15 protein having at least 80% identity to SEQ ID NOs: 1, 4, 8,14, 84 or 86 and wherein the Cas9 protein is fused to a histone demethylase, a transcriptional activator, or to a deaminase.

21b. The engineered, non-naturally occurring Cas9 fusion protein of claim 21 further  
comprising a nuclear localization sequence (NLS) and/or a FLAG, HIS or HA tag.

21c. The engineered, non-naturally occurring Cas9 fusion protein of claim 22 having at  
20 least 80% identity to SEQ ID NOs: 2, 5, 9, 15, 85, 87, 95 or 96.

22. The Cas9 protein of claim 21, wherein the Cas9 protein is fused to a cytosine  
deaminase or to an adenosine deaminase.

23. The Cas9 protein of claim 22, wherein the Cas9 protein is fused to a adenosine  
deaminase and has an amino acid sequence at least 80% identical to

25 (a)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPGAVLVLNNRVIGEGWNRAIGLH  
DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA

GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
 TSESATPESSGPKKKRKGKVPYSIGLAIGTNSVGWAVVTDDYKVPKMKVKGNTDKQSIK  
 KNLLGALLFDSGETAEATRLKRTARRRYTRRKNRLRYLQEI FTGEMNKVDENFFQRLDDSF  
 VDEDKRGEHHP IFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHF  
 5 LYEGDLKAENTDVQALFKDFVEEYDKTIEESHLS EITVDALSILTEKVS KSSRLENLIAHYP  
 TEKNTLFGNLIALS LDLPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGD EYADLFAS  
 AKNLYDAILLSGILTVDDNSTKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYN AIFKDK  
 NKKGYASYIESGVKQDEFYKYLK GILLKINGS GDFLDKIDREDFLRKQRTFDNGSIPHQIHL  
 QEMHAILRRQGEHY PFLKENQDKIEKILTFRIPIYYVGPLARKGSRFAWA EYKADEKITPWNF  
 10 DDILDKEKSAEKFITRMTLNDLYLP EEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFD  
 TNMKQEIFDHVFKENRKVTKDKLLNYLNKEFEFRIVNLTGLDKENKAFNSSLGTYHDLRKI  
 LDKSFLDDKAN EKTIEDIIQTLTLFEDREMIRQRLQKYS DI FTKAQLKKLERRHYTGWGRLS  
 YKLINGIRNKENKKTILDYLI DDGYANRNF MQLINDDALSFKEEIARAQIIDDVDDIANVVH  
 DLPGPSAIIKKGILQSVKIVDELVKVMGHN PANII IEMARENQTTDKGRNSQQRLKLLQDSL  
 15 KNLDNPVNIKNVENQQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHII PQAFIKDNS  
 LDNRVLTRS DKNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGLTEEDK  
 AGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKS NLVSNFRKEFELYKV  
 REINDYHHAHDAYLNAVVG NALLKYPQLEPEFVYGEYPKYNSYRSRKSATEKFLFYSN ILR  
 FFKKEDIQTNE DGEIAWNKEKH I KILRKVLSYPQVNI VKKTEEQTGGFSKESILPKGESDKL  
 20 IPRKTKNSYWDPKKYGGFDS PVVAYSILVFADVEK GSKKLRKVQDMVGITIMEKKRFEKNP  
 VDFLEQRGYRNVRL EKI I KLPKYSLFELENKRRRL LASAKELQGNELVIPQRFTTLLYHSY  
 RIEKDYEP EPREYVEKHKDEFKELLEYISVFSRKYVLADNNLT KIEMLFSKNKDAEVSSLAK  
 SFISLLTFTAFGAPAAFNFFGENIDRKRYTSVTECLNATLIHQ SITGLYETRIDL SKLGEDG  
 KRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 20);

25 (b)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH  
 DPTAHAEIMALRQGLVMQNYRLYDATLYVT FEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA  
 GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
 TSESATPESSGPKKKRKGAKNKDIRYSIGLAIGTNSVGWAVMDEHYELLKKNHMMWGSRL  
 30 FDAAEPAATR RASRSIRRRYNKRRERIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQK  
 NLGNDYKDNYNLFIEKDFNDKTYDYKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGN  
 FLKEGQSFAKVYEDIEEKLDNTLKKFMSLNDLNDL FVDNDINSMITVLSKIYQRSKKADDLL  
 KIMNPTKEERAAYKEFTKALVGLKFNVSKMILAQEVKKDDKDIELDFSNVDYDSTVDGLQAE

LGEYIEFIEMLHSINSWVELQDILGNNSTISAAMVERYEEHKNDLRVLKVKVIREELPKYNE  
 VFREDNPKLHNYLGYIKYPKNTPVEEFYIYIKRLLAKVDTGEAREILERIDLEKFMKQNSR  
 TNGSIPYQMOKDEMIQIIDNQSVYYPQLKENREKLISILEFRIPIYFGLNTHSEFAWIKKF  
 EDKQKERILPWNVDQIVDIDATAEGFIERMQNTGTYFPDKPVMKNSLTVSKFEVLNENLKI  
 5 RINGKLI PVETKKELLSDFMKNKTITDKKLDWLVTHQYYDTNEELKIEGYQKDLQFSTSL  
 APWIDFTKIFGEINASNYQLIEKIYDISIFEDKKILKRRLKVKYQLDDLLVDKILKLNITG  
 WSRLSEKLLTGIKSKNSKETILSILENSNMNLMEIINDESLGFKQIIEESNKKDIEGPFYD  
 EVKLAGSPAIRGIWQALLVVQEITKFMKHEPSHIYIEFAREEQEKVRTESRIAKLQKIYK  
 DLNLQTKEDQLVYESLKKEDAKKIDTDALYLYYLQMGKSMYSGKPLDIDKLSTYHIDHILP  
 10 RSLIKDSDLDNRVLPKENEKLDSETVPEIRNKMMGFQKLEHENGLEMSNKKFFSLIRTD  
 FNEKDKKRFINRQLVETRQIIKNVAVIINDHYTNTNVTVRAELSHQFRERYKIYKNRDLND  
 LHHADAYIACILGQFIHQNFNMDVNMIYGQYKKNYKQDVQEHNNYGFILNSMNIHFND  
 NSVIWDPSYIGKIKSCFCYKDVYVTKKLEQNDAKLFDLTILPSDKNSENGVTAKAPIVKNYK  
 KDVNKYGGFSGDAPIMLAIEADKGGKHVRQVIAFPLRLKNYNDEERIKFIEKEKNLKNVKIL  
 15 TEVKNQLILINHQYFFITGTNELVNATQLKLSAKNTKNLFDANKHNKLESIDDANFNE  
 VIQELICKLQEPYISRYNSIGKEFEDSYEKINAVTKQDKLYIIEYLIAIMSAKATQGYIKPE  
 LAREIGTNGKNKGRIKSFTIDLNKTTFISTSVTGLFSKKYKLGKRPAAATKAGQAKKKKGSY  
 PYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 6);

(c)

20 MPAARVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPGAVLVLNNRVIGEGWNRAIGLH  
 DPTAHAEIMALRQGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA  
 GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
 TSESATPESSGPKKKRKGVSIIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLAIGTSSV  
 GWAVTNKAYELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRLKLEELFADAM  
 25 AQVDPTFFMRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTD  
 DIRKLFLAVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFCNSAIISSIGQIL  
 MEAGKTKSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDL  
 FGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPGLTIS  
 ESKVKA FNKHKDDLA I LKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFY  
 30 KYTKKIVEGLSDSKDKEYILSQIELQILLPLQRIKDNQVPIPYQLHLEELKAILAKCGPKFPF  
 LNEVADGFSVAEKLIKMLEFRIPIYVGPLNTHHNVNDGGFAWAVRKASGRVTWNFDDKIDR  
 EKSAAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNVRIDGKPLEKVVKEHLIEAV  
 FKQDHKKMTKNRIEQFLKDNQYISETHKHEITGLDGEIKNDLASYRDMVRI LGDGFDRSMAE



EIITDITIFGESKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAG  
 DGT PET I I I LMRNFSYNLMELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKR  
 AVWQALRIVDEVAHIKKALPARI FVEVTRS NKNEKKKDSRQKRLSDLYAAIKKDDVLLNGL  
 NNEIFGELKSSLAKYDDAALRSK KLYLYYTQMGRCAYTGEI I ELSLLNTDNYDIDHIYPRSL  
 5 TKDDSFDNLVLCRRTANAQKSDAYPISEEIQKTQKPFWTFLKQQGLISERKYERLTRITPLT  
 ADDLSGFIARQLVETNQSVKAATLLRRLYPGVDVVFVKAENVTD FRHDN NFIKVRSLNHHH  
 HAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKD GKAWDVKTS  
 MDTVKKMMSDNDVRVTKRLL EQTGALADATIYKATVAGKAKDGAYIGMKT KSSVFADVSKYG  
 GMTKIKNAYSIIVQYTGKKGEVIKEIVPLPIYL TNRTTDQDLINVASIIPQAKDISIIYG  
 10 KLCINQLVKVNGFY YLGGKTNSKFCIDNAIQVIVSNEWIPYLVLEKFNNMRKDNKDLKAN  
 VVSTRALDNKHTIEVRIVEEKNI EFFDYLVSKLMP IYQMKGNKAAELSEKGYGLFKKMSL  
 EEQSIHLIELLNL TNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTI  
 VGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 10);

(d)

15 MPKKRKRKVS I INFQRRGLMETQASNQLISSHLKGYPIKDYFVGLAIGTSSVGWAVTNKAYEL  
 LKFRSHKMWGSRLFDEGESAVARRGFRSMRRLERRKLRLKLLLEELFADAMAQVDPTFFMRL  
 RESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHH  
 ILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFCNSAIISSIGQILMEAGKTKSDKA  
 KAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEDL  
 20 KKLQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAI ILLSIKEPGLTISESKVKA FNKHK  
 DDLAILKSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETS CNREDFYKYTKKIVEGLS  
 DSKDKEYILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLNEVADGF SVA  
 EKLIKMLEFRIPIYYVGPLNTHHNVDNGGFAVA VRKASGRVTPWNFDDKI DREKSAAAFIKNL  
 TNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTKN  
 25 RIEQFLKDNNGYISETHKHEITGLDGEIKNDLAS YRDMVRILGDGFDRSMAEEIITDITIFGE  
 SKKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGT PET I I I LM  
 RNFSYNLMELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVDE  
 VAHIKKALPARI FVEVTRS NKNEKKKDSRQKRLSDLYAAIKKDDVLLNGLNNEIFGELKSS  
 LAKYDDAALRSK KLYLYYTQMGRCAYTGEI I ELSLLNTDNYDIDHIYPRSLTKDDSFDNLVL  
 30 CKRTANAQKSDAYPISEEIQKTQKPFWTFLKQQGLISERKYERLTRITPLTADDLSGFIARQ  
 LVETNQSVKAATLLRRLYPGVDVVFVKAENVTD FRHDN NFIKVRSLNHHHHAKDAYLNIVV  
 GNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKD GKAWDVKTSMDTVKKMMSDN  
 DVRVTKRLL EQTGALADATIYKATVAGKAKDGAYIGMKT KSSVFADVSKYGGMTKIKNAYS I

IVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISIIYGKLCINQLVKVN  
 GFYYLGGKTNSKFCIDNAIQVIVSNEWIPYLKVLKFNMRKDNKDLKANVSTRALDNKH  
 TIEVRIVEEKNIEFFDYLVSKLKMPIYQMKGNKAAELSEKGYGLFKKMSLEEQSIHLIELL  
 NLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIVKRPAATKKAG  
 5 QAKKKKSGSETPGTSESATPESSGSEVEFSHEYWMRHALTLAKRARDEREVPVGA VLVLNLR  
 VIGEGWNRAIGLHDPTAHAEIMALRQGG LVMQNYRLYDATLYVT FEPCVMCAGAMIHSRIGR  
 VVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQ  
 SSTDPAAKRVKLDGSPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 11);

(e)

10 MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGA VLVLNLRVIGEGWNRAIGLH  
 DPTAHAEIMALRQGG LVMQNYRLYDATLYVT FEPCVMCAGAMIHSRIGR VVFGVRNAKTGA  
 GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
 TSESATPESSGPKKKRKGVTKVVDYIIGLAIGTSSVGWAVTDEAYNVLKFNSKMMWGVRLFD  
 DAKTAEERRGQRGARRRLDRKKERLSLLQDFFAEVAKVDPNFFLRDLNDSLIMEDKDQKLL  
 15 SKYTLFNDKDFKDNFHKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFI FEGQK  
 FDTKSSFENSLNELKVHLNDEYGLDLEFDNENLINILTPKLNKTAKKELKSVIGDTKFLK  
 AVSAIMIGSSQKLVDLFENPEDFDDSAIKSVDFSTTSFDDKYSYELALGDKIALVNILKEI  
 YDSSILENLLKEADKSKDGNKYISNAFVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKV  
 NDNYVSYTKSSISNNKRVKANKFTDQEA FYKFAKKHLETIKYKINKVNGSKADLELIDGMLR  
 20 DMEFKNFMPKIKSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFR  
 IPYVYVGPLNPN SKYAWIKKQKDSEITPWNFKDVVDLSSREEFIDSLIGRCTYLKDEKVLPK  
 ASLLYNEYMVLNELLNKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEI  
 LLSGTDGDFKQGLNSYNDFKAI VGDKVDSDDYRDKIEEIIKLVLYGDDKSYLQKKIKAGYG  
 KYFTDSEIKKMAGLNYKDWGRLSKLLTGLEGANKITGERGSI IHFMREYNLNLME LMSASF  
 25 TFTEEIQKLN PVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIE MAR  
 GKEEVKARKESRKNQLLK FYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQ  
 LGRCMYSLEPIDISELSSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPIPDEI  
 LNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTI  
 CKNSEIVYSKAENASRFRQEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKDP MNFVKK  
 30 QEKARSYNLENMFKYDVKRGGYTAWIADDEKGT VKNASIKRIRKELEGTNYRFTRMNYIESG  
 ALFNATLQRKNKGSRPLKDKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLMPVEREI  
 YAKQKNDKLSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSYFITGGS DKTLELRS  
 ALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSF LNL

FQDDKIENIDFKFKSMDFKKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGRLLS SKTNNIGN  
YQEFKIINQSI TGLFENEVDLLKLGKRPAATKKAGQAKKKKGSYPYDVPDYAYPYDVPDYAY  
PYDVPDYA (SEQ ID NO: 16);

(f)

5 MPKKKRKVTKVVDYYIGLAIGTSSVGVAVTDEAYNVLKFNSKKMWGVRLFDDAKTAEERRGQ  
RGARRRLDRKKERLSLLQDFFAEVAVKVDPNFFLRDNLSDLYMEDKDQKLKSKYTLFNDKDF  
KDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFI FEGQKFDTKSSFENSL  
NELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQ  
KLVDLFENPEDFDDSAIKSVDFSTTSFDDKYSYELALGDKIALVNILKEIYDSSILENLLK  
10 EADKSKDGNKYISNAFVKYKNGHQDLKEFKRLVRQYHKSAYFDIFRSEKVNNDYVSYTKSS  
ISNNKRVKANKFTDQEAIFYKFAKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKI  
KSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDYGSVCDKIASIMEFRIPIYVGPLNPN  
SKYAWIKKQKDEITPWNFKDVVDLSDSREEFIDSLIGRCTYLKDEKVLPKASLLYNEYMVL  
NELNNLKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGDFKQ  
15 GLNSYNDFKAI VGDKVSDDYRDKIEEIIKLVLYGDDKSYLQKKIKAGYGYFTDSEIKKM  
AGLNYKDWGRLSKKLLTGLEGANKITGERGSI IHFMREYNLNMELMSASFTFTEEIQKLN  
VDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFEMARGKEEVKARKES  
RKNQLLFYKDGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQLGRCMYSLEPI  
DISELSSKNIYDQDHIYPKSKIYDDSIENRVLVKKDLNSKKGNSYPIPDEILNKNCYAYWKI  
20 LYDKGLIGQKQYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTICKNSEIVYSKA  
ENASRFRQEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKDPMNFVKKQEKARSYNLEN  
MFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRKN  
KGSRPLKDKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLMVVEREIIYAKQKNDKCLS  
DEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITGGSDKTELELSALQLILPKNE  
25 WAIKQIDKSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFNLNFQDDKIENIDF  
KFKSMDFKKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGRLLS SKTNNIGNYQEFKIINQSI  
TGLFENEVDLLKLGKRPAATKKAGQAKKKKSGSETPGTSESATPESSGSEVEFSHEYWMRHAL  
TLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMALRQGGGLVMQNYRLYDA  
TLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADE  
30 CAALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPYDVPDYAYPYDVPDYAYPYDVP  
DYA (SEQ ID NO: 17);

(g)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLH

DPTAHAEIMALRQGGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA  
 GSLMDVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG  
 TSESATPESSGPKKKRKGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGK  
 TAAERRTFRTTRRRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEPDAKNQAFIG  
 5 KLLFPDLLKKNERGYPTLIKMRDELVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAVH  
 HIVKYRGHFLNNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLD  
 TKMRKLDLDRQKAVAKLLEVKVADKEETKRNKQIATAMSKLVLGYKADFATVAMANGNEWKIDL  
 SSETSEDEIEKFREELESDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWOTHERQLA  
 EVKEYLATQPASARKEFDQVYNKYIGQAPKEKGFDFLEKGLKKILSKKENWKEIDELLKAGDF  
 10 LPKQRTSANGVIHQMHQQLDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSRIFRI  
 PYYVGPLVTPPEVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKDITYLL  
 NEDVLPANSLLYQKYNVLNELNVRVNGRRLSVGKQDIYTELFKKKTKVAGDVASLVMK  
 TRGVNKPVSVEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFA  
 DKLTEVEWLTDEQRSALVKKRYKGGWRLSKLLTGIVDENGQRIIDLWNTDQNFMQIVNQP  
 15 VFKEQIDQLNQKAITNDGMTLRERVESVLD DAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKS  
 ISIEFARNEGKGEITRSRRTQLQKLFEDQAHELVKDTSLTEELEKAPDLSDRYYFYFTQGG  
 KDMYTGDPINFDEISTKYDIDHILPQS FVKDDSLDNRVLVSR AENNKSDRVPKLYAAKMK  
 PYWNQLLKQGLITQRKFENLTMDVDQTIKYRSLGFVKRQLVETRQVIKLTANILGSMYQEAG  
 TDIETRAGLTKQLREEFDLPKVREVNDYHHAVDAYLTFAGQYLNRYPKLRSEFFVYGEYM  
 20 KFKHGSDDLKLRNFNFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWINLKVMLISNE  
 TYEETGKYFDASHESLKYLNQNKSKLVLVPLKKNLQPEYYGAYTGITQGYMVLKLLDKK  
 GFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKSEVVVPRVLKGTFLVDGEEK  
 FILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLNEMQTKKLEIYDNI LRQVDKYKLY  
 DINKFRAKLHDGRSKFVELDDFGQDASKEKVIKILRGLHFGSDLQNLKEIGFGTTPLGQFQ  
 25 VSEAGIRLSNTAFIIFKSPTGLFNRKLYLKNLGKRPAAATKKAGQAKKKKGSYPYDVPDYAYP  
 YDVPDYAYPYDVPDYA (SEQ ID NO: 88);

(h)

MPKKKRKVGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAAERRTFRT  
 TRRRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEPDAKNQAFIGKLLFPDLLKK  
 30 NERGYPTLIKMRDELVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAVHHIVKYRGHFL  
 NNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLD TKMRKLDLDRQK  
 AVAKLLEVKVADKEETKRNKQIATAMSKLVLGYKADFATVAMANGNEWKIDLSSETSEDEIE  
 KFREELESDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWOTHERQLAEVKEYLATQP

ASARKEFDQVYNKYIGQAPKEKGFLEKGLKKILSKKENWKEIDELLKAGDFLPKQRTSANG  
 VI PHQM HQQELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPYVVGPLVTP  
 EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEAFIKRMTVKDTYLLNEDVLPANSL  
 LYQKYNVLNELLNNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMKTRGVNKPVSVE  
 5 GLSDPKKFNENLATYLDLKSIVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVEWLT  
 DEQRSALVKKRYKGGWRLSKKLLTGIVDENGQRIIDL MWNTDQNF MQIVNQPVFKEQIDQLN  
 QKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSSISIEFARNEG  
 NKGEITRSRRTQLOKLFEDQAHELVKDTSLTEELEKAPDLSDRYFYFTQGGKDMYTGD PIN  
 FDEISTKYDIDHILPQS FVKDDSLDNRVLVSRAENNKSDRVPAKLYAAKMKPYWNQLLKQG  
 10 LITQRKFENLTMDVDQTIKYRSLGFVKRQLVETRQVIKLTANILGSMYQEAGTDIIETRAGL  
 TKQLREEFDLPKRVENDYHHAVDAYLTTFAGQYLNRRYPKLRSEFFVYGEYMKFKHGS DLKL  
 RNFNFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLISNETYEETGKYFD  
 ASHESSSLYLKNQNKSKLVVPLKNKLQPEYYGAYTGITQGYMVLKLLDKKGGFGVYRIPR  
 YAADILNKCHDEVAYRNKIAEIISSDPRAPKSEVVVPRVLKGTFLVDGEEKFILSSYRYKV  
 15 NATQLILPVSDIKLIQDNFKALKKLNEMQTKKLIIEIYDNILRQVDKYYKLYDINKFRAKLH  
 DGRSKFVELDDFGQDASKEKVIKILRGLHFGSDLQNLKEIGFGTTP LGQFQVSEAGIRLSN  
 TAFIIFKSP TGLFNRKLYLKNLKRPAATKAGQAKKKKSGSETPGTSESATPESGSEVEFS  
 HEYWMRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMALRQGGVL  
 MQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVE  
 20 ITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPYDVPDYAYPYDVP  
 DYAYPYDVPDYA (SEQ ID NO: 89); or

(i)

MSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMA  
 LRQGGVLVMQNYRLYDATLYSTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHP  
 25 GMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDSGGSSGGSSGSETPGTSES  
 ATPESGSSGGSGKPYSIGLAIGTNSVGWAVVTDDYKVPKMKV LGNTDKQSIKKNLLGA  
 LLFDSGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKR  
 GEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDL  
 KAENTDVQALFKDFVEEYDKTIEESHLSEITVDALSILTEKVS KSSRLENLIAHYPT EKNT  
 30 LFGNLIALS LLDLHPNFKTNFQ LSEDAKLQFSKDTYEEDLEGFLGEVGD EYADLFASAKNLYD  
 AILLSGILTVDNSTKAPLSASMVKRYEEHQKDLKCLKDFIKVNAPDQYN AIFKDKNKKGYA  
 SYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAI  
 LRRQGEHY PFLKENQDKIEKILTFRIPYVVGPLARKGSRFAWAEYKADEKITPWNFDDILDK

EKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQE  
 IFDHVFKENRKVTKDKLLNYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKIILDKSFL  
 DDKANEKTIEDI IQTLTLFEDREMIRQLQKYSDI FTKAQLKKLERRHYTGWGRLSYKLING  
 IRNKENKKTILDYLI DDGYANRNFQMLINDDALS FKEEIARAQI IDDVDDIANVVHDLPGSP  
 5 AIKKGILQSVKIVDELVKVMGHN PANI I IEMARENQT TDKGRNSQQRLKLLQDSLKNLNDP  
 VNIKNVENQQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHI IPQAFIKDNSLDNRVL  
 TRSDKNRGKSDDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKR  
 QLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKSNLVSNFRKEFELYKVREINDY  
 HHAHDAYLNAVVGNALLLKYPQLEPEFVYGEYPKYNSYRSRKSATEKFLFYSNILRFFKED  
 10 IQTNEDGEIAWNKEKH IKILRKVLSYPQVNIVKKTTEEQTGGFSKESILPKGESDKLIPRKT  
 NSYWDPKKYGGFMQPVVAYSILVFADVEKGKSKKLRKVQDMVGITIMEKKRFEKNPVDFLEQ  
 RGYRNVRLKIKL PKYSLFELENKRRRLASAKFLQKGNELVIPQRFTTLLYHSYRIEKDY  
 EPEHREYVEKHKDEFKELLEYSVFSRKYVLADNNLTKIEMLFSKNKDAEVSSLAKSFISLL  
 TFTAFGAPRAFNFFFGENIARKEYRSVTECLNATLIHQSI TGLYETRIDL SKLGEDGEGADKR  
 15 TADGSEFESPKKKRKV (SEQ ID NO: 98);

(j)

MSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVI GEGWNRAI GLHDPTAHAE IMA  
 LRQGGGLVMQNYRLYDATLYSTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHP  
 GMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDSGGSSGGSSGSETPGTSES  
 20 ATPESSGGSSGGSGKPYSIGLAIGTNSVGWAVVTDDYKVPKMKVVLGNTDKQS I KKNLLGA  
 LLFDSGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDS FLVDEDKR  
 GEHHP IFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDL  
 KAENTDVQALFKDFVEEYDKTIEESHLS EITVDALSILTEKVS KSSRLENLIAHYPT EKNT  
 LFGNLI ALSLDLHPNFKTNFQ LSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYD  
 25 AILLSGILTVDN STKAPLSASMVKRYEEHQKDLKKLKD FIKVNAPDQYNAIFKDKNKKGYA  
 SYIESGVKQDEFYKYLK GILLKINGSGDFLDKIDREDFLRKQRTFDNGIIPHQIHLQEMHAI  
 LRRQGEHY PFLKENQDKIEKILTFRI PYYVGPLARKGSRFAWAEYKADEKITPWNFDDI LDK  
 EKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQE  
 IFDHVFKENRKVTKDKLLNYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKIILDKSFL  
 30 DDKANEKTIEDI IQTLTLFEDREMIRQLQKYSDI FTKAQLKKLERLHYTGWGRLSYKLING  
 IRNKENKKTILDYLI DDGYANRNFQMLINDDALS FKEEIARAQI IDDVDDIANVVHDLPGSP  
 AIKKGILQSVKIVDELVKVMGHN PANI I IEMARENQT TDKGRNSQQRLKLLQDSLKNLNDP  
 VNIKNVENQQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHI IPQAFIKDNSLDNRVL

TRSDKNRGKSDVPSIEVVHEMKS FWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKR  
 QLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKSNLVSNFRKEFELYKVREINDY  
 HHAHDAYLNAVVGNALLLKYPQLEPEFVYGEYPKYNSYRSRKSATEKFLFYSNILRFFKED  
 IQTNEDGEIAWNKEKH IKILRKVLSYPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKT  
 5 NSYWDPKKYGGFMQPVVAYSILVFADVEKGGKSKLKRKQDMVGITIMEKKRFEKNPVDFLEQ  
 RGYRNVRLKIKIKLPKYSLFELENKRRRLASAKFLQKGNELVIPQRFTTLLYHSYRIEKDY  
 EPEHREYVEKHKDEFKELLEYSVFSRKYVLADNNLTKIEMLF SKNKDAEVSSLAKSFISLL  
 TFTAAGAPRAFNFNGENIARKEYRSVTECLNATLIHQSI TGLYETRIDL SKLGEDGEGADKR  
 TADGSEFESPKKKRKV (SEQ ID NO: 99).

10 24. The Cas9 protein of claim 22, wherein the Cas9 protein is fused to a cytosine  
 deaminase and has an amino acid sequence at least 80% identical to

(a)

MPAARVKLDTSEKGPSTGDPTLRRRIESWEFDVFDPRELRKETCLLYEIKWGMSRKIWR  
 SGKNTTNHVEVNFIIKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVII  
 15 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYPPLWML  
 YALELHCIIISLPPCLKISRRWQNHLAFFRLHLQNCYQTI PPHILLATGLIHPSVTWRLKS  
 GGSSGGSSGSETPGTSESATPESSGGSSGSPKKKRKVGKPYSIGLAIGTNSVGWAVVTD  
 YKVPAKKMKVLGNTDKQSIKKNLLGALLFDSGETAEATRLKRTARRRYTRRKNRLRYLQEIF  
 TGEMNKVDENFFQRLDDSFLVDEDKRGEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSK  
 20 ADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFKDFVVEYDKTIEESHLSSEITVDALS  
 ILTEKVS KSRLENLIAHYPTKNTLFGNLI ALSLDLHPNFKTNFQLSEDAKLQFSKDTYE  
 EDLEGFLGEVGDYADLFASAKNLYDAILLSGILTVDDNSTKAPLSASMVKRYEEHQDLKK  
 LKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEFYKYLKGI LLKINGS GDFLDKI DRE  
 DFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHY PFLKENQDKIEKILTFRIPYVYVGLARK  
 25 GSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTV  
 YNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTKDKLLNYLNKEFEEFRIVNLTGL  
 DKENKAFNSLGTYHDLRKI LDKSFLDDKAN EKTI EDIIQTTLTFEDREMIRQLQKYS DIF  
 TKAQLKKLERRHYTGWGRLSYK LINGIRNKENKKTILDYLI DDGYANRNF MQLINDDALSFK  
 EEIARAQIIDDVDDIANVVHDLPGSPA I KKGILQSVKIVDELVKVMGHNPANII IEMARENQ  
 30 TTDKGRNSQQLRLLQDSLKNLDNPVNIKNVENQQQLQNDRLFLYYIQNGKDMYTGETLDIN  
 NLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSDVPSIEVVHEMKS FWSKLLSVKLIT  
 QRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKI  
 ITLKS NLVSNFRKEFELYKVREINDYHHAHDAYLNAVVGNALLLKYPQLEPEFVYGEYPKYN

SYRSRKSATEKFLFYSNILRFFKKEDIQTNEDEGIAWNKEKHIKILRKVLSYPQVNIVKKTE  
 EQTGGFSKESILPKGESDKLI PRKTKNSYWDPKKYGGFDSPVVAYSILVFADVEKGKSKKLR  
 KVQDMVGITIMEKKRFEKNPVDFLEQRGYRNVRLKIIKLPKYSLELENKRRRLLASAKEL  
 QKGNELVIPQRFTTLLYHSYRIEKDYEPHREYVEKHKDEFKELLEYISVFSRKYVLADNNL  
 5 TKIEMLF SKNKDAEVSSLAKSFISLLTFTAAGAPAAAFNFFGENIDRKRYTSVTECLNATLIH  
 QSITGLYETRIDL SKLGEDGKRPAATKKAGQAKKKKGSSGGSSGGSGGSTNLSDIIEKETGKQ  
 LVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNG  
 ENKIKMLSGSSGGSSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTA  
 YDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAYPYDVPDYAYPYDVP  
 10 DYA (SEQ ID NO: 21);

(b)

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVIFYDPREL RKETCLLYEIKWGMSRKI WRS  
 SGKNTTNHVEVNFIIKFKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIIY  
 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYPPLWMMML  
 15 YALELHCII LSLPCLKISRWRQNHLAFFRLHLQNCYQTI PPHILLATGLIHPSVTWRLKS  
 GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKKRKVGSIINFQRRGLMETQASNQLISSH  
 LKGYPIKDYFVGLAIGTSSVGWAVTNKAYELLKFRSHKMWGSRLFDEGESAVARRGFRSMRR  
 RLERRKLRLKLEELFADAMAQVDPTFFMRLRESKYHYEDKTTGHSKHI LFI DKNYNDQDY  
 FKEYPTVYHLRSELMKSGTDDIRKLFLAVHHILKYRGNFLYEGATFDSNASTLDDVIKQALE  
 20 NITFNCFCDCNSAIISSIGQILMEAGKTKSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDK  
 KRLKAFANLVLGLNASLIDLFGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDDCC  
 KSVYDAI ILLSIKEPGLTISESKVKAFNKHKDDLAILKSLKSDRSIYNTMFKVDEKGLHNY  
 VHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEYIILSQIELQILLPLQRIKDNQVPIPY  
 QLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKMLEFRIPIYVGPLNTHHNVDNNGGFAW  
 25 AVRKASGRVT PWNFDDKIDREKSAAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNN  
 VRIDGKPLEKVVEHLIEAVFKQDHKMTKNRIEQFLKDNGYISETHKHEITGLDGEIKNDL  
 ASYRDMVRI LGDGFDRSMAEEIITDITIFGESKMLRET LRKKFASCLDDEAIKKLTKLRYS  
 DWGRLSQKLLNGIEGCDKAGDGPETII IILMRNFSYNLMELLGDKFSFMERIQEINAKLTEG  
 QIVNPHDIIDDLALSPAVKRAVWQALRIVDEVAHIKKALPARI FVEVTRSNKNEKKKKDSRQ  
 30 KRLSDLYAAIKKDDVLLNGLNNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGRCAYTGEII  
 ELSLLNTDNYDIDHIYPRSLTKDSDFDNLVLCRRTANAQKSDAYPISEEIQKTQKPFWTFLK  
 QQGLISERKYERLTRITPLTADDLSGFIARQLVETNQSVKAATTLLRRLYPGVVVVKAEN  
 VTDFRHDNNFIKVRSLNHHHHAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAKMF



NYDVNCTNAKDGKAWDVKTSMDTVKKMMSNDVRVTKRLLLEQTGALADATIYKATVAGKAKD  
 GAYIGMKTSSVFADVSKYGGMTKIKNAYSIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQD  
 LINYVASIIPQAKDISIIYGKLCINQLVKVNGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPY  
 LKVLEKFNNMRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLVSCLKMPIYQKMK  
 5 GNKAAELSEKGYGLFKMSLEEQSIHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQ  
 DEFKVINESITGLYSNEVTIVGKRPAATKKAGQAKKKKSSGGSSGGSGGSTNLSDIIEKETG  
 KQLVIQESILMLPEEVEEVIIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDS  
 NGENKIKMLSGGSSGGSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIIGNKPESDILVH  
 TAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAYPYDVPDYAY

10 (SEQ ID NO: 12);

(c)

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFDPRELRKETCLLYEIKWGMSRKIWS  
 SGKNTTNHVEVNFIIKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIIY  
 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHPQYPPPLWMLL  
 15 YALELHCIIISLPPCLKISRWRQNHHLAFFFFRLHLQONCHYQTIPPHILLATGLIHPSVTWRLKS  
 GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKKRKVGTKVKDYIIGLAIGTSSVGWAVTD  
 EAYNVLFKNSKKMWGVRLFDDAKTAEERRGQRGARRRLDRKKERLSLLQDFFAEVAKVDPN  
 FFLRLDNSDLYMEDKDQKLKSKYTLFNDKDFKDKNFHKYPTIHLLMDLIEDDSKKDIRLV  
 YLACHYLLKNRGHFI FEGQKFDTKSSFENSLNELKVHLNDEYGLDLEFDNENLINILTDPKL  
 20 NKTAKKKELKSVIGDTKFLKAVSAIMIGSSQKLVLDLFENPEDFDDSAIKSVDFSTTSFDDKY  
 SDYELALGDKIALVNILKEIYDSSILENLLKEADKSKDGNKYISNAFVKKYNKHGQDLKEFK  
 RLVRQYHKSAYFDIFRSEKVNNDYVSYTKSSISNNKRVKANKFTDQEAIFYKFAKKHLETIKY  
 KINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNGVIPPYQLKLMELNKILENQSKEHFEFL  
 NVSDEYGSVCDKIASIMEFRIPIYVGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLDSREE  
 25 FIDSLIGRCTYLKDEKVLPKASLLYNEYMVLNELNKLNDLPI TEEMKKKIFDQLFKTRKK  
 VTLKAVANLLKKEFNINGEILLSGTDGDFKQGLNSYNDFKAIIVGDKVDSDDYRDKIEEIIKL  
 IVLYGDDKSYLQKKIKAGYGKYFTDSEIKKMAGLNYKDWGRLSKLLTGLEGANKITGERGS  
 I IHFMREYNLNLMELEMSASFTFTEEIQKLN PVDDRKLSYEMVDELYLSPSVKRMLWQSLRIV  
 DEIKNIMGTDSKKIFIE MARGKEEVKARKE SRKNQLLFYKDGKKA FISEIGEERYSYLLSE  
 30 IEGEEENKFRWDNLYLYTQLGRCMYSLEPIDISELSSKNIYDQDHIYPKSKIYDDSIENRV  
 LVKKDLNSKKGNSYPI PDEILNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDELVQFIS  
 RQIVETRQATKETANLLKTI CKNSEIVYSKAENASRFRQEFDIVKCRAVNDLHHMHDAYINI  
 IVGNVYNTKFTKDPMNFVKKQEKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRI

RKELEGTNYRFTRMNYIESGALFNATLQRKNKGSRPLKDKGPKSSIEKYGGYTNINKACFAV  
 LDIKSKNKIERKLMPPERIEIYAKQKNDKKLSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLK  
 IDGSYYFITGGSDKTLELRSAQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNT  
 FDIIVNKFKTSVFKKSFNLNFQDDKIENIDFKFKSMDFKKCKTLLMLVKAIRASGVRQDLK  
 5 SIDLKS DYGR LSSKTNNIGNYQEFKIINQSITGLFENEVDLLKLGKRPAATKKAGQAKKKK  
 SSGSGSGSGSTNLSDIIEKETGKQLVIQESILMLPEEVVEEVIGNKPESDILVHTAYDESTD  
 ENVMLLTS DAPEYKPWALVIQDSNGENKIKMLSGSGSGSGSTNLSDIIEKETGKQLVIQES  
 ILMPEEVVEEVIGNKPESDILVHTAYDESTDENVMMLTS DAPEYKPWALVIQDSNGENKIKM  
 LYPYDVPDYAYPYDVPDYAY (SEQ ID NO: 18);

10 (d)

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFDYDPRELKRETCLLYEIKWGMSRKIWRS  
 SGKNTTNHVEVNFIIKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIIY  
 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYPPPLWMLL  
 YALELHCIILSLPPCLKISRRWQNHLAFFRLHLQONCHYQTIPPHILLATGLIHPSVTWRLKS  
 15 GGSSGGSSGSETPGTSESATPESSGGSSGSPKKKRKVGKEYHIGLAIGTSSIGWAVTDSQF  
 KLMRIK GKTAIGVRLFEEGKTAERRTFRTRRRRLKRRKWRLHYLDEIFAPHLQEVDENFLR  
 RLKQSNIHPEDPAKNQAFIGKLLFPDLLKNERGYPTLIKMRDEL PVEQRAHYPVTNIYKLR  
 EAMINEDRQFDLREVYLAVHHIVKYRGHFLNNASVDKFKVGRIDFDKSFNVLNEAYEELQNG  
 EGSFTIEPSKVEKIGQLLLDTKMRKLDROKAVAKLLEVKVADKEETKRNKQIATAMSKLVLG  
 20 YKADFATVAMANGNEWKIDLSSETSEDEIEKFREELESDAQNDILTEITSLSFSQIMLNEIVPN  
 GMSISESMMDRYWOTHERQLAEVKEYLATQPASARKEFDQVYNKYIGQAPKEKGFLEKGLKK  
 ILSKKENWKEIDELLKAGDFLPKQRTSANGVI PHQMHQQELDRIIEKQAKYYPWLATENPAT  
 GERDRHQAKYELDQLVSFRIPYYVGPLVTPEVQKATSGAKFAWAKRKEDGEITPWNLWDKID  
 RAESAEAFIKRMTVKD TYLLNEDVLPANSLLYQKYNVLNELNNVRVNGRRLSVGIKQDIYTE  
 25 LFKKKKT VKAGDVASLVMAKTRGVNKPSVEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQ  
 MDLENIIEWRSVFEDEI FADKLTEVEWLTDEQRSALVKKRYKVGRLSKLLTGIVDENGQ  
 RIIDLWNTDQNFMQIVNQPVFKEQIDQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAI  
 WQVVRVVEDIVKAVGNAPKSI SIEFARNEGKGEITRSRRTQLQKLFEDQAHELVKDTSLTE  
 ELEKAPDLSDRYYFYFTQGGKDMYTGDPI NFEI STKYDIDHILPQS FVKDDSLDNRVLVSR  
 30 AENNKSDRVP AKLYAAKM KPYWNQLLKQLITQRKFENLTMDVDQTIKYRSLGFVQRQLVE  
 TRQVIKLTANILGSMYQEAGTDIIE TRAGLT KQLREEFDLPKVREVNDYHHA VDAYLTT FAG  
 QYLNRRYPKLR SFFVYGEYMKFKHGS DLKLRNFNFFHELMEGDKS QGKVVDQQT GELITTRD  
 EVADYFDWINLKVMLISNETYEETGKYFDASHSSSLYLKNQNKSKLVVPLKNKLQPEYY

GAYTGITQGYMVILKLLDKKGGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKS  
 FEVVVPRVLKGTFLVDGEEKFILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLNEMQTK  
 KLIEIYDNI LRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEKVI I KILRGLHFG  
 SDLQNLKEIGFGTTP LGQFQVSEAGIRLSNTAFI I FKSPTGLFNRKLYLKNLGRPAATKKA  
 5 GQAKKKKGSSGGSGGSGGSTNLSDIIEKETGKQLVIQESILMLPEEVVEEVIGNKPESDILVH  
 TAYDESTDENVMLLTS DAPEYKPWALVIQDSNGENKIKMLSGGSGGSGGSGGSTNLSDIIEKETG  
 KQLVIQESILMLPEEVVEEVIGNKPESDILVHTAYDESTDENVMLLTS DAPEYKPWALVIQDS  
 NGENKIKMLYPYDVPDYAYPYDVPDYAY (SEQ ID NO: 90);

(e)

10 MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFDPRELRKETCLLYEIKWGMSRKIWRS  
 SGKNTTNHVEVNFIIKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIIY  
 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEY YHCWRNFVNYPPGDEAHWPQYPPLWMLL  
 YALELHCI ILSLPPCLKISRWRQNH LAFFRLHLQONCHYQTI PPHILLATGLIHPSVTWRLKS  
 GGSSGGSSGSETPGTSESATPESGGSSGGSPKKKRKVGEKKTNYTIGLAIGTDSVGWAVVK  
 15 DDLELVKKRMKVLGNTETNYIKKNLWGSLLFESGQTAKDRRLKRVARRRYERRRNRLTELQK  
 IFAPAIDEVDENFFFRLNESFLVPEDKAFSKNPIFGTLGEDKTYKYTYPTIYHLRQHLADSE  
 EKADVRLIYLALAHMIKYRGHFLEIGKLDTEHIAINENLEQFFESYNALFSEPIELRKEEL  
 IAIENILREKNSRTVKEKRITSFLKDIGRANKQSPMAFITLIVGKKAKFKAAFNLEEEIISL  
 NLTDDSYDENLEILLNTIGSDFADLFDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYE  
 20 RHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKDLKGYAGYIDGNMSQDSFYKFVKDQLKE  
 VPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVHLAEMEA ILDRQENYYPWLKENREKIISLL  
 TFRIPYYVGPLADGQSEFAWLERKSDEKIKPWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEY  
 VLPKKS LIYQKYMVFNELTKIAYLDERQKRMNLS SVEKKEIFETLFKKRSKVTEKQLVKFFE  
 NYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMDDPDNEDLMEEIVKILT VFEDRKMR  
 25 RKQLEKYKERLSPEQIKELAKKHGTGWGRLSKLLV GIRDKETQKTILDYLVEDDNHSGGRQ  
 HLNRLMQLINDRRLSFKKTI AELQMIDPSADLYAQVQEIAGSPA I KKGILLGLKIVDEIIR  
 VMGEKPENIV IEMARENQTTARGKALSKRREAKIKEGLAALGSSLLKENLPGNADLSQRKIY  
 LYQTQNGKDIYLDEPLDFDRLSQYDEDHII PQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSL  
 EVVNRQLAYWRS LK DAGLMTQRKFDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKLL  
 30 DMRLNDKKDEAGNKIRETNIVLLKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAAIAINL  
 LALYPYMADDFVYGEFRYKKKPQAEKATYEKLRQWNLIKRFGEKQLFTP DHEDCWNKERDIK  
 TIKKVMGYRQVNVVKAERTGMLFKETINGKTNKGSRIPIKKDLDP SKYGGYIEEKMAYYA  
 VISYEDKKKKPGKTI VGISIMDKKEFEYDSISYLGKLGFSNPV VQIILKNYSLIAYPDGRRR

YITGATKTTKGKVELQKANQIAMEQDLVNFYIHLKNYDEISHPESYAFVQSHTDYFDRLFDS  
 IEHYTRRFLDAETNINRLRRIYEEEEKKKDPVDIEALVASFIELLKLTSAGAPADFI FMGEAI  
 SRRRYNSMTGLFDGQVIYQSLTGLYETRMRFEDGKRPAATKKAGQAKKKKSGSSGGSSGGSS  
 TNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTS DAP  
 5 EYKPWALVIQDSNGENKIKMLSGSSGGSSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEV  
 IGNKPESDILVHTAYDESTDENVMLLTS DAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAY  
 PYDVPDYAY (SEQ ID NO: 93);

(f)

MPAAKRVKLDTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDE  
 10 NVMLLTS DAPEYKPWALVIQDSNGENKIKMLSGSSGGSSGGSTNLSDIIEKETGKQLVIQESI  
 LMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTS DAPEYKPWALVIQDSNGENKIKML  
 SGGSSGGSSGSPKKKRKVEKKTNYTIGLAIGTDSVGVAVKDDLELVKKRMKVLGNTETNYIK  
 KNLWGSLLFESGQTAKDRRLKRVARRRYERRRNLTELQKIFAPAI DEVDENFFRNLNESFL  
 VPEDKAFSKNPIFGTLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHF  
 15 LIEGKLDTEHIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITS  
 FLKDIGRANKQSPMAFITLIVGKKAKFKAAFNLEEEISLNLTDSDYDENLEILLNTIGSDF  
 ADLFDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYD  
 MTFVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRS  
 FYNGVIPNQVHLAEMEAILDRQENYYPWLKENREKISLLTFRI PYYVGPLADGQSEFAWLE  
 20 RKSDEKIKPWNFSDVDLDRSAEKFIEQLIGRDTYLPDEYVLPKKS LIYQKYMVFNELTKIA  
 YLDERQKRMNLS SVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYS  
 TYVELAKVPGMKSMDDPDNEDLMEEIVKILTVFEDRKMRRKQLEKYKERLSPEQIKELAKK  
 HYTGWGRLSKLLVGIRDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDRRLSFKKTIA  
 ELQMI DPSADLYAQVQEIAGSPA I KKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTAR  
 25 GKALSKRREAKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLS  
 QYDEDHIIPQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEVVNRLAYWRS LKDAGLMTQR  
 KFDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKKDEAGNKIRETNIVL  
 LKSAMASEFRKMFRLYKVRELN DYHHAHDAYLNAAIAINLLALY P YMADDFVYGEFRYKKKP  
 QAEKATYEKLRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNVVKKAEERTG  
 30 MLFKETINGKTNKGSRIPIKKDLDP SKYGGYIEEKMAYYAVISYEDKKKKPGKTIVGISIMD  
 KKEFEYDSISYLGKLGFSNPVVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKANQIA  
 MEQDLVNFYIHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIY  
 EEEKKKDPVDIEALVASFIELLKLTSAGAPADFI FMGEAISRRRYNSMTGLFDGQVIYQSLT

GLYETRMRFEDKRPAATKKAGQAKKKKGSSGGSSGGSSGSETPGTSESATPESSGGSSGGST  
 SEKGPSTGDPTLRRRIESWEFDVFDYDPRELKRETCCLLYEIKWGMRSRKIWRSSGKNTTNHVEV  
 NFIKKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIYVARLFWHMDQR  
 NRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPPGDEAHWPQYPPLWMMMLYALELHCIIIS  
 5 LPPCLKISRWRQNHLLAFFRLHLQNCYQTI PPHILLATGLIHPSVTWRYPYDVPDYAYPYDV  
 PDYAYPYDVPDYA (SEQ ID NO: 94).

25. The Cas9 protein of claim 2, wherein the Cas9 protein recognizes a PAM sequence comprising 5' - NGG - 3'.
26. The Cas9 protein of claim 3, wherein the Cas9 protein recognizes a PAM sequence  
 10 comprising 5' - NAGHC - 3', wherein H is adenine, cytosine, or thymine.
27. The Cas9 protein of claim 4, wherein the Cas9 protein recognizes a PAM sequence comprising 5' - NRHRRH - 3', wherein H is adenine, cytosine or thymine, and R is adenine or guanine.
28. The Cas9 protein of claim 5 or claim 7, wherein the Cas9 protein recognizes a PAM  
 15 sequence comprising 5' - NGG - 3'.
29. The Cas9 protein of claim 6, wherein the Cas9 protein recognizes a PAM sequence comprising 5' - NNAAA - 3'.
- 29b. The Cas9 protein of claim 10b or 10c, wherein the Cas9 protein recognizes a PAM sequence comprising 5' - NGG - 3'.
- 20 30. A nucleic acid encoding the Cas9 protein of any one of the preceding claims.
31. The nucleic acid of claim 30, wherein the nucleic acid is codon-optimized for expression in mammalian cells.
32. The nucleic acid of claim 31, wherein the nucleic acid is codon-optimized for expression in human cells.
- 25 33. A eukaryotic cell comprising the Cas9 protein of any one of claims 29.
34. The eukaryotic cell of claim 33, wherein the cell is a human cell.
35. A method of cleaving a target nucleic acid in a eukaryotic cell comprising:

contacting the cell with a Cas9 of any one of claims 1-29, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and

wherein the Cas9 protein is capable of binding to the RNA guide and of causing a  
5 break in the target nucleic acid sequence complementary to the RNA guide.

36. A method of altering expression of a target nucleic acid in a eukaryotic cell comprising:

contacting the cell with a Cas9 of any one of claims 1-29, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat  
10 sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and

wherein the Cas9 protein is capable of binding to the RNA guide and of causing a break in the target nucleic acid sequence complementary to the RNA guide.

37. A method of altering expression of a target nucleic acid in a eukaryotic cell comprising:

15 contacting the cell with a Cas9 of any one of claims 1-29, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and

wherein the Cas9 protein is capable of binding to the RNA guide and editing the target nucleic acid sequence complementary to the RNA guide.

20 38. A method of modifying a target nucleic acid in a eukaryotic cell comprising:

contacting the cell with a Cas9 of any one of claims 1-29, and an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to the target nucleic acid, and

25 wherein the Cas9 protein is capable of binding to the RNA guide and editing the target nucleic acid sequence complementary to the RNA guide.

39. The method of claim 37 or 38, wherein the Cas9 protein is an inactive Cas9 (dCas9).

40. The method of claim 39, wherein the dCas9 is fused to a deaminase.

41. The method of any one of claims 35-40, wherein the RNA guide comprises a crRNA and a tracrRNA.

42. The method of any one of claims 35-39, wherein the RNA guide comprises a sgRNA.

43. The method of claim 42, wherein the sgRNA for use with *Streptococcus constellatus*  
5 Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

GUUUUAGAGCUGUGCUGUUUAAACAACACAGCAAGUUAAAUAAGGCUUUGU  
CCGUACUCAAGCUUGCAAAGCGUGCACCGAUUCGGUGCU-3' (SEQ ID NO: 3).

44. The method of claim 42, wherein the sgRNA for use with *Sharpea* Cas9 comprises a  
10 scaffold comprising a sequence having at least about 80% identity to

5'-

GUUUUAGAGUUGUGUUUUUGAAAAUAACACAACGAGUUAAAUAAGCUUA  
UGCUUAAAUGCCAGCUUUGCUGGUGUCAUUUAGAUGACUUUACUAAGGUUGC  
UUCGGCAACCUUUUU-3' (SEQ ID NO: 7).

15 45. The method of claim 42, wherein the sgRNA for use with *Veillonella parvula* Cas9  
comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

GUUUGAGAGUAGUGUGAAAACAUUACGAGUUCAAAUACAAUUAUUUACAA  
UGCCUUCGGGCUGCCCGACGUAGGGCACCUACUCUCAAUUCUUCGGAAUUGAG  
20 UU-3' (SEQ ID NO: 13).

46. The method of claim 42, wherein the sgRNA for use with *Ezakiella peruensis* Cas9  
comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

GUUUGAGAGUUAUGUAAUUGAAAAUUACAUGACGAGUUCAAAUAAAAUUU  
25 AUUCAACCGCCUAUUUAUAGGCCGCAGAUGUUCUGCAUUAUGCUUGCUAUU  
GCAAGCUU-3' (SEQ ID NO: 19).

47. The method of claim 42, wherein the sgRNA for use with *Lactobacillus fermentum* strain AF15-40LB Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

5 GUCUUGGAGUGUGUGAAAACACUCAUAGUCAAGAUCAAACGAGUGGUUUUC  
CACGAGUUAUUACUUUUGAGGUCUUAUAUGGCCCAUACAUAAAAGGAGUCG  
GAAUUCCGGCUCCUUUUCUU-3' (SEQ ID NO: 95)

48. The method of claim 42, wherein the sgRNA for use with *Peptoniphilus sp. Marseille-P3761* Cas9 comprises a scaffold comprising a sequence having at least about 80%  
10 identity to

5'-

GUUUUAGAGCCAUGUAGAAUACAUUGCAAGUUA AAAU AAGGCUUUGUCCGU  
AAUCAACUUGAAAAAGUGGCGCUGUUUCGGCGCUUU-3' (SEQ ID NO: 96)

49. The method of claim 41, wherein the crRNA comprises a guide sequence of between  
15 about 16 and 26 nucleotides long.

50. The method of claim 49, wherein the crRNA comprises a guide sequence between 18 and 24 nucleotides long.

51. The method of claim 35 or 36, wherein the break in the target nucleic acid is a single-stranded or double-stranded break.

20 52. The method of claim 51, wherein the break in the target nucleic acid is a single-stranded break.

53. The method of claim 34 or 35, wherein the Cas9 protein is a nuclease that cleaves both strands of the target nucleic acid sequence, or is a nickase that cleaves one strand of the target nucleic acid sequence.

25 54. The method of any one of claims 34-53, wherein the target nucleic acid is 5' to a protospacer adjacent motif (PAM) sequence.



55. The method of any one of claims 34-54, wherein the Cas9 is operably linked to a promoter sequence for expression in a eukaryotic cell, and wherein the guide RNA is operably linked to a promoter sequence for expression in a eukaryotic cell.

56. The method of claim 55, wherein the eukaryotic cell is a human cell.

5 57. The method of claim 55, wherein the promoter sequence is a eukaryotic or viral promoter.

58. An engineered, non-naturally occurring CRISPR-Cas system comprising:

an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide comprises a direct repeat sequence and a spacer sequence capable of hybridizing to a target  
10 nucleic acid; and

a codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to SEQ ID NOs: 1, 4, 8, 14, 84 or 86, and wherein the Cas protein is capable of binding to the RNA guide and of causing a break in the target nucleic acid sequence complementary to the RNA guide.

15 58b. The engineered, non-naturally occurring CRISPR-Cas system of claim 58 where the codon-optimized CRISPR-associated (Cas) protein further comprising a nuclear localization sequence (NLS) and/or a FLAG, HIS or HA tag.

58c. The engineered, non-naturally occurring CRISPR-Cas system of claim 59 where the codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to  
20 SEQ ID NOs: 2, 5, 9, 15, 85, 87, 95 or 96, and wherein the Cas protein is capable of binding to the RNA guide and of causing a break in the target nucleic acid sequence complementary to the RNA guide.

59. An engineered, non-naturally occurring CRISPR-Cas system comprising:

an RNA guide or a nucleic acid encoding the RNA guide, wherein the RNA guide  
25 comprises a direct repeat sequence and a spacer sequence capable of hybridizing to a target nucleic acid; and

a codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to SEQ ID NOs: 1, 4, 8, 14, 84 or 86;

wherein the Cas protein is fused to a deaminase, and wherein the Cas protein fusion is capable of binding to the RNA guide and of editing the target nucleic acid sequence complementary to the RNA guide.

5 59b. The engineered, non-naturally occurring CRISPR-Cas system of claim 59 where the codon-optimized CRISPR-associated (Cas) protein further comprising a nuclear localization sequence (NLS) and/or a FLAG, HIS or HA tag.

10 59c. The engineered, non-naturally occurring CRISPR-Cas system of claim 59b where the codon-optimized CRISPR-associated (Cas) protein having at least 80% sequence identity to SEQ ID NOs: 2, 5, 9, 15, 85, 87, 95 or 96, wherein the Cas protein is fused to a deaminase, and wherein the Cas protein fusion is capable of binding to the RNA guide and of editing the target nucleic acid sequence complementary to the RNA guide.

60. The system of claim 59, wherein the Cas9 protein is an inactive Cas9 (dCas9).

61. The system of claim of any one of claims 58-60, wherein the RNA guide comprises a crRNA and a tracrRNA.

15 62. The system of any one of claims 58-60, wherein the RNA guide comprises an sgRNA.

63. The system of claim 62, wherein the sgRNA for use with *Streptococcus constellatus* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

20 GUUUUAGAGCUGUGCUGUUUAAACAACACAGCAAGUUAAAUAAGGCUUUGU  
CCGUACUCAAGCUUGCAAAGCGUGCACCGAUUCGGUGCU-3' (SEQ ID NO: 3).

64. The system of claim 62, the sgRNA for use with *Sharpea* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

25 GUUUUAGAGUUGUGUUUUGAAAAUAACACAACGAGUUAAAUAAGCUUA  
UGC UAAAUGCCAGCUUUGCUGGUGUCAUUUAGAUGACUUUACUAAGGUUGC  
UUCGGCAACCUUUUUU-3' (SEQ ID NO: 7).

65. The system of claim 62, wherein the sgRNA for use with *Veillonella parvula* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

GUUUGAGAGUAGUGUGAAAACAUUACGAGUUCAAAUACAAUAAUUUACAA  
 UGCCUUCGGGUGCCCGACGUAGGGCACCUACUCUCAAUUCUUCGGAAUUGAG  
 UU-3' (SEQ ID NO: 13).

- 5 66. The system of claim 62, wherein the sgRNA for use with *Ezakiella peruensis* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

GUUUGAGAGUUAUGUAAUUGAAAAAUACAUGACGAGUUCAAAUAAAAUUU  
 AUUCAACCGCCUAUUUAUAGGCCGCAGAUGUUCUGCAUUAUGCUUGCUAUU  
 10 GCAAGCUU-3' (SEQ ID NO: 19).

67. The system of claim 62, wherein the sgRNA for use with *Lactobacillus fermentum* strain AF15-40LB Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

15 GUCUUGGAGAGUGUGAAAACACUCAUAGUCAAGAUCAAACGAGUGGUUUUC  
 CACGAGUUAUUACUUUUGAGGUCUUAUAUGGCCCAUACAUAAAAGGAGUCG  
 GAAUUCCGGCUCCUUUUCUU-3' (SEQ ID NO: 95).

68. The system of claim 62, wherein the sgRNA for use with *Peptoniphilus sp. Marseille-P3761* Cas9 comprises a scaffold comprising a sequence having at least about 80% identity to

20 5'-

GUUUUAGAGCCAUGUAGAAAUACAUUGCAAGUUA AAAUAAGGCUUUGUCCGU  
 AAUCAACUUGAAAAGUGGCGCUGUUUCGGCGCUUU-3' (SEQ ID NO: 96).

69. The system of any one of claims 58-68, wherein the Cas protein is operably linked to a promoter sequence for expression in a eukaryotic cell, and wherein the guide RNA is  
 25 operably linked to a promoter sequence for expression in a eukaryotic cell.

70. The system of claim 69, wherein the eukaryotic cell is a human cell.

71. The system of claim 70, wherein the promoter sequence is a eukaryotic promoter sequence.

72. A nucleic acid encoding the system of any one of claims 58-71.
73. A vector comprising the system of any one of claims 58-72.
74. The vector of claim 73, wherein the vector is a plasmid vector or a viral vector.
75. The vector of claim 74, wherein the viral vector is an adeno associated virus (AAV)  
5 vector or a lentiviral vector.
76. The vector of claim 75, wherein the viral vector is an AAV vector.
77. The vector of claim 76, wherein more than one AAV vector is used for packaging the system of claims 59-71.
78. A method of treating a disorder or a disease in a subject in need thereof, the method  
10 comprising administering to the subject a system of any one of claims 58-71,  
wherein the guide RNA is complementary to at least 10 nucleotides of a target nucleic acid associated with the condition or disease;  
wherein the Cas protein associates with the guide RNA;  
wherein the guide RNA binds to the target nucleic acid;  
15 wherein the Cas protein causes a break in the target nucleic acid, optionally wherein the Cas9 is an inactive Cas9 (dCas9) fused to a deaminase and results in one or more base edits in the target nucleic acid, thereby treating the disorder or disease.
79. The method of claim 78, wherein the guide RNA is complementary to about 18-24 nucleotides.
- 20 80. The method of claim 79, wherein the guide RNA is complementary to 20 nucleotides.
81. A base editor comprising the fusion protein of any one of claims 16-19.
82. The base editor of claim 81 comprising an adenosine deaminase domain or a cytidine deaminase domain.
- 82b. The base editor of claim 81 comprising an adenosine deaminase domain and a  
25 cytidine deaminase domain.
83. A method of editing a nucleobase of a polynucleotide, the method comprising contacting the polynucleotide with the base editor of claim 81 in complex with one or more

guide RNAs, wherein the base editor comprises an adenosine deaminase domain and wherein the one or more guide RNAs target the base editor to effect an A•T to G•C alteration in the polynucleotide.

84. A method of editing a nucleobase of a polynucleotide, the method comprising  
5 contacting the polynucleotide with the base editor of claim 81 in complex with one or more guide RNAs, wherein the base editor comprises a cytidine deaminase domain, and wherein the one or more guide RNAs target the base editor to effect an C•G to T•A alteration in the polynucleotide.

85. The method of claim 83 or 84, wherein the editing results in less than 50% indel  
10 formation in the target polynucleotide sequence.

86. The method of any one of claims 83-85, wherein the editing generates a point mutation.

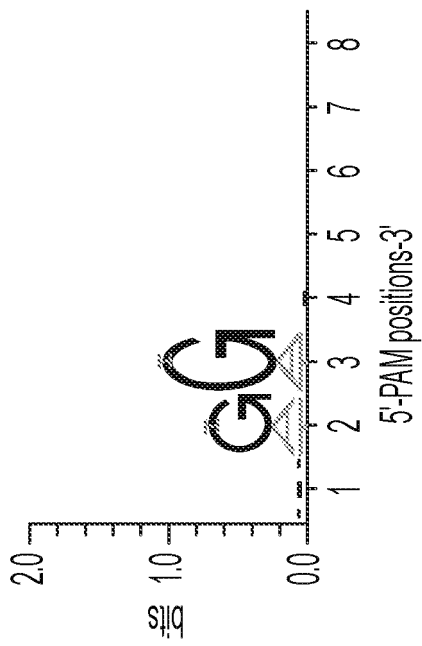


FIG. 1A

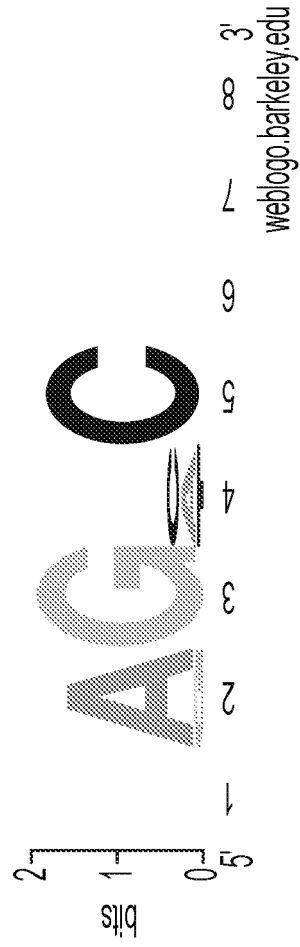


FIG. 1B

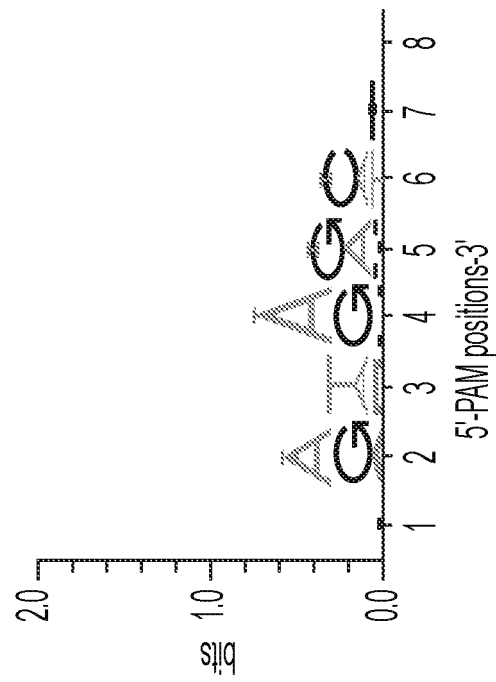


FIG. 1C

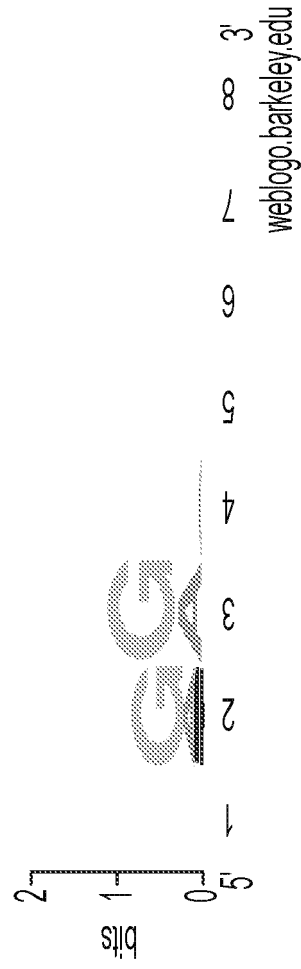


FIG. 1D

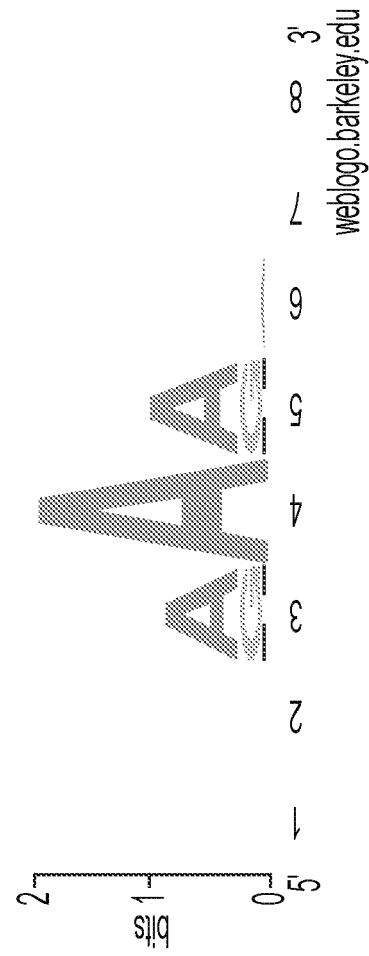


FIG. 1E

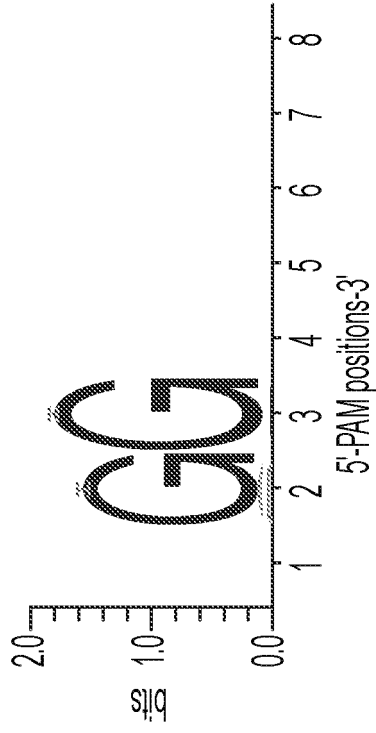


FIG. 1F

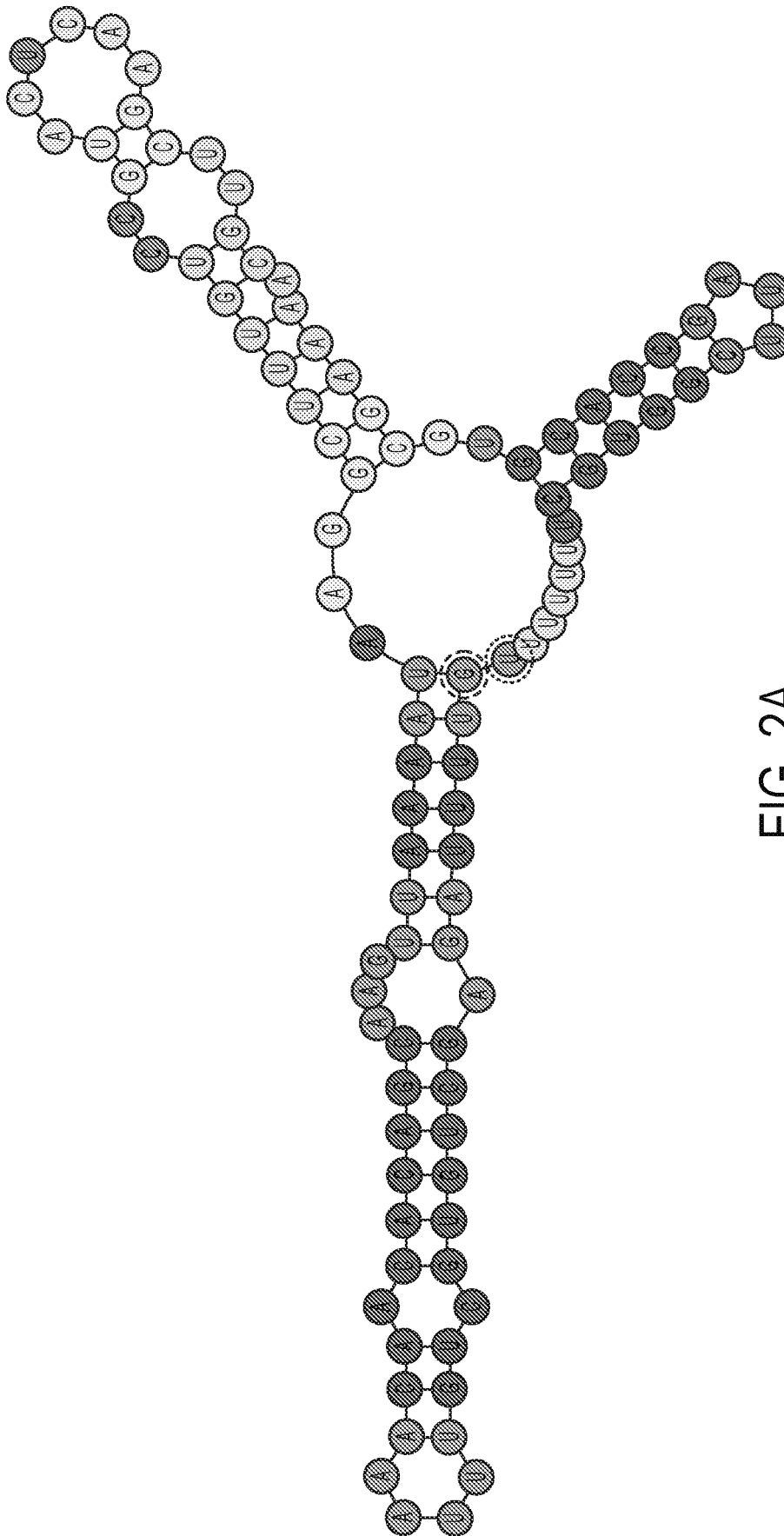


FIG. 2A



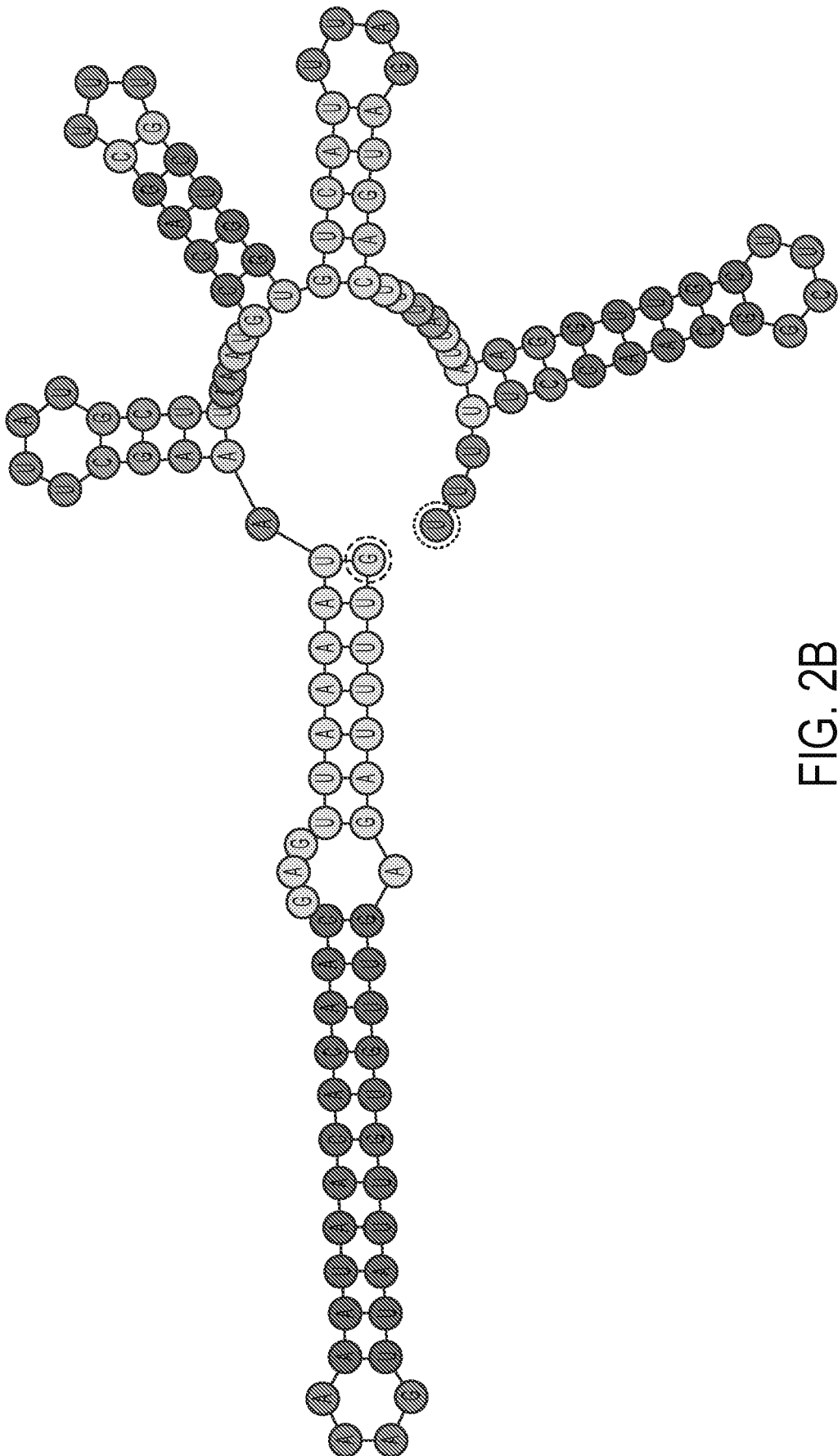


FIG. 2B

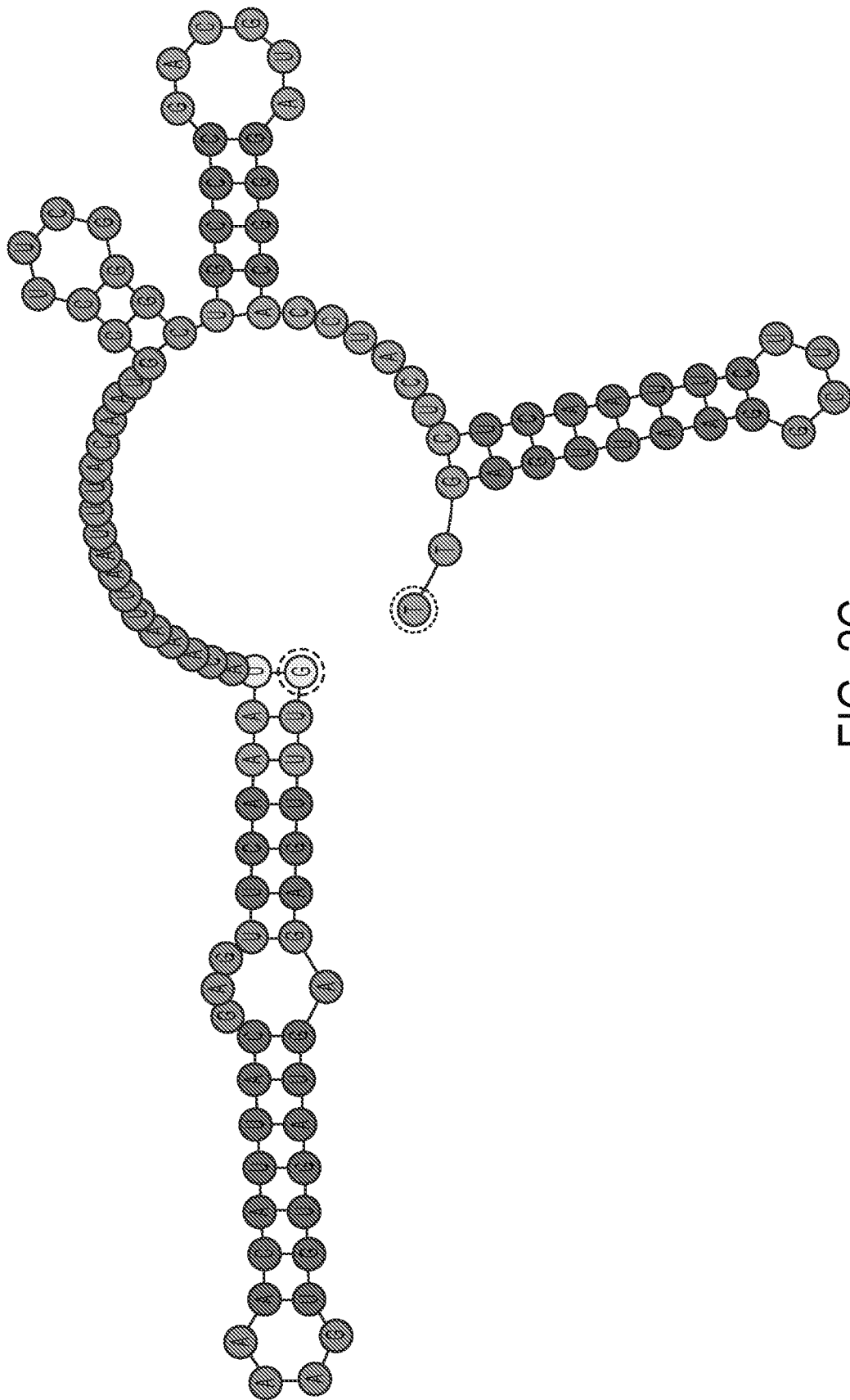


FIG. 2C

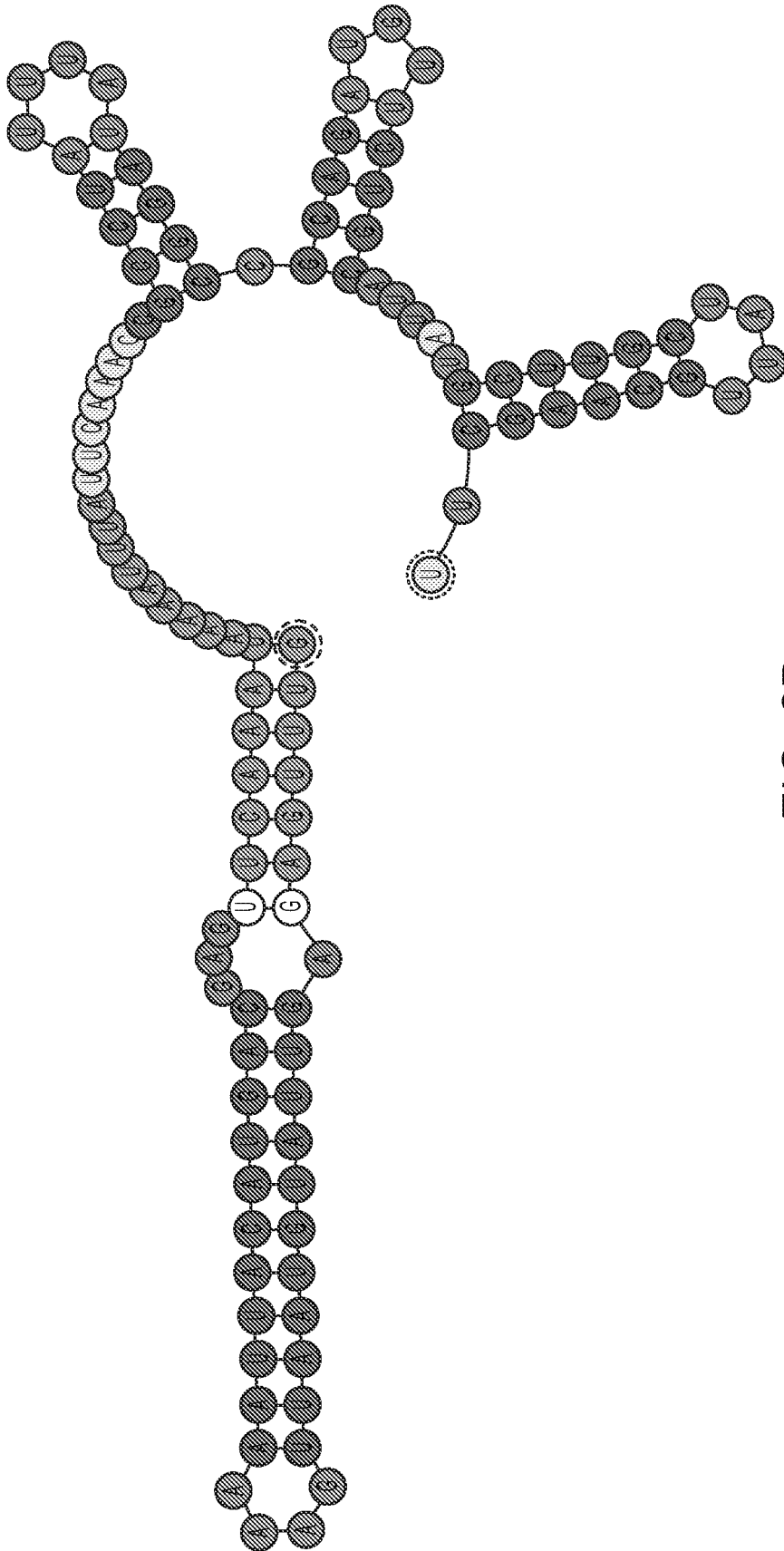


FIG. 2D

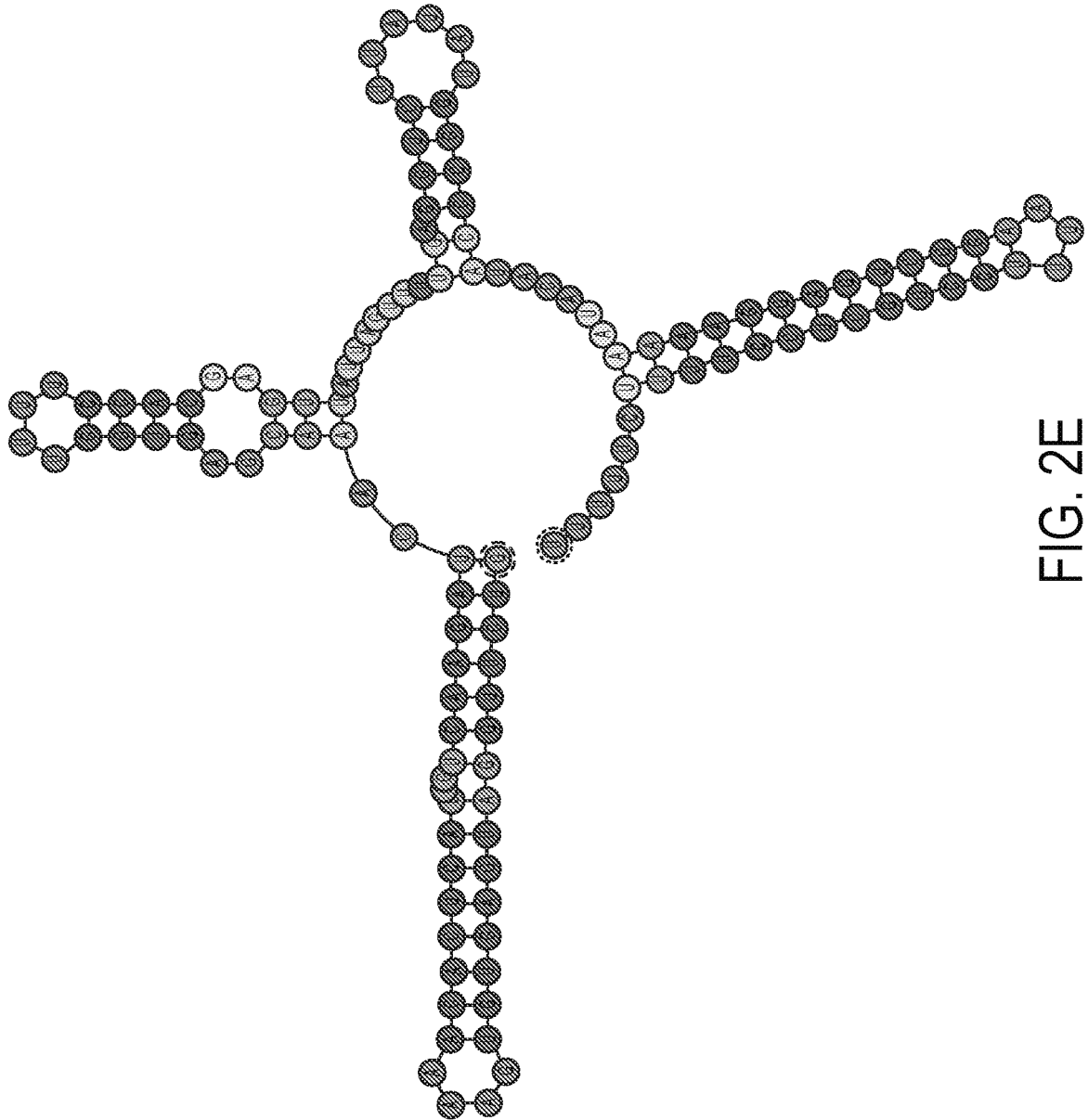


FIG. 2E

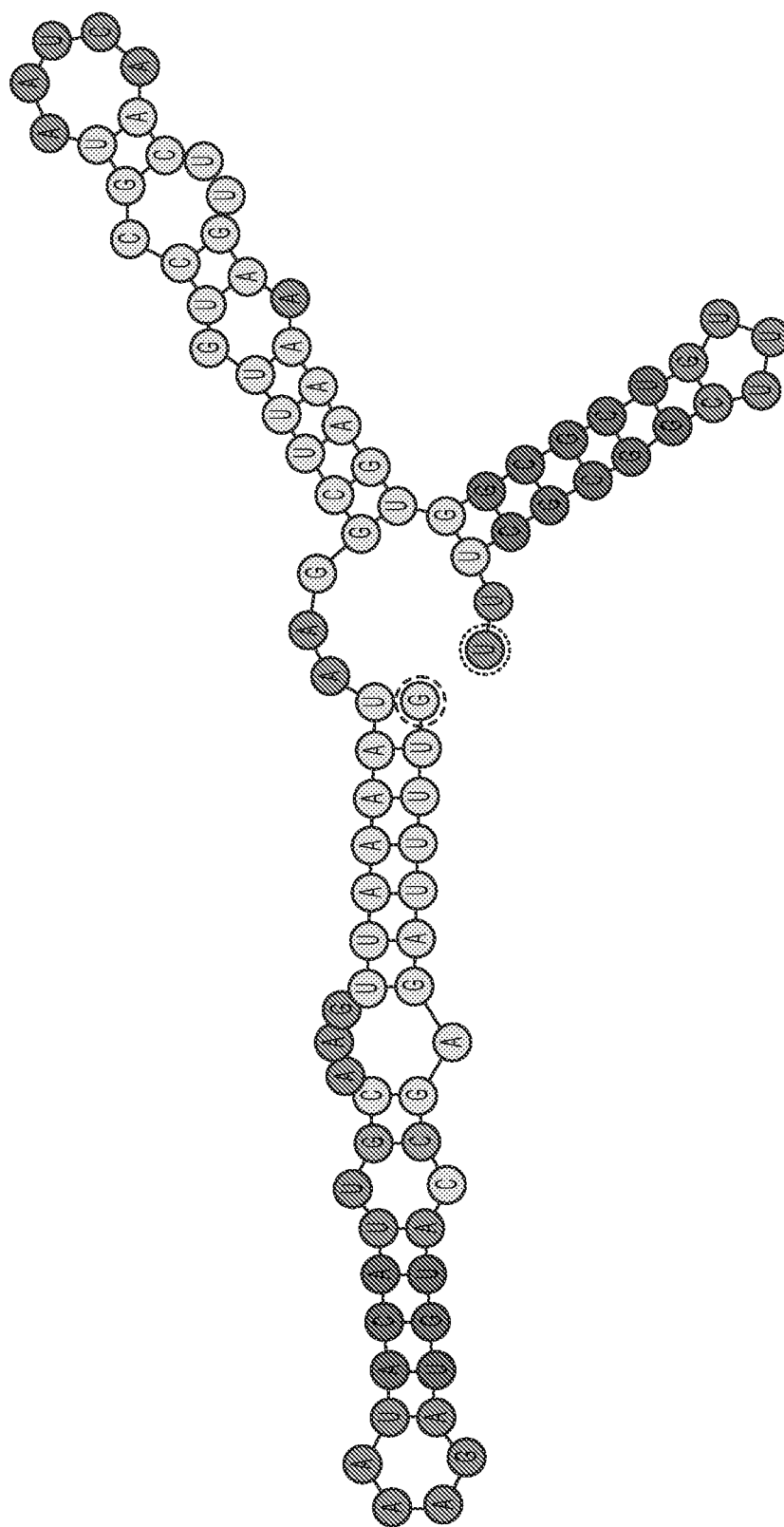


FIG. 2F

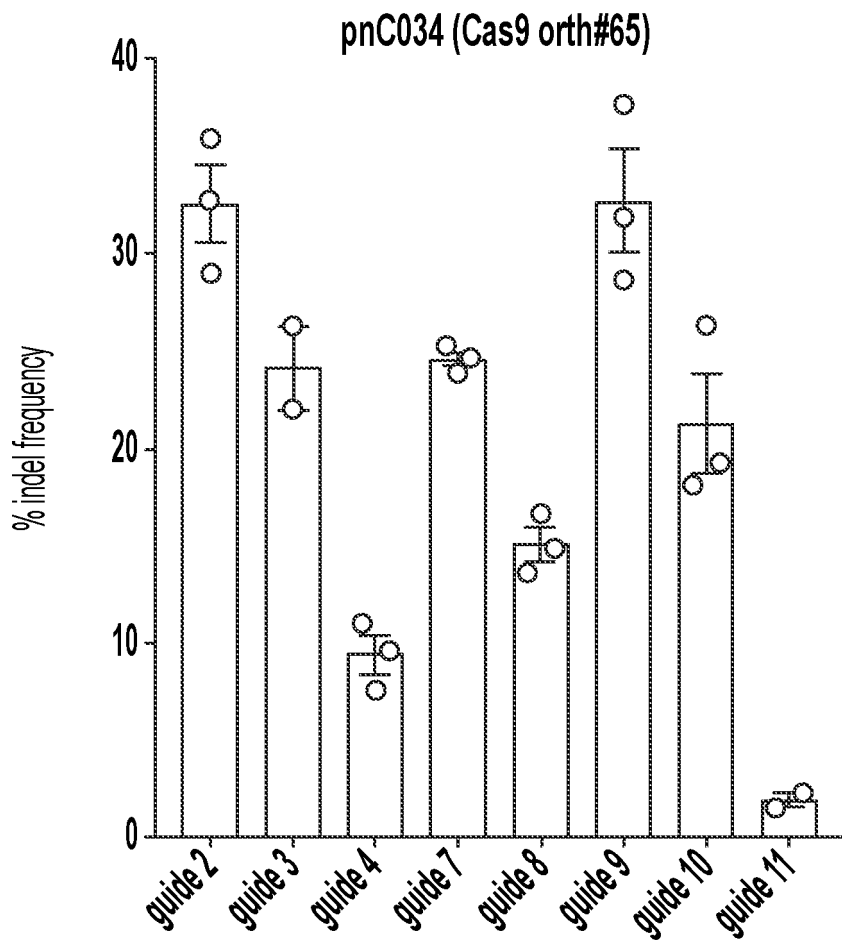


FIG. 3

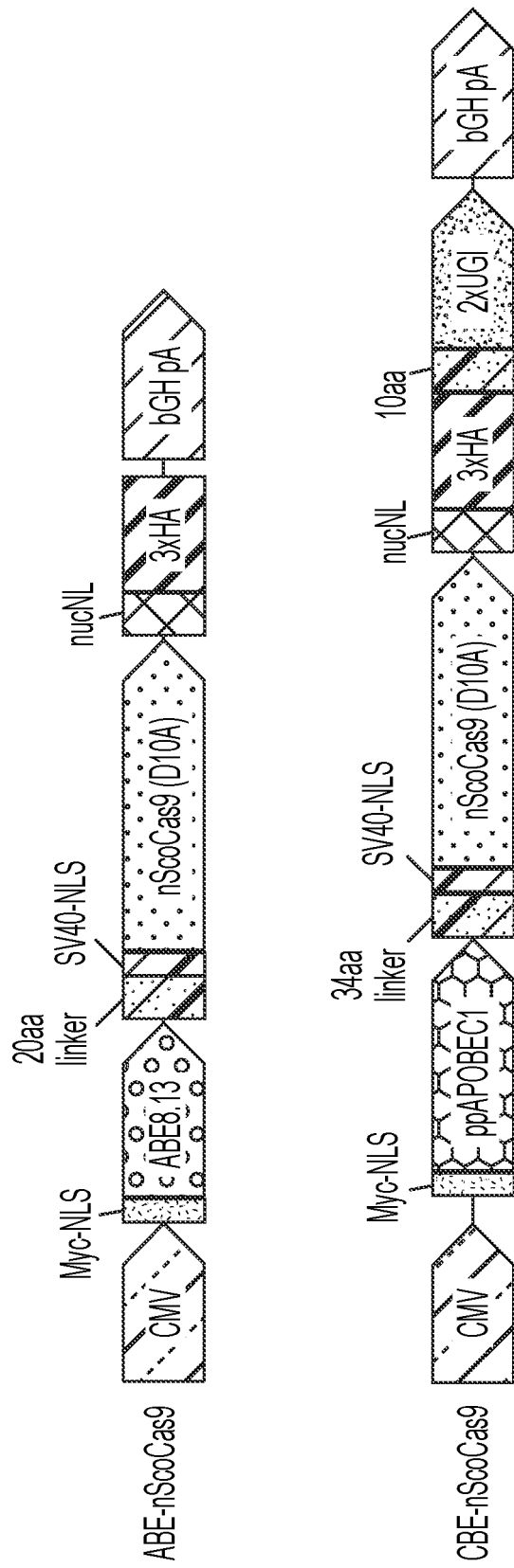


FIG. 4A

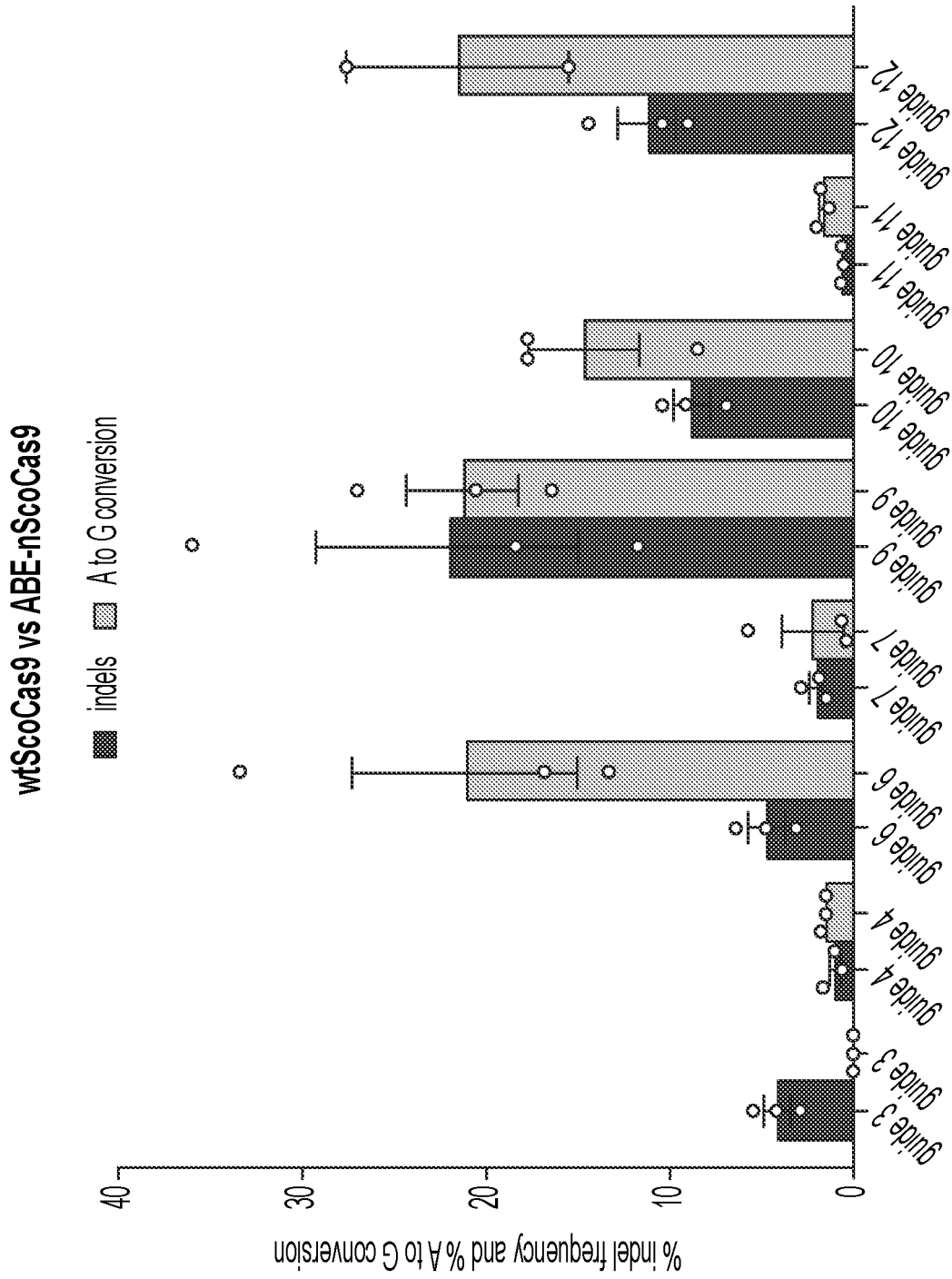


FIG. 4B



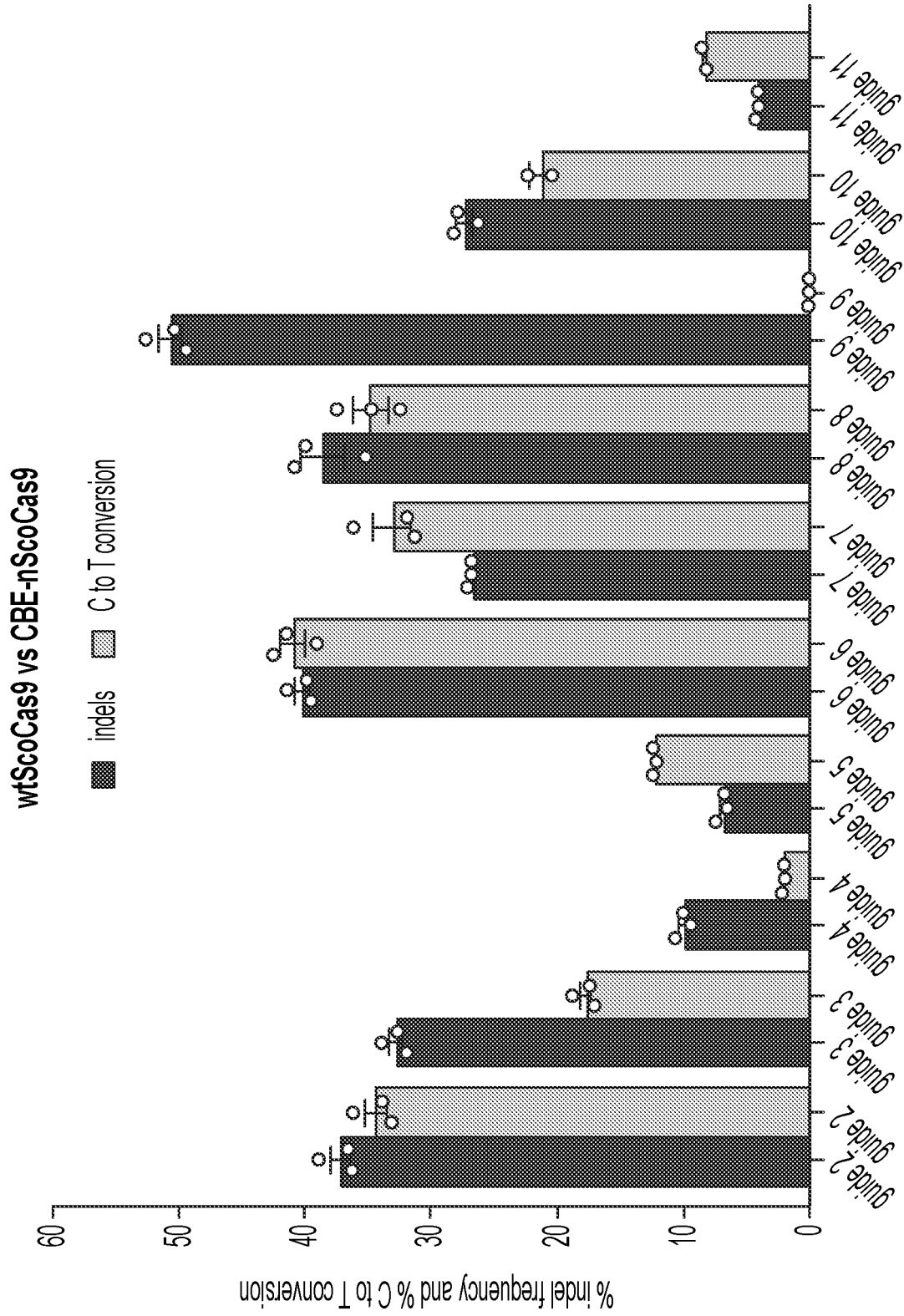


FIG. 4C

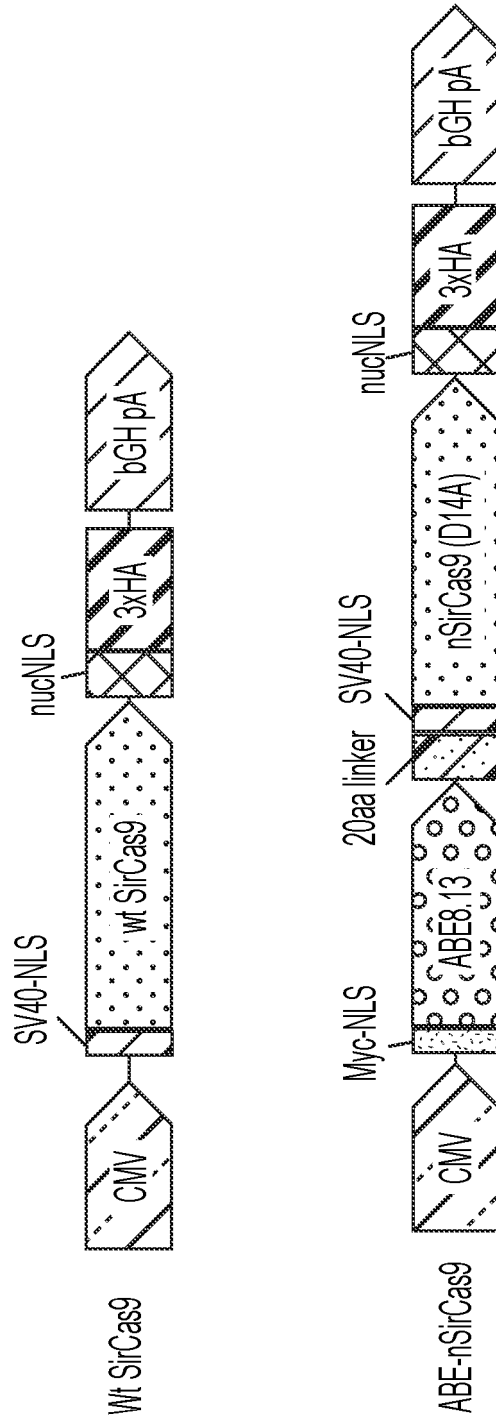


FIG. 5A

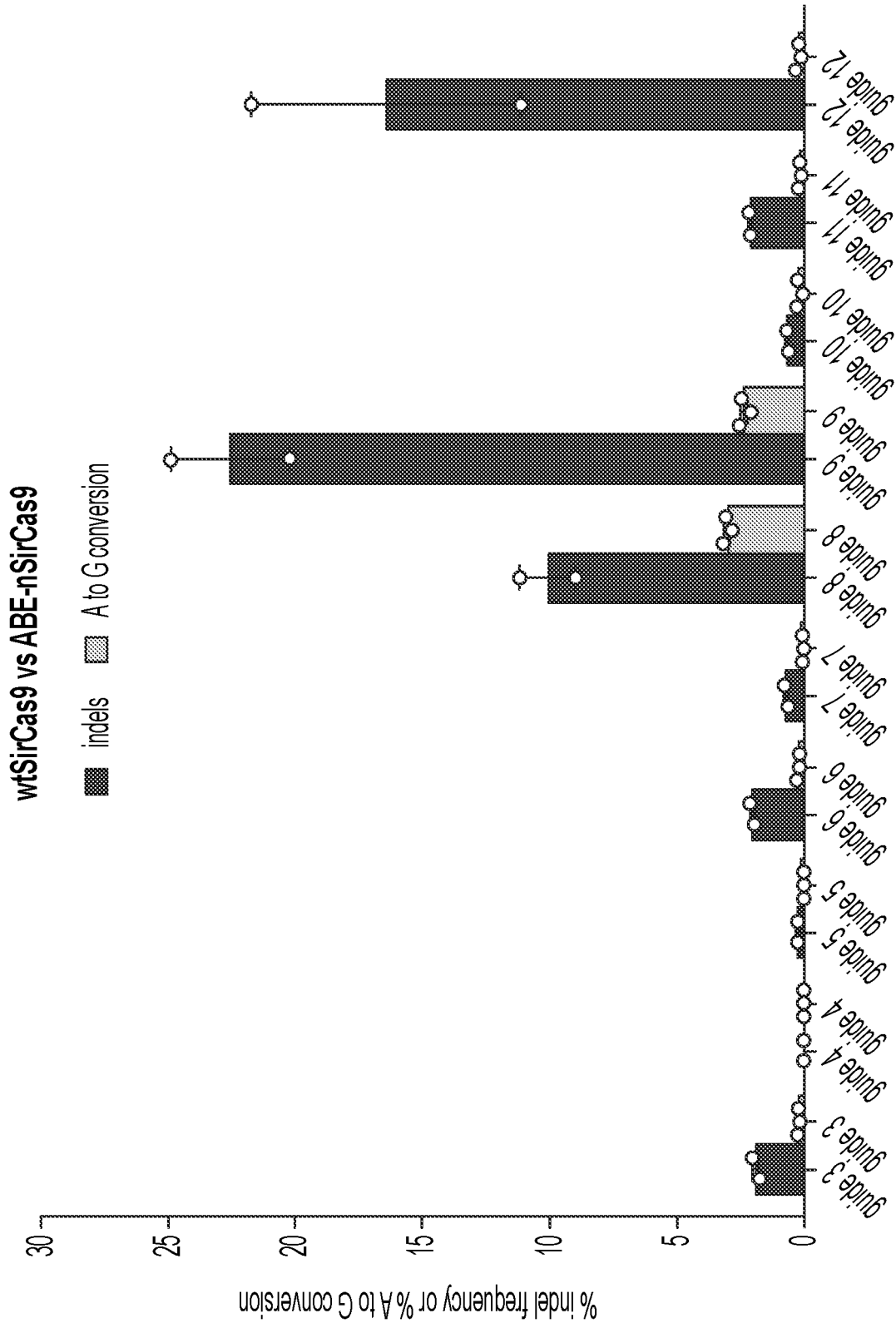


FIG. 5B

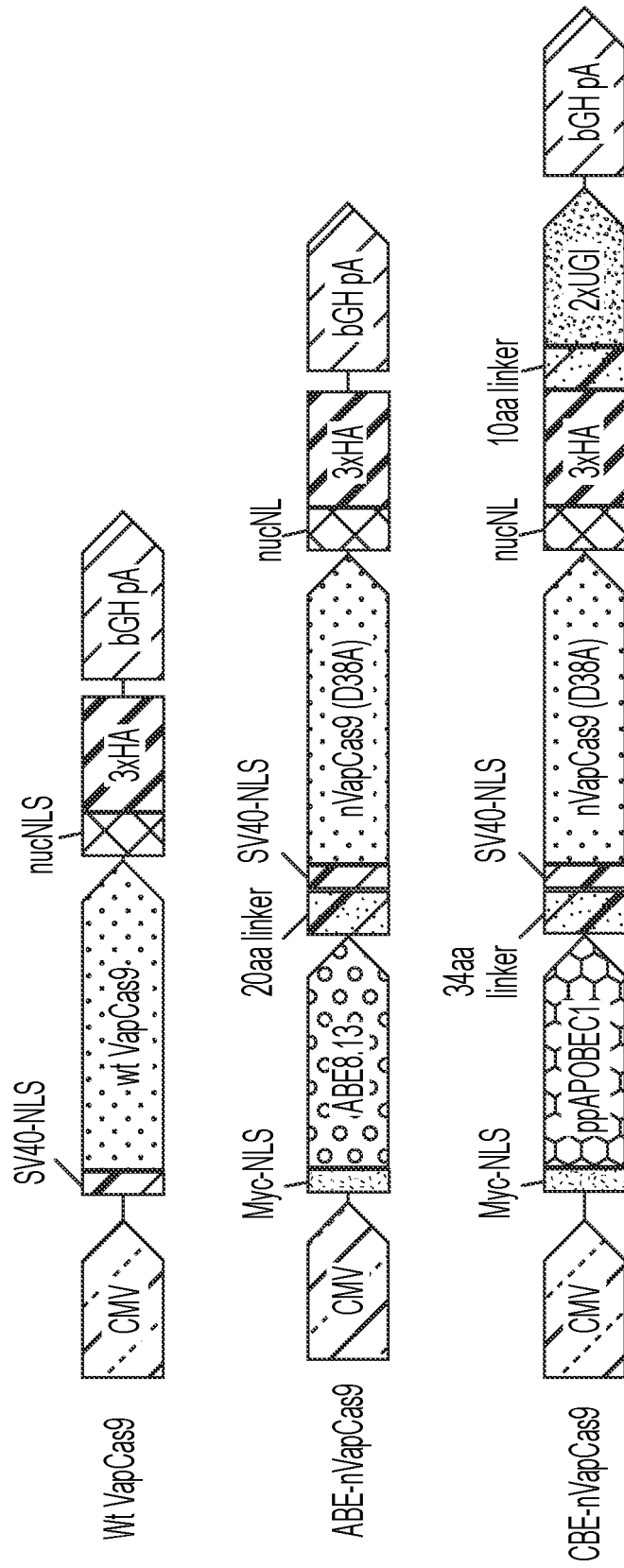


FIG. 6A

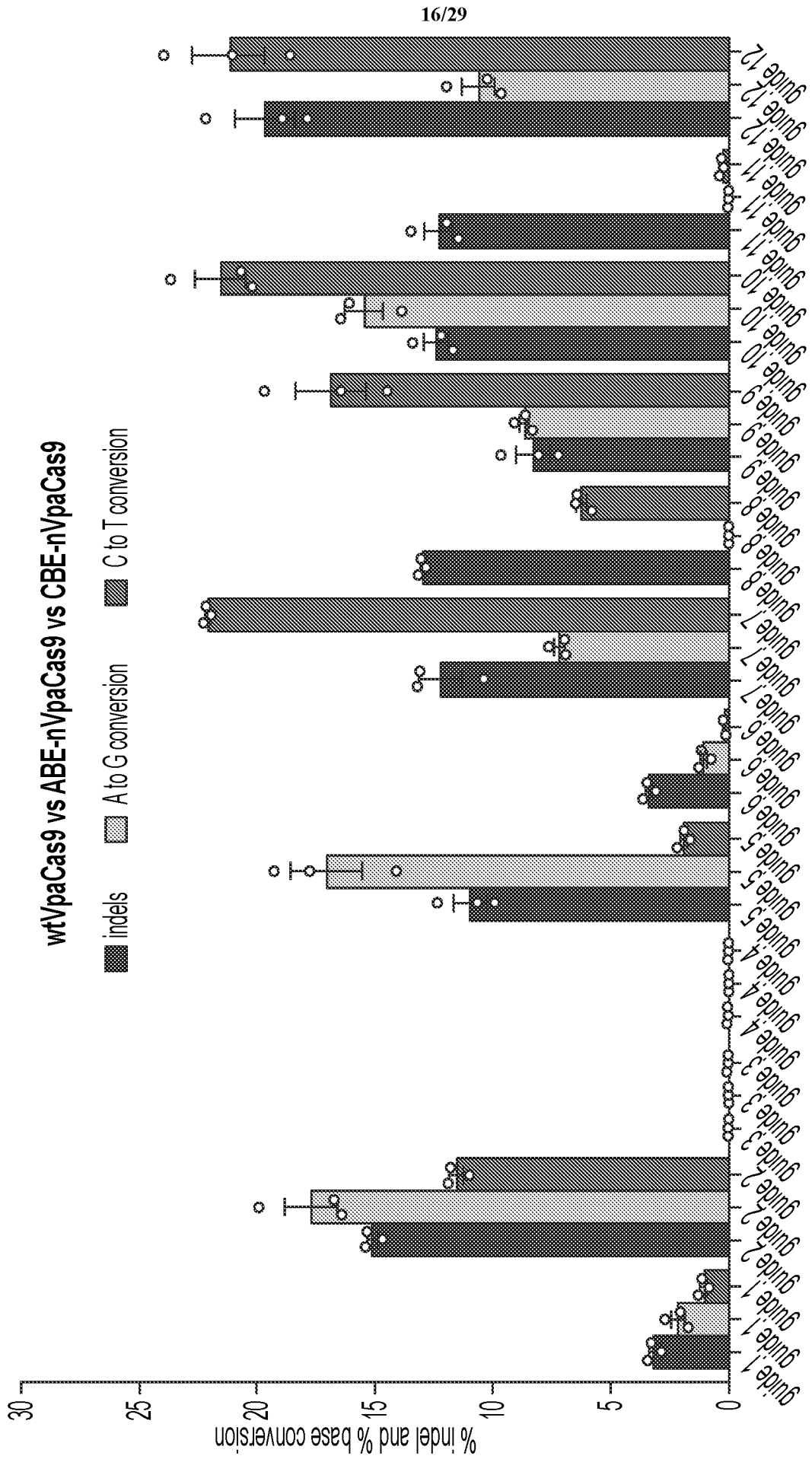


FIG. 6B

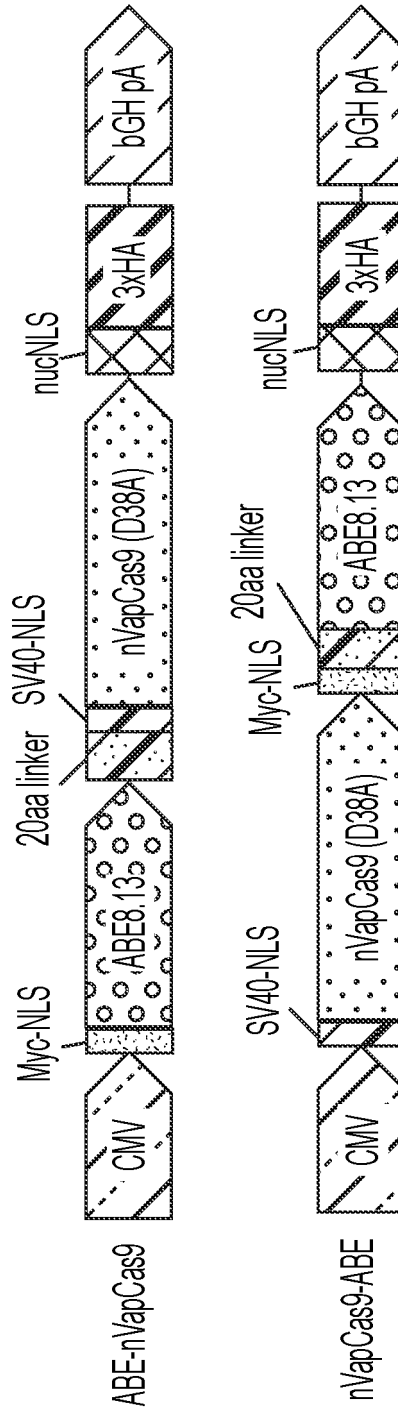


FIG. 7A

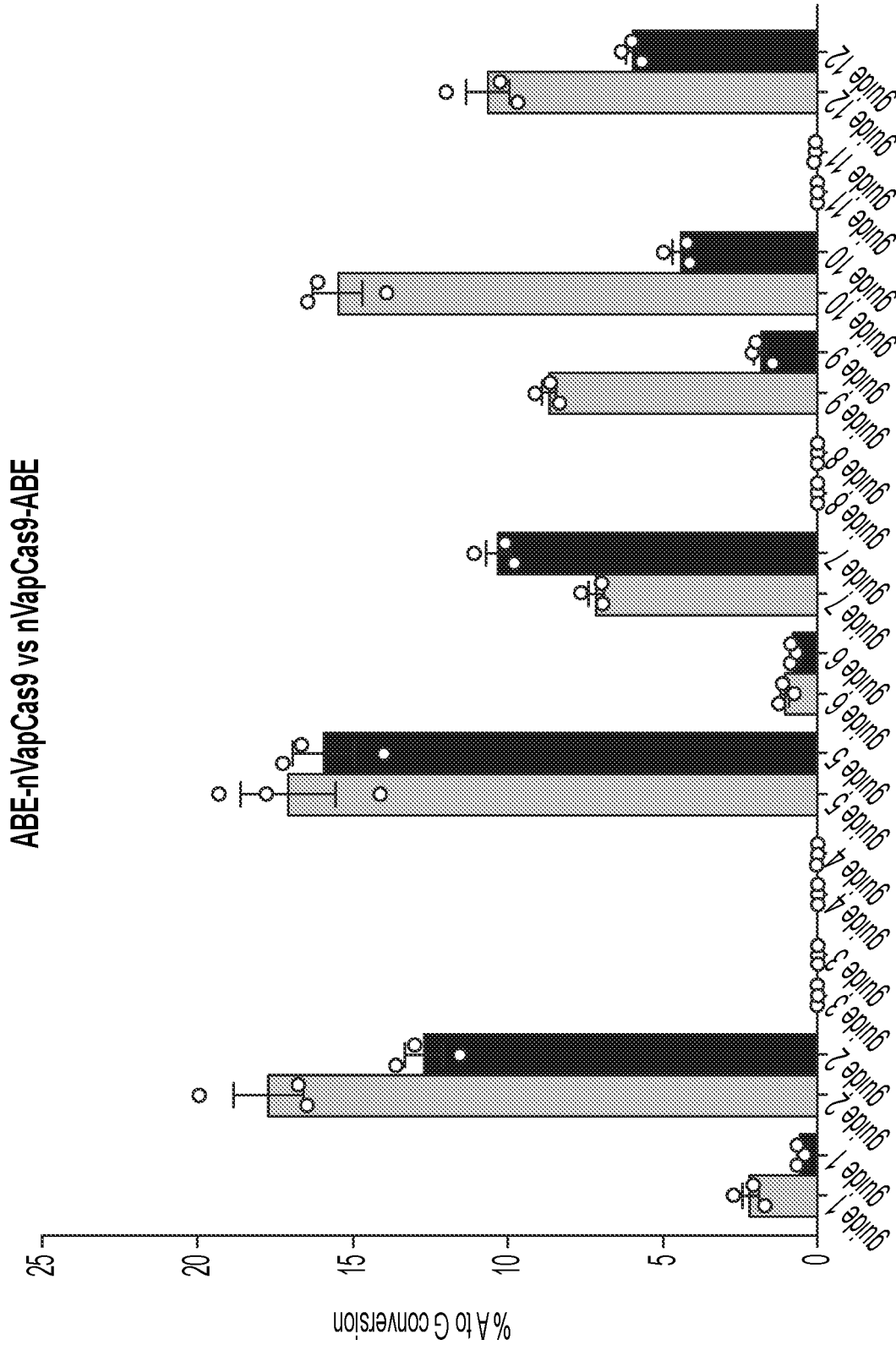


FIG. 7B

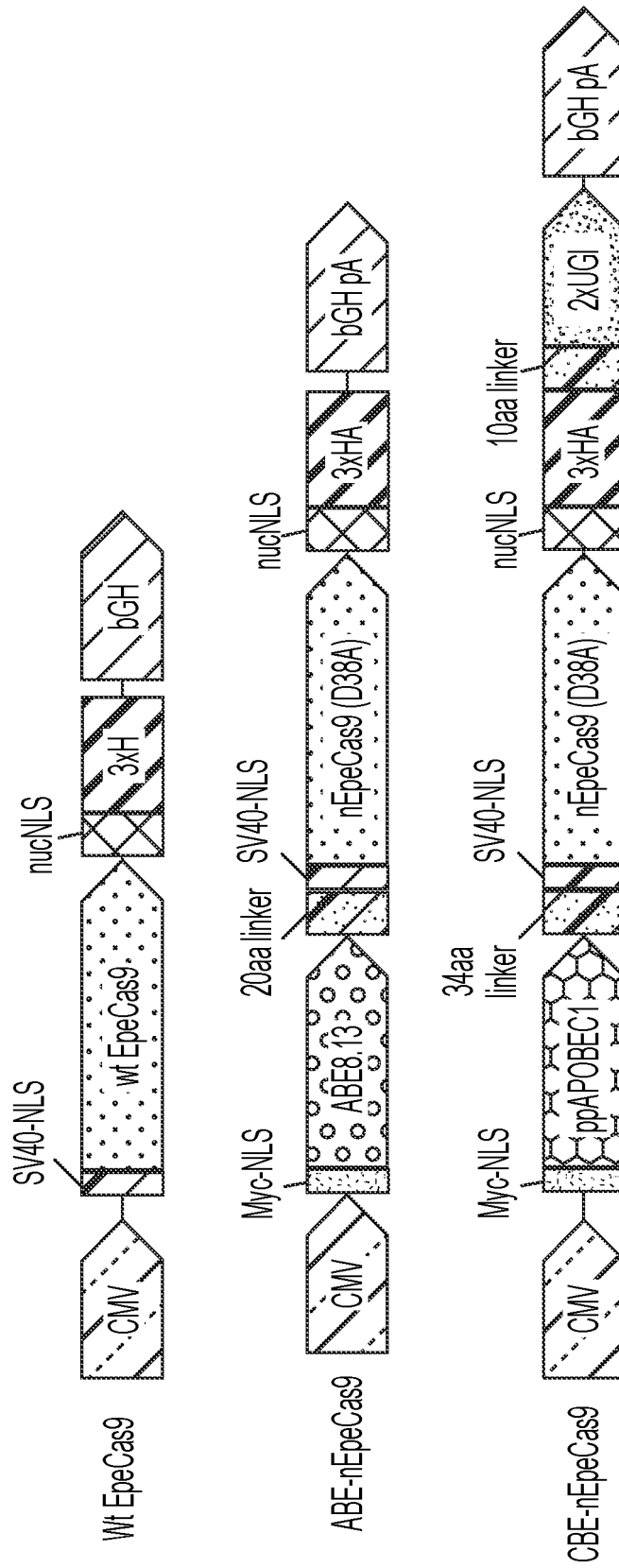


FIG. 8A



wtEpeCas9 ABE-nEpeCas9 CBE-nEpeCas9

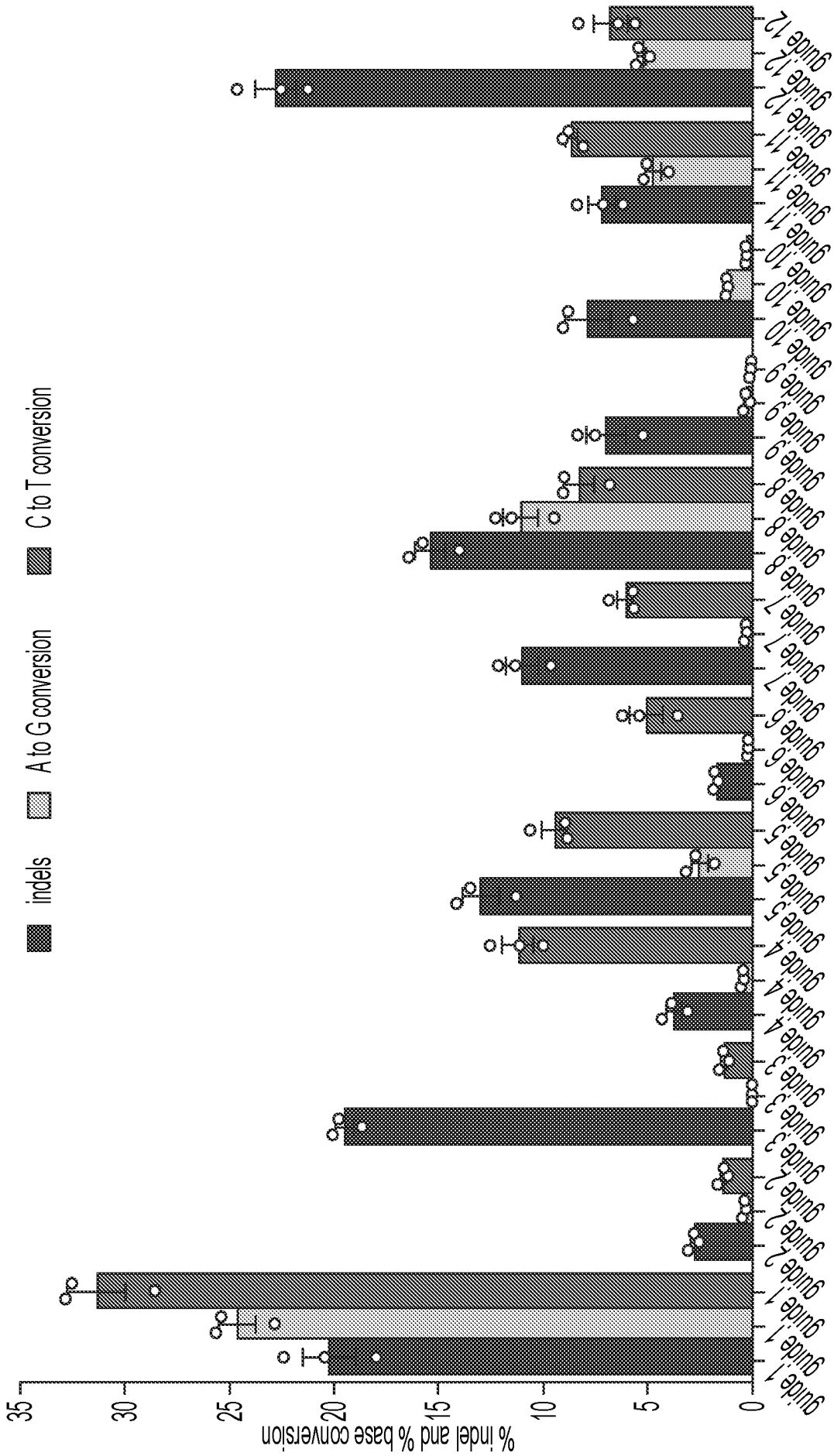


FIG. 8B

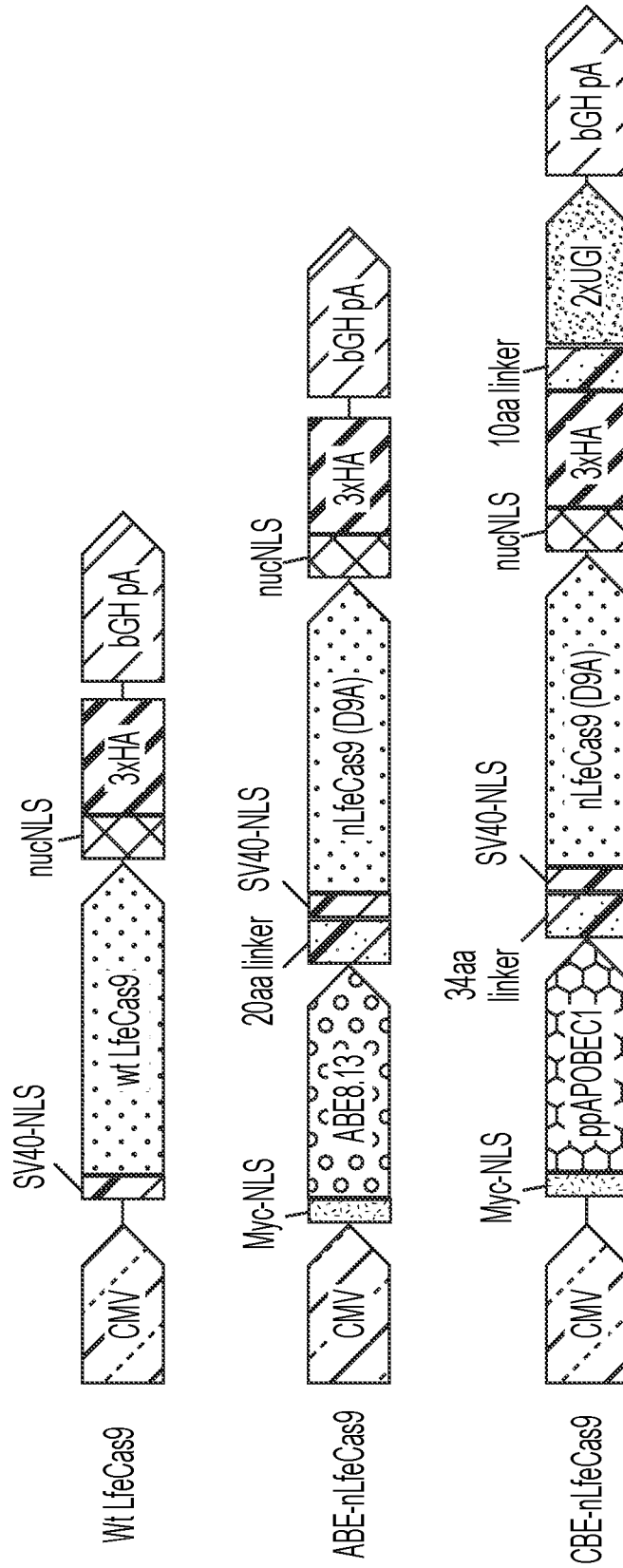


FIG. 9A

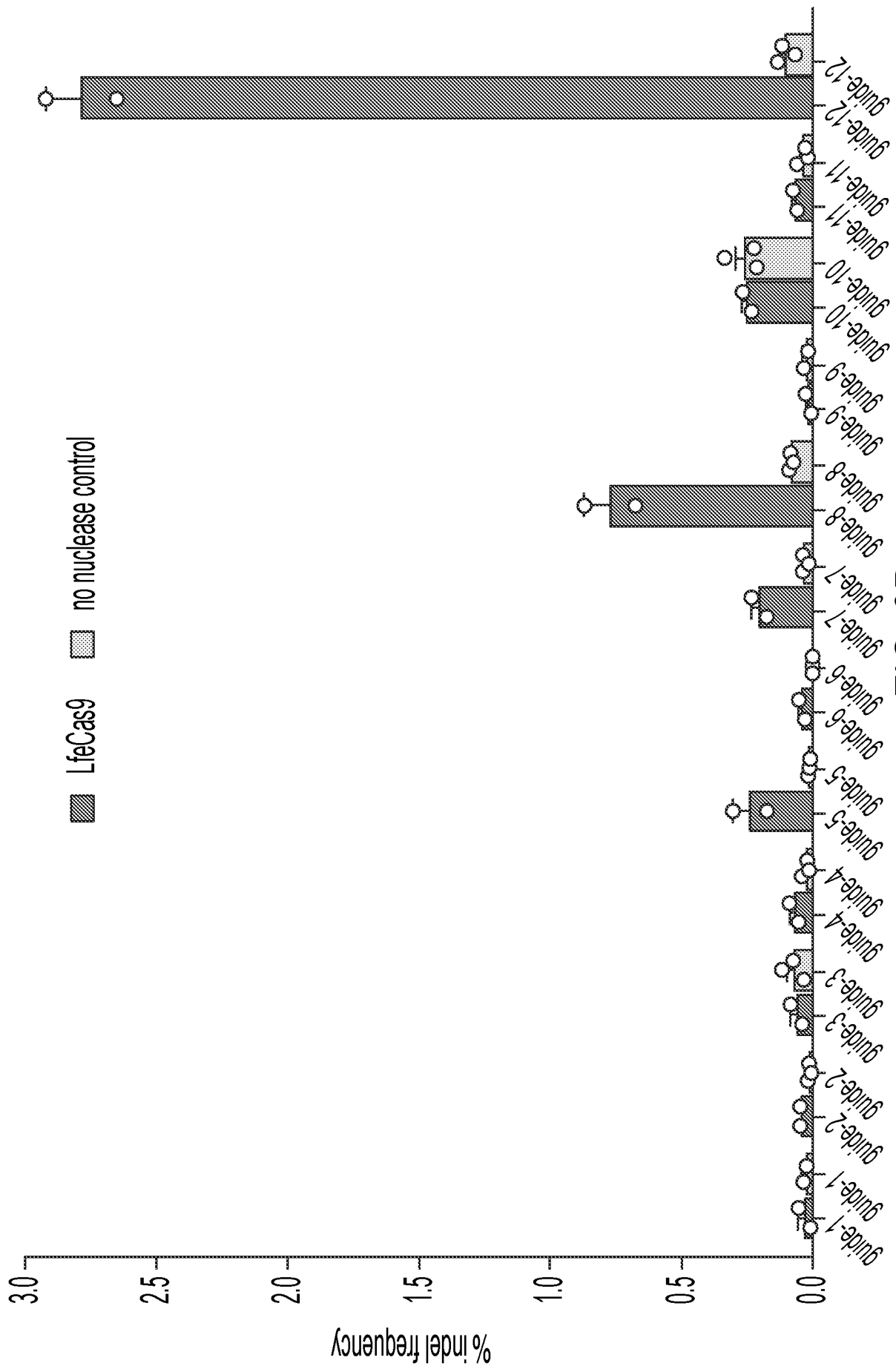


FIG. 9B

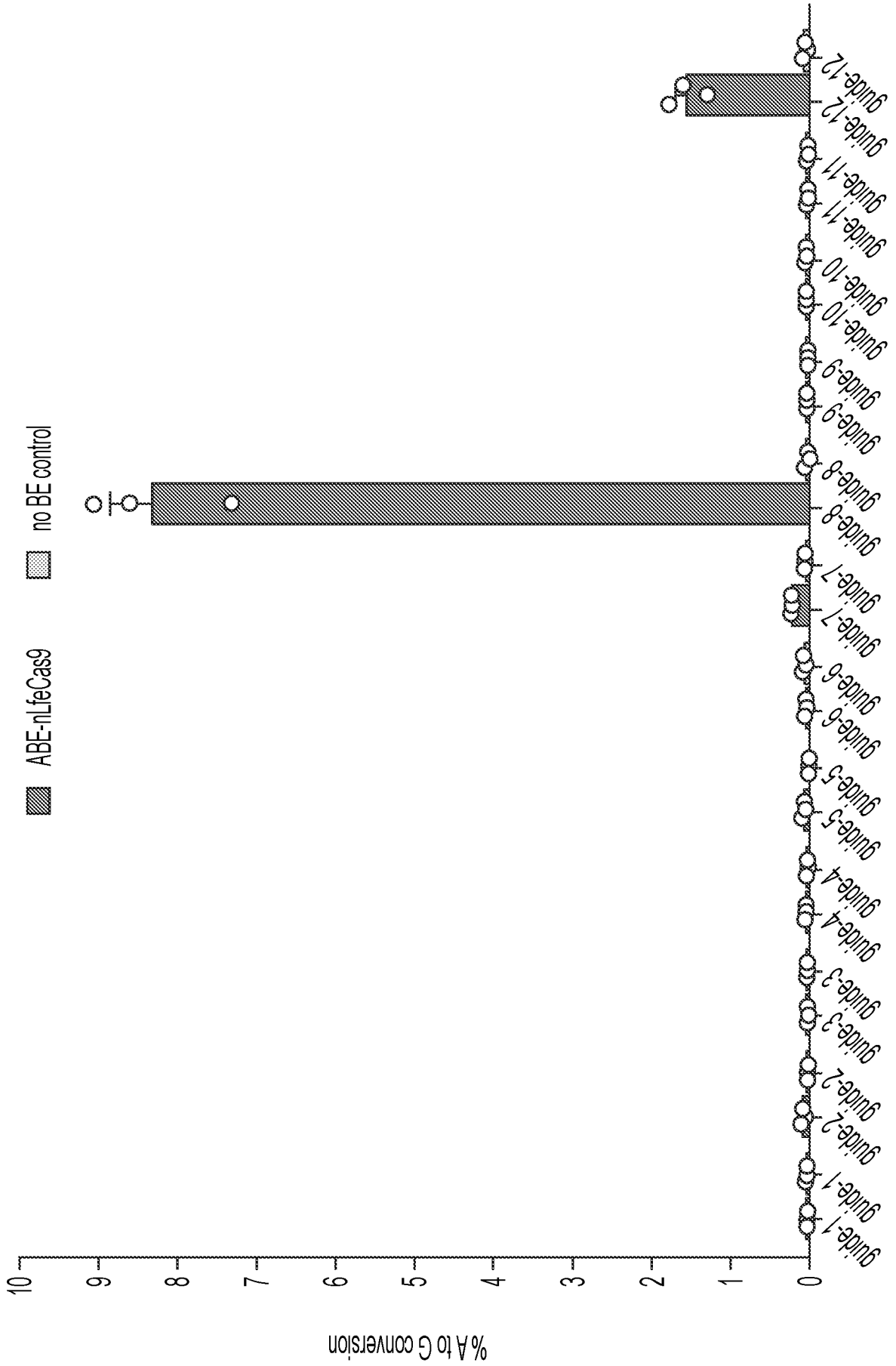


FIG. 9C

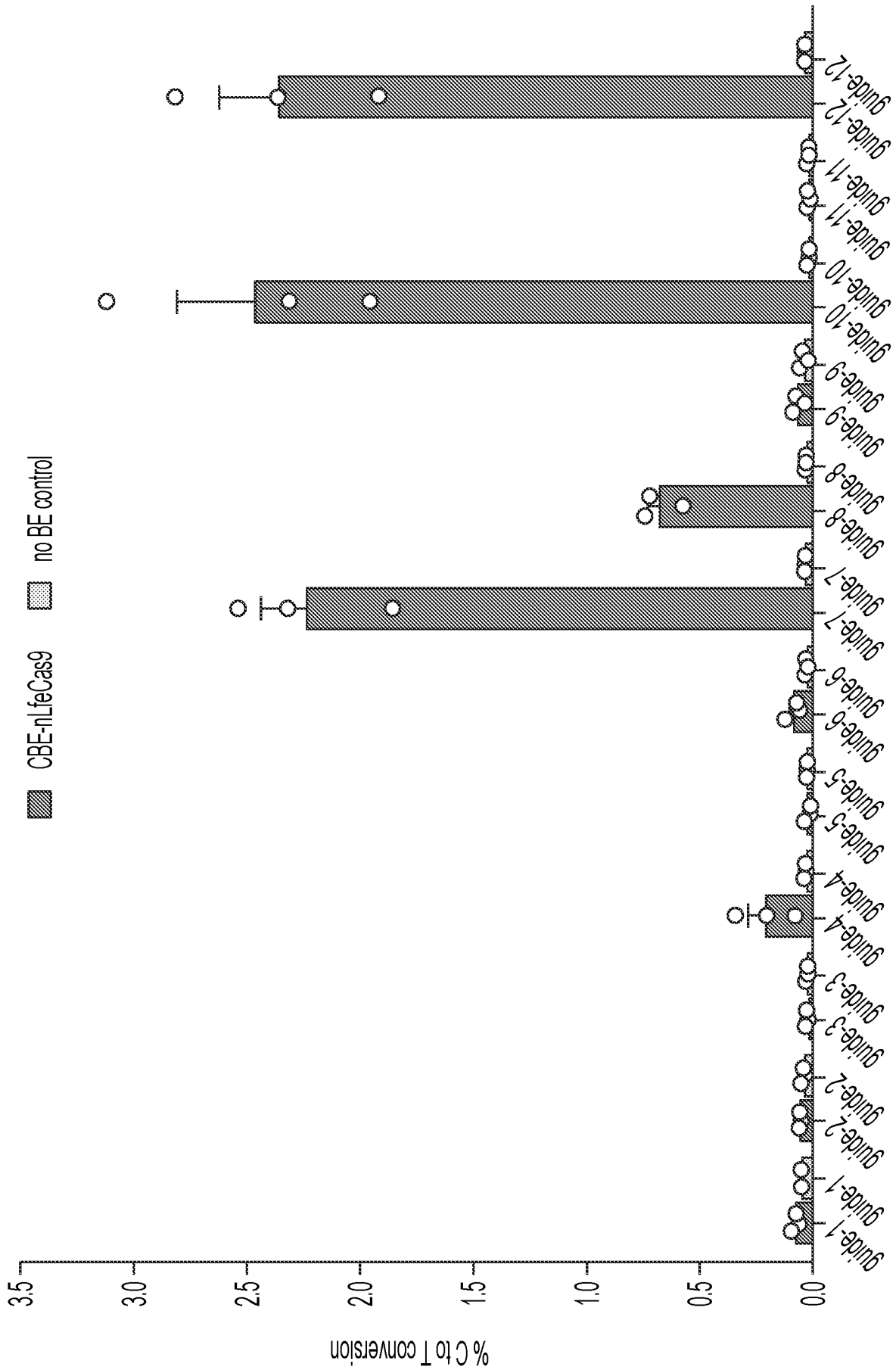


FIG. 9D

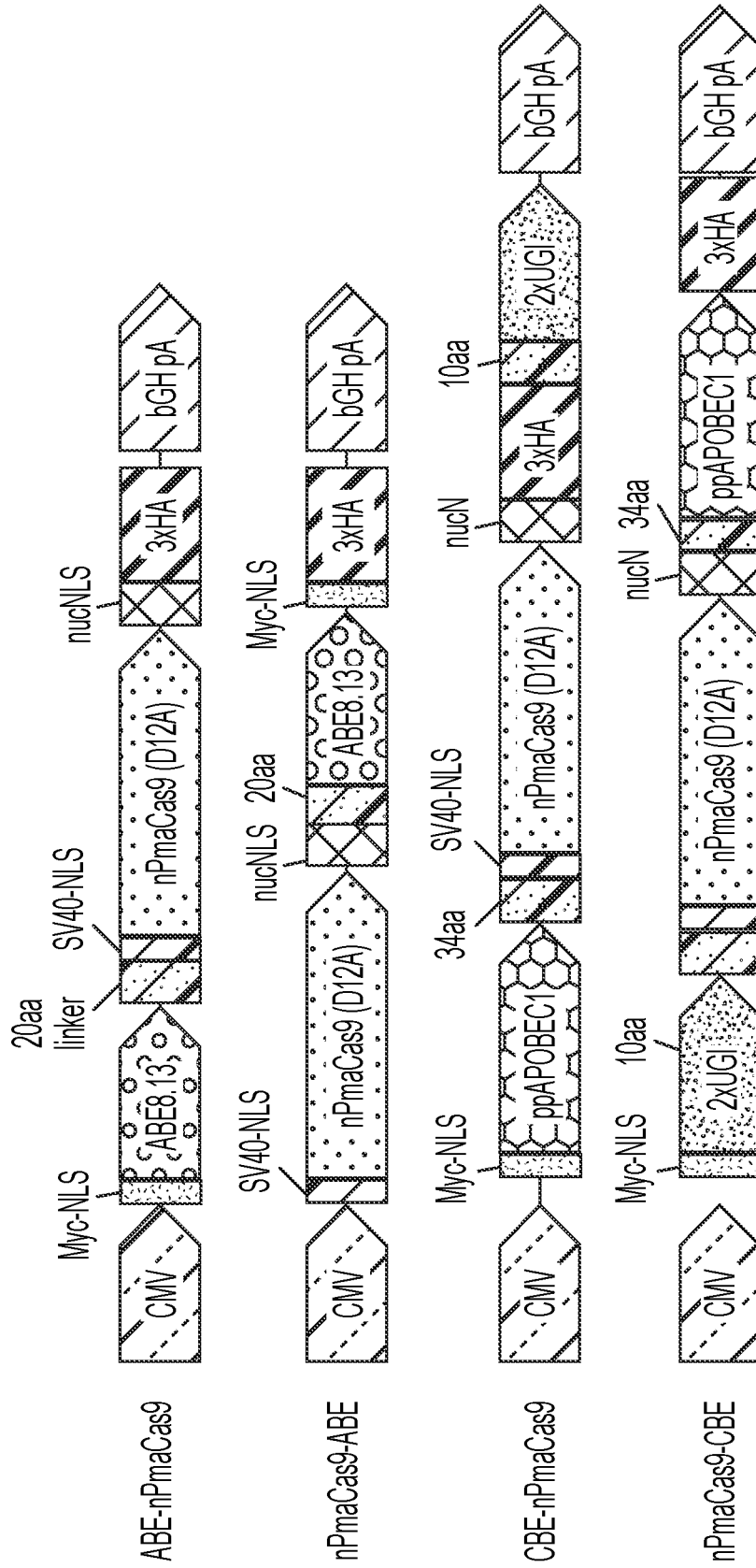


FIG. 10A

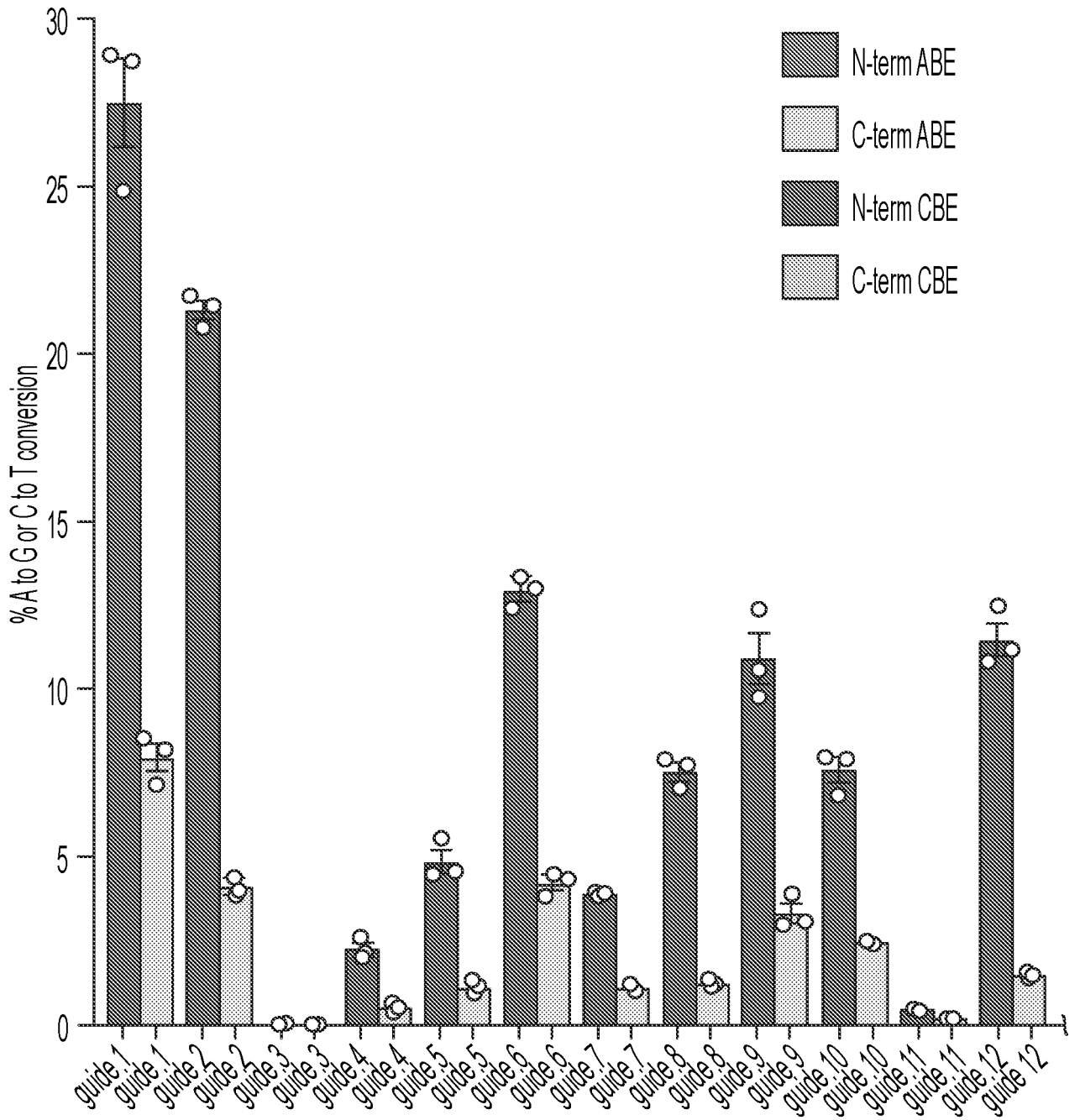


FIG. 10B

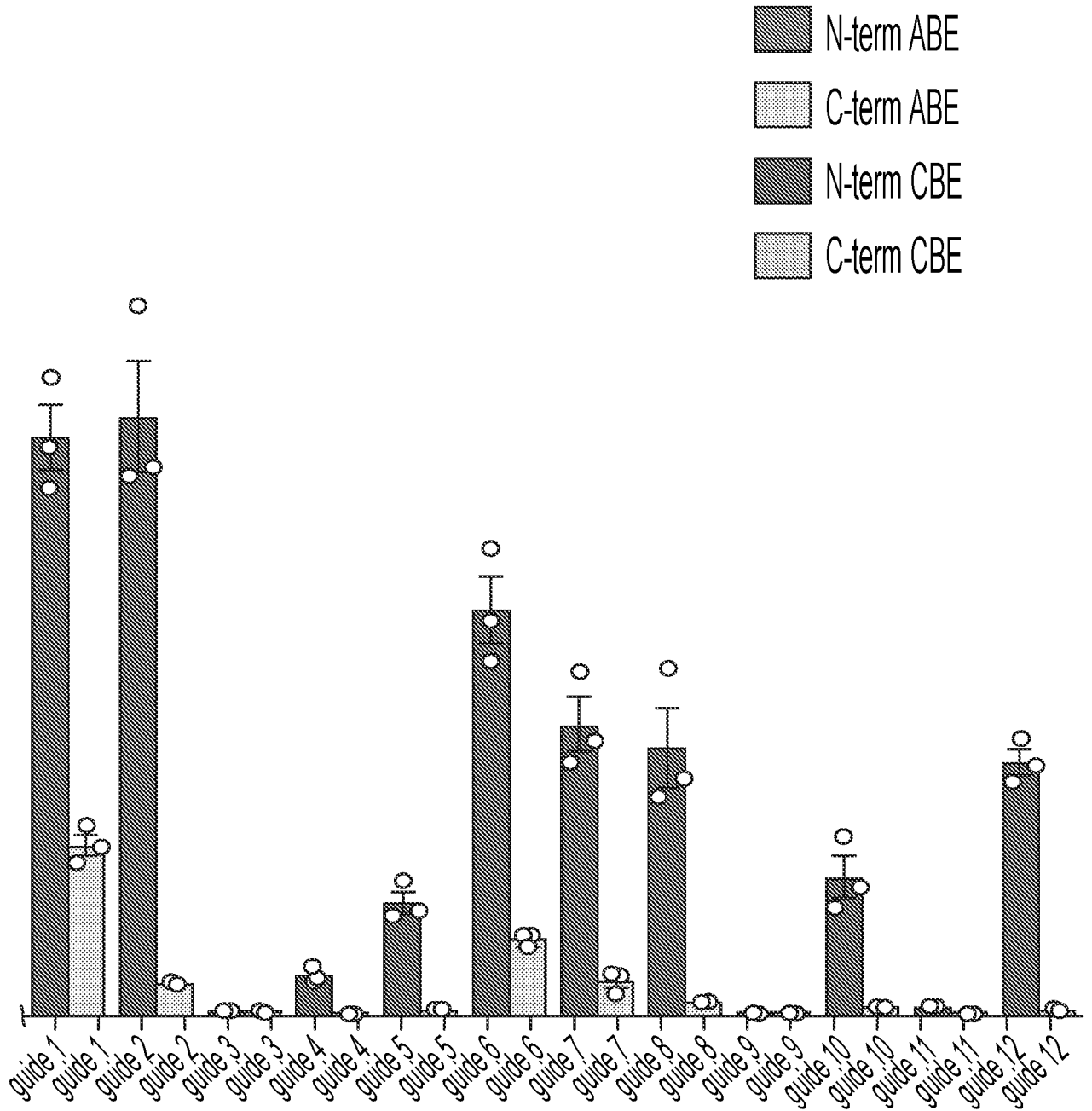


FIG. 10B  
CONTINUED



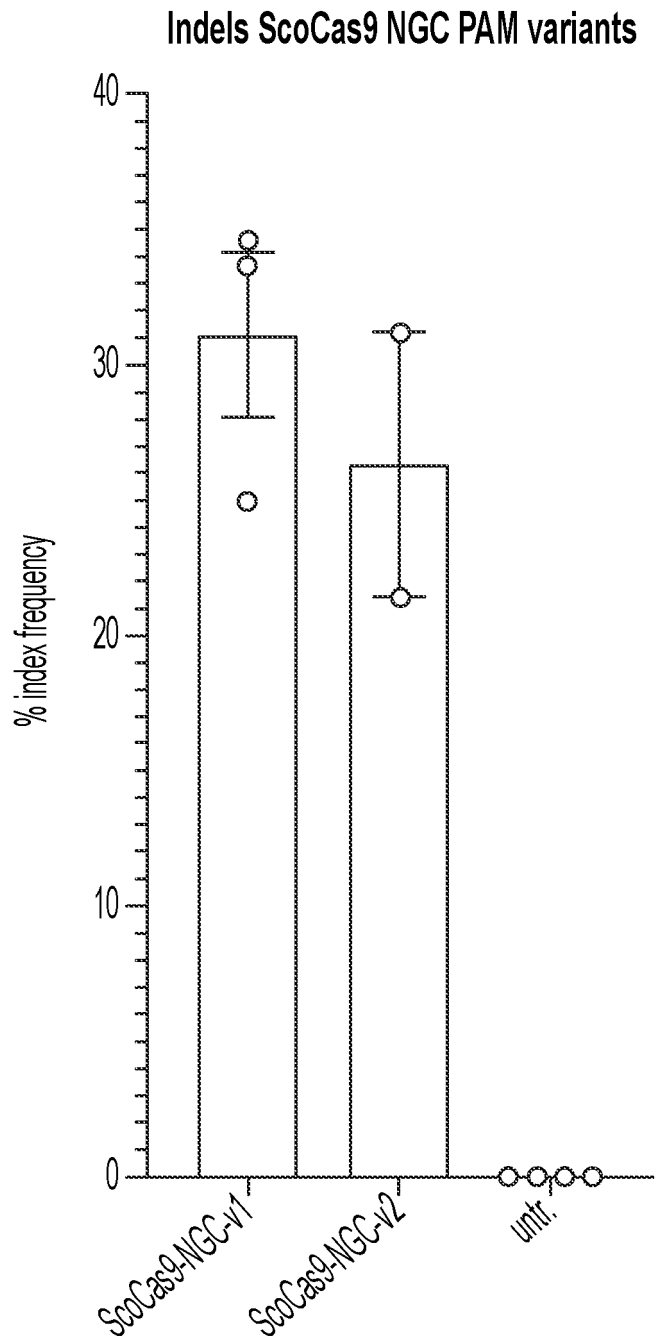


FIG. 11A

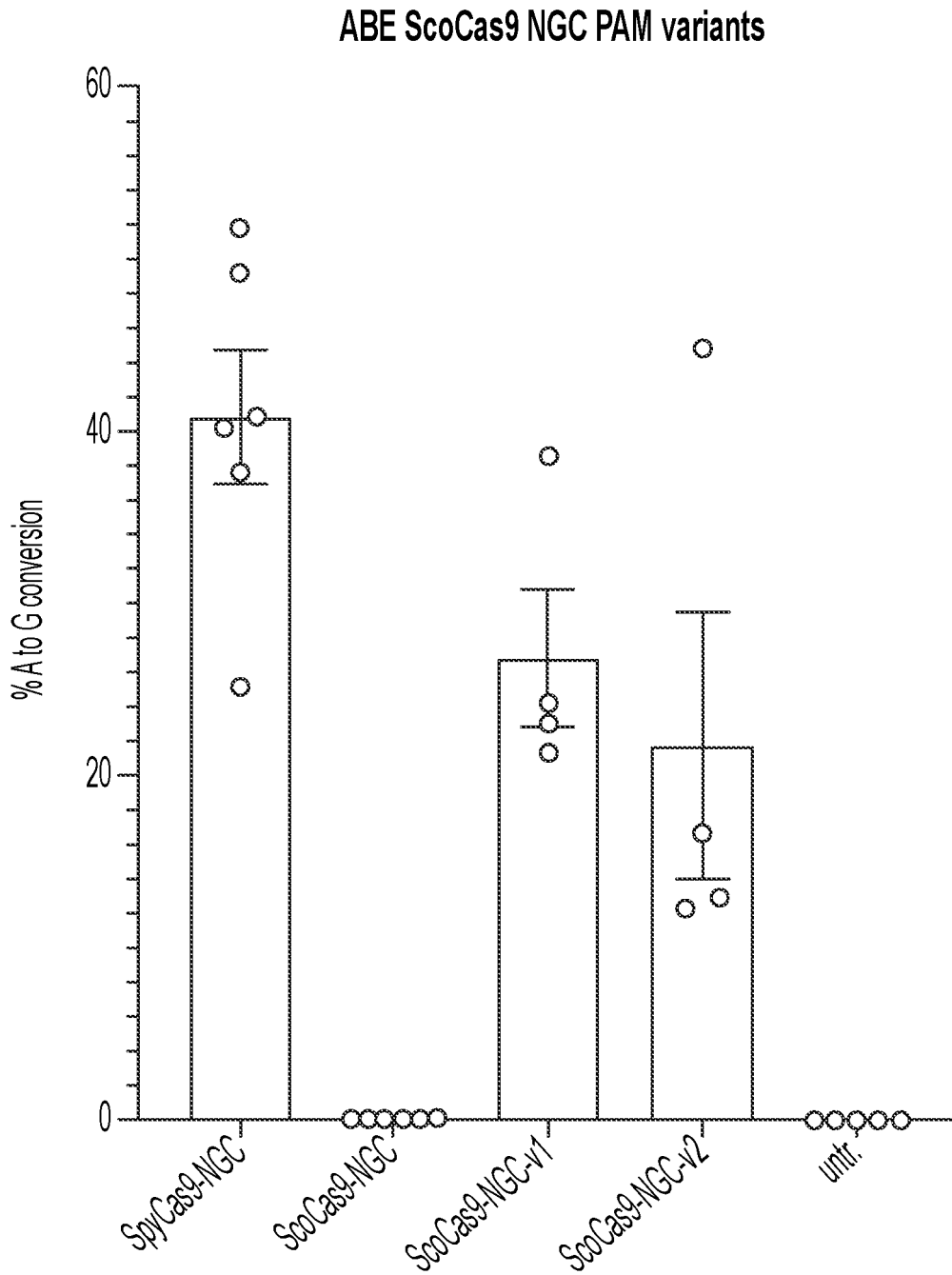


FIG. 11B