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

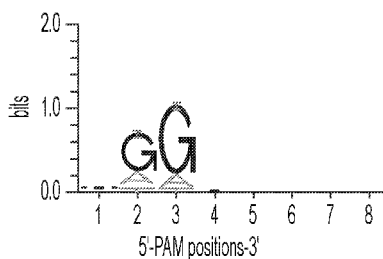


FIG. 1A

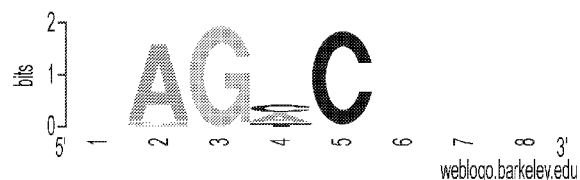


FIG. 1B

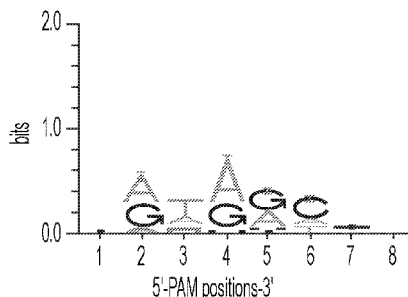


FIG. 1C

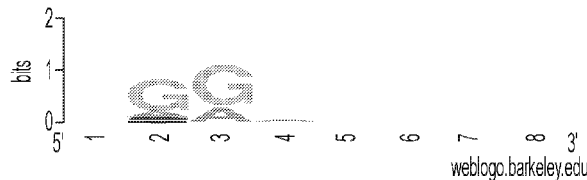


FIG. 1D

(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 peruvensis*, *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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20 October 2022 (20.10.2022)

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

5 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
10 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

MGKPYSIGLDIGTNSVGWAVTDDYKVPAKKMKVLGNTDKQS IKKNLLGALLFDGETAEAT
RLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRGEHHPIFGNIAA
20 EVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFK
DFVEEYDKTIEESHLSEITVDALSILTEKVSKSSRLENLIAHYPTEKKNTLFGNLIALSLDL
HPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAILLSGILTVD
NSTKAPLSASMVKRYEEHQDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEF
YKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHYPFLK
25 ENQDKIEKILTFRIPYYGPLARKGSRFAWEYKADEKITPWNFDDILDKEKSAEKFITRMT
LNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKV
TKDKLNYLNKEFEEFRIVNLTGLDENKAFNSSLGTYHDLRKILDKSFLDDKANEKTIEDI
IQTTLFEDREMIRQRLQKYSIFTKAQLKKLERRHYTGWRSLSYKLINGIRNKENKKTILD
YLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPAICKGILQSVKI
30 VDELVKVMGHNPANIIEMARENQTTDKGRRNSQQRLLQDSLKNLDNPVNICKNVENQQLQ
NDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSDD
VPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKQLVETRQITKHV
AQILDERFNTEDFGNKRRIRNVKIIITLKSNLVSNFRKEFELYKVREINDYHHAHDAYLNAV

GNALLLKYPQLEPEFVYGEYPKNSYRSRKSATEKFLFYSNILRFFKKEDIQTNEDGEIAWN
KEKHIKILRKVLSPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKTKNSYWDPKKYGGF
DSPVVAYSILVFADVEKGKSKLRKVQDMVGITIMEKKRFEKNPVDFLEQRGYRNVRLEKII
KLKPYSILFELENKRRLLASAKELQKGNELVIPQRFTTLYHSYRIEKDYEPHREYVEKHK
5 DEFKELLEYISVFSRKYVLADNNLTKEIMLFSKNKDAEVSSLAKSFISLLTFTAFGAPAAFN
FFGENIDRKRYTSVTECLNATLIHQSIITGLYETRIDLSKLGED (SEQ ID NO: 1).

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

10 NDKTYYDKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQSFAKVYEDIEEK
LDNTLKKFMSLNDLDNLFVDNDINSMITVLSKIYQRSKKADDLLKIMNPTKEERAAYKEFTK
ALVGLKFNVSKMILAQEVKDDKDIELDFSNDYDSTVDGLQAELEYIEFIEMLHSINSWV
ELQDILGNNSTISAAMVERYEEHKNDLRLVKKVIREELPDKYNEVFREDNPKLHNLYLGYIKY
PKNTPVEEFYEYIKRLLAKVDTGEAREILERIDLEKFMLKQNSRTNGSIPYQMOKDEMIQII

15 DNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFEDKQKERILPWNYDQIVD
IDATAEGFIERMQNTGYFPDKPVMAKNSLTISKFEVNLNELNKIRINGKLIPVETKKELLSD
LFMKNKTITDKKLKDWLVTHQYYDTNEELKIEGYQKDLQFSTSAPWIDFTKIFGEINASNY
QLIEKIIYDISIFEDKKILKRRLKVKYQLDDLVDKILKLNWTGWSRLSEKLLTGIKSNSK
ETILSILENSNMNLMEIINDESLGFQIIEESNKKDIEGPFRYDEVKKLAGSPAIKRGIWQA

20 LLVVQEITKFMKHEPSHIYIEFAREEQEKVRTESRIAKLQKIYKDLNLQTKEDQLVYESLKK
EDAOKKIDTDALYLYLQMGKSMYSGKPLDIDKLSTYHIDHILPRSLIKDDSLDNRVLVLPK
ENEWKLDSETVPFEIRNKMMGFWQKLHENGLMSNKKFFSLIRTDFNEKDKKRFINRQLVETR
QIIKNVAIINDHYTNVVTVRAELSHQFRERYKIYKNRDLNLLHAAHDAYIACILGQFIH
QNFGNMDVNMIYGQYKKNYKKDVQEHNNYGFILNSMNHIHFNDDNSVIWDPSYIGKIKSCFC

25 YKDVYVTKLEQNDAKLFDLTLIPSDKNSENGVTAKAIPVNKYRKDVNKYGGFSGDAPIMLA
IEADKGKKHVRQVIAFPLRLKNYNDEERIKFIEKEKNLKNVKILTEVKKNQLILINHQYFFI
TGTNELVNATQLKLSAKNTKNLFNLVDANKHNLLESIDDANFNEVIQELICKLQEPIYSRYN
SIGKEFEDSYEKINAUTQDKLYIEYLIAIMSAKATQGYIKPELAREIGTNGKNKGRIKSFT
TIDLNKTTFISTSVTGLFSKKYKL (SEQ ID NO: 4).

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

MSIINFQRRGLMETQASNQLISSHLKGYPYKDYFVGLDIGTSSVGWAVTNKAYELLKFRSHK
MWGSRLFDEGESAVARRGFRSMRRRLERRKLRLKLLEELFADAMAQVDPTFFMRLRESKYHY
EDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKGSTDDIRKLFLAVHHILKYRGN
FLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGTKSDKAKAIEHLV
5 DTYIATDTVDTSSKTQKDQVKEDKKRLKAFANVLGLNASLIDLFGSVEELEDLKKLQITG
DTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPGLTISESKVKA
FNFNKHKKMTKNRIEQFLK
SLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEY
ILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLNEADGFSVAEKLIKML
EFRIPIYYVGPLNTHHNVDNGGFAWRKASGRVT
10 PWNFDDKIDREKSAAAFIKNLTNKCTYL
LGEDVLPKSSL
LYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTKNRIEQFLK
DNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGESKKMLRE
TLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGT
PETIIILMRNFSYNL
MELLGDKFSFMERIQUEINAKLTEQQIVNPHDIIDDLALS
PAVKRAVWQALRIVDEVAHIKKA
L
15 LPARI
FVEVTRS
NKNEKKKDSRQKRLSDL
YAAIKDDVLLNGLNNEIFGELKSSLAKYDDA
ALRSKKLYLYYTQMGRCA
YTGEIIELS
LLNTDNYDIDHIYPRSL
TKDDSFDNLVLCKRTANA
QKS
DAYPI
SEEI
QKTQKPFWTFLKQQGL
ISERKYERL
TRITPLTADDLS
GFIARQL
VETNQS
VKAATT
LRR
LYPGVDVV
FVKA
ENVTDFRHDNN
FIKVRS
LNHH
HAKDAY
LNIV
VG
NVY
HER
FTRNFRAFF
KKNGAN
RTYNL
AKMFNY
DV
NCTNA
DGKAW
DV
KTS
SMD
TVK
KMM
DS
ND
VR
VT
KR
L
LEQT
GALA
DATIY
KATVAG
KAKD
GAY
IGM
KTK
SSV
FADV
SKYGG
MTKI
KNAY
SI
IV
QYT
TGK
20 KGEV
IKEIV
PLPI
YLT
NRNT
DQD
LIN
YVAS
IIP
QAK
DIS
IIY
GKLC
CIN
QLV
KVNG
F
YY
LG
GKT
NSKFC
CID
NAIQ
VIVS
NEW
I
PYL
K
V
LE
KF
NN
MR
KD
NL
K
AN
VV
STR
AL
DN
K
HT
IE
VR
IV
EEK
NIE
FF
D
Y
LV
SK
L
K
MP
IY
Q
KM
KG
N
K
AA
E
L
SE
K
GY
GL
FK
K
MS
L
EE
QS
I
H
L
I
E
L
N
L
L
T
N
Q
K
TT
FEV
K
PL
GIT
ASR
STV
GSK
ISN
Q
DEF
KV
INES
IT
GLY
S
NE
VT
IV
(SEQ ID NO: 8).

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

MTKV
KDYY
IGLD
IGTSS
VGWAVT
DEAY
NVLF
NSKK
MWGV
RLF
DDAK
TAEERR
GQRG
ARRRL
DRKK
KERL
SLLQ
DFFA
EEVA
KVD
PNFL
RLDN
SDLY
MEDK
DQKL
KSKY
TLF
NDK
DFKD
KNFH
K
YPTI
HLL
MDL
IED
DSK
KD
DIRL
VY
LACHY
LL
KNRG
HF
FEGQ
KFDT
KSS
FENS
L
NEL
KVHL
N
DEY
GLD
LEFD
NEN
LIN
ILTDP
KLN
TAK
KKEL
K
SVI
GDT
KFL
KAV
SAIM
IGSS
QKL
VDL
FE
30 NP
EDF
DSA
IKS
VDF
STTS
FDD
K
SDY
E
LA
GDK
K
IAL
VN
IL
KEI
YD
SS
I
LEN
L
KEAD
KSKD
GN
KY
ISNA
FVK
KYN
KHG
QDL
KEF
KRL
VRQ
YH
KSAY
FD
IF
RSE
K
V
ND
N
Y
V
SY
TK
S
S
NN
K
RV
K
ANK
FTD
QEAF
Y
KFA
KKH
LET
I
KY
K
INK
VNG
SKAD
LE
L
IDG
MLR
D
MEF
K
NF
MP
K
IK
SS
D
NG
V
I
PY
QL
K
L
M
E
LN
K
I
LEN
Q
SK
H
E
FL
N
VS
DEY
GS
V
CD
K
IAS
IME
F
RI
PI
YY
VG
PL
NP
NS
SKY
AW
IK

KQKDSEITPWNFKDVVDLDSREEFIDSLIGRCTYLKDEKVLPKASLLYNEYMVNLNELNNLK
LNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGDFKQGLNSYND
FKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGKYFTDSEIKKMAGLNYKD
WGRLSKKLLTGLEGANKITGERGSIIHFMREYNLNLMELMSASFTFTEEIQKLNPVDDRKLS
5 YEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIEMARGKEEVKARKESRKNQLLK
FYKDGKKAFISEIGEERYSYLLSEIEGEEENKFRWDNLYLYTQLGRCMYSLEPIDISELSS
KNIYDQDHYPKSKIYDDSIENRVLVKDKLNSKKGNSYPIPDEILNKNCYAYWKILYDKGLI
GQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTICKNSEIVYSAENASRFR
QEFDIVKCRAVNLDLHHMHDAYINIIVGNVYNTKFTKDPMNFKVKKQEKARSYNLENMFKYDVK
10 RGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRKNKGSRPLK
DKGPKSSIEKYGGYTNINKACFAVLDIKSKNKIERKLMPVEREIYAKQKNDKKLSDEIFSKY
LKDRFGIEDYRVVYPVVKMRTLKIDGSYYFITGGSDKTLELRSALQLILPKKNEWAIKQID
KSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLFQDDKIENIDFKFKSMDF
KEKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGRLOSSKTNIGNYQEFKIINQSITGLFENE
15 VDLLKL (SEQ ID NO: 14)

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

MKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAERRTFRTTRRLKRR
KWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEDPAKNQAFIGKLLFPDLLKKNERGYPTL
20 IKMRDELPVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAHHIVKYRGHFLNNASVDKF
KVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLDRQKAVAKLLEV
KVADKEETKRNKQIATAMSKLVLYKADFATVAMANGNEWKIDLSSETSEDEIEKFREELSD
AQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWTHERQLAEVKEYLATQPASARKEFD
25 QVYNKYIGQAPKEKGFDLEKGLKKILSKKENWKEIDELLKAGDFLPKQRTSANGVIPHQMHQ
QELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPIYYGVPLVTPEVQKATSG
AKFAWAKRKEDGEITPWNLWDKIDRAESAFAFIKRMTVKDTYLLNEDVLPANSLLYQKYNVL
NELNNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMAKTRGVNKPSVEGLSDPKKF
30 NSNLATYDLKSIVGDKVDDNRYQMDLENIEWRSVFEDGEIFADKLTEVEWTDEQRSALV
KKRYKGWGRLSKKLLTGIVDENQRIIDLWNTDQNFMQIVNQPVFKEQIDQLNQKAITNDG
MTLRERVESVLDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSISIEFARNEGNGEITRS
RRTQLQKLFEDQAHELVKDTSLTEELEKAPDLSDRYYFYFTQGGKDMYTGDPIINFDEISTKY
DIDHILPQSFVKDDS LDNRVLVSRAENNKKSDRVPALKYAAKMKPYWNQLKQGLITQRKFE
NLTMDVDQTICKYRSLGFVKRQLVETRQVIKTANILGSMYQEAGTDIETRAGLTQLREEF

DLPKVREVNDYHHAVDAYLTTFAGQYLNRYPKLRSFFVYGEYMKFKHGSDLKLRNFNFHE
LMEGDKSQGKVVDQQTGELITRDEVADYFDWVINLKVMLISNETYEETGKYFDASHESSSL
YLKNQNKKSKLVVPLKNKLQPEYYGAYTGITQGYMVLKLLDKGGFGVYRIPRYAADILNK
CHDEVAYRNKIAEIISSDPRAPKSFEVVVPRVLKGTFLVGEEKFILSSYRYKVNATQLILP
5 VSDIKLIQDNFKALKLNEMQTKKLIEIYDNILRQVDKYYKLYDINKFRAKLHDGRSKFVE
LDDFGQDASKEKVIIKILRGLHFGSDLQNLKEIGFGTTPLGQFQVSEAGIRLSNTAFIIFKS
PTGLFNRKLYLKNL (SEQ ID NO: 84).

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

10 MEKKTNYTIGLDIGTDSVGWAVVKDDLELVKKRMKVILGNTEINYIKKNLWGSLLFESGOTAK
DRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFFRLNESFLVPEDKAFSKNPIFGTL
GEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIAINEN
LEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDGRANKQSPMMA
FITLIVGKKAKFKAAFNLEEEISLNLTDDSYDENLEILLNTIGSDFADLFDHAQRVYNAVEL
15 AGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKKDLKG
YAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVHLAEME
AILDROENYYPWLKENREKIISLLTFRIPIYYVGPLADGQSEFAWLERKSDEKIKPWNFSDVV
DLDRSAEKFIEQLIGRDTYLPDEYVLPKKSLIYQKYMVFNELTKIAYLDERQKRMNLSSVEK
KEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMM
20 DPDNEDLMEEIVKILTVFEDRKMRKQLEKYKERLSPEQIKEAKHYTGWGRLSKKLLVGI
RDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDDRLSFKKTIAELQMDPSADLYAQVQ
EIAGSPAICKGILLGLKIVDEIIRVMGEKOPENIVIEMARENQTTARGKALSKRREAKIKEGL
AALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHIIPQSFTVDN
SLDNLVLTNSSQNRGNKKDDVPSLEVNRQLAYWRSLKAGLMTQRKFDNLTKAMRGGLTDK
25 DRERFIQRQLVETRQITKNAKLLDMRLNDKDEAGNKIRETNIVLLKSAMASEFRKMFRLY
KVRELNDYHHAHDAYLNAAIAINLLALYPYMADDFVYGEFRYKKKPQAEKATYEKLRQWNLI
KRFGEKQLFTPDHEDCWNKERDIKITIKKVMGYRQVNVVKKAEERTGMLFKETINGKTNKGSR
IPIKKDLDPSKYGGYIEEKMAYYAVISYEDKKKPGKTIVGISIMDKFEYDSISYLGKLG
FSNPVVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKANQIAMEQDLVNFIYHLKNYD
30 EISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIYEEEKKDPVDIEALVA
SFIELLKLTSAGAPADFIFMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMRFED (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.

5 In some embodiments, the Cas9 protein further comprises a nuclear localization 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

MPKKKRKVGGKPYSIGLDIGTNSVGAVVTDDYKVPACKMKVLGNTDKQSIKKNLLGALLFD
SGETAETRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSLVDEDKRGEHH
10 PIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAEN
TDVQALFKDFVEEYDKTIEESHLEITVDALSILTEVKVSSRLENLIAHYPTEKKNTLFGN
LIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAILL
SGILTVDNSTKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYIE
SGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQ
15 GEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRFWAEYKADEKITPWNFDDILDKEKSA
EKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKVYVNEQGEAKFFDTNMKQEIFDH
VFKENRKVTDKLNYLNKEFEEFRIVNLTGLDENKAFNSSLGTYHDLRKILDKSFLDDKA
NEKTIEDIIQTLTFEDREMIRQLQKSDIFTKAQLKKLERRHYTGWGRRLSYKLINGIRNK
ENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPAICKK
20 GILQSVKIVDELVKVMGHNPANIIEMARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNIK
NVENQQLQNDRFLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRSD
KNRGKSSDVPSIEVVHEMKSFWSKLLSVKLITQRKF DNLTKAERGGLTEEDKAGFIKRQLVE
TRQITKHVAQILDERFNTEDGDNKRRIRNVKIITLKSNLVSFRKEFELYKVREINDYHHAH
DAYLNAVVGNNALLKYPQLEPEFVYGEYPKNSYRSRKSATEKFLFYSNILRFFKKEDIQTN
25 EDGEIAWNKEKHIKILRKVLSYPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKTKNSYW
DPKKYGGFDSPVVAYSILVFADVEKGKSKKLRKVQDMVGITIMEKKRFEKNPVDLFEQRGYR
NVRLEKIIKLPKYSLFELENKRRRLLASAKELQKGNELVIPQRFETLLYHSYRIEKDYEP
REYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMLFSKNKDAEVSSLAKSFISLLFTA
FGAPAAFNFFGENIDRKRYTSVTECLNATLIHQSTITGLYETRIDLSKLGEDGKRP
30 QAKKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 2).

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

MPKKKRKGAKNNDIRYSIGLDIGTNSVGWAVMDEHYELLKKGNHHMWGSRLFDAEPAATR
RASRSIRRKYRKRRERIRLLRDLLGDMVMEVDPTFFIRLLNVSLDEEDKQKNLGNDYKDNY
NLFIEKDFNDKTYDKYPTIYHLRKELCENKEADPRLIYLALHHIVKYRGNFLKEGQSFAK
VYEDIEEKLDNTLKKFMSLNLDNLFVDNDINSMITVLSKIYQRSKKADDLKIMNPTKEER
5 AAYKEFTKALVGLKFNVSKMILAQEVKDDKIELDFSNDYDSTVDGLQAELEYIEFIEM
LHSINSWVELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPDKYNEVFREDNPKLH
NYLGYIKYPNTPVEEFYEYIKRLLAKVDTGEAREILERIDLEKFMLQONSRTNGSIPYQM
KDEMIQIIDNQSVPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFEDKQKERILP
WNYDQIVDIDATAEGFIERMQNTGTYFPDKPVMAKNSLTFSKFEVLNELNKIRINGKLIPVE
10 TKKELLSDLFMKNKTITDKKLKDWLVTHQYYDTNEELKIEGYQKDLQFSTSAPWIDFTKIF
GEINASNYQLOIEIIYDISIFEDKKILKRRLLKVVQLODDLLVDKILKLNWTGWSRLSEKLLT
GIKSKNSETILSILENSNMNLMEIINDES LGFKQIIEESNKKDIEGPFRYDEVKKLAGSPA
IKRGIWQALLVVQEITKFMKHEPSHIYIEFAREEQEKRATESRIAKLQKIYKDLNLQTKEDQ
LVYESLKKEDAKKIDTDALYLYLQMGKSMYSGKPLDIDKLSTYHIDHILPRSLIKDDSLD
15 NRVLVLPKENEWKLDSETVPFEIRNKMMGFWQKLHENGLMSNKKFFSLIRTDFNEKDKKRFI
NRQLVETRQIINKNAVIINDHTNTNVVTVRAELSHQFRERYKIYKNRDLNDLHHADAYIA
CILGQFIHQNFGNMDVNMIYGQYKKNYKKDVQEHNNYGFILNSMNHIHFNDNSVIWDPSYI
GKIKSCFCYKDVYVTKKLEQNDAKLFDLTILPSDKNSENGVTKAKIPVNKYRKDVNKYGGFS
GDAPIMLAIEADKGKKHVRQVIAFPLRLKNYNEERIKFIEKEKNLKNVKILTEVKKNQLIL
20 INHQYFFITGTNELVNATQLKLSAKNTKNLFNLVDANKHNLESIDDANFNEVIQELICKLQ
EPIYSRYNSIGKEFEDSYEKINAUTQDKLYIIEYLIAIMSAKATQGYIKPELAREIGTNGK
NKGRIKSFTIDLNTTFISTSVTGLFSKKYKLGKRPAATKKAGQAKKKGSYPYDVPDYAYP
YDVPDYAYPYDVPDYA (SEQ ID NO: 5).

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

MPKKKRKGSIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLDIGTSSVGWAVTNKAYE
LLKFRSHKMWGSRLFDEGESAVARRGFRSMRRLERRKLRLKLEELFADAMAQVDPTFFMR
LRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVH
HILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGTKSDK
30 AKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANVLGLNASLIDLFGSVEELED
LKKLQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPLTISESKVKAFNKH
KDDLAILKSLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGL
SDSKDKEYIILSQIELQILLPLQRIKDNGVIYQLHLEELKAILAKCGPKFPFLNEVADGFSV

AEKLIKMLEFRIPIYYVGPLNTHHNVNDNGGFAWAVRKASGRVT PWNFDDKIDREKSAAAFIKN
LTNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKHEHLIEAVFKQDHKKMTK
NRIEQFLKDNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEI ITDITIFG
ESKKMLRETLRKKFASCLDEAIKKLTKRYRDWGRLSQKLLNGIEGCDKAGDGTPETIIIL
5 MRNFSYNLMELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVD
EVAHIKKALPARIFVEVTRSNKNEKKKDSRQKRLSDLYAIIKKDDVLLNGLNNEIFGELKS
SLAKYDDAALRSKKLYLYTQMGRCAYTGEIIELSLLNTDNYDIDHIYPRSLTKDDSFDNLV
LCKRTANAQKSDAYPISEEIQKTQKFWTFLKQQGLISERKYERLTRITPLTADDLSGFIAR
QLVETNQSVKAATLLRRLYPGVDVVFVKAENVTDFRHNNFIKVRSLNHHHHAKDAYLNIV
10 VGNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKGKAADVKTSMDTVKKMMD
NDVRVTKRLLEQTGALADATIYKATVAGKAKDGAYIGMKTKSSVFADVSKYGGMTKIKNAYS
IIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISIYGKLCINQLVKV
NGFYYYLGGKTNSKFCIDNAIQVIWSNEWIPYLVLEKFNNMRKDNKDLKANVVSTRALDNK
HTIEVRIVEEKNIEFFDYLVSKLKMPIYQKMGNKAAELSEKGYGLFKKMSLEEQSIIHLIEL
15 LNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIVGKRPAATKK
AGQAKKKGSYPYDVPDYAPYDVPDYAPYDVPDYA (SEQ ID NO: 9).

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

MPKKKRKVGTKV DYYIGLDIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAEERRG
20 QRGARRRLDRKKERLSLLQDFFAEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKD
FKDKNFHKKYPTIHLLMDLIEDDSKKDIRIVYLACHYLLKNRGHFIFEGQKFDTKSSFENS
LNELKVLNDEYGLDLEFDNENLINILTDPKLNKTAKKKELKSVIGDTKFLKAVSAIMIGSS
QKLVDLFENPEDFDDSAIKSVDFSTTSFDDKYS DYELALGDKIALVNIIKEIYDSSILENLL
KEADKSKDGNKYISNAFKVVKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNDNYVSYTKS
25 SISNNKRVKANKFTDQEAFYKFAKKLETI KYKINKVNGSKADLELIDGMLRDMEFKNFMPK
IKSSDNGVI PYQLKLMELNKILENQSKHHEFLNVSD EYGSVCDKIASIMEFRIPIYYVGPLNP
NSKYAWIKKQKDSEITPWNFKDVVDLDSSREEFIDS LIGRCTYLKDEKVLPKASLLYNEYMV
LNELNNLKLNDLPITEEMKKIFDQLFKTRKKVTLKAVANLLKEFNINGEILLSGTGDFK
QGLNSYNDFKAIVGDKVDSDDYRDKIEEIIK LIVLYGDDKS YLQKKIKAGYGKYFTDSEIKK
30 MAGLNYKDWGRSLSKKLLTGLEGANKITGERSIIFMREYNLNLMSASFTFTEEIQKLN
PVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSSKKIFIEMARGKEEVKARKE
SRKNQLLKFYKDGGKKA FISEIGEERYSYLLSEIEGEEENKFRWDNLYLYTQLGRCMYSLEP
IDISELSSKNIYDQDH IYPKSKIYDDS IENRVLVKKDLNSKKGNSYPIPDEILNKNCYAYWK

I LYDKGLIGQKKYTRLTRRTGFTDELVQFISRQIVETRQATKETANLLKTICKNSEIVYSK
AENASRFRQEFDIVKCRAVNDLHHMH DAYINIIVGNVYNTKFTKDPMN FVKKQE KARSYNLE
NMFKYDVKRGGYTAWIADDEKGTVKNASI KIRKELEG TNYRFTRMNYIESGALFNATLQRK
NKGSRPLKDKGPKSSIEKYGGYTNI NKACFAVLDIKSKNKIERKLMPVERE IYAKQKNDKKL
5 SDEIFSKYLKDRFGIEDYRVVYPVVKMRTL LKIDGSYYFITGGSDKTLELRSALQLILPKKN
EWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDITVNKFKTSVFKKSFLNL FQDDK IENID
FKFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGR LSSKTNNIGNY QEFKIINQS
ITGLFENEVDLLKLGKRPAATKKAGQAKKKGSYPYDVPDYA PYDVPDYA PYDVPDYA
(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

MPKKKRKVGKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAERRTFR T
TRRLKRRKWRHLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEDPAKNQAFIGKLLFPDLLKK
NERGYPTLI KMRDEL PVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLA VHHIVKYRGHFL
15 NNASVDFKVG RIDFDKS FNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLD RQK
AVAKLLEVKVADKEETKRNKQIATAMS KLV LGYKADFATVAMANGNEWKIDLSSETSEDEIE
KFREELS DAQNDILTEITS LFSQIMLNEIVPNGMSI SESMMDRYWTHERQLAEVKEYLATQP
ASARKEFDQVYNKYIGQAPKEKGFDLEKG LKKILSKKENWKEI DELLKAGDFLPKQRTSANG
VIPHQMHQQELDRIIEKQAKYY PWL ATENPATGERDRHQAKYELDQ LVS FRIPYYVGPLVTP
20 EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESA EAFIKRMTVKDTYLLNEDVLPANS L
LYQKYNVLNELNNVRVN GRRLSVG IKQDIYTELFKKKKTVKAGDV ASLVMAKTRGVN KPSVE
GLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENII EWR SVFEDGEI FADKL TEVEWL T
DEQR SALVKKRYKGWGR LSKKLLTGIVDENGQRIIDL MWNTDQN FMQIVN QPVFKEQIDQLN
QKA ITNDGMTLRERVESVLLDAYTSPQNK KAIWQV VRVVEDIVKAVGNAPKSISIEFARNEG
25 NKGEITRS RRTQLQKL FEDQA HELVKDTSLTEELEKAPDLS DRYYFYFTQGGKDMYTGD PIN
FDEISTKYDIDHILPQS FVKDDSLDN RVL VSRAENN KKS DRVPAKLYAAKMKP YWNQ LLKQ G
LITQRKFENLTMDQTIKYRSLGFV KRQLVETRQVI LTAN ILGS MYQEAGTDIETRAGL
TKQLREEFDLPKV REVNDYH AVDAYLTT FAGQYLN RRYPKLRSFFVYGEYMKFKHGSDLKL
RNFNFFHELMEGDKSQGKVV DQQT GELIT TRDEVADYFDWVINLKVMLISNETYEETGKYFD
30 ASHESSSLYLNQNKSKL VVPLK NKLQPEYYGAYT GITQGYM VILK LLDKGGFGVYR IPR
YAADI LNKCHDEV AYRN KIAEIISSD PRAPKS FEVV VPRVLKGTFLV DGE EK FILSSY RY KV
NATQLI LPVSDIKL IQDNF KALK KLN VEMQ TKKL IEIYDNILRQVDKYYKLYDINKFRAKL H
DGRSKFVELDDFGQDASKEK VIIK IRLGLHFGS DLQNLKEIGFGTTPLGQFQVSEAGIRLSN

TAFIIIFKSPTGLFNRKLYLKNLGKRPAATKKAGQAKKKGSYPYDVPDYAYPDVDVPDYAYPD
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 MPKKRKVGEKKTNYTIGLDIGTDSVGAVVKDDLELVKKRMKVLGNTE NYIKKNLWGSLL
FESGQTAKDRRLKRVARRRYERRNRNLTELQKIFAPAIDEVDENFFFRLNESFLVPEDKA
FSKNPIFGTLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDT
EHIAINENLEQFFESYNALFSEEPIELKEELIAENILREKNSRTVKEKRITSFLKD
GINKQS PMMAFITLIVGKKAKFKAAFNLEEEISLNLTDDSYDENLEILLNTIGSDFADLF
DHAQ
10 RVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAP
KA
AQKKDLKGYAGYIDGNMSQDFYKFKVDQLKEVPGSEKF
LDSIEKEDFLRKQRSFYNGVIPN
QVHLAEMEAILDRQENYYPW
LKENREKIISLLTFRI
PYVGPLADGQSEFAWLERKSDEKIK
PWNFS DVVDLDRSAEKF
IEQLIGRDTYLPDEYVLP
KKSLIYQKYMVFNE
LT
KIA
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ERQKR
MNLSSVEKKEI
FETLFKKRS
KVTE
QLV
KF
FENYL
QIDN
PTI
FG
IEDA
FNAD
YSTY
VELAKV
15 PGMKSMMDDPDNEDLMEEIVKILT
VFEDRK
MRRK
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(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 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)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVNNRVIGEGWNRAIGLH
DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA
GSIMDVLLHPGMNHRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG
TSESATPESSGPKKRKVGGKPYSIGLAIGTNVGWAVTDDYKVPACKMKVLGNTDKQSIK
10 KNLLGALLFDGETAETRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQLDDFL
VDEDKRGEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHF
LYEGDLKAENTDVQALFKDFVEEYDKTIEESHLEITVDALSIILTEKVSKSSRLENLIAHYP
TEKKNTLFGNLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDEGLGEVGDEYADLFAS
AKNLYDAILLSGILTVDNSTKAPLSASMVKRYEEHQDLKKLKDFIKVNAPDQYNAIFKDK
15 NKKGYASYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHL
QEMHAILRRQGEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRFAWAELYKADEKITPWNF
DDILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFD
TNMKQEIFDHVFKENRKVTDKLNYLNKEEEFRIVNLTGLDENKAFFNSSLGTYHDLRKI
LDKSFLDDKANEKTIEDITIQTTLFEDREMIRQRQLQKSDIFTKAQLKKLERRHYTGWRGLS
20 YKLINGIRNKENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVVH
DLPGSPAIIKKGILQSVKIVDELVKVMGHNPAIIIEMARENQTDKGRRNSQQLKLLQDSL
KNLDNPVNIKVENQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNS
LDNRVLTRSDKNRGKSDDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGGLTEEDK
AGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKSNLVSFRKEFELYKV
25 REINDYHHAHDAYLNAVGNALLKYPQLEPEFVYGEYPKNSYRSRKSATEKFLFYSNILR
FFKKEDIQTNEDEIAWNKEKHIKILRKVLSPQVNIVKKTEEQTGGFSKESILPKGESDKL
IPRKTKNSYWDPKKYGGFDSPVVAYSILVFADVEKGKSKLRKVQDMVGITIMEKKRFEKNP
VDFLEQRGYRNVRLEKIIKLPKYSLFELENKRRLLASAKELQKGNELVIPQRFTTLLYHSY
RIEKDYEPEHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKEMLFSKNKDAEVSSLAK
30 SFISLLTFTAFCGAPAAFNFFGENIDRKRYTSVTECLNATLIHQSITGLYETRIDLSKLGEDG
KRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 20);

(b)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH
DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA
GSLMDVLHHPGMNRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG

5 TSESATPESSGPKKRKVGAKNDIRYSIGLAIGTNSVGWAVMDEHYELLKGHNHHMWGSRL
FDAAEPAACTRASRSIRRYNKRERIRLLRDLLGDMVMEDPTFFIRLLNVSFLDEEDKQK
NLGNDYKDNYNLFIEKFNDKTYYDKYPTIYHLRKELCENKEADPRLIYLALHHIVKYRGN
FLKEGQSFAKVYEDIEEKLDNTLKFKMSLNLDNLFVDNDINSMITVLSKIYQRSKKADDLL

KIMNPTEERAAYKEFTKALVGLKFNVS KMLAQEVKKDDKIELDFS NVDYDSTVDGLQAE

10 LGEYIEFIEMLHSINSWVELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPDKYNE
VFREDNPKLHNLYLGVIKYPKNTPVEEFYEYIKRLLAKVDTGEAREILERIDLEKFMLKQNSR
TNGSIPYQMOKDEMIOIIDNQSVYPQLKENREKLISILEFRIPIYYFGPLNTHSEFAWIKKF
EDKQKERILPWNYDQIVDIDATAEGFIERMQNTGTYFPDKPVMAKNSLTISKFEVNLNELNKI

RINGKLI PVETK KELLSDL FMKNKTITDKKLKDWLVT HQYYDTNEELKIEGYQKDLQFSTS L

15 APWIDFTKIFGEINASNYQ LIEKIIYDISIFEDKKILKRRKKVYQLDDLVDKILKLN YTG
WSRLSEKLLTGIKS KNSKETILSILENSNMNLMEIINDES LGFKQIIEESNKKDIEGPFRYD
EVKKLAGSPA IKRGWI QALLVVQEITKFMKHEPSHIYIEFA REEQE KV R TESRIAKLQKIYK
DLNLQTKEDQLVYESLK KEDAKKKIDTD ALYLYLQMGKSMYSGKPLDIDKLSTYHIDHILP

RSLIKDDSLDNRVLVLPKENEWKLDSETVPFEIRNKMMGFWQKLHENGLMSNKKFFSLIRT D

20 FNEKDKKRFINRQLVETRQIIKNVAIINDHYTNVNTVRAELSHQFRERYKIKYKNRDLND
LHHAH DAYIACILGQFIHQNGNMDVNMIYGQYKKNYKKDVQEHNNYGFILNSMNHIHFNDD
NSVIWDPSYIGKIKSCFCYKD VYVTKLEQNDAKLFDLTILPSDKNSENGVT KAKIPVNKYR
KDVKKYGGFSGDAPIMLAIEADKGKKHVRQVIAFPLRLKNYNE DEERIKFIEKEKNLKNVKIL

TEVKKNQLILINHQYFFITGTNELVNATQLLSAKNTKNLFNLVDANKHNKLESIDDANFNE

25 VIQELICKLQEPIYSRYNSIGKEFEDSYEKINA VTQDKLYIIEYLIAIMS AKATQGYIKPE
LAREIGTNGKNKGRIKSFTIDLNTTFIST SVTGLFSKKYKL GKRPAATKKAGQAKKKGSY
PYDVPDYA PYDVPDYA PYDVPDYA (SEQ ID NO: 6);

(c)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH

30 DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA
GSLMDVLHHPGMNRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG

TSESATPESSGPKKRKVGSIINFQRRLMETQASNQLISSHLKGYP IKDYFVGLAIGTSSV
GWAVTNKAYELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRLERKLRKLLEELFADAM

AQVDPTFFMRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKGTD
DIRKLFLAVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQIL
MEAGKTSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANVLGLNASLIDL
FGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPLTIS
5 ESKVKAFNKHKDDLAILKSLLKSDRSIYNTMFVDEKGLHNYVHYIKQGRTEETSCNREDFY
KYTKKIVEGLSDSKDKEYIILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPF
LNEVADGFSVAEKLIKMLEFRIPYYVGPLNTHHNVNDNGGFAWAVRKASGRVT PWNFDDKIDR
EKSAAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAV
FKQDHKKMTKNRIEQFLKDNGYISETHKHEITGLGEIKNDLASYRDMVRILGDGFDRSMAE
10 EIITDITIFGESKKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAG
DGT PETIIIILMRNFSYNLMELLGDKFSFMERIQEINAALKTEGQIVNPHDIIDDLALSPAVKR
AVWQALRIVDEVAHIKKALPARIFVEVTRSNKNEKKKDSRQKRLSDLYAIAKKDDVLLNGL
NNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGRCAVTGEIIIELSLLNTDNYDIDHIYPRSL
TKDDSF DNVLVLCRTANAQKSDAYPISEEIQKTQKFWTFLKQQGLISERKYERLTRITPLT
15 ADDLSGFIARQLVETNQSVKAATTLLRRLYPGDVVFVKAENVTDFRHDNNFIKVRSLNHHH
HAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKDGKA DVKTS
MDTVKKMMDSNDVRVTKRLLEQTGALADATIYKATVAGKADGAYIGMKTSSFADVSKYG
GMTKIKNAYSIIIVQYTGKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIIPQAKDISIIYG
KLCINQLVKVNGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPYLVLEKFNNMRKDNLKAN
20 VVSTRALDNKHTIEVRIEEKNIEFFDYLVSKLKMPIYQKMKGNAEELSEKGYGLFKKMSL
EEQSIHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTI
VGKRPAATKKAGQAKKKGSYPDVDPDYAYPYDVDPYAYPYDVDPYAY (SEQ ID NO: 10);

(d)

MPKKKRKVSIINFQRRGLMETQASNQLISSHLKGYPIKDYFVGLAIGTSSVGWAVTNKAYEL
25 LKFRSHKMWGSRLFDEGEESAVERRGFRSMRRLERKRLRLKLEELFADAMAQVDPTFFMRL
RESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKGTD DIRKLFLAVHH
ILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGKTSDKA
KAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANVLGLNASLIDLFGSVEELEEDL
KKLQITGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPLTISESKVKAFNKH
30 DDLAILKSLLKSDRSIYNTMFVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLS
DSKDKEYIILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLNEVADGFSVA
EKLIKMLEFRIPYYVGPLNTHHNVNDNGGFAWAVRKASGRVT PWNFDDKIDREKSAAFIKNL
TNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTKN

RIEQFLKDNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGE
SKKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGTETIIILM
RNFSYNLMEELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALS PAVKRAVWQALRIVDE
VAHIKKALPARIFVEVTRSNKNEKKKDSRQKRLSDLYAAIKDDVLLNGLNNEIFGELKSS
5 LAKYDDAALRSKLYLYTQMGRCAYTGEIELSLLNTDNYDIDHIYPRLTKDDSF DN LVL
CKRTANAQKS DAYPISEEIQKTQKFWTFLKQQGLISERKYERLTRITPLTADDLSGFIARQ
LVETNQSVKAATTLLRRLYPGVDVFVKAENVTDFRHNNFIKVRSLNHHHAKDAYLNIVV
GNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCNAKDGKAWDVKTSMDTVKKMMDSN
DVRVTKRLLEQTGALADATIYKATVAGKAKDGAYIGMKTKS SVFADVS KYGGMTKIKNAYS
10 IVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISIIFYGKLCINQLVKVN
GFYYYLGGKTNSKFCIDNAIQVIVSNEWI PYLK VLEKFNNMRKDNKDLKANVVSTRALDNKH
TIEVRIEKNIEFFDYLVS KLKMPIYQKMKGNKAAELSEKGYGLFKKMSLEEQS IHLIELL
NLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIVKRPAATKKAG
QAKKKKSGSETPGTSESATPESSGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNR
15 VIGEGWNRAIGLHDPTAHAEIMALRQGGLVMQNYRLYDATLYVT FEP CVMCAGAMIHSRIGR
VVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADEC AALLCRFFRMPRRVFNAQKKAQ
SSTDPAAKRVKLDGSYPYDVPDYAPYDVPDYAPYDVPDYA (**SEQ ID NO: 11**);

(e)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH
20 DPTAHAEIMALRQGGLVMQNYRLYDATLYVT FEP CVMCAGAMIHSRIGR VFGVRNAKTGAA
GSLMDVLHHPGMNHRVEITEGILADEC AALLCRFFRMPRRVFNAQKKAQ SSTDGSSGSETPG
TSESATPESSGPKKRKVGT KVKDYYIGLAIGTSSVGWAVTDEAYNVLFNSKKMWGVRLFD
DAKTAEERRGQRGARRRLDRKKERLSLLQDFFAEEVAKVDPNFFLRLDNSDLYMEDKDQKLK
SKYTLFNDKDFKDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYLA CHYLLKNRGHFIFEGQK
25 FDTKSSFENSNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLK
AVSAIMIGSSQKLVDLFENPEDFDDSAIKSVDFSTTSFDDKYS DYELALGDKIALVNILKEI
YDSSILENLLKEADSKDGNKYISNAFKVVKYNGQDLKEFKRLVRQYHKSAYFDIFRSEKV
NDNYVSYTKSSISNNKRVKANKFTDQEAFYKFAKKLETIKYKINKVNGSKADLELIDGMLR
DMEFKNFMPKIKSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFR
30 IPYYVGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLDSSREEFIDS LIGRCTYLKDEKVLPK
ASLLYNEYMVLNELNNLKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKEFNINGEI
LLSGTDGDFKQGLNSYNDKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGY
KYFTDSEIKK MAGL NYKDWGRSLSKLLTGLEGANKITGERGSIIHFMREYNLNLMELMSASF

TFTEEIQKLNPVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSSKKIFIEMAR
GKEEVKARKESRKKNQLLKFYKDGKKAFISEIGEERYSYLLSEIEGEENKFRWDNLLYYTQ
LGRCMYSLEPIDISELSSKNIYDQDHIPKSKIYDDSIENRVLVKKDLNSKGNSYPIPDEI
LNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKTI
5 CKNSEIVYNSKAENASRFRQEFDIVKCRAVNDLHHMDAYINIIVGNVYNTKFTKDPMNFVKK
QEKAWSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESG
ALFNATLQRKNKGSRPLDKGPKSIEKYGGYNINKACFAVLDIKSKNKIERKLMPVEREI
YAKQKNDKKLSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITGGSDKTLELRS
ALQLILPKKNEWAIKQIDKSENDYLTIERIQDLTEELVYNTFDITVNKFKTSVFKKSFLNL
10 FQDDKIEINIDFKFKSMDFKEKCTLMLVKAIRASGVRQDLKSIDLKSDYGRLLSKTNNIGN
YQEFKIINQSITGLFENEVDLLKLGKRPAAKKAGQAKKKGSYPYDVPDYAYPYDVPDYAY
PYDVPDYA (SEQ ID NO: 16);

(f)

MPKKKRKVTKVKDYYIGLAIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAERRGQ
15 RGARRRLDRKKERLSLLQDFFAEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKDF
KDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIGEQKFDTKSSFENSL
NELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQ
KLVDLFENPEDFDDSAIKSVDFSTTSFDDKSYDYEALGDKIALVNILKEIYDSSILENLLK
EADKS KDGNEYISNAFKVKKYNGQDLKEFKRLVRQYHKSAYFDIFRSEKVNDNYVSYTQSS
20 ISNNKRVKANKFTDQEAFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKI
KSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRIPIYYVGPLNPN
SKYAWIKKQKDSEITPWNFKDVVDLDSSREEFIDSЛИGRCYTLKDEKVLPKASLLYNEYMVL
NELNNLKLNDLPITEEMKKIFDQLFKTRKKVTLKAVANLLKEFNINGEILLSGTGDFKQ
GLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGKYFTDSEIKKM
25 AGLNYKDWGRLSKKLLTGLEGANKITGERSIIFMREYNLNLMELMSASFTFTEEIQKLNP
VDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSSKKIFIEMARGKEEVKARKES
RKNQLLKFYKDGKKAFISEIGEERYSYLLSEIEGEENKFRWDNLLYYTQLGRCMYSLEPI
DISELSSKNIYDQDHIPKSKIYDDSIENRVLVKKDLNSKGNSYPIPDEIINKNCYAYWKI
LYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKICKNSEIVYNSKA
30 ENASRFRQEFDIVKCRAVNDLHHMDAYINIIVGNVYNTKFTKDPMNFVKKQEKARSYNLEN
MFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRKN
KGSRPLDKGPKSIEKYGGYNINKACFAVLDIKSKNKIERKLMPVEREIYAKQKNDKKLS
DEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITGGSDKTLELRSALQLILPKKNE

WAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLFQDDKIEINIDF
KFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGRILLSKTNNIGNYQEJKIINQSI
TGLFENEVDLLKLKRPAATKKAGQAKKKSGSETPGTSESATPESSGSEVEFSHEYWMRHAL
TLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDPTAHAEIMALRQGGLVMQNYRLYDA
5 TLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADE
CAALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPYDVPDYAYPYDVPDYAYPYDVP
DYA (SEQ ID NO: 17);

(g)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH
10 DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA
GSIMDVLHHPGMNHRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG
TSESATPESSGPKKRKVGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGK
TAAERRTFRTTRRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEDPAKNQAFIG
KLLFPDLLKKNERGYPTLIKMRDEL PVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAVH
15 HIVKYRGHFLNNASVDKFVGRIDFDKS FNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLD
TKMRKLDQKAVAKLLEVKVADKEETKRNKQIATAMS KLVLYKADFATVAMANGNEWKIDL
SSETSEDEIEKFREELSDAQNDILTEITSLSQIMLNEIVPNGMSISESMMDRYWTHERQLA
EVKEYLATQPASARKEFDQVYNKYIGQAPKEKGFDLEKGKLLKILSKKENWEIDEELLKAGDF
LPKQRTSANGVI PHQMHQQUELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVS FRI
20 PYYVGPLVTPEVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESA EAFIKRMTVKDTYLL
NEDVLPANSILYQKYNVLNE LNNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMAK
TRGVNKPSVEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENII EWSVFEDGEIFA
DKLTEVEWLTDEQRSA LVKKRYKGWGRLSKKLLTGIVDENGQRIIDL MWNTDQNFMQIVNQP
VFKEQIDQLNQKAITNDGMTLRERVESVLDAYSPQNKKAIWQVVRVVEDIVKAVGNAPKS
25 ISIEFARNEGKGEITRS RRTQLQKL FEDQAHELVKDTSLTEELEKAPDLSDRYYFYFTQGG
KDMYT GDPINFDEISTKYDIDHILPQS FVKDDSLDNRLV SRAENN KSDRVP A KLYAAKMK
PYWNQLLKQGLITQRKFENLTMDVDQTIKYRSLGFVKRQLVETRQVI LTANILGS MYQEAG
TDIIETRAGLTQLREEFDLPKVREVNDYH HAVDAYLTTFAGQYLNRRYPLRSFFVYGEYM
KFKHGSDLKLRNFNFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLISNE
30 TYEETGKYFDASHESSSLYLKNQNKKSKLVVPLKNKLQPEYYGAYTGITQGYM VILKLLDKK
GGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKS F EVVVPRVLKGTFLVDGEEK
FILSSYRYKV NATQLILPVSDIKLIQDNFKALKLN VEMQTKKLIEIYDNILRQVDKYYKLY
DINKFRAKLHDGRSKFVELDDFGQDASKEKVI IKILRGLHFGSDLQNLKEIGFGTTPLGQFQ

VSEAGIRLSNTAIFIIFKSPTGLFNRKLYLKNLGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 88);

(h)

MPKKKRKVGVKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAERRTFRT

5 TRRRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEDPAKNQAFIGKLLFPDLLKK

NERGYPTLIKMRDELPVEQRAHYPTVNIYKLREAMINEDRQFDLREVYLAVHHIVKYRGHFL

NNASVDFKFVKVGRIDFDKSFNVLNEAYEELONGEGSFTIEPSKVEKIGQQLLDTKMRKLDRQK

AVAKLLEVKVADKEETKRNKQIATAMSKVLGYKADFATVAMANGNEWKIDLSSETSEDEIE

KFREELSDAQNDILTEITSLSQIMLNEIVPNGMSISESMMDRYWTHERQLAEVKEYLATQP

10 ASARKEFDQVYNKYIGQAPKEKGFDLEKGILKKILSKKENWKEIDELLKAGDFLPKQRTSANG

VIPHQMHQELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPYYGPLVTP

EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEEFIKRMTVKDTYLLNEDVLPANSI

LYQKYNVLNELNNVRVNRRGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMAKTRGVNPKSVE

GLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIEWRSVFEDGEIFADKLTEVEWLT

15 DEQRSLVKKRYKGWGRSLKKLLTGIVDENGQRIIDLWNTDQNFMQIVNQPVFKEQIDQLN

QKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSISIEFARNEG

NKGEITRSRRTQLQKLFEQAHELVKDTSLTEELEKAPDLSDRYFYFTQGGKDMYTGDPIN

FDEISTKYDIDHILPQSFVKDDSLDNRVLVSRRAENNKSDRVPAKLYAAKMKYWNQLLKQG

LITQRKFENLTMDVDQTICKYRSILGFVKRQLVETRQVIKTANILGSMYQEAGTDIETRAGL

20 TKQLREEFDLPKVREVNDYHHAVDAYLTTFAGQYLNRRYPKLRFFVYGEYMKFKHGSDLKL

RNFNFFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINLKVMLISNETYEETGKYFD

ASHESSSLYLNQNKKSKLVPPLKNKLQPEYYGAYTGITQGYMVLKLLDKGGFGVYRIPR

YAADIINKCHDEVAYRNKIAEIISSDPRAPKSFEVVVPRVLKGTFLVDGEEKFILSSYRYKV

NATQLILPVSDIKLIQDNFKALKLNVEMQTKKLIEIYDNILRQVDKYYKLYDINKFRAKLH

25 DGRSKFVELDDFGQDASKEKVIIKILRGLHFGSDLQNLKEIGFGTTPLGQFQVSEAGIRLSN

TAFIIFKSPTGLFNRKLYLKNLKRPAATKKAGQAKKKSGSETPGTSESATPESSGSEVEFS

HEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIDEGWNRAIGLHDPTAHAEIMALRQGGLV

MQNYRLYDATLYVTFEPVCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVE

ITEGILADECALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPYDVPDYAYPYDVP

30 DYAYPYDVPDYA (SEQ ID NO: 89);

(i)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH
DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA
GSIMDVLHHPGMNRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG
TSESATPESSGPKKRKVGEKNTYTIGLAIGTDSVGAVVKDDLELVKKRMKVLGNTETNY
5 IKKNLWGSLLFESGQTAKDRLKRVARRRYERRNRNLTTELQKIFAPAIDEVDENFFFRLNES
FLVPEDKAFAKNPIFGTILGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRG
HFLIEGKLDTEHIAINENLEQFFESYNALFSEEPIELKEELIAIENILREKNSRTVKEKRI
TSFLKDGRANKQSPMMAFITLIVGKKAKFKAAFNLEEEISLNLTDDSYDENLEILLNTIGS
DFADLFDAQRVYNAVLAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQ
10 YDMTFVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQ
RSFYNGVIPNQVHLAEMEAILDRQENYPWLKENREKIISLLTFRIPYYVGPLADGQSEFAW
LERKSDEKIKPWNFSDVVDLDRSAEKFIEQLIGRDTYLPDEYVLPKSLIYQKYMVFNELTK
IAYLDERQKRMNLSSVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNAD
YSTYVELAKVPGMKSMMDDPDNEDLMEEIVKILTVFEDRKMRKQLEKYKERLSPEQIKEA
15 KKHYTGWRSLSKLLVGIRDKETQKTILDYLVEDDNHSGGRQHNRNLMQLINDDRLSFKKT
IAELQMIDPSADLYAQVQEIAGPSAIKKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTT
ARGKALSKRREAKIKEGLAALGSSLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDR
LSQYDEDHII PQSFTVDNSLDNLVLTNSQN RGNKKDDVPSLEV VNRQLAYWRS LKDAGLMT
QRKFDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKDEAGNKIRETN
20 VLLKSAMASEFRKMFRKYVRELNDYHHADAYLNAIAIAINLLALYPYMADDFVYGEFRYKK
KPQAEKATYEKLQWNLIKRFGEKQLFTP DHEDCWNKERDIKTIKKVMGYRQVN VVKAEER
TGMLFKETINGKTNKGSRIPIKKLDPSKYGGYIEEKMAYYAVISYEDKKKPGKTIVGISI
MDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIAYPDGRRYITGATKTTKGKVELQKANQ
IAMEQDLVNFYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRR
25 IYEEEKKDPVDIEALVASFIELLKLT SAGAPADFI FMGEAISRRRYNSMTGLFDGQVIYQS
LTGLYETRMRFEDGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA
(SEQ ID NO: 91);

(j)

MPKKKRKVEKNTYTIGLAIGTDSVGAVVKDDLELVKKRMKVLGNTETNYIKKNLWGSLLF
30 ESGQTAKDRLKRVARRRYERRNRNLTTELQKIFAPAIDEVDENFFFRLNESFLVPEDKAFA
NPIFGTILGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTE
HIAINENLEQFFESYNALFSEEPIELKEELIAIENILREKNSRTVKEKRITSFLKDGRAN
KQSPMMAFITLIVGKKAKFKAAFNLEEEISLNLTDDSYDENLEILLNTIGSDFADLFDAQR

VYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDA
QKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQ
VHLAEMEAILDRQENYYPWLKRNREKIISLLTFRIPYYVGPLADGQSEFAWLERKSDEKIKP
WNFSDVVLDLRSACKFIEQLIGRDTYLPDEYVLPKSLIYQKYMVFNELTKIAYLDERQKRM
5 NLSSVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVP
GMKSMMDDPDNEDLMEEIVKILTVFEDRKMRKQLEKYKERLSPEQIKELAKKHYTGWGRSL
KKLLVGIRDKETQKTILDYLVEDDNHSGGRQHLNRLNQLINDDRLSFKKTIAELQMIDPSA
DLYAQVQEIAAGSPAICKGILLGLKIVDEIIIRVMGEKOPENIVIEMARENQTTARGKALSKRRE
AKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHIIP
10 QSFTVDNSLDNLVLTNSSQNRGNKDDVPSLEVNRQLAYWRSLKDAGLMTQRKFDNLTKAM
RGGLTDKDRERFIQRQLVETRQITKNVAKLDMRLNDKKDEAGNKIRETNIVLLKSAMASEF
RKMFRLYKVRELNDYHHAHDAYLNAIAINLLALYPYMADDFVYGEFRYKKPQAEKATYEK
LRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTICKVMGYRQVNVVKAERTGMLFKETING
15 KTNKGSRIPIKKDLDPSKYGGYIEEKMAYYAVISYEDKKKPGKTIVGISIMDKKEFEYDSI
SYLGKLGFSNPVVQIILKNYSLIYPDGRRRYITGATTTKGKVELQKANQIAMEQDLVNFI
YHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFIDAETNINRLRRIYEEEKKDPV
DIEALVASFIELLKLTSAAGAPADFIFMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMRF
EDKRPAATKKAGQAKKKSGSETPGTSESATPESSGSEVEFSHEYWMRHALTAKRARDERE
20 VPVGAVLVLNNRVIGEGWNRAIGLHDPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVM
CAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADECACALLCRFFRM
PRRVFNAQKKAQSSTDPAAKRVKLDGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (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)
MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR
SGKNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLV
VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPLWMML
YALELHCIILSLPPCLKISRWRQNHIAFFRLHLQNCHYQTIPPHILLATGLIHP
30 GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKRKVGGKPYSIGLAIGTNSVGWAVVTDD
YKVPAAKKMKVLGNTDKQSIKKNLLGALLFDGETAEATRLKRTARRRYTRRKNRLRYLQEIF
TGEMNKVDENFFQLDDSLVDEDKRGEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKK
ADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFKDFVEEYDKTIEESHLEITVDALS

ILTEKVKSSRLENLIAHYPTEKKNTLFGNLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYE
EDLEGFLGEVGDEYADLFASAKNLYDAILLSGILTVDNSTKAPLSASMVKRYEEHQKDLKK
LKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDRE
DFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHYPFLKENQDKIEKILTFRIPYYVGPLARK
5 GSRAFAWEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTV
YNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTKDKLLNYLNKEFEERIVNLTGL
DKENKAFNSSLGTYHDLRKILDKSFLDDKANEKTIEDIIQTLTFEDREMIRQLQKYSdif
TKAQLKKLERRHYTGWGRLSYKLINGIRNKENKKTILDYLIDDGYANRNFMQLINDDALSKF
EEIARAQIIDDVDDIANVHDLPGSPAICKGILQSVKIVDELVKVMGNPANIIEMARENQ
10 TTDKGRRNSQQRLKLLQDSLKNLDNPVNICKNVENQQLQNDRLFLYYIQNGKDMYTGETLDIN
NLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSDDVPsieVVHEMKSFWSKLLSVKLIT
QRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKI
ITLKSNLVSNFRKEFELYKREINDYHHADAYLNAVGNALLKYPQLEPEFVYGEYPKYN
SYRSRKSATEKFLLFYSNILRFFKKEDIQTNEGEIAWNKEKHKILRKVLSYPQVNIVKKTE
15 EQTGGFSKESILPKGESDKLIPRKTNSYWDPKKYGGFDSPVVAYSILVFADVEKGKSKKLR
KVQDMVGITIMEKKRFEKNPVDFLEQRGYRNRLEKIIKLPKYSIFELENKRRLLASAKEL
QKGNELVIPQRFTTLLYHSYRIEKDYEPHREYVEHKDEFKELLEYISVFSRKYVLADNNL
TKIEMLFSKNDAEVSSLAKSFISLLTFTAFGAPAAFNFFGENIDRKRYTSVTECLNATLIH
QSITGLYETRIDLSKLGEDGKRPAATKKAGQAKKKKGSSGGSGGSTNLSDIIIEKETGKQ
20 LVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNG
ENKIKMLSGGSGGSGGSTNLSDIIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTA
YDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAYPYDVPDYAYPYDVP
DYA (SEQ ID NO: 21);

(b)

25 MPAAKRVKLDTSEKG PSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR
SGKNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWPCWECSQAIREFLSQHPGVTLVIY
VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMML
YALELHCIILSLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHPSTWRLKS
GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKRKVGSIINFQRRGLMETQASNQLISSH
30 LKGYPIKDYFVGLAIGTSSVGWAVTNKAYELLKFRSHKMWGSRLFDEGESEAVARRGFRSMRR
RLERRKLRKLLEELFADAMAQVDPTFFMRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDY
FKEYPTVYHLRSELMKSGTDDIRKLFLAVHILKYRGNFLYEGATFDSNASTLDDVIKQALE
NITFNCFDCNSAIISSIGQILMEAGTKSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDK

KRLKAFANVLGLNASLIDLFGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDC
KSVYDAIILLSIKEPLTISESKVKAFNKHKKDLAILKSLLKSDRSIYNTMFVDEKGLHNY
VHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEYILSQIELQILLPLQRIKDNGVIPY
QLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKMLEFRIPIYYVGPLNTHHNVDNGGFAW
5 AVRKASGRVT PWNFDDKIDREKSAAFIKNLTNKCTYLLGEDVLPKSSLSEFMNLNN
VRIDGKPLEKVVKHEHLIEAVFKQDHKKMTKNRIEQFLKDNGYISETHKHEITGLDGEIKNDL
ASYRDMVRILGDGFDRSMAEEIITDITIFGESKKMLRETLRKKFASCLDDEAIKKLTKLRYR
DWGRLSQKLLNGIEGCDKAGDGTETIIILMRNFSYNLMELLGDKFSFMERIQEINAKLTEG
QIVNPHDITDDLALSPAVKRAVWQALRIVDEVAHIKKALPARIFVEVTRSNKNEKKKDSRQ
10 KRLSDLYAAIKKDDVLLNGLNNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGRCAUTGEII
ELSLLNTDNYDIDHIYPRSLTKDDSFDNLVLCRTANAQKSDAYPISEEIQKTQKPFWTFLK
QQGLISERKYERLTRITPLTADDLSGFIARQLVETNQSVKAATLLRRLYPGVVVVFVKAEN
VTDFRHDDNNFIKVRSLNHHHHAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAKMF
NYDVNCNTNAKGKADEVKTSMDTVKKMMDSDNRVTKRLLEQTGALADATIYKATVAGKAKD
15 GAYIGMKTSSVFADVKYGGMTKIKNAYSIIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQD
LINYVASIIPQAKDISIIYGKLCINQLVKVNGFYYLGKTNFKCIDNAIQVIVSNEWIPY
LKVKLEKFNNMRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLVSKLKMPIYQKMK
GNKAAEILSEKGYGLFKKMSLEEQSILHIELLNLTNQKTTFEVKPLGITASRSTVGSKISNQ
DEFKVINESITGLYSNEVTIVGKRPAAKKAGQAKKKGSSGGSGGGSTNLSDIIIEKETG
20 KQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDS
NGENKIKMLSGGGSGGGSTNLSDIIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVH
TAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAYPYDVPDYAY
(SEQ ID NO: 12);

(c)

25 MPAAKRVKLDSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR
SGKNTTNHVEVNFIKKFTSERRFHSSISCSITWFWSWPCWECSQAIREFLSQHPGVTLV
VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMML
YALELHCIILSLPPCLKISRWRQNHLAFFRLHLQNCHYQTIPPHILLATGLIHP
GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKRKVGTKVKDYYIGLAIGTSSVG
30 EAYNVLFNSKKMWGVRLFDDAKTAEERRGQRGARRRLDRKKERLSLLQDFFAEVAKVDP
FFLRLDNSDLYMEDKDQKLKSKYTLFNDKDFKDNFKHKKYPTIHLLMDLIEDDSKKDIRLV
YLACHYLLKNRGHFIFEGQKFDTKSSFENSNELKVHLNDEYGLDLEFDNENLINIL
NKTAKKKEKLSVIGDTKFLKAVSAIMIGSSQKLVDLFENPEDFDDSAIKSVDFSTTSFDDKY

SDYELALGDKIALVNILKEIYDSSILENLLKEADSKDGNKYISNAFKVKKYNKHGQDLKEFK
RLVRQYHKSAYFDIFRSEKVNDNYVSYTKSSISNNKRVKANKFTDQEAFYKFAKKHLETIKY
KINKVNNGSKADLELIDGMLRDMEFKNFMPKIJKSSDNGVI PYQLKLMELNKILENQSKHHEFL
NVSDEYGSVCDKIASIMEFRIPYVGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLDSSREE
5 FIDSLIGRCTYLKDEKVLPKASLLYNEYMLNELLNLNDLPITEEMKKIFDQLFKTRKK
VTLKAVANLLKKEFNINGEILLSGTGDFKQGLNSYNDFKAIVGDKVDSDDYRDKIEEIKL
IVLYGDDKSYLQKKIKAGYGKYFTDSEIKMAGLNYKDWGRLSKLLTGLEGANKITGERGS
IIHFMRREYNLNLMELMSASFTFTEEIQKLNPVDDRKLSYEMVDELYLSPSVKRMLWQSLRIV
DEIKNIMGTDSKKIFIEMARGKEEVKARKESRKRNQLLFYKDGKKAFISEIGEERYSYLLSE
10 IEGEENEKFRWDNLYLYYTQLGRCMYSLEPIDISELSSKNIYDQDHIPKSKIYDDSIEENRV
LVKKDLNSKKGNNSYPIPDEILNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDELVQFIS
RQIVETRQATKETANLLKTICKNSEIVYSAENASRFRQEFDIVKCRAVNDLHHMHDAYINI
IVGNVYNTKFTKDPMN FVKKQEKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASI KRI
RKELEGTNYRFTRMNYIESGALFNATLQRKNKGSRPLDKGPKSSIEKYGGYNINKACFAV
15 LDIKSKNKIERKLMPVERE IYAKQKNDKKSDEIFSKYLDKDRFGIEDYRVVYPVVKMRTLLK
IDGSYYFITGGSDKTLELRSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNT
FDIIVNKFKTSVFKKSFLNLQDDKNIENIDFKFKSMDFKEKCKTLLMLVKAIRASGVRQDLK
SIDLKSDYGRSSKTNIGNYQEFKIINQSITGLFENEVDLLKLGKRPAAKKAGQAKKKKG
SSGGSGGSGGSTNLSDIIIEKETGKQLVIQESIILMLPEEEVIGNKPESDILVHTAYDESTD
20 ENVMLLTSDAPEYKPWALVIQDSNGENKIKMLGGSGGSGGSTNLSDIIIEKETGKQLVIQES
IILMLPEEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKM
LYPYDVPDYAYPYDVPDYAY (SEQ ID NO: 18);

(d)

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR
25 SGKNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLV
VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGPDEAHWPQYPPLWMML
YALELHCIILSLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHPSTWRLKS
GGSSGGSSGSETPGTSESATPESSGGSSGSPKKRKVGKEYHIGLAIGTSSIGWAVTDSQF
KLMRIKGKTAIGVRLFEEGKTAERRTFRTTRRLKRRKWRLHYLDEIFAPHLQEVDENFLR
30 RLKQSNIHPEDPAKNQAFIGKLLFPDLLKKNERGYPTLIKMRDELPVEQRAHYPTNIYKLR
EAMINEDRQFDLREVYLAHHIVKYRGHFLNNASVDKFKVGRIDFDKSFNVLNEAYEELQNG
EGSFTIEPSKVEKIGQLLDTKMRKLDRQKAVAKLLEVKVADKEETKRNKQIATAMSKLVLG
YKADFATVAMANGNEWKIDSSETSEDEIEKFREELSDAQNDILTEITSLFSQIMLNEIVPN

GMSISESMMDRYWTHERQLAEVKEYLATQPASARKEFDQVYNKYIGQAPKEKGFDFLEKGLKK
ILSKKENWEIDEELLKAGDFLPKQRTSANGVIPHQMHQQELDRIIEKQAKYYPWLATENPAT
GERDRHQAKYELDQLVSFRIPIYYVGPLVTPEVQKATSGAKFAWAKRKEDGEITPWNLWDKID
RAESAEEAFIKRMTVKDTYLLNEDVLPANSLLYQKYNVLNELNNVRVNGRRLSVGIKQDIYTE
5 LFKKKKTVKAGDVASLVMAKTRGVNKPSVEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQ
MDLENIEWRSVFEDGEIFADKLTEVEWLTDQRSALVKKRYKGWGRSLKLLTGIVDENGQ
RIIDLWMNTDQNFMQIVNQPVFKEQIDQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAI
WQVVRVVEDIVKAVGNAPKSISIEFARNEGNKGEITRSRRTQLQKLFEDQAHELVKDTSLTE
ELEKAPDLSDRYYFYFTQGGKDMYTGDPINFDEISTKYDIDHILPQSFKVDDSLDNRVLVSR
10 AENNKKSDRVPACKYAAKMKPWNQLLQGLITQRKFENLTMDVDQTIKYRSLGFVVKRQLVE
TRQVIKLTANILGSMYQEAGTDIETRAGLTQLREFDLPKVREVNDYHHAVDAYLTTFAG
QYLNRRYPKLRFFVYGEYMKFKHGSDLKLRNFNFFHELMEGDKSQGKVVDDQQTGELITTRD
EVADYFDWVINLKVMLISNETYEETGKYFDASHESSSLYLNQNKKSCLVVPLKNKLQPEYY
GAYTGITQGYMVLKLLDKGGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKS
15 FEVVVPRVLKGTFLVDGEEKFILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLNVEMQTK
KLIEIYDNILRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEVIIKILRGLHFG
SDLQNLKEIGFGTTPLGQFQVSEAGIRLSNTAIFIIFKSPTGLFNRKLYLKNLGKRPAAKKA
GQAKKKKGSSGGSGGSTNLSDIIIEKETGQLVIQESILMLPEEEVIGNKPESDILVH
TAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSGGSTNLSDIIIEKETG
20 KQLVIQESILMLPEEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDS
NGENKIKMLYPYDVPDYAYPYDVPDYAY (SEQ ID NO: 90);

(e)

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR
SGKNTTNHVEVNFIKKFTSERRHSSISCSITWFLSWPCWECSQAIREFLSQHPGVTLVIY
25 VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMML
YALELHCIILSLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHPSTWRLKS
GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKRKVGEEKTNYTIGLAIGTDSVGWAVVK
DDLELVKKRMKVLGNTETNYIKKNLWGSLLFESGQTAKDRRLKRVARRRYERRRNRLTELQK
IFAPAIDEVDENFFFRLNESFLVPEDKAFSKNPIFGTLGEDKTYKTYPTIYHLRQHLADSE
30 EKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIAINENLEQFFESYNALFSEEPIELRKEEL
IAIENILREKNSRTVKEKRITSFLKDGRANKQSMMAFITLIVGKKAKFKAASNLEEEISL
NLTDSDSYDENLEILLNTIGSDFADLFDAQRVYNAVLAGILSGDVKNTHAKLSAQMVAMYE
RHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFKVDQLKE

VPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVHLAEMEAILDRQENYYPWLKENREKIISLL
TFRIPYYVGPLADGQSEFAWLERKSDEKIKPWNFSVVLDRSAEKFIEQLIGRDTYLPDEY
VLPPKSLIYQKYMVFNELTKIAYLDERQKRMNLSSVEKKEIFETLFKKRSKVTEKQLVKFFE
NYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMMDDPDNEDLMEEIVKILTVFEDRKMR
5 RKQLEKYKERLSPEQIKEALKHYTGWGRLSKKLLVGIRDKETQKTILDYLVEDDHSGGRQ
HLNRNLMQLINDDRLSFKKTIAELQMIDPSADLYAQVQEAGSPAIIKGILLGLKIVDEIIR
VMGEKPENIVIEMARENQTTARGKALSKRREAKIKEGLAALGSSLKENLPGNADLSQRKIY
LYYTQNGKDIYLDEPLDFDRLSQYDEDHII PQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSL
EVVNRQLAYWRSLKDAGLMTQRKFDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKLL
10 DMRLNDKKDEAGNKIRETNIVLLKSAMASEFRKMFRILYKVRELNDYHHADAYLNAAIAINL
LALYPYMADDFVYGEFRYKKPQAEKATYEKLQWNLIKRFGEKQLFTP DHEDCWNKERDIK
TIKKVMGYRQVNVVKAERTGMLFKETINGKTNKGSRIPIKKLDPSKYGGYIEEKMAYYA
VISYEDKKKPGKTIVGISIMDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIAYPDGRRR
YITGATKTTKGKVELQKANQIAMEQDLVNFIYHLKNYDEISHPESYAFVQSHTDYFDRLFDS
15 IEHYTRRFLDAETNINRLRRIYEEEKKDPVDIEALVASFIELLKLTSA GAPADFI FMGEAI
SRRRYNSMTGLFDGQVIYQSLTGLYETRMRFEDGKRPAATKKAGQAKKKKGSSGGSGSGGS
TNLSDIIEKETGKQLVIQESI LMLPEEEVIGNKPESDILVHTAYDESTDENVMLLTSADP
EYKPWALVIQDSNGENKIKMLSGGSGGSGSTNLSDIIEKETGKQLVIQESI LMLPEEEV
IGNKPESDILVHTAYDESTDENVMLLTSAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAY
20 PYDVPDYAY (SEQ ID NO: 93); or

(f)

MPAAKRVKLDTNLSDIIIEKETGKQLVIQESI LMLPEEEVIGNKPESDILVHTAYDESTDE
NVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSGGSGSTNLSDIIIEKETGKQLVIQESI
LMLPEEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKML
25 SGGSGGSGGSPKKRKVEKKNTYTIGLAIGTDSVGWAVVKDDLELVKKRMKVLGNTETNYIK
KNLWGSLLFESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFRLNESFL
VPEDKAFSKNPIFGTLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHF
LIEGKLDTEHIAINENLEQFFESYNALFSEEPIELKEELIAIENILRENSRTVKEKRITS
FLKDIGRANKQSPMMAFITLIVGKKAKFKA FNLEEEISLNLTDDSYDENLEILLNTIGSDF
30 ADLFDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYD
MTFVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVQDQLKEVPGSEKFLDSIEKEDFLRKQRS
FYNGVIPNQVHLAEMEAILDRQENYYPWLKENREKIISLLTFRIPYYVGPLADGQSEFAWLE
RKSDEKIKPWNFSVVLDRSAEKFIEQLIGRDTYLPDEYVLPKSLIYQKYMVFNELTKIA

YLDERQKRMNLSSVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYS
TYVELAKVPGMKSMMDPDNEDLMEEVKILTVFEDRKMRKQLEKYKERLSPEQIKELAKK
HYTGWGRLSKLLVGIRDKETQKTILDYLVEDDNHSGGRQHLNRNLMQLINDDRLSFKKTIA
ELQMIDPSADLYAQVQEIAGSPAIIKGILLGLKIVDEIIRVMGEKOPENIVIEMARENQTTAR
5 GKALSKRREAKIKEGLAALGSSLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLS
QYDEDHIIIPQSFTVDNSLDNLVLTNSSQRGNKKDDVPSLEVVRNQLAYWRSLKDAGLMTQR
KFDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKDEAGNKIRETNIVL
LKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAIAINLLALYPYMADDFYGEFRYKKKP
QAEKATYEKLROWNLIKRGFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQNVVKAAERTG
10 MLFKETINGKTNKGSRIPIKKDLDP SKYGGYIEEKMAYYAVISYEDKKKKPGKTIVGISIMD
KKEFEYDSISYLGKLGFSNPVVQIILKNYSLIAYPDGRRYITGATKTTKGKVELQKANQIA
MEQDLVNFIYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRIY
EEEKKKDPVDIEALVASFIELLK LTSAGAPADFI FMGEAISRRYNSMTGLFDGQVIYQSLT
GLYETRMRFEDKRPAATKKAGQAKKKGSSGGSSGGSSGSETPGTSESATPESSGGSSGGST
15 SEKG PSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWRSSGKNTTNHVEV
NFIKKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIYVARLFWHMDQR
NRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPILWMMLYALELHCIILS
LPPCLKISR RWQNHLAFLRLHLQNCHYQTIPPHILLATGLIHP SVTWRYPYDVPDYAYPYDV
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.

30 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

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 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' -

GUUUUAGAGCUGUGCUGUUUAACAAACACAGCAAGUUAAAUAAGGCUUUGUCCGUACUCAA
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' -

GUUUUAGAGUUGUGUUAUUGAAAAAAUACACAACAGAGUUAAAUAAGCUUAUGCUUAAAUG
CCAGCUUUGCUGGUGUCAUUAGAUGACUUUACUAAGGUUGCUUCGGCAACCUUUUU-3'
(SEQ ID NO: 7).

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

5' -

GUUUGAGAGUAGUGUGAAAACAUUACGAGUUCAAAUACAAAUUAAAUAACAAUGCCUUCGGG
CUGCCCGACGUAGGGCACCUACUCUCAAUUCUUCGGAAUUGAGUU-3' (SEQ ID NO: 13).

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

5' –
GUUUGAGAGUUAAUGUAUUGAAAAAUUACAUGACGAGUUCAAAUAAAUAUUCAAACCG
5 CCUAUUUAUAGGCCGCAGAUGUUCUGCAUUAUGCUCAUUGCAAGCUU-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 GUCUUGGAUGAGUGUGAAAACACUCAUAGUCAAGAUCAAACGAGUGGUUUCCACGAGUUAU
UACUUUUGAGGUCUUAUAUGGCCAUACAUAAAAGGAGUCGGAAUUCGGCUCUUUCU
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' –
GUUUUAGAGCCAUGUAGAAAUAUCAUUGCAAGUUAAAUAAGGCUUUGUCCGUAAUCAACUUG
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 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 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.

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.

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 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 5 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 10 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.

15 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.

20 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 25 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 adenine (A) within DNA. In some

embodiments, the base editor is capable of deaminating a cytosine (C) and an adenine (A) within DNA. In some embodiments, the base editor is a cytidine base editor (CBE). In some embodiments, the base editor is an adenine base editor (ABE). In some embodiments, the base editor is an adenine 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 adenine 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 15 cleavage” *Nature* 533, 420-424 (2016); Gaudelli, N.M., *et al.*, “Programmable base editing of 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:eaao4774 (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 adenine 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 adenine 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 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 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, 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, 5 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. 10

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 15 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 20 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 25 (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 30 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.

10 *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 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.

15 *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.

20 *Functional equivalent or analog:* As used herein, the term "functional equivalent" or "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 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.

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

30 *Improve, increase, or reduce:* As used herein, the terms "improve," "increase" or "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, Hoogstein 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 5 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.

10 *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.

15 *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.

20 *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 25 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.

30 *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.

15 *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).

25 *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 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 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, 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 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. A target nucleic acid may be interspersed with non-nucleic acid components. A target nucleic 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 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 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) 5 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; 10 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 15 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 20 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

25 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 30 *Veillonella parvula* Cas9. FIG. 1D is a graph that shows a consensus PAM motif recognized by human codon-optimized *Ezakiella peruvensis*. 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. 5 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 peruvensis* 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. 2D 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. 2D 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 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 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 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 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-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 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 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 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 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-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 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 herein), *Ezakiella peruvensis* (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: 5 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

Wild Type <i>Streptococcus constellatus</i> Cas9
MGKPYSIGLDIGTNSVGWAVTDDYKVPACKMKVLGNTDKQSIIKKNLLGALLFDSGETAEA TRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSLVDEDKRGEHHPIFGNI AAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQA LFKDFVEEYDKTIEESHLEITVDALSILTEKVSKSSRLENLIAHYPTEKNTLFGNLIAL SLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAILLSGI LTVDDNSTKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYIESG VKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQG EHYPFLKENQDKIEKILTFRIPYYGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSA EKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFD HVFKENRKVTDKLNNYLNKEFEEFRIVNLTGLDENKA FNSSLGTYHDLRKILDKSFLDD KANEKTIEDIIQTTLFEDREMIRQRLQKYSIFTKAQLKKLERRHYTGWGRLSYKLINGI RNKENKKTILDYLIDDGYANRNFMLINDDALSFKEEIARAQIIDDVDDIANVVHDLPGSP AIKKGILQSVKIVDELVKVMGNHPANIIEMARENQTTDKGRRNSQQRLKLLQDSLKNLDN PVNIKNVENQQLQNDRLFYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNR VLTRSDKNRGKSDDVPSIEVVHEMKSFW SKLLSVKLITQRKF DNLTKAERGGLTEEDKAGF IKRQLVETRQITKHVAQILDERFNTEDFGNKRRIRNVKIITLKSNLVSFRKEFELYKVRE INDYHHAHDAYLNAVVGNA LLKYPQLEPEFVYGEYPKNSYRSRKSATEKFLFYSNILRF FKKEDIQTNEEDGEIAWNKEKH KIILRKVLSYPQVNIVKKTEEQTGGFSKESILPKGESDKL IPRKTKNSYWDPKKYGGFDSPVVAYSILVFADVEKGKSKKLRKVQDMVGITIMEKKRFEKN PVDFLEQRGYNRVRL EKIIKLPKYSLELENKRRLLASAKELQKGNE LVI PQRFTTLLYH SYRIEKDYEPHREYVEKHDEFKELLEYISVFSRKYVLADNNLT KIEMLFSKNKDAEVSS LA KS FISLLTFTAFGAPAAF NFFGENIDRKRYTSVTECLNATLIHQ SITGLYETRIDLSKL GED (SEQ ID NO: 1).
<i>Streptococcus constellatus</i> Cas9 with Nuclear Localization Signal (NLS) and Linker

MPKKKRKVGGKPYSIGLDIGTNSVGWAVTDDYKVPACKMKVLGNTDKOSIKKNLLGALLF
DSGETAEATRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRGE
HHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLK
AENTDVQALFKDFVEEYDCTIEESHLEITVDALSILTEKVSKSSRLENLIAHYPTEKKNT
LFGNLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLY
DAILLSGILTVDNSTKAPLSASMVKRYEEHQDLKCLKDFIKVNAPDQYNAIFKDKNKKG
YASYIESGVQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEM
HAILRRQGEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRAWAEYKADEKITPWNFDD
IILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKVYVNEQGEAKFFDT
NMKQEIFDHVFKENRKVTDKLNYLNKEFEFRIVNLTGLDENKAFFNSSLGTYHDLRKI
LDKSFLDDKANEKTIEDIIQTTLFEDREMIRQRLQKYSIFTKAQLKKLERRHYTGWRGL
SYKLINGIRNKENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANV
VHDLPGSPAICKGILQSVKIVDELVKVMGHNPNANIIEMARENQTTDKGRRNSQQLKLLQ
DSLKNLDNPVNICKVENQQLQNDRFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFI
KDNSLDNRVLTRS DKNRGKSDDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGGL
TEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKSNLVSFRKE
FELYKVREINDYHHAHDAYLNAVVGNNALLLKYPQLEPEFVYGEYPKNSYRSRKSATEKFL
FYSNILRFFKKEDIQTNEDGEIAWNKEHKIKLRKVLSPQVNIVKKTEEQTGGFSKESIL
PKGESDKLIPRKTNSYWDPKKYGGFDSPVVAYSILVFADVEKGKSKKLRKVQDMVGITIM
EKKRFEKNPVDFLEQRGYRNVRLEKIIKLPKSLFELENKRRRLLASAKELQKGNELVIPQ
RFTTLLYHSYRIEKDYEPHEHREYVEKHDEFKELLEYISVFSRKYVLADNNLTKEMLFSK
NKDAEVSSLAKSFISLLFTAFTGAPAAFFGENIDRKRYTSVTECLNATLIHQSIITGLYE
TRIDLSKLGEDGKRPAATKKAGQAKKKGS YPYDVPDYAYPYDVPDYAYPYDVPDYA
(SEQ ID NO: 2).

Wild Type Sharpea Cas9

MAKNKDIRYSIGLDIGTNSVGWAVMDEHYELLKKGNHHMWGSRLFDAEPAATRARSIR
 RRYNKRRERIRLLRDLLGDMVMEVDPFFIRLLNVSFLDEEDKQKNLGNDYKDNYNLIEK
 DFNDKTYDKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQSFAKVYEDI
 EEKLDNTLKKFMSLNDLDNLFVDNDINSMITVLSKIYQRSKKADDLLKIMNPTKEERAAYK
 EFTKALVGLKFNVSKMILAQEVKKDDKDIELDFSNVDYDSTVDGLQAELEYIEFIEMLHS
 INSWVELQDILGNNSTISAAMVERYEEHKNDLRLKKVIREELPDKYNEVFREDNPKLHNY
 LGYIKYPKNTPVEEFYEWIKRLLAKVDTGEAREILERIDLEKFMLKQNSRTNGSIPYQMOK
 DEMIQIIDNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFEDKQKERILP

WNYDQIVDIDATAEGFIERMQNTGTYFPDKPVMAKNSLTISKFEVLNELNKIRINGKLIPV
ETKKELLSDFMKNKTITDKKLKDVLVTHQYYDTNEELKIEGYQKDLQFSTSLAPWIDFTK
IFGEINASNYQLIEKIIYDISIFEDKKILKRRLKVVYQLDDLLVDKILKLNWTGWSRLSEK
LLTGIKSNSKETILSILENSNMNLMEIINDESLGFKQIEESNKKDIEGPFRYDEVKKLA
GSPAIIKRGIIWQALLVVQEITKFMKHEPSHIYIEFAREEQEKRVTESRIAKLQIYKDLNLQ
TKEDQLVYESLKKEDAKKIDTDALYLYYLQMGKSMYSGKPLDIDKLSTYHIDHILPRSLI
KDDSLDNRVLVLPKENEWKLDSETVPFEIRNKMMGFQKLHENGMSNKKFFSLIRTDFNE
DKKKRFINRQLVETRQIIKNVAVIINDHYTNTNVVTVRAELSHQFRERYKIYKNRDNLH
HAHDAYIACILGQFIHQNGNMDVNMIYGQYKKNYKKDVQEHNNYGFILNSMNHIHFNDNN
SVIWDPYIGKIKSCFCYKDVYVTKLEQNDAKLFDLTILPSDKNSENGVTKAIPVNKYR
KDVNKYGGFSGDAPIMLAIEADKGKKHVRQVIAFPLRLKNYNDEERIKFIEKEKNLKNVKI
LTEVKKNQLILINHQYFFITGTNELVNATQLLSAKNTKNLFNLVDANKHNLESIDDANF
NEVIQELICKLQEPIYSRYNSIGKEFEDSYEKINAVTKQDKLYIEYLIAIMSAKATQGYI
KPELAREIGTNGKNKGRIKSFTIDLNTTFISTSVTGLFSKKYKL (SEQ ID NO: 4).

Sharpea Cas9 with Nuclear Localization Signal (NLS) and Linker

MPKKKRKVGAKNkdirySIGLDIGTNsvgwAvMdehyellkkgnhHMwgsrlfdaaepaat
RRASRSIRRynkrerirllrdllgdmvmevdptffirllnvsfldeedkqknlgndykd
nynlifiekdndktyydkyptiyhlrkelcenkekadprliylalhhivkyrgnflkegqs
fakv yedieekldntlkfmslnldnlfvndinsmitvlskiyorskkaddlkimnpt
keeraaykeftkalvglkfnnvskmilaqevkdddkieldfsndydstvdglqaelgeyi
efi emlhsinswvelqdilgnnstisaamveryeekndlrvlkkvireelpdkynevfre
dnpklhnlylgikypkntpveefyeyikrllakvdtgeareileridlekfmlkqnsrtng
sipyqmqkdemiqiidnqsvypqlkenrekliislefripyyfgplnthsefawikkfed
kokerilpwnydqivdidataegfiermqntgtyfpdkpvmaknsltiskfevlnelnkir
ingklipvetkkellsdfmknktitdkklkdvlvthqyydtneelkiegyqkdlqfstsl
apwidftkifgeinasyqliekiiydisifedkkilkrrlkvvyqlddllvdkilknwt
gwsrlseklltgiksnsketilsilensnmnlmeiindestgfkqieesnkkdiegpf
ydevkklagspaikrgiiwqallvvqeitkfmkhepshiyiefaareeqekrvtessriaklq
iykdlnlqtkedqlvyeslkkedakkidtdalylyylqmgksmysgkpldidklstyhid
hilprslilikdssldnrvlvlpknenewkldsetvpfeirnkmmgfqklhengmsnkkffs
lirtdfnekdkkrfinrqlvetrqiiknnavviindhytnvvtvraelshqfrerykiy
nrndlndlhhadayiacilgqfihqngnmdvnmiygqykknykkdvqehnnygfilnsmn
hihfnddnsviwdpsyigkikscfcykdvyvttklevqndaklfdltilpsdknsengvtka

KIPVNKYRKDVNKYGGFSGDAPIMLAIEADKGKKHVRQVIAFPPLRLKNYNDEERIKFIEKE
KNLKNVKILTEVKKNQLILINHQYFFITGTNELVNATQLKLSAKNTKNLFNLVDANKHNKL
ESIDDANFNEVIQELICKLQEPIYSRYNSIGKEFEDSYEKINAUTQDKLYIIEYLIAIMS
AKATQGYIKPELAREIGTNGKNKGRIKSFTIDLNTTFISTSVTGLFSKKYKLGKRPAATK
KAGQAKKKKGS YPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 5).

Wild Type *Veillonella parvula* Cas9

MSIINFQRRGLMETQASNQLISSHLKGYPPIKDYFVGLDIGTSSVGWAVTNKAYELLKFRSH
KMWGSLRFDEGESAVARRGFRSMRRLERKRLRLKLLLEELFADAMAQVDPTFFMRLRESKY
HYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHHILKY
RGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGTKSDKAKAI
EHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEEDLKK
LQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPGLTISESKVKA
DNKHD DLAILKSLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLS
DSKDKEYILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLNEVADGFSV
AEKLIKMLEFRIPIYYVGPLNTHHNVNDNGGFAWA
RKASGRVT PWN FDDKIDREKSAAFIK
NLTNKCTYLLGEDVLPKSSL
LYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKM
TKNRIEQFLKDNGYISETHKHEITGLDGEIKNDLASYRDMV
RILGDGFDRSMAEEIITDIT
IFGESKKMLRETLRKKFASCLDDEAIKKLT
KLRYRDWGRLSQKLLNGIEGCDKAGDGT PET
IIILMRNFSYNLMELLGDKFSFMERIQEINAKTEGQIVNPHDIIDDLALS
PAVKRAVWQA
LRIVDEVAHIKKALPARIFVEVTRSNKNE
KKKDSRQKRLSDLYAAIKKDDVLLNGLNNEI
FGELKSSLAKYDDAALRSKKLYLYYTQMGR
CAYTGEIIELSLLNTDNYDIDHIYPRSLTKD
DSFDNVLCKRTANAQKSDAYPISEEI
QKTQKPFWTFLKQQGLISERKYERL
TRITPLTAD
DLSGFIARQLVETNQSVKAATTLLRRLY
PGVDVVFVKAENVTDFRH
DNNFIKVRSLNHHH
AKDAYLNIVVGNVYHERFTRNFRAFFKNG
GANRTYNLAKMFNYDV
NCTNAKDGKA
WDV
KTS
MDTVKKM
MDSNDVR
VTKRLLEQT
GALADATIYK
ATVAGK
AKDGAY
IGMKT
KSSV
FADV
SKY
GGMT
KIKNAYS
II
IVQYT
GKK
GEVI
KEIV
PLPI
YLT
NRNTT
DQDL
INY
VASI
IIP
QAK
DISII
YGKLC
INQLV
KVNG
FYY
LGG
KTN
SKFC
IDNA
I
QV
IVS
NEW
I
PYL
KV
LEKF
NNMR
KD
N
K
DL
KAN
VV
STR
ALDN
KHT
IEV
RIV
EEK
NIE
FFD
YLV
SKL
KM
PI
YQ
KM
GN
KAA
ELSE
K
GY
GL
FK
KMS
LEE
QS
I
H
L
I
E
L
N
L
T
N
Q
K
TT
FEV
K
PL
GIT
A
R
S
T
V
G
S
K
I
S
N
Q
D
E
F
K
V
I
N
E
S
I
T
G
L
Y
S
NE
VT
I
V
(SEQ ID NO: 8).

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

MPKKKRKVGSIINFQRGLMETQASNQLISSHLKGYPIKDYFVGLDIGTSSVGWAVTNKAY
 ELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRLERKLRLKLLEELFADAMAQVDPTFF
 MRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFL
 AVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGKT
 KSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANVLGLNASLIDLFGSVE
 ELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPLTISESKV
 KAFNKHKKDDLAILKSLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYT
 KKIVEGLSDSKDKEYILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLN
 EVADGFSVAEKLICKMLEFRIPIYYVGPLNTHHNVDNGGFAAWRKASGRVT PWN FDDKIDRE
 KSAAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAV
 FKQDHKKMTKNRIEQFLKDNGYISETHKHEITGLGEIKNDLAS YRDMVRILGDGFDRSMA
 EEEITDITIFGESKKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDK
 AGDGT PET IIILMRNFSYNLMELLGDKFS FMERIQUEINAKLTEGQIVNPHDIIDDLALSPA
 VKRAVWQALRIVDEVAHIKKALPARIFVEVTRSNKNEKKKDSRQKRLSDLYAAIKDDVL
 LNGLNNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGRCAVTGEIIELSLLNTDNYDIDHI
 YPRSLTKDDSFDNLVLCKRTANAQKSDAYPISEEIQKTQKPFWTFLKQQGLISERKYERLT
 RITPLTADDLSGFIARQLVETNQSVKAATTLLRRLYPGVDVVFVKAENVTDFRHNNFIKV
 RSLNHHHHAKDAYLNIVVGNVYHERFTRNFRAFFKNGANRTYNLAKMFNYDVNCTNAKDG
 KAWDVKTSMDTVKKMMD SNDVRVTKRLLEQTGALADATIYKATVAGKAKDGAYIGMKTKSS
 VFADVSKYGGMTKIKNAYSIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIP
 QAKDISIIYGKLCINQLVKVNGFYYYLGGKTN SKFCIDNAIQVIVSNEWIPYLVLEKFNN
 MRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLVSKLKMPIYQKMKGNKAAELS
 EKGYGLFKKMSLEEQS IHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVIN
 ESITGLYSNEVTIVG**KRPAAKKAGQAKKKGS** YPYDVPDYAYPYDVPDYAYPYDVPDYA
 (SEQ ID NO: 9).

Wild Type *Ezakiella peruensis* Cas9

MTKVKDYYIGLDIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAERRGQRGARRR
 LDRKKERLSLLQDFFAEEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKDFKDKNF
 HKKYPTIHLLMDLIEDDSKKDIRLVLYACHYLLKNRGHFIFEGQKFDTKSSFENSNELK
 VHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQKLV
 DLFENPEDFDDSAIKSVDSTTSFDDKYS DYELALGDKIALVNILKEIYDSSILENLKEA
 DKSKDGNKYISNAFKVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNNDNYVSYTKSSI
 SNNKRVKANKFTDQEAFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKI

KSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSCDKIASIMEFRIPIYYVGPLNP
 NSKYAWIKKQKDSEITPWNFKDVVDLDSSREEFIDSЛИGRCYLKDEKVLPKASILLYNEYM
 VLNELNKLNLNDLPITEEMKKIFDQLFKTRKKVTLKAVANLLKEFNINGEILLSGTGD
 FKQGLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGYFTDSE
 IKKMAGLNYKDWGRSLSKLLTGLEGANKITGERGSIIHFREYNLNLMELMSASFTFTEEI
 QKLNPVDDRKLSEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIEMARGKEEV
 KARKESRKRNQLKFYKDGKKAFISEIGEERYSYLLSEIEGEEENKFRWDNLYLYTQLGRC
 MYSLEPIDISELSSKNIYDQDHYPKSKIYDDSIENRVLVKKDLNSKKGNSYPIPDEILNK
 NCYAYWKILYDKGLIGQKKYTRLRTGFTDDELVQFISRQIVETRQATKETANLLKTICK
 NSEIVVSKAENASRFRQEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKDPMNFVKKQ
 EKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESG
 ALFNATLQRKNKGSRPLKDKGPSSIEKYGGYTNIINKACFAVLDIKSKNKIERKLMPVERE
 IYAKQKNDKKLSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITGGSDKTLEL
 RSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSF
 LNLFQDDKIENIDFKFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGRLLSKTN
 NIGNYQEFKIINQSITGLFENEVDLLKL (SEQ ID NO: 14).

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

MPKKKKRKVGTVKDYYIGLDIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAEERR
 GORGARRRLDRKKERLSSLQDFFAAEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFND
 KDFKDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIFEGQKFDTKSSF
 ENSLNELKVHLNDEYGLDLEFDNENLINILTDPKLNKAKKELKSVIGDTKFLKAVSAIM
 IGSSQKLVDLFENPEDFDDSAIKSVDFSTTSFDDKYSDYELALGDKIALVNILKEIYDSSI
 LENLLKEADKSKDGNKYISNAFKVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNDNY
 VSYTKSSISNNKRVKANKFTDQEAFYKFAKKLETIKYKINKVNGSKADLELIDGMLRDME
 FKNFMPKIKSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSCDKIASIMEFRIP
 YYVGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLDSSREEFIDSЛИGRCYLKDEKVLPKA
 SLLYNEYMVLNELNKLNLNDLPITEEMKKIFDQLFKTRKKVTLKAVANLLKEFNINGEI
 LLSGTGDFKQGLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGY
 GKYFTDSEIKKMAGLNYKDWGRSLSKLLTGLEGANKITGERGSIIHFREYNLNLMELMSA
 SFTFTEEIQKLNPVDDRKLSEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFI
 MARGKEEVKARKESRKRNQLKFYKDGKKAFISEIGEERYSYLLSEIEGEEENKFRWDNLYL
 YYTQLGRCMYSLEPIDISELSSKNIYDQDHYPKSKIYDDSIENRVLVKKDLNSKKGNSYPI
 PDEILNKNCYAYWKILYDKGLIGQKKYTRLRTGFTDDELVQFISRQIVETRQATKETA

NLLKTICKNSEIVYNSKAENASRFRQEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKD
PMNFVKKQEKAWSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGNYRFT
RMNYIESGALFNATLQRKNKGSRPLDKGPKSSIEKYGGYNINKACFAVLDIKSKNKIER
KLMPVEREIYAKQKNDKKLSDEIFSKYLDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITG
GSDKTLELRSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFK
TSVFKKSFLNLQDDKNIENIDFKFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKSDY
GRLSSKTNNIGNYQEFKIINQSITGLFENEVDLLKLGKRPAATKKAGQAKKKGS YPYDVP
DYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 15).

Wild Type Lactobacillus fermentum strain AF15-40LB Cas9

MKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAERRTFRTTRRLKR
RKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEDPAKNQAFIGKLLFPDLLKKNERGYP
TLIKMRDEL PVEQRAHY PVTNIYKLREAMINEDRQFDLREVYLAHHIVKYRGHFLNNASV
DKFKVGRIDFDKS FNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLDTKMRKLDRQKAVAK
LLEVKVADKEETKRNKQIATAMSKVLGYKADFATVAMANGNEWKIDLSSETSEDEIEKFR
EELSDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWTHERQLAEVKEYLATQPAS
ARKEFDQVYNKYIGQAPKEKGFDLEKGLKKILSKKENWEIDEELLKAGDFLPKORTSANGV
IPHQMHQELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPYYVGPLVTP
EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEEFIKRMTVKDTYLLNEDVLPANS
LLYQKYNVLNELNNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVMAKTRGVNKPS
VEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIIIEWRSVFEDGEIFADKLTEVE
WLTDEQRSALVKKRYKGWGRLSKKLLTGIVDENGQRIIDLWNTDQNFMQIVNQPVFKEQI
DQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSISIEF
ARNEGNGEITRSRRTQLQKLFDQAH ELVKDTSLTEELEKAPDLSDRYYFYFTQGGKDMY
TGDPINFDEISTKYDIDHILPQSFVKDDS LDNRVLVSRAENNKKSDRVPAKLYAAKMKPW
NQLLKQGLITQRKFENLTMDVDQT IKYRS LGFVKQLVETRQVIKLTANILGSMYQEAGTD
IIETRAGLTQQLREEFDL PKVREVNDYHHAVDAYLTT FAGQYLNRRYPKLRSFFVYGEYMK
FKHGS DLKLRNFNFFHELMEGDKSQGKVV DQQTGELITTRDEVADY FDWVINLKVMLISNE
TYEETGKYFDASHES SSSL YLKNQNKKSKL VVPLKNKLQPEYYGAYT GITQGYMVILKLLDK
KGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKS FEV VVPRVLKGTFLVDGE
EKFILSSYRYKVNATQ LILPVSDIKLIQDNFKALKKLN VEMQT KKLIEIYDNILRQVDKYY
KLYDINKFRAKLHDGRSKFVELDDFGQDASKEVII KILRGLHFGSDLQNLKEIGFGTTPL
GQFQVSEAGIRLSNTAIFI FKSP TGLFNRKLYLKNL (SEQ ID NO: 84).

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

MPKKRKVGKEYHIGLDIGTSSIGWAVTDQFKLMRIKGKTAIGVRLFEEGKTAERRTFR
TTRRLKRRKWRLHYLDEIFAPHIQLQEVDENFLRRLQSNIHPEDPAKNQAFIGKLLFPDLL
KKNERGYPTLIKMRDELPEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAVHHIVKYRG
HFLNNASVDKFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLDTKMRKL
DRQKAVAKLLEVVKVADKEETKRNKQIATAMSKLVLYGKADFATVAMANGNEWKIDLSSETS
EDEIEKFREELSDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRYWTHERQLAEVKE
YLATQPASARKEFDQVYNKYIGQAPKEGFDEGLKKILSKKENWKEIDELLKAGDFLPK
QRTSANGVIPHQMHQQELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPY
YVGPLVTPEVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAFAIKRMTVKDTYLLN
EDVLPANSLLYQKYNVLNELNNVRVNGRRLSVGIKQDIYTELFKKKKTVKAGDVASLVM
TRGVNKPSVEGLSDPKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIEWRSVFEDGEIF
ADKLTEVEWLTDQRSALVKKRYKGWGRSLKKLLTGIVDENGQRIIDLWMNTDQNFMQIVN
QPVFKEQIDQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNA
PKSISIEFARNEGKGEITRSRRTQLQKLFEDQAHELVKDTSLTEELEKAPDLSDRYYFYF
TQGGKDMYTGDPINFDEISTKYDIDHILPQSFKDDSLDNRVLVSRAENNKKSDRVPAKLY
AAKMKPYWNQLLQGLITQRKFENLTMDVDQTICKYRSLGFVKRQLVETRQVIKLTANILGS
MYQEAGTDIIETRAGLTKQLREEFDLPLKVREVNDYHHAVDAYLTTFAGQYLNRRYPKLR
FVYGEYMKFKHGSDLKLRNFNFFHELMEGDKSQGKVVDQQTGELITTRDEVADYFDWVINL
KVMLISNETYEETGKYFDASHESSSLYLNQNKSKLVVPLKNKLQPEYYGAYTGITQGYM
VILKLLDKGGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKSFEVVVPRVLK
GTFLVDGEEKFILSSYRYKVNATQLILPVSDIKLIQDNFKALKLNEMQTKKLIEIYDNI
LRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEVIIKILRGLHFGSDLQNLKE
IGFGTTPLGQFQVSEAGIRLSNTAIFIIFKSPTGLFNRKLYLKLN
GKRPAATKKAGQAKKKGS **YPYDVPDYAYPYDVPDYAYPYDVPDYA** (SEQ ID NO: 85).

Wild Type *Peptoniphilus* sp. Marseille-P3761 Cas9

MEKKNTYTIGLDIGTDSVGWAVVKDDLELVKKRMKVLGNTEINYIKKNLWGSLLFESGQTA
 KDRRLKRVARRRYERRNRNRLTELQKIFAPAIDEVDENFFFRLNESFLVPEDKAFSKNPIFG
 TLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIAI
 NENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDGRANKQS
 PMMAFITLIVGKKAKFKAANLEEEISLNLTDDSYDENLEILLNTIGSDFADLFDHAQRVY

NAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQ
 KKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQ
 VHLAEMEAIILDRQENYYPWLKRNREKIISLLTFRIPYYVGPLADGQSEFAWLERKSDEKIK
 PWNFSDVVDLDRSAEKFIEQLIGRDTLPDEYVLPKKSLIYQKYMVFNELTKIAYLDERQK
 RMNLSSVEKKEIFETLFKKRSKVTEQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELA
 KVPGMKSMMDDPDNEDLMEEIVKILTVFEDRKMRKQLEKYKERLSPEQIKELAKKHYTGW
 GRLSKLLVGIRDQETQKTILDYLEDNHSGGRQHNRNLMQLINDDRLSFKKTIAELQM
 IDPSADLYAQVQEIAGSPIKKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTTARGKA
 LSKRREAKIKEGLAALGSSLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQY
 DEDHIIIPQSFTVDNSLDNLVLTNSSQNRCGNKKDDVPSLEVNRQLAYWRSLKDALMTQRK
 FDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNAVAKLLDMRLNDKKDEAGNKIRETNIVL
 LKSAMASEFRKMFRKYKVRELNDYHHADAYLNAAIAINLLALYPYMADDFVYGEFRYKKK
 PQAEKATYEKLQRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQVNWKKAER
 TGMLFKETINGKTNKGSRIPIKKDLDPSKYGGYIEEKMAYYAVISYEDKKKPGKTIVGIS
 IMDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIYPDGRRRYITGATKTTKGKVELQKA
 NQIAMEQDLVNFIYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINR
 LRRIYEEKKDPVDIEALVASFIELDLKLTSAGAPADFIFMGEAISRRRYNSMTGLFDGQV
 IYQSLTGLYETRMRFED (SEQ ID NO: 86).

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

MPKKKRKVGEKKTNYTIGLDIGTDSVGAWAVVKDDLELVKKRMKVLGNTETNYIKKNLWGSL
 LFESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFFRLNESFLVPEDKA
 FSKNPIFGTLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGK
 LDTEHIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKD
 IGRANKQSPMMAFITLIVGKKAFAFNLEEEISLNLTDDSYDENLEILLNTIGSDFADL
 FDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMT
 FVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSF
 YNGVIPNQVHLAEMEAIILDRQENYYPWLKRNREKIISLLTFRIPYYVGPLADGQSEFAWLE
 RKSDEKIKPWNFSDVVDLDRSAEKFIEQLIGRDTLPDEYVLPKKSLIYQKYMVFNELTKI
 AYLDERQKRMNLSSVEKKEIFETLFKKRSKVTEQLVKFFENYLQIDNPTIFGIEDAFNAD
 YSTYVELAKVPGMKSMMDDPDNEDLMEEIVKILTVFEDRKMRKQLEKYKERLSPEQIKEL
 AKKHYTGWGRSLSKLLVGIRDQETQKTILDYLEDNHSGGRQHNRNLMQLINDDRLSFK
 KTIAELQMIDPSADLYAQVQEIAGSPIKKGILLGLKIVDEIIRVMGEKPENIVIEMAREN

QTTARGKALSKRREAKIKEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPL
DFDRLSQYDEDHIIPQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEVVRQLAYWRSLK
AGLMTQRKFNDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKLLDMRLNDKDEAGNK
IRETNIVLLKSAMASEFRKMFRILYKVRELNDYHHAHDAYLNAAIAINLLALYPYMADDFVY
GEFRYKKKPQAEKATYEKLQRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTICKVMGYRQVN
VVKKAAEERTGMLFKETINGKTNKGSRIPIKKLDPSKYGGYIEEKMAYYAVISYEDKKKP
GKTIIVGISIMDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIYPDGRRRYITGATKTTK
GKVELQKANQIAMEQDLVNFYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFL
DAETNINRLRRIYEEEKKDPVDIEALVASFIELLKLTSGAPADFI 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.

20 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
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 aspartic acid-to-alanine substitution (D12A) in the RuvC domain of EpeCas9. In some 10 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.

15 In some embodiments, the mutation is an aspartic acid-to-glycine substitution (D10G) 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 20 embodiments, 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.

25 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 Cas9 protein comprises one or more mutations that inhibits the ability of Cas9 to cleave both strands of a DNA duplex.

30 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 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 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

Sequence of <i>ScoCas9</i> with Nuclear Localization Signal (NLS) and Linker
MPKKKRKV<u>GGKPYSIGLDIGTN</u>SVGWAVVTDDYKVPAAKKMKVLGNTDKQSIKKNILLGALLFDSGETAEATRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRGEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFKDFVEEYDKTIEESHLEITVDALSILTEKVSKSSRLENLIAHYPTEKKNT

LFGNLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLY
 DAIILSGILTVDNSTKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDKNKKG
 YASYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEM
 HAILRRQGEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRFAWAEYKADEKITPWNFDD
 ILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDT
 NMKQEIFDHVFKENRKVTKDKLLNYLNKEFEFRIVNLTGLDENKAFLNSSLGTYHDLRKI
 LDKSFLDDKANEKTIEDIIQTTLFEDREMIRQLQKSDIFTKAQLKKLERRHYTGWGRL
 SYKLINGIRNKENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDVDDIANV
 VHDLPGSPAICKGILQSVKIVDELVKVMGHNPANIIEMARENQTTDKGRNSQQLKLLQ
 DSLKNLDNPVNICKNVENQQLQNDRFLYIYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFI
 KDNSLDNRVLTRSDKNRGKSSDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGGL
 TEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKSNLVSFRKE
 FELYKVREINDYHHAHDAYLNAVVGNNALLKYPQLEPEFVYGEYPKNSYRSRKSATEKFL
 FYSNILRFFKKEDIQTNEEDGEIAWNKEKHIKILRKVLSYPQVNIVKKTEEQTGGFSKESIL
 PKGESDKLIPRKTNSYWDPKKYGGFDSPVVAYSILVFADVEKGKSKLRKVQDMVGITIM
 EKKRFEKNPVDFLEQRGYNRVRLEKIIKLPKYSLFELENKRRRLLASAKELQKGNELVIPQ
 RFTTLLYHSYRIEKDYEPHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMLFSK
 NKDAEVSSLAKSFISLLTFTAFGAPAAFNFFGENIDRKRTSVTECLNATLIHQSIITGLYE
 TRIDLSQLGEDKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA
 (SEQ ID NO: 2).

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

MPKKKRKVGAKNKDIRYSIGLDIGTNSVGWAVMDEHYELLKKGNHHMWGSRLFDAAEPAAT
 RRASRSIRRINKRERIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQKNLGNDYKD
 NYNLFIEKDFNDKTYDKYPTIYHLRKELCENKEADPRLIYIYALHHIVKYRGNFLKEGQS
 FAKVYEDIEEKLDNTLKKFMSLNLDNLFVDNDINSMITVLSKIYQRSKKADDLLKIMNPT
 KEERAAYKEFTKALVGLKFNVSkmilaQEVKKDDKDIELDFSNVDYDSTVDGLQAELEYI
 EFIEMLHSINSWVELQDILGNNSTISAAMVERYEEHKNDLRLKKVIREELPDKYNEVFRE
 DNPKLHNLYLGYIKYPKNTPVEEFYFYIKRLLAKVDTGEAREILERIDLEKFMLKQNSRTNG
 SIPYQMOKDEMIIQIIDNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFED
 KQKERILPWNYDQIVDIDATAEGFIERMQNTGTYFPDKPVMAKNSLTISKFEVLNELNKIR
 INGKLIPVETKHELLSDLFMKNKTITDKKLKDWLVTHQYYDTNEELKIEGYQKDLQFSTSL
 APWIDFTKIFGEINASNYQLIEKIIYDISIFEDKKILKRRLKVKYQLDDLVDKILKLNYT
 GWSRLSEKLLTGIKSNSKETILSILENSNMNLMEIINDESLGFKQIIEESNKKDIEGPFR

YDEVKKLAGSPAIKRGIGWQALLVVQEITKFMKHEPSHIYIEFAREEQEKRTERIAKLQK
 IYKDLNLQTKEQLVYESLKKEDAKKIDTDALYLYLQMGKSMYSGKPLDIDKLSTYHID
 HILPRSLIKDDSLDNRVLVLPKENEWKLDSETVPFEIRNMMGFWQKLHENGLMSNKKFFS
 LIRTDFNEKDKKRFINRQLVETRQIIKNVAVIINDHYTNVVTVRRAELSHQFRERYKIYK
 NRDLNDLHHADAYIACILGQFIHQNFGNMDVNMIYGQYKKNYKKDVQEHNNGFILNSMN
 HIHFNDDNSVIWDPSYIGKIKSCFCYKDVTKKLEQNDAKLFDLTILPSDKNSENGVTKA
 KIPVNKYRKDVNKYGGFSGDAPIMLAIEADKGKKHVRQVIAFPLRLKNYNDERIKFIEKE
 KNLKNVKILTEVKKNQLILINHQYFFITGTNELVNATQLKLSAKNTKNLFNLVDANKHNKL
 ESIDDANFNEVIQELICKLQEPIYSRYNSIGKEFEDSYEKINAUTQDKLYIIEYLIAIMS
 AKATQGYIKPELAREIGTNGKNKGRIKSFTIDLNTTFISTSVTGLFSKKYKL**KRPAATK**
KAGQAKKKGS YPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 5).

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

MPKKRKVGS INFQRRLMETQASNQLISSHLKGYPKDYFVGLDIGTSSVGWAVTNKAY
 ELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRLERKLRKLLEELFADAMAQVDPTFF
 MRLRESKYHYEDKTTGHSSKHILFIDKNYNQDYFKEYPTVYHLRSELMKSGTDDIRKLFL
 AVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGKT
 KSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANVLGLNASLIDLFGSVE
 ELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPGLTISESKV
 KAFNKHKKDDLAILKSSLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREFYKYT
 KKIVEGLSDSKDKEYILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLN
 EVADGSVAEKLIKMLEFRIPIYYVGPLNTHHNVDNGGFAWRKASGRVT PWNFDDKIDRE
 KSAAAFIKNLTKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAV
 FKQDHKKMTKNRIEQFLKDNGYISETHKHEITGLGEIKNDLASYRDMVRILGDGFDRSMA
 EEEITDITIFGESKKMLRETLRKKFASCLDDEAIKKLTLYRDWGRLSQKLLNGIEGCDK
 AGDGTPETIIILMRNFSYNLMEELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPA
 VKRAVWQALRIVDEVAHIKKALPARIFVEVTRSNKNEKKKDSRQKRLSDLYAAIKDDVL
 LNGLNNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGRCAVTGEIIIELSLLNTDNYDIDHI
 YPRSLTKDDSFNLVLCKRTANAQKSDAYPISEEIQKTQKPFWTFLKQQGLISERKYERLT
 RITPLTADDLSGFIARQLVETNQSVKAATTLLRRLYPGVVVFVKAENVTDFRHDDNFIKV
 RSLNHHHHAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKDG
 KAWDVKTSMDTVKKMMDSDNRVTKRLLEQTGALADATIYKATVAGKAKDGAYIGMKTKSS

VFADVSKYGGMTKIKNAYSIIIVQYTGKKGEVIKEIVPLPIYLTLNRNTTDQDLINYVASIIP
QAKDISIIYGKLCINQLVKVNGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNN
MRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLVSKLKMPIYQKMKGNKAAELS
EKGYGLFKKMSLEEQSIHLLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVIN
ESITGLYSNEVTIVGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA
(SEQ ID NO: 9).

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

MPKKKRKVGTVKDYYIGLDIGTSSVGWAVTDEAYNVLFNSKKMWGVRLFDDAKTAEERR
GQRGARRRLDRKKERLSSLQDFFAEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFND
KDFKDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIGEGQKFDTKSSF
ENSLNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIM
IGSSQKLVDLFENPEDFDSAIAKSVDFSTTSFDDKYSDYELALGDKIALVNLKEIYDSSI
LENLLKEADKSKDGNKYISNAFKVKKYNHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNDNY
VSYTKSSISNNKRVKANKFTDQEAFYKFAKKLETIKYKINKVNGSKADLELIDGMLRDME
FKNFMPKIKSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRI
YYVGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLDSSREEFIDSILGRCTYLDEKVLPKA
SLLYNEYMVLNELNNLKLNLPITEEMKKIFDQLFKTRKKVTLKAVANLLKEFNINGEI
LLSGTDGDFKQGLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGY
GKYFTDSEIKKMAGLNYKDWGRSLSKLLTGLEGANKITGERGSIIHFMREYNLNLMSA
SFTFTEEIQKLNPVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIE
MARGKEEVKARKESRKNQLLKFYKDGGKAFIGEIGEERYSYLLSEIEGEEENKFRWDNLYL
YYTQLGRCMYSLEPIDISELSSKNIYDQDHIPKSKIYDDSIENRVLVKKDLNSKGNSYP
IPDEILNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETA
NLLKTICKNSEIVYASKENASRFRQEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKD
PMNFVKKQEKAWSYNLENMFYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFT
RMNYIESGALFNATLQRKNKGSRPLDKGPKSSIEKYGGYTINKACFAVLDIKSKNKIER
KLMPVEREITYAKQKNDKLSDEIFSKYLCDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITG
GSDKTLELRSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFK
TSVFKKSFLNLQDDKNIENIDFKFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKSDY
GRLSSKTNNIGNYQEJKIINQSITGLFENEVDLLKLGKRPAATKKAGQAKKKGSYPYDVP
DYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 15).

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

MPKKKRKVGKEYHIGLDIGTSSIGWAVTD**SQFKLMRIKGKTAIGVRLFEEGKTA**ERRTFR
 TTRRRLKRRKWR~~LHYLDEI~~FAPH~~LQEVDENFL~~RR~~LKQSNIHPEDPAKNQAFIGKLLFP~~DLL
 KKNERGYPT**L**I**KMRDEL**PVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLVHIVKYRG
 HFLNNASV**D**KFKVGR**I**DFDKSFNVLNEAYEELQN~~GEGSFTIEPSKVEKIGQLLDTKMRKL~~
 DRQKAVAKLLEV**V**KVADKEETKRNQ**IATAMS**KLVLGYKADFATVAMANGNEWKIDLSSETS
 EDEIEKFREELSDAQNDILTEITS**LFSQIMLNEIVPNGMSI**SESMMDRYW~~THERQLAEVKE~~
 YLATQPASARKEFDQVYNKYIGQAPKEGF**DLEKGLKKILSKKENWKEI**DELLKAGDFLPK
 QRTSANGVI**PHQMHQ**QELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPY
 YVGPLVTPEVQKATSGAKFAWAKRKEDGEIT**PWNLWDKIDRAESA**EAFIKRMTVKDTYLLN
 EDVLPANSLLYQKYNVLNELNNVRVNGRRLSVGI**KQDIYTEL**FKKKKTVKAGDVASLVMAK
 TRGVNKPSVEGLSDPKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIEWRSVFEDGEIF
 ADKLTEVEWLTD**EQRSALVKKRY**KGWGR~~L~~SKLLTGIVDENGQRIIDLMWNTDQNFMQIVN
 QPVFKEQ**IDQLNQKAIT**NDGMTLRERVESVL~~DAYTSPQNKKAIWQVVRVVEDIVKAVGNA~~
 PKSISIEFARNEGKGEITRS~~RRTQLQKL~~FEDQAHELVKDTSLTEELEKAPDLSDRYYFYF
 TQGGKDMYT**GDPINFDE**ISTKYDIDHILPQS**FVKDDSLDNRVLVSRAENNKKSDRVPAKLY**
 AAKMKPYWNQLLQGLITQRKFENLTMDVDQT**IKYRSLGFV**KRQLVETRQVIKLTANILGS
 MYQEAGTDIIETRAGLT**KQLREFDLPKV**REVNDYHHAVDAYLTTFAGQYLNRRYPKLR~~SF~~
 FVYGEYM**KFKHGSDLKLRNFNFFHELMEGDKSQGKVVDQQTGELIT**TRDEVADYFDWVINL
 KVMLISNETYEETGKYFDASHES~~SSLYLKNQNKKSKLVVPLKNKLQPEYYGAYTGITQGYM~~
 VILKLLDKGGFGVYRIPRYAADILNKCHDEVAYRNKIAEI**ISSDPRAPKS**FEVVVPRVLK
 GTFLVDGEEKF**ILSSYRYKV**NATQLILPVSDIKLIQDNFKALKLNVEMQTKKLIEIYDNI
 LRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEVIIKILRGLHFGSDLQNLKE
 IGF~~GTTPLGQFQVSEAGIRLSNTAIFI~~FKSPTGLFN~~RKLYLK~~N
GKRPAATKKAGQAKKKGS YPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 85).

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

MPKKKRKVGEKKNYTIGLDIGTDSVGWAVVKDDLELVKKRMKVLGNTE~~NYIKKNLWGSL~~
 LFESGQTAKD**RRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFFRLNESFLVPEDKA**
 FSKNP~~I~~FGTLGEDKTYKTYPTIYHLRQH**LA**SEEKADVRLIYLALAHMIKYRGHFLIEGK
 LDTEHIAINENLEQFFESYNALFSEEPIELKEELIAENILREKNSRTVKEKRITSFLKD

I GRANKQS PMMAFITLIVGKKAKFKA AFNLEEEISLNLTDDSYDENLEILLNTIGSDFADL
 FDHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMT
 FVAPKDAQKKDLKG YAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSF
 YNGVI PNQVHLAEMEAILDRQENYY PWLKENREKIISLLTFRIPYYVGPLADGQSEFAWLE
 RKSDEKIKPWNFS DVVDLDRSAEKFIEQLIGRTYLPDEYVLPKKSLIYQKYMVFNELTKI
 AYLDERQKRMNLSSVEKKEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNAD
 YSTYVELAKVPGMKSMMDDPDNE DLMEEIVKILT FEDRKMRKQLEKYKERLSPEQIKEL
 AKKHYTGWGRSLSKLLVGIRDKETQKTILDYLVEDDNHSGGRQHNRNLMQLINDDRLSFK
 KTIAELQMIDPSADLYAQVQEIA GSPA IKKGILLGLKIVDEITIRVMGEKPENIVIEMAREN
 QTTARGKALSKRREAKIKEGLA ALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPL
 DFDRLSQYDEDHIIPQSFTVDNSLDNLVLTNSSQNRGNKKDDVPSLEVNRQLAYWRSLK
 AGLMTQRKFDNLTKAMRGGLTDKRERFIQRQLVETRQITKNVAKLLDMRLNDKDEAGNK
 IRETNIVLLKSAMASEFRKMFR LYKVRELNDYHHADAYLNAAIAINLLALYPYMADDFVY
 GEFRYKKKPQAEKATYEKL RQWNLI KRFGEKQLFTP DHEDCWNKERDIKTIKKVMGYRQVN
 VVKKAEERTGMLFKETINGKTNKGSRIPIKKLDPSKYGGYIEEKMAYYAVISYEDKKKP
 GKTIVGISIMDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIYPDGR RRYITGATKTTK
 GKVELQKANQIAMEQDLVNFIYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFL
 DAETNINRLRRIYEEEKKDPVDIEALVASFIELLKLT SAGAPADFIFMG EAISRRRYNSM
 TGLFDGQVIYQSLTGLYETRMRFEDGKRPAATKKAGQAKKKGS YPYDVPDYAYPYDVPDY
 AYPYDVPDYA (SEQ ID NO: 87).

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

MPKKKRKVGMGKYSIGLDIGTNSVGWAVVTDDYKVPACKMKVLGNTDKQS IKKNLLGALL
 FDSGETAETRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRG
 EHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDL
 KAENTDVQALFKDFVEEYDKTIEESHLS EITVDALSILTEKVSKSSRLENLIAHYPTEKKN
 TLFGNLIALS LDH PNF KTNFQLSEDAKLQFSKDTYEE DLEGFLGEVGDEYADLFASAKNL
 YDAI LLSGILT VDDNSTKAPLSASMVKRYEEHQKDLKKLDFIKVNAPDQYNAIFKDKNKK
 GYASYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQE
 MHAI LRRQGEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRF AWA EYKADEKITPWNFD
 DILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKV KYVNEQGEAKFFD
 TNMKQEIFDHVFKENRKVT KDKL NYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRK
 ILDKSFLDDKANEKTIEDIIQTTLFEDREMIRQRLQKYS DIFTKAQLKKLERRHYTGWGR

LSYKLINGIRNKENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIAN
VVHDLPGSPAICKGILQSVKIVDELVKVMGHNPNANIIEMARENQTTDKGRRNSQQRLKLL
QDSLKNLDNPVNICKNVENQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAF
IKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGG
LTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKITITLKSNLVSNFRK
EFELYKVREINDYHHAHDAYLNAVVGNAALLKYPQLEPEFVYGEYPKYNYSYRSRKSATEKF
LFYSNILRFFKKEDIQTNEDGEIAWNKEKHIKILRKVLSYPQVNIVKKTEEQTGGFSKESI
LPKGESDKLIPRKTKNSYWDPKKYGGFMQPVVAYSILVFADVEKGKSKLRKVQDMVGITI
MEKKRFEKNPVDFLEQRGYRNVRLEKIILPKYSLFELENKRRRLLASAKFLQKGNELVIP
QRFTTLLYHSYRIEKDYPEHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKEMLFS
KNKDAEVSSLAKSFISLTFATFGAPRAFNFFGENIARKEYRSVTECLNATLIHQSITGLY
ETRIDLSKLGEDGE~~GADKRTADGSEFESP~~**KKKRKV** (SEQ ID NO: 95)

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

MPKKKRKVGMGKPYSIGLDIGTNSVGWAVVTDDYKVPACKMKVLGNTDKQSIIKKNLLGALL
FDSGETAETRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRG
EHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDL
KAENTDVQALFKDFVEEYDKTIEESHlseITVDALSILTEKVSKSSRLENLIAHYPTEKKN
TLFGNLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNL
YDAILLSGILTVDNSTKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDKNKK
GYASYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGIIPHQIHLQE
MHAILRRQGEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRFOWAEYKADEKITPWNFD
DILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKVYVNEQGEAKFFD
TNMKQEIFDHVFKENRKVTDKLLNYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRK
IILDKSFLDDKANEKTIEDIITQTLFEDREMIRQRLQKYSIFTKAQLKKLERLHYTGWR
LSYKLINGIRNKENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIAN
VVHDLPGSPAICKGILQSVKIVDELVKVMGHNPNANIIEMARENQTTDKGRRNSQQRLKLL
QDSLKNLDNPVNICKNVENQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAF
IKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGG
LTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKITITLKSNLVSNFRK
EFELYKVREINDYHHAHDAYLNAVVGNAALLKYPQLEPEFVYGEYPKYNYSYRSRKSATEKF

LFYSNILRFFKKEDIQTNEDGEIAWNKEKHIKILRKVLSYPQVNIVKKTEEQTGGFSKESI
LPKGESDKLIPRKTKNSYWDPKKYGGFMQPVVAYSILVFADVEKGKSKKLRKVQDMVGITI
MEKKRFEKNPVDFLEQRGYRNRLEKIIKLPKYSLFELENKRRLLASAKFLQKGNELVIP
QRFTTLLYHSYRIEKDYEPHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKEMLFS
KNKDAEVSSLAKSFISLLTFTAFCAPRAFNFFGENIARKEYRSVTECLNATLIHQSI
ETRIDLSKLGEDGE**GADKRTADGSEFESPKKKRKV** (SEQ ID NO: 96)

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) of the native sequence with codons that are more frequently used or most frequently used in 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 peruvensis* 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 peruvensis* 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)

MDKKYSIGLDIGTNSVGAVITDDYKVPSSKKFKVLGNTDRHSIKKNLIGALLFDGETAEAT
RLKRTARRRYTRRKNRICYLQEIFSNEAKVDDS FFHRLEESFLVEEDKKHERHPIFGNIVD
EVAYHEKYPTIYHLRKKLVDSTDKADLRLIYLALAHMIKFRGHFLIEGDLNPDNSDVDKLFI
QLVQTYNQLFEENPINASGVDAKAILSARLSKSRRLENLIAQLPGEKKNGLFGNLIALSLGL
5 TPNFKSNFDLAEDAKLQLSKDTYDDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNT
EITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEF
YKFIKPILEKMDGTEELLVKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYPFLK
DNREKIEKILTFRIPYYVGPLARGNSRFAMTRKSEETITPWNFEVVVDKGASAQSFIERTMT
NFDKNLPNEKVLPKHSSLYEYFTVYNELTKVKVYVTEGMRKPAFLSGEQKKAIVDLLFKTNRK
10 VTVKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILEDIV
LTTLTFEDREMIEERLKYAHLFDDKVMQQLKRRRTGWRGRLSRKLINGIRDQSGKTILDF
LKSDGFANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAICKGILQTVKVV
DELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEGIKELGSQLKEHPVENTQL
QNEKLYLYLQNGRDMYVDQELDINRLSDYDWDHVIPQSFNKDDSIDNKVLTRS DKNRGKSD
15 NVPSEEVVKMKNYWRQLLNAKLITQRKF DNLTKAERGGI SELDKAGFIKRQLVETRQITKH
VAQILD SRMNTKYDENDKLIREVKVITLKS KLVDFRKDFQFYKVREINNYHHAHDAYLNAV
VGTALIKKYPKLESEFVYGDYKVDVRKMIAKSEQEIGKATAKYFFYSNIMNFFKTEITLAN
GEIRKRPLIETNGETGEIVWDKGRDFATVRKVLSMPQVNIVKKTEVQTGGFSKESILPKRNS
DKLIARKKDWDPKYGGFDSPTVAYSVLVVAKVEKGKSKKLKSVKELLGITIMERSSFEKNP
20 IDFLEAKGYKEVKKDLIIKLPKYSIFELENGRKMLASAGELQKGNELALPSKYVNFLYLAS
HYEKLKGSPEDNEQKQLFVEQHKHYLDEIIEQISEFSKRVILADANLDKVL SAYNKHRDKPI
REQAENIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQSI TGlyETRIDLSQ
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 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 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.

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 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 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 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 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 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 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 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 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. 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.

10 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 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.

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

20 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, 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.

25 In some embodiments, the loop forming sequences are 3, 4, 5 or more nucleotides in length. In some embodiments, the loop has the sequence GAAA, AAAG, CAAA, AAAC, UUUU, UUAUAU, 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 UUAUAU. 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.

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

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

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

10 5'-

GUUUUAGAGUUGUGUUAUUGAAAAAAUACACAACACAGAGUAAAAUAAGCUUA
UGCUUAAAUGCAGCUUUGCUGGUGUCAUUAGAUGACUUUACUAAGGUUGC
UUCGGCAACCUUUUU-3' (SEQ ID NO: 7) for SirCas9,

5'-

15 GUUUGAGAGUAGUGUGAAAACAUUACGAGUCAAAUACAAUUAUUUACAA
UGCCUUCGGGCUGCCCACGUAGGGCACCUACUCUCAAUUCUUCGGAAUUGAG
UU-3' (SEQ ID NO: 13) for VapCas9,

5'-

20 GUUUGAGAGUUAUGUAUUGAAAAAUUACAUGACGAGUCAAAUAAAAAUU
AUUCAAACGCCUAUUUAUAGGCCGCAGAUGUUCUGCAUAUGCUUGCUALU
GCAAGCUU-3' (SEQ ID NO: 19) for EpeCas9,

5'-

25 GUCUUGGAUGAGUGUGAAAACACUCAUAGUCAAGAUCAAACCGAGUGGUUUC
CACGAGUUAUUACUUUUGAGGUCUUAUAGGCCAUACAUAAAAGGAGUCG
GAAUUUCCGGCUCCUUUCUU-3' (SEQ ID NO: 95) for LfeCas9, and

5'-

GUUUUAGAGCCAUGUAGAAAACAUUGCAAGUAAAAUAAGGCUUUGUCCGU
AAUCAACUUGAAAAAGUGGCGCUGUUUCGGCGCUUU-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
5 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
10 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
15 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
20 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,
25 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.

30 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 activity, kinase activity, phosphatase activity, ubiquitin ligase activity, deubiquitinating activity, adenylating activity, deadenylation activity, SUMOylating activity, deSUMOylating activity, ribosylation activity, deribosylation activity, myristylation activity, demyristylation 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 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 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 a adenine and cytosine base editor (ACBE or CABE), wherein ACBE or CABE is generated by fusing a heterodimer of TadA and an activation-induced cytidine deaminase (AID) to the N- and C-terminals of Cas9 nickase (nCas9). In some embodiments, the ACBE or CABE 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 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%, 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 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, 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 some embodiments, a deaminase incorporated into a fusion protein comprises all or a portion of 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 an APOBEC3B deaminase. In some embodiments, a deaminase incorporated into a fusion 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 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 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 deaminases are provided below.

5 pmCDA1 (*Petromyzon marinus*)
MTDAEYVRIHEKLDIYTFKKQFFNNKKSVSHRCYVLFELKRRGERRACFWGYAVNK
PQSGTERGIHAEIFSIRKVEEYLDRNPGQFTINWYSSWSPCADCAEKILEWYNQELRG
NGHTLKIWACKLYYEKNARNQIGLWNLRDNGVGLNVMVSEHYQCCRKIFIQSSHNQ

10 LNENRWLEKTLKRAEKRSELIMIQVKILHTTKSPAV (SEQ ID NO: 22)

Human AID:

MDSLLMNRRKFLYQFKNVRWAKGRRETYLCYVVKRRDSATSFSLDFGYLRNKNGC
HVELLFLRYISDWLDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGNPNLSLRIFTAR
LYFCEDRKAEP EGLRRLH RAGVQIAIMTFKAPV (SEQ ID NO: 23)

15 Human AID:

MDSLLMNRRKFLYQFKNVRWAKGRRETYLCYVVKRRDSATSFSLDFGYLRNKNGC
HVELLFLRYISDWLDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGNPNLSLRIFTAR
LYFCEDRKAEP EGLRRLH RAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHEN
SVRLSRQLRILLPLYEVDDL RDAFR TLGL (underline: nuclear localization sequence;

20 double underline: nuclear export signal) (SEQ ID NO: 24)

Mouse AID:

MDSLLMKQKKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSCSLDFGHLRNKSGC
HVELLFLRYISDWLDLDPGRCYRVTWFTSWSPCYDCARHVAEFLRWNPNLSLRIFTAR
LYFCEDRKAEP EGLRRLH RAGVQIGIMTFKDYFYCWNTFVENRERTFKAWEGLHEN

25 SVRLTRQLRILLPLYEVDDL RDAFR MLGF (underline: nuclear localization sequence;
double underline: nuclear export signal) (SEQ ID NO: 25)

Canine AID:

MDSLLMKQRKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSFSLDFGHLRNKSGC
HVELLFLRYISDWLDLDPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSLRIFAAR
30 LYFCEDRKAEP EGLRRLH RAGVQIAIMTFKDYFYCWNTFVENREKTFKAWEGLHEN
SVRLSRQLRILLPLYEVDDL RDAFR TLGL (underline: nuclear localization sequence;
double underline: nuclear export signal) (SEQ ID NO: 26)

Bovine AID:

MDSLLKKQRQFLYQFKNVRWAKGRHETLYCYVVKRRDSPTSFSLDFGHLRNKAGC
HVELLFLRYISDWLDLPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSRIFTAR
LYFCDKERKAEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHE
NSVRLSRQLRRILLPLYEVDDLRAFRTLGL (underline: nuclear localization sequence;

5 double underline: nuclear export signal) (SEQ ID NO: 27)

Rat AID:

MAVGSKPKAALVGPHWERERIWCFCLSTGLGTQQTGQTSRWLPAATQDPVSPPRS
LLMKQRKFLYHFKNVRWAKGRHETLYCYVVKRRDSATSFSLDFGYLRNKSGCHVE
LLFLRYISDWLDLPGRCYRVTWFTSWSPCYDCARHVADFLRGNPNLSRIFTARLTG
10 WGALPAGLMSPARPSDYFYCWNTFVENHERTFKAWEGLHENSVRLSRRLLRILLPL
YEVDDLRAFRTLGL (SEQ ID NO: 28)

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

clAID (*Canis lupus familiaris*):

MDSLLMKQRKFLYHFKNVRWAKGRHETLYCYVVKRRDSATSFSLDFGHLRNKSGC
15 HVELLFLRYISDWLDLPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSRIFAAR
LYFCEDRKAEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENREKTFKAWEGLHEN
SVRLSRQLRRILLPLYEVDDLRAFRTLGL (SEQ ID NO: 29)

btAID (*Bos taurus*):

MDSLLKKQRQFLYQFKNVRWAKGRHETLYCYVVKRRDSPTSFSLDFGHLRNKAGC
20 HVELLFLRYISDWLDLPGRCYRVTWFTSWSPCYDCARHVADFLRGYPNLSRIFTAR
LYFCDKERKAEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHE
NSVRLSRQLRRILLPLYEVDDLRAFRTLGL (SEQ ID NO: 30)

mAID (*Mus musculus*):

MDSLLMNRRKFLYQFKNVRWAKGRRETYLCYVVKRRDSATSFSLDFGYLRNKNGC
25 HVELLFLRYISDWLDLPGRCYRVTWFTSWSPCYDCARHVADFLRGNPNLSRIFTAR
LYFCEDRKAEPEGLRRLHRAGVQIAIMTFKDYFYCWNTFVENHERTFKAWEGLHEN
SVRLSRQLRRILLPLYEVDDLRAFRTLGL (SEQ ID NO: 31)

rAPOBEC-1 (*Rattus norvegicus*):

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELKETCLLYEINWGGRHSIWRHTSQNT
30 NKHVEVNIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSPSNEAHWPRYPHLW
VRLYVLELYCIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLK (SEQ
ID NO: 32)

maAPOBEC-1 (*Mesocricetus auratus*):

MSSETGPVVVDPTLRRRIEPIHEFDAFFDQGELRKETCLLYEIRWGGRHNIWRHTGQN
TSRHVEINFIEKFTSERYFYPSTRCSIVWFLSWSPCGECSKAITEFLSGHPNVTLFYAA
RLYHHTDQRNRQGLRDLISRGVTIRIMTEQEYCYCWRNFVNYPPEVYWPYPNL
WMRLYALELYCIHLGLPPCLKIKRRHQYPLTFFRLNLQSCHYQRIPPHILWATGFI

5 (SEQ ID NO: 33)

ppAPOBEC-1 (*Pongo pygmaeus*):

MTSEKGPSGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWRSSGKN
TTNHVEVNFIKKFTSERRFHSSISCSITWFLSWSPCWECSSQAIREFLSQHPGVTLVIYV
ARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYP

10 PLWMMLYALELHCIILSLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIH
PSVTWR (SEQ ID NO: 34)

ocAPOBEC1 (*Oryctolagus cuniculus*):

MASEKGPSNKDYTLRRRIEPIWEFEVFFDPQELRKEACLLYEIKWGASSKTWRSSGKN
TTNHVEVNFKLTSEGRLGPSTCCSITWFLSWSPCWECSSMAIREFLSQHPGVTLIIFV
15 ARLFQHMDRRNRQGLKDLVTSGVTVRVMSVSEYCYCWNFVNYPGKAAQWPRY
PPRWMLMYALELYCIILGLPPCLKISRRHQKQLTFFSLTPQYCHYKMIPPYILLATGLL
QPSVPWR (SEQ ID NO: 35)

mdAPOBEC-1 (*Monodelphis domestica*):

MNSKTGPSVGDAVLRRRIKPWEFVAFFNPQELRKETCLLYEIKWGNQNIWRHSNQN
20 TSQHAEINFMEKFTAERHFNSSVRCSSITWFLSWSPCWECSSKAIRKFLDHYPNVTLAIFI
SRLYWHMDQQHRQGLKELVHSGVTIQIMSYSEYHYCWRNFVDYPQGEEDYWPKYP
YLWIMLYVLELHCIILGLPPCLKISGSHSNQLALFSDLQDCHYQKIPYNVLVATGLV
QPFVTWR (SEQ ID NO: 36)

ppAPOBEC-2 (*Pongo pygmaeus*):

25 MAQKEEEAAAATEAASQNGEDLENLDDPEKLKELIELPPFEIVTGERLPANFFKFQFRN
VEYSSGRNKTFLCYVVEAQGKGGQVQASRGYLEDEHAAAHAEEAFFNTILPAFDPA
LRYNVTWYVSSPCAACADRIIKTLSKTKNLRLLILVGRLFMWEELEIQDALKKLKE
AGCKLRIMKPQDFEYVWQNFVEQEEGESKAFQPWEDIQENFLYYEKLADILK (SEQ
ID NO: 37)

30 btAPOBEC-2 (*Bos taurus*):

MAQKEEEAAAAEPASQNGEEVENLEDPEKLKELIELPPFEIVTGERLPAHYFKFQFRN
VEYSSGRNKTFLCYVVEAQSKGGQVQASRGYLEDEHATNHAEEAFFNSIMPTFDPA
LRYMVTWYVSSPCAACADRIVKTLNKTKNLRLLILVGRLFMWEEPEIQAALRKLKE

AGCRLRIMKPQDFEYIWQNFVEQEEGESKAFEPWEDIQENFLYYEEKLADILK (SEQ ID NO: 38)

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

MQPQRLGPRAGMGPCLGCSHRKCYSPIRNLISQETFKFHKNLGYAKGRKDTFLCY

5 EVTRKDCDSPVSLHHGVFKNKDNIHAEICFLYWFHDKVVKVLSPREEFKITWYMSW

SPCFECAEQIVRFLATHHNLSDLIFSSRLYNVQDPETQQNLCRLVQEQAQVAAMDLY

EFKKCWKKFVDNGGRRFRPKRLLTNFRYQDSKLQEILRPCYISVPSSSSTLSNICL

TKGLPETRFWVEGRRMDPLSEEEFYSQFYNNQRVKHLCYYHRMKPYLCYQLEQFNG

QAPLGCLLSEKGKQHAEILFLDKIRSMELSQVTITCYLTWSPCPNCAWQLAAFKRD

10 RPDYLILHIYTSRLYFHWKRPFKGLCSLWQSGILVDVMDLPQFTDCWTNFVNPKRPF

WPWKGLEIISRRTQRRLRIKESWGLQDLVNDFGNLQLGPPMS (SEQ ID NO: 39)

Mouse APOBEC-3-(2):

MGPFCLGCSHRKCYSPIRNLISQETFKFHKNLGYAKGRKDTFLCYEVTRKDCDSPV

SLHHGVFKNKDNIHAEICFLYWFHDKVVKVLSPREEFKITWYMSWSPCFECAEQIVRFL

15 ATHHNLSLDIFSSRLYNVQDPETQQNLCRLVQEQAQVAAMDLYEFKKCWKKFVDN

GGRRFRPKRLLTNFRYQDSKLQEILRPCYIPVPSSSSTLSNICLTKGLPETRFCVEG

RRMDPLSEEEFYSQFYNNQRVKHLCYYHRMKPYLCYQLEQFNGQAPLGCLLSEKGK

QHAEILFLDKIRSMELSQVTITCYLTWSPCPNCAWQLAAFKDRDPDLILHIYTSRLYFHW

KRPFQKGLCSLWQSGILVDVMDLPQFTDCWTNFVNPKRPFWPWKGLEIISRRTQRRL

20 RRIKESWGLQDLVNDFGNLQLGPPMS (italic: nucleic acid editing domain) (SEQ ID

NO: 40)

Rat APOBEC-3:

MGPFCLGCSHRKCYSPIRNLISQETFKFHKNRRLRYAIDRKDTFLCYEVTRKDCDSPV

SLHHGVFKNKDNIHAEICFLYWFHDKVVKVLSPREEFKITWYMSWSPCFECAEQVLRFL

25 ATHHNLSLDIFSSRLYNIRDOPENQQNLCRLVQEQAQVAAMDLYEFKKCWKKFVDNG

GRRFRPKKLLTNFRYQDSKLQEILRPCYIPVPSSSSTLSNICLTKGLPETRFCVERR

RVHLLSEEEFYSQFYNNQRVKHLCYYHGVPKYLCYQLEQFNGQAPLGCLLSEKGKQ

HAEILFLDKIRSMELSQVIITCYLTWSPCPNCAWQLAAFKDRDPDLILHIYTSRLYFHWK

RPFQKGLCSLWQSGILVDVMDLPQFTDCWTNFVNPKRPFWPWKGLEIISRRTQRRLH

30 RRIKESWGLQDLVNDFGNLQLGPPMS (italic: nucleic acid editing domain) (SEQ ID NO:

41)

hAPOBEC-3A (*Homo sapiens*):

MEASPASGPRHLMDPHIFTSNFNNNGIGRHKTLCYEVERLDNGTSVKMDQHRGFLH

NQAKNLLCGFYGRHAELRFLDLVPSLQLDPAQIYRVTFISWSPCFSWGCAGEVRAF

LQENTHVRRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIMTYDEFKHCWDTFVDHQ
GCPFQPWDGLDEHSQALSGRLRAILQNQGN (SEQ ID NO: 42)

hAPOBEC-3F (*Homo sapiens*):

MKPHFRNTVERMYRDTFSYNFYNRPILSRRNTVWL CYEVTKGPSRPLDAKIFRGQ

5 VYSQPEHHAEMCFLSWFCGNQLPAYKCFQITWFVSWTPCPDCVAKLAEFLAEHPNV
TLTISAARLYYYWERDYRRALCRLSQAGARVKIMDDEFAYCWENFVYSEGQPFMP
WYKFDDNYAFLHRTLKEILRNPMEMAMYPHIFYFHFKNLRKAYGRNESWLCFTMEV
VKHHSPVSWKRGVFRNQVDPETHCHAERCFLSWFCDDILSPNTNEYVTWYTSWSPC
PECAGEVAEFLARHSNVNLTIIFTARLYYFWDTDYQEGLRSLSQEGASVEIMGYKDFK

10 YCWENFVYNDDEPKPWKGLKYNFLFLDSKLQEILE (SEQ ID NO: 43)

Rhesus macaque APOBEC-3G:

MVEPMDPRTFVSNFNNRPILSGLNTVWL CCEVTKDPSGPPLDAKIFQGKVYSKAKY

HPEMRFLRFHKWRQLHHDQEYKVTWYVWSPECTRCANSVATFLAKDPKVTLTIF
VARLYYFWKPDYQQALRILCQKRGPHATMKIMNYNEFQDCWNKFVDGRGKPFKP

15 RNNLPKHYTLLQATLGELLRHLMDPGTFTSNFNNKPWVSGQHETYL CYKVERLHND
TWVPLNQHRGFLRNQAPNIHGFPKGRHAELCFLDLIPFWKLDGQQYRVTCTSWSPC
FSCAQEMAKFISNNEHVSLCIFAARIYDDQGRYQEGLRALHRDGAKIAMMNYSEFEY
CWDTFVDRQGRPFQPWDGLDEHSQALSGRLRAI (italic: nucleic acid editing domain;
underline: cytoplasmic localization signal) (SEQ ID NO: 44)

20 Chimpanzee APOBEC-3G:

MKPHFRNPVERMYQDTFSDNFYNRPILSHRNTVWL CYEVTKGPSRPLDAKIFRGQ

VYSKLKYHPEMRFFHWFSKWRKLHRDQEYEVTWYISWSPCTKTRDVATFLAEDPKV
TLTIFVARLYYFWDPDYQEALRSLCQKRDGPRATMKIMNYDEFQHCWSKFVYSQRE
LFEPWNNLPKYYILLHIMLGEILRHSMDPPTFTSNFNNELWVRGRHETYL CYEVERL

25 HNDTWVLLNQRRGFLCNQAPHKHGFLEGRHAELCFLDVIPFWKLDLHQDYRVTCTFS
WSPCFSCAQEMAKFISNNKHVSLCIFAARIYDDQGRQCQEGLRTLAKAGAKISIMTYSE
FKHCWDTFVDHQGCPFQPWDGLEEHSQALSGRLRAILQNQGN (SEQ ID NO: 45)

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

Green monkey APOBEC-3G:

30 MNPQIRNMVEQMEPDIFVYYFNNRPILSGRNTVWL CYEVTKDPSGPPLDANIFQGK

LYPEAKDHPEMKFLHWFRKWRQLHRDQEYEVTWYVWSPECTRCANSVATFLAEDPKV
TLTIFVARLYYFWKPDYQQALRILCQERGGPHATMKIMNYNEFQHCWNEFVDGQQ
KPFKPRKNLPKHYTLLHATLGELLRHVMMDPGTFTSNFNNKPWVSGQRETYLCYKVE
RSHNDTWVLLNQHRGFLRNQAPDRHGFPKGRHAELCFLDLIPFWKLDDQQYRVTCTFT

SWSPCFSCAQKMAKFISNNKHVSLCIFAARIYDDQGRCQEGLRTLHRDGAKIAVMNY
SEFEYCWDTFVDRQGRPFQPWDGLDEHSQALSGRLRAI (SEQ ID NO: 46)

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

Human APOBEC-3G:

5 MKPHFRNTVERMYRDTFSYNFYNRPILSRRNTWLCYEVKTKGPSRPLDAKIFRGQ
VYSELKYHPEMRFFHWFSKWRKLHRDQEYEVTWYISWSPCTKCTRDMATFLAEDPKV
TLTIFVARLYYFWDPDYQEALRSLCQKRDGPRATMKIMNYDEFQHCWSKFVYSQRE
LFEPWNNLPKYYILLHIMLGEILRHSMDDPPTFTFNNEPWVRGRHETYLCYEVERM
HNDTWVLLNQRRGFLCNQAPHKHGFLEGRHAELCFLDVIPFWKLDLQDYRVTFCFTS

10 *WSPCFSCAQEMAKFISKNKHVSLCIFTARIYDDQGRCQEGLRTLAEAGAKISIMTYSE*
FKHCWDTFVDHQGCPFQPWDGLDEHSQDLSGRLRAILQNQEN (SEQ ID NO: 47)

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

Human APOBEC-3F:

MKPHFRNTVERMYRDTFSYNFYNRPILSRRNTWLCYEVKTKGPSRPLDAKIFRGQ
15 VYSQPEHHAEMCFLSWFCGNQLPAYKCFQITWFVSWTPCPDCVAKLAEFLAEHPNVTL
TISAARLYYYWERDYRRALCRLSQAGARVKIMDDEEFAYCWENFVYSEGQPFMPW
YKFDDNYAFLHRTLKEILRNPMEAMYPHIFYFHFKNLRKAYGRNESWLCFTMEVVK
HHSPVSWKRGVFRNQVDPETHCHAERCFLSWFCDDILSPNTNEYVTWYTSWSPCPECA
GEVAEFLARHSNVNLTIIFTARLYYFWDTDYQEGLRSLSQEGASVEIMGYKDFKYCW
20 ENFVYNNDEPFKPWKGKLYNFLFLDSKLQEILE (SEQ ID NO: 48)

(italic: nucleic acid editing domain)

Human APOBEC-3B:

MNPQIRNPMERMYRDTFYDNFENEPILYGRSYTWLCYEVKIKRGRSNLLWDTGVFR
GQVYFKPQYHAEMCFLSWFCGNQLPAYKCFQITWFVSWTPCPDCVAKLAEFLSEHPN
25 VTLTISAARLYYYWERDYRRALCRLSQAGARVTIMDYEEFAYCWENFVYNEGQQF
MPWYKFDENYAFLHRTLKEILRYLMDPDTFTFNFFNNDPLVRRRQTYLCYEVERLD
NGTWVLMDQHMGFLCNEAKNLLCGFYGRHAELRFLDVPSLQLDPAQIYRTWFIWS
PCFSWGCAGEVRAFLQENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIMTY
DEFEYCWDTFVYRQGCPFQPWDGLEEHSQALSGRLRAILQNQGN (SEQ ID NO: 49)

30 (italic: nucleic acid editing domain)

Rat APOBEC-3B:

MQPQGLGPNAGMGPVCLGCSHRRPYSPIRNPLKKLYQQTFYFHFKNVRYAWGRKN
NFLCYEVNGMDCALPVPLRQGVFRKQGHIHAELCFIYWFHDKVLRVLSPMEEFKVT
WYMSWSPCSKCAEQVARFLAAHRNLSLAIFSSRLYYLRNPNYQQKLCRLIQEGVH

VAAMDLPEFKKCWNKFVDNDGQPFRPMRLRINF SYDCKLQEIFSRMNLLREDVF
YLQFNNSHRVKPVQNRYYRRKSYLCYQLERANGQEPLKGYLLYKKGEQHVEILFLE
KMRSMELSQVRITCYLTWSPCPNCARQLAAFKKDHPDLLIRIYTSL YFWRKKFQKG
LCTLWRSGIHVDVMDLPQFADCWTNFVN PQRPF RPWNELEKNSWRIQRLRRIKES

5 WGL (SEQ ID NO: 50)

Bovine APOBEC-3B:

MDGWEVAFRSGTVLKAGVLGVSMTEGWAGSGHPGQGACVWTPGTRNTMNL REV
LFKQQFGNQPRVPAPYYRRKTYLCYQLKQRNDLTLDRGCFRNKKQRHAERFIDKIN
SLDLNPSQSYKIICYITWSPCPNCANELVN FITRNNHLKLEIFASRLYFHWIKSF KMGL

10 QDLQNAGISVAVMTHTEFEDCWEQFVDNQS RPFQPWDKLEQYSASIRRRLQRILTAP
I (SEQ ID NO: 51)

Chimpanzee APOBEC-3B:

MNPQIRNPMEMW MYQRTFY YNFENEPI LYGRSYTWLCYEVKIRRGHSNLLWDTGVFR
GQMY SQPEHHAEMCFLSWFCGNQLSAYKCFQITWFVSWTPCPDCVAKLAKFLAEH

15 PNVTLTISAARLYYYWERDYRRALCRLSQAGARVKIMDDEEFAYCWENFVYNEGQP
FMPWYKFDDNYAFLHRTLKEIIRHLM DPDTFTFN FNN DPLVLRRHQ TYLCYEVERLD
NGTWVLMQDHMGFLCNEAKNLLCGFYGRHAELRF LDLVPSLQLDPAQIYRV TWFIS
WSPCF SWGCAGQVRAFLQENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIM
TYDEF EYCWDTFVYRQGCPFQPWDGLEEH SQALSGRLRAILQVRASSLCMVPHRPPP
20 PPQSPGPCLPLCSEPPLG SLLPTGRPAPSLPFLLTASF SF PPPASLPLPSLSLSPG HLP VP
SFHSLTSCSIQPPCSSRIRETEG WASVSKEGRDLG (SEQ ID NO: 52)

Human APOBEC-3C:

MNPQIRNPMKAMYPGT FYFQFKNLWEANDRNETWL CFTVEGIKRRSVVSWKTGVF
RNQVDSETHCHAERCFLSWFCDDILSPNTKYQVTWY TS WSPCPDCAGEVAEFLARHSN

25 VNLTIFTARLYYFQYPCYQEGLRSLSQEGVAVEIMDYEDFKYC WENFVYNDNEPFKP
WKGLKTNFRLLKRRRLRESLQ (SEQ ID NO: 53)

(italic: nucleic acid editing domain)

Gorilla APOBEC-3C

MNPQIRNPMKAMYPGT FYFQFKNLWEANDRNETWL CFTVEGIKRRSVVSWKTGVF

30 RNQVDSETHCHAERCFLSWECDDILSPNT NYQVTWY TS WSPCP ECAGEVAEFLARHSN
VNLTIFTARLYYFQDTDYQEGLRSLSQEGVAVKIMDYKDF KYC WENFVYNDDEPFK
PWKGLKYNFRFLKRRRLQEILE (SEQ ID NO: 54)

(italic: nucleic acid editing domain)

Human APOBEC-3A:

MEASPASGPRHLMRPHIFTSNFNNGIGRKTYLCYEVERLDNGTSVKMDQHRGFLH
NQAKNLLCGFYGR*HAE*LRFLDVPSLQLDPAQIYRVTFISWSPCFSGCAGEVRAFLQ
ENTHVRLRIFAARIYDYDPLYKEALQMLRDAGAQVSIMTYDEFKHCWDTFVDHQGC
PFQPWDGLDEHSQALSGLRAILQNQGN (SEQ ID NO: 55)

5 (italic: nucleic acid editing domain)

Rhesus macaque APOBEC-3A:

MDGSPASRPRHLMRPHIFTSNFNNDLSVRGRHQTYLCYEVERLDNGTWVPMDER
GFLCNKAKNVPCGDYGCHVELRFLCEVPSWQLDPAQTYRVTFISWSPCFRRGCAGQ
VRVFLQENKHVRLRIFAARIYDYDPLYQEALRTLDAQVSIMTYEEFKHCWDTF

10 VDRQGRPFQPWDGLDEHSQALSGLRAILQNQGN (SEQ ID NO: 56)

(italic: nucleic acid editing domain)

Bovine APOBEC-3A:

MDEYTFTENFNNQGWPSKTYLCYEMERLDGDATIPLDEYKGFVRNKGLDQPEKPC*H*
AELYFLGKIHSWNLDRNQHYRLTCFISWSPCYDCAQKLTTFLKENHISLHILASRIYTH

15 NRFGCHQSGLCELQAAGARITIMTFEDFKHCWETFVDHKGPQWPEGLNVKSQAL
CTELQAILKTQQN (SEQ ID NO: 57)

(italic: nucleic acid editing domain)

Human APOBEC-3H:

MALLTAETFRLQFNKRRRLRPYYPRKALLCYQLTPQNGSTPTRGYFENKKK*CHAEI*
20 *CFINEIKSMGLDETQCYQVTCYLWSPCSSCAWELVDFIKAHDHLNLGIFASRLYYHWC*
KPQQKGLRLLCGSQVPVEVMGFPKFADCWENFVDHEKPLSFNPYKMLEELDKNSRA
IKRRLERIKIPGVRAQGRYMDILCDAEV (SEQ ID NO: 58)

(italic: nucleic acid editing domain)

Rhesus macaque APOBEC-3H:

25 MALLTAKTFSLQFNKRRVNKPYYPRKALLCYQLTPQNGSTPTRGHLKNKKDHAE
IRFINKIKSMGLDETQCYQVTCYLWSPCPSCAGELVDFIKAHRHNLRIFASRLYYH
WRPNYQEGLLLCGSQVPVEVMGLPEFTDCWENFVDHEKPLSFNPSEKLEELDKNS
QAIKRRLERIKRSRSVDVLENGRLSQLGPVTPSSIRNSR (SEQ ID NO: 59)

Human APOBEC-3D:

30 MNPQIRNPMERMYRDTFYDNFENEPILYGRSYTWLCYEVKIKRGRSNLLWDTGVFR
GPVLPKRQSNHRQEYFRFENHAEMCFLSWFCGNRLPANRRFQITWFVSWNPCLPCVV
KVTKFLAEHPNVTLTISAARLYYYRDRDWWRVLLRLHKAGARVKIMDYEDFAYCW
ENFVCNEGQPFMPWYKFDDNYASLHRTLKEILRNPMEAMYPHIFYFHFKNLLKACG
RNESWLCTFMEVTKHSAVFRKRGVFRNQVDPETHCHAERCFLSWFCDDILSPNTNY

EVTWYTSWSPCPECAGEVAEFLARHSNVNLTIFTARLCYFWDTDYQEGLCSLSQEGAS
VKIMGYKDFVSCWKNFVYSDEPFKPWKGLQTNFRLKRRRLREILQ (SEQ ID NO:
60)

(italic: nucleic acid editing domain)

5 Human APOBEC-1:

MTSEKGPSTGDPTLRRRIEPWEFDVFYDPRELRK**EACLLYEIKWGMSRKIWRSSGKN**
TTNHVEVNFIKKFTSERDFHPSMCSITWFLSWSPCWECSQAIREFLSRHPGVTLVIV
ARLFWHMDQQNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQY
PPLWMMLYALELHCIILSLPPCLKISRRWQNHLTFFRLHLQNCHYQTIPPHILLATGLI

10 HPSVAWR (SEQ ID NO: 61)

Mouse APOBEC-1:

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELRK**EETCLLYEINWGGRHSVWRHTSQN**
TSNHVEVNLEKFTTERYFRPNTRCSITWFLSWSPCGECSRAITEFLSRHPYVTLFIYIA
RLYHHTDQRNRQGLRDLISSGVTIQIMTEQEYCYCWRNFVNYPSPSNEAYWPRYPHL

15 WVKLYVLELYCIILGLPPCLKILRRKQPQLTFFTITLQTCHYQRIPP~~H~~LLWATGLK
(SEQ ID NO: 62)

Rat APOBEC-1:

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELRK**EETCLLYEINWGGRHSIWRHTSQNT**
NKHVEVNIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVT~~L~~FIYIAR
LYHHADPRNRQGLRDLISSGVTIQIMTEQESGYCWRNFVNYS~~P~~SNEAHWPRYPHLW
VR~~LY~~VLELYCIILGLPPCLNILRRKQPQLTFFTIALQSCHYQRLPP~~H~~ILWATGLK (SEQ
ID NO: 63)

Human APOBEC-2:

MAQKEEA~~AA~~ATEAASQNGEDLENLDDPEKLKELIELPPFEIVTGERLPANFFKFQFRN
25 VEYSSGRNKTFLCYVVEAQGKGQVQASRGY~~E~~DEHAAAHAEEAFFNTILPAFDPA
LRYNVTWYVSSPCAACADRIIKTLSKTKNLRLLILVGRLFMWEEPEIQAALKKLKE
AGCKLRIMKPQDFEYVWQNFVEQEEGESKA~~F~~QPWEDIQENFLYYE~~K~~LA~~D~~ILK (SEQ
ID NO: 64)

Mouse APOBEC-2:

30 MAQKEEA~~AA~~APASQNGDDLENLEDPEKLKELIDLPPFEIVTGVRLPVNFFKFQFR
NVEYSSGRNKTFLCYVVEVQSKGGQAQATQGY~~E~~DEHAGAHAEEAFFNTILPAFD~~P~~
ALKYNVTWYVSSPCAACADRILTLSKTKNLRLLILVSRLFMWEEPEVQAALKKLKE
EAGCKLRIMKPQDFEYIWQNFVEQEEGESKA~~F~~EPWEDIQENFLYYE~~K~~LA~~D~~ILK (SEQ
ID NO: 65)

Rat APOBEC-2:

MAQKEEEAAEAAAAPASQNGDDLENLEDPEKLKELIDLPPFEIVTGVRLPVNFFKFQFR
NVEYSSGRNKTFLCYVVEAQSKGGQVQATQGYLEDEHAGAHAEAFFNTILPAFDP
ALKYNVTWYVSSSPCAACADRILKTLSKNLRLLILVSRLFMWEEPEVQAALKKLK

5 EAGCKLRIMKPQDFEYLWQNFVEQEEGESKAFEPWEDIQENFLYYEELADILK
(SEQ ID NO: 66)

Bovine APOBEC-2:

MAQKEEEAAAAAEPASQNGEEVENLEDPEKLKELIELPPFEIVTGERLPAHYFKFQFRN
VEYSSGRNKTFLCYVVEAQSKGGQVQASRGYLEDEHATNHAEEAFFNSIMPTFDPA

10 LRYMVTWYVSSSPCAACADRIVKTLKNLRLLILVGRLFMWEEPEIQAALRKLKE
AGCRLRIMKPQDFEYIWQNFVEQEEGESKAFEPWEDIQENFLYYEELADILK (SEQ
ID NO: 67)

Petromyzon marinus CDA1 (pmCDA1):

MTDAEYVRIHEKLDIYTFKKQFFNNKKSVSHRCYVLFELKRRGERRACFWGYAVNK
15 PQSGTERGIHAEIFSIRKVEEYLRDNPGQFTINWYSSWSPCADCAKILEWYNQELRG
NGHTLKIWACKLYYEKNARNQIGLWNLRDNGVGLNMVSEHYQCCRKIFIQSSHQNQ
LNENRWLEKTLKRAEKRRSELSFMIQVKILHTTKSPA (SEQ ID NO: 68)

Human APOBEC3G D316R D317R:

MKPHFRNTVERMYRDTFSYNFYNRPILSRRNTVWL CYEVTKGPSRPLDAKIFRGQ
20 VYSELKYHPEMRFFHWFSKWRKLHRDQEYEVTWYISWSPCTKTRDMATFLAEDP
KVTLTIFVARLYYFWDPDYQEALRSCLCQKRDGPRATMKFNYDEFQHCWSKFVYSQ
RELFEPWNNLPKYYILLHFMLGEILRHSMDPPTFTFNNEPWVRGRHETYL CYEVE
RMHNDTWVLLNQRRGFLCNQAPHKHGFLEG RHAELCFLDVIPFWKLDLDQDYRVT
CFTSWSPCFSCAQEMAKFISKKHVSLCIFTARIYRRQGRCQEGLRTLAEAGAKISFTY
25 SEFKHCWDTFVDHQGCPFQPWDGLDEHSQDLSGRLRAILQNQEN (SEQ ID NO: 69)

Human APOBEC3G chain A:

MDPPTFTFNNEPWGRHETYL CYEVERMHNDTWVLLNQRRGFLCNQAPHKHGF
LEGRHAELCFLDVIPFWKLDLDQDYRVT CFTSWSPCFSCAQEMAKFISKNKHVSLCIF
TARIYDDQGRCQEGLRTLAEAGAKISFTYSEFKHCWDTFVDHQGCPFQPWDGLD

30 EHSQDLSGRLRAILQ (SEQ ID NO: 70)

Human APOBEC3G chain A D120R D121R:

MDPPTFTFNNEPWVRGRHETYL CYEVERMHNDTWVLLNQRRGFLCNQAPHKHG
FLEG RHAELCFLDVIPFWKLDLDQDYRVT CFTSWSPCFSCAQEMAKFISKNKHVSLCI

FTARIYRRQGRCQEGLRTLAEAGAKISFMTYSEFKHCWDTFVDHQGCPFPWDGLD
EHSQDLSGRLRAILQ (SEQ ID NO: 71)

hAPOBEC-4 (*Homo sapiens*):

MEPIYEEYLANHGTIVKPYYWLSFSLDCSNCPYHIRTGEEARVSLTEFCQIFGFPYGT

5 FPQTKHLYELKTSSGSLVQKGHASSCTGNYIHPESMLFEMNGYLDASIYNND SIRH
IILYSNNSPCNEANHCCISKMYNFLITYPGITLSIYFSQLYHTEMDFPASAWNREALRS
LASLWPRVVLSPISGGIWHSLHSFISGVSGSHVFQPILTGRALADRHNAYEINAITGV
KPYFTDVLLQTKRNPNTKAQEAELEYPLNNAFPGQFFQMPSGQLQPNLPPDLRAPVV

FVLVPLRDLPPMHMGQNPNKPRNIVRHLNMPQMSFQETKDLGRLPTGRSVEIVEITE

10 QFASSKEADEKKKKKGKK (SEQ ID NO: 72)

mAPOBEC-4 (*Mus musculus*):

MDSLLMKQKKFLYHFKNVRWAKGRHETYLCYVVKRRDSATSCSLDFGHLRNKSGC

HVELLFLRYISDWLDLPGRCYRTWFTSWSPCYDCARHVAEFLRWNPNLSLRIFTAR
LYFCEDRKAEP EGLRRLH RAGVQIGIMTFKD YFYCWNTFVENRERTFKAW EGLHEN

15 SVRLTRQLRILLPLYEVDDLRAFRMLGF (SEQ ID NO: 73)

rAPOBEC-4 (*Rattus norvegicus*):

MEPLYEEYLTHSGTIVKPYYWLSVSLNCTNC PYHIRTGEEARVPYTEFHQTGFPWST
YPQTKHLYELRSSSGNLIQKGLASNCTGSHTHPESMLFERDGYLDLSIFHDSNIRHI
IILYSNNSPCDEANHCCISKMYNFLNYPEVTLSVFFSQLYHTENQFPTSAWNREALR

20 GLASLWPQVTL SAISGGIWQSILETFVSGISEGLTA VRPFTAGRTLTDRYNAYEINCIT
EVKPYFTDALHSWQKENQDQKVWAASENQPLHNTTPAQWQPDMSQDCRTPAVFM
LV PYRDLPIHVNPSPQPKRTVVRHLN TQLSASKVKALRKSPSGRPVKKEARKGS
TRSQEANETNKS KWKKQTLFIKS NICHLLEREQKKIGILSSWSV (SEQ ID NO: 74)

mfAPOBEC-4 (*Macaca fascicularis*):

25 MEPTYEEYL ANHGTIVKPYYWLSFSLDCSNCPYHIRTGEEARVSLTEFCQIFGFPYGT
TYPQTKHLYELKTSSGSLVQKGHASSCTGNYIHPESMLFEMNGYLDASIYNND SIRH
HILYCNSNP CNEANHCCISKVYNFLITYPGITLSIYFSQLYHTEMDFPASAWNREALRS
SLASLWPRVVLSPISGGIWHSLHSFISGVSGSHVFQPILTGRALTD RYNAYEINAITGV

VKPFFTDVLLHTKRNPNTKAQMALESYPLNNAFPGQSFQMTSGIPPD LRAPVV FVLL

30 PLRDLPPMHMGQDPN KPRNIIRHLNMPQMSFQETKDLERLPTRRSVETVEITERFASS
KQAEKTKKKKGKK (SEQ ID NO: 75)

pmCDA-1 (*Petromyzon marinus*):

MAGYECVRVSEKLD FDT FEFQFENLHYATERHRTYVIFDVKPQSAGGRS RRLWGYII
NNPNVCHAE LILMSMIDRHLESNPGVYAMTWYMSWSPCANCSSKLN PWLKNLLEE

QGHTLTMHFSRIYDRDREGDHRLRGLKHSNSFRMGVVGRAEVKECLAEYVEAS
RRTLTWLDTTESMAAKMRRKLFILVRCAGMRESGIPLHLFTLQTPLLSGRVVWR
V (SEQ ID NO: 76)

pmCDA-2 (*Petromyzon marinus*):

5 MELREVVDICALASCVRHEPLSRVAFLRCFAAPSQKPRGTVILFYVEGAGRVTGGH
AVNYNKQGTSIHAEVLLSAVRAALLRRRCEDGEATRGCTLHCYSTYSPCRDCVE
YIQEFGASTGVRVVIHCCRLYELDVNRRSEAEGLRSLSRLGRDFRLMGPRAIA
LLGGRLANTADGESGASGNAWVTETNVVEPLVDMTGFDEDLHAQVQRNKQIREA
YANYASAVSLMLGELHVDPDKFPFLAEFLAQTSVEPSGTPRETRGRPRGASSRGPEIG
10 RQRPADFERALGAYGLFLHPRIVSREADREEIKRDLIVVMRKHNYQGP (SEQ ID NO:
77)

pmCDA-5 (*Petromyzon marinus*):

MAGDENVRVSEKLDFTFQFENLHYATERHRTYVIFDVKPQSAGGRSRRLWGYII
NNPNVCHAELILMSMIDRHLESNPGVYAMTWYMSWSPCANCSSKLNWLKNLEE
15 QGHTLMMHFSRIYDRDREGDHRLRGLKHSNSFRMGVVGRAEVKECLAEYVEAS
RRTLTWLDTTESMAAKMRRKLFILVRCAGMRESGMPLHLFT (SEQ ID NO: 78)

yCD (*Saccharomyces cerevisiae*):

MVTGGMASKWDQKGMDIAYEEALGYKEGGVPIGGCLINNKDGSVLGRGHNMRF
QKGSATLHGEISTLENCGRLEGKVYKDTLYTTLSPCDMCTGAIIMYGIPRCVVGEN
20 VNFKSKEKYLQTRGHEVVVVDERCKKIMQFIDERPQDWFEDIGE (SEQ ID NO:
79)

rAPOBEC-1 (delta 177-186):

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELRKETCLLYEINWGGRHSIWRHTSQNT
NKHVEVNIEKFTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
25 LYHHADPRNRQGLRDLISSLGVTIQIMTEQESGYCWRNFVNYSPSNEAHWPRYPHLW
VRGLPPCLNILRRKQPQLTFTIALQSCHYQRLPPHILWATGLK (SEQ ID NO: 80)

rAPOBEC-1 (delta 202-213):

MSSETGPVAVDPTLRRRIEPHEFEVFFDPRELRKETCLLYEINWGGRHSIWRHTSQNT
NKHVEVNIEKFTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIAR
30 LYHHADPRNRQGLRDLISSLGVTIQIMTEQESGYCWRNFVNYSPSNEAHWPRYPHLW
VRLYVLELYCIILGLPPCLNILRRKQPQHYQRLPPHILWATGLK (SEQ ID NO: 81)

Mouse APOBEC-3:

MGPFCLGCSHRKCYSPIRNLISQETFKFHKNLGYAKGRKDFTLCYEVTRKDCDSPV
SLHHGVFKNDNIHAEICFLYWFHDKVLKVLSREEFKITWYMSWSPCFECAEQIVRFL

ATHHNLSDLIFSSRLYNVQDPETQQNLCRLVQEQAQVAAMDLYEFKKCWKKFVDN
 GGRRFRPKRLLTNFRYQDSKLQEILRPCYIPVPSSSSTLSNICLTKGLPETRFCVEG
 RRMDPLSEEEFYSQFYNQRVKHLCYYHRMKPYLCYQLEQFNGQAPLKGCLLSEKGK
 QHAEILFLDKIRSMELS^{QVTITCYLTWSPCPNC}AWQLAAFKRDRPDYLILHIYTSLRYFHW
 5 KRPFQKGLCSLWQSGILVDVMDLPQFTDCWTNFVNPKRPFWPWKGLEIISRTQRRL
 RRIKESWGLQDLVNDFGNLQLGPPMS (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

Adenosine Deaminase	Adenosine Deaminase Description
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

Adenosine Deaminase	Adenosine Deaminase Description
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	WMRHALTLLAK	<u>RARDE</u> REVPV	GAVLVLNNRV	I GEGWNRAIG
5					
	60	70	80	90	100
	LHD <u>P</u> TAHAEI	MALRQGGLV <u>M</u>	<u>QNY</u> RLIDATL	YVTFEPCVMC	AGAM <u>IHSRIG</u>
	110	120	130	140	150
	RVVFGVRNAK	TGAAGSLMDV	LHY <u>P</u> GMNHRV	E <u>I</u> <u>T</u> EGILADE	CAALL <u>C</u> YFFR
	160				
10	MPRQVFNA <u>Q</u> K	KAQSSTD	(SEQ ID NO: 83)		

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.

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 β 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 degron 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 degron sequence is controlled at least in part by the degron sequence. In some cases, a suitable degron is constitutive such that the degron exerts its influence on protein stability independent of experimental control (i.e., the degron is not drug inducible, temperature inducible, etc.) In some cases, the degron 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 degron is a temperature sensitive degron, 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 degron is a drug inducible degron, 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 degron is derived from the FKBP12 protein. The stability of the degron is controlled by the presence or absence of a small molecule that binds to the degron.

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 degron: 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 degron; Barbour et al., Biosci Rep. 2013 Jan 18;33(1).: Characterization of the bipartite degron 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 5 fluorescent protein (GFP)-based reporter protein; all of which are hereby incorporated in their entirety by reference).

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

10 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 degron 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 15 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 20 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 25 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. 30 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 5 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 10 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 15 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* 20 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, 25 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 30 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 10 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 15 (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) - 20 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 25 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 30 (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^{2+} -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 adiponectin 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 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 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 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.

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 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 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, adenylate cyclase 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 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
10 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
15 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 silence genes. In some embodiments, base editing results in altered protein function by
20 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
25 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 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 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 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 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 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:eaao4774 (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 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 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 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 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 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 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 embodiments, the CRISPR methods or systems described herein comprise a nucleobase 10 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 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 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 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 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, 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. 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.

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 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-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 exonuclease 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 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 amides, 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 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, 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, 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 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 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, 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 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.

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 1×10^3 cells will be administered, for example 5×10^3 cells, 1×10^4 cells, 5×10^4 cells, 1×10^5 cells, 1×10^6 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 µm 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-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 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 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 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 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 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-associated proteins, or RNA guides, can be delivered by various delivery systems such as 10 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.

Similar to wt AAV, recombinant AAV (rAAV) utilizes the *cis*-acting 145-bp ITRs to 30 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
embodiments, a portion or fragment of a fusion protein is fused to an intein and fused to an
30 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 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 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 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 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 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 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); Sommnerfelt *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).

Retroviral vectors, especially lentiviral vectors, can require polynucleotide sequences 5 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 10 into a target cell via a retroviral vector. In some cases, a Cas9 is of a size so as to allow 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 15 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 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 20 *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 (1994); Muzychka, 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 & Muzychka, PNAS 81:6466-6470 (1984); and Samulski *et al.*, J. Virol. 63:3822-3828 (1989).

A CRISPR system (e.g., including the Cas9 disclosed herein) described herein can 25 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 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-Dioleyloxy)propyl]N,N,N-trimethylammonium chloride	DOTMA	Cationic
1,2-Dioleyloxy-3-trimethylammonium-propane	DOTAP	Cationic
Dioctadecylamidoglycylspermine	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-Dioleyloxypropyl)-2,4,6-trimethylpyridinium	2Oc	Cationic
2,3-Dioleyloxy-N-[2(sperminecarboxamido-ethyl]-N,N-dimethyl-1-propanaminium trifluoroacetate	DOSPA	Cationic
1,2-Dioleyl-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
3β-[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-Diodeoxy-2-(6-carboxy-spermyl)-propylamide	DOSPER	Cationic
Dimethyloctadecylammonium bromide	DDAB	Cationic
Dioctadecylamidoglycylspermidin	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
Ethyldimyristoylphosphatidylcholine	EDMPC	Cationic
1,2-Distearyloxy-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-tetradecylaminopropionamidine	diC14-amidine	Cationic
Octadecenolyl[ethyl-2-heptadecenyl-3 hydroxyethyl]imidazolinium chloride	DOTIM	Cationic
N1 -Cholesteryloxycarbonyl-3,7-diazanonane-1,9-diamine	CDAN	Cationic
2-(3-[Bis(3-amino-propyl)-amino]propylamino)-N-ditetradecylcarbamoylme-ethyl-acetamide	RPR209120	Cationic
1,2-dilinoleyloxy-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(β -aminoester)	
Poly(4-hydroxy-L-proline ester)	PHP
Poly(allylamine)	
Poly(α -[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	

 Polymers Used for Gene Transfer

Polymer	Abbreviation
Dextran-spermine	D-SPM

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	(e.g., 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
Non-Viral	Herpes Simplex Virus	YES	Stable	NO	DNA
	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.*

10 *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).

20 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 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, 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 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 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, 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. 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 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 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 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, 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 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 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 µl Plus reagent). After 6 hours, the media is changed to antibiotic-free DMEM with 10% fetal 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 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 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* 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 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 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.

5 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/ (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.

30 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

5 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 poly anhydrides; (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.

10 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.

15 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 administrating 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, intraosseus, 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 ah, 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 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 ah, Gene Ther. 1999, 6: 15 1438-47). Positively charged lipids such as N-[*l*-(2,3-dioleyloxi)propyl]-N,N,N-trimethyl-amoniummethylsulfate, 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.

20

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 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 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 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 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 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 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 provided 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 (ScoCas9, 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 peruvensis*, ,
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 5
10 *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 peruvensis* 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-15 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 peruvensis, Lactobacillus fermentum strain AF15-40LB and Peptoniphilus sp. Marseille-P3761 bacteria

20 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 25 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, 30 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	GTGGCCCTGTGCCAGCCC (SEQ ID NO: 98)	TGG
guide 4	GTCCCCAAATATGTAGCTGTT (SEQ ID NO: 99)	TGG
guide 6	GCTCCCATCACATCAACCGG (SEQ ID NO: 100)	TGG

guide 7	GATGTCACCTCCAATGACTA (SEQ ID NO: 101)	GGG
guide 9	GTTGAAGATGAAGGCCAGAG (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 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

5 (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 10 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 15 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	GTGGCCCTGTGCCAGCCC (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	GTGCCAGAACAGGGGTGAC (SEQ ID NO: 116)	GGG

FIG. 5A shows a schematic diagram of constructs of WT *SirCas9* as well as *SirCas9*

5 (“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).

10 **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	AGAGAAAATGAAACTTCAA (SEQ ID NO: 121)	AAGCC
guide 6	CCAAACCCAACCTCCATCTAC (SEQ ID NO: 122)	CAGCC
guide 7	GGTCCTTGAATTGCAGTATC (SEQ ID NO: 123)	TAGCC
guide 8	GCATAGACTGCGGGCGGGC (SEQ ID NO: 124)	CAGCC
guide 9	GGAAACTGGAACACAAAGCA (SEQ ID NO: 125)	TAGAC
guide 10	GACAGCATGTGGTAATTTC (SEQ ID NO: 126)	CAGCC
guide 11	GCCCCCGGAAACTCTGTCCA (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	GAATCCTGCCATACACTTG (SEQ ID NO: 135)	AATAGC
guide 8	CTGCGGGCGGGCCAGCCTG (SEQ ID NO: 136)	AATAGC
guide 9	ACATTGTCAGAGGGACACAC (SEQ ID NO: 137)	TGTGGC
guide 10	AGCAACTCCAGTCCAAATA (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	GTCCCCAATATGTAGCTGTT (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	GCCAACACCAACCAGAACCTT (SEQ ID NO: 150)	GGG
guide 11	TGCTGCACACAGCAGGCCTT (SEQ ID NO: 151)	TGG
guide 12	GTGCCAGAACAGGGGTGAC (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	AATGAGAGAAAATGAAACTT (SEQ ID NO: 157)	TCAAA
guide 6	GGCCATCAAGGATGCCACG (SEQ ID NO: 158)	AGAAA
guide 7	AAATTGTCCAGCCCCATCTG (SEQ ID NO: 159)	TCAAA
guide 8	CCTGTAAAGGAAACTGGAAC (SEQ ID NO: 160)	ACAAA
guide 9	TACATGAAGCAACTCCAGTC (SEQ ID NO: 161)	CCAAA
guide 10	AAACTCCCCCACCCCCTT (SEQ ID NO: 162)	CCAAA
guide 11	GAGTTGGTTGGTGCTCAA (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	GTTGAAGATGAAGGCCAGAG (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
Amino Acid Sequence of Adenine Deaminase, TadA8.13m-nickase fused to the N-terminal of nickase <i>ScoCas9</i> (ABE-nScoCas9, D10A mutant)	
<u>MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPGAVLVLNNRVIGEGWNRAIGLHDP</u> <u>TAHAEIMALRQGLVMQNYRLYDATLYVTFEPVCVMCAGAMIHSR1GRVVFGVRNAKTGAAGSLM</u> <u>DVLHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESAT</u> <u>PESSGPKKKRKVGGKPYSIGLAIGTNSVGAVVTDDYKVPACKMKVLGNTDKQSIKKNLLGALL</u> <u>FDSGETAEATRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRGEHH</u> <u>PIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTD</u> <u>VQALFKDFVEEYDKTIEESHLEITVDALSILTEKVSKSSRLENLIAHYPTEKKNTLFGNLIAL</u> <u>SLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAILLSGILT</u> <u>DDNSTKAPLSASMVKRYEEHQDLKKLDFIKVNAPDQYNAIFKDKNKGYASYIESGVKQDEF</u> <u>YKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHYPLKEN</u> <u>QDKIEKILTFRIPYYVGPLARKGSRFWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDL</u> <u>YLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTDKLL</u> <u>NYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSFLDDKANEKTIEDIIQTTLFE</u> <u>DREMIRQRLQKYSDIFTKAQLKKLERRHYTGWGRLSYKLINGIRNKENKTIILDYLIIDDGYANR</u> <u>NFMQLINDDALSFKEEIARAQIIDDVDDIANVVHLPSPAIIKKGILQSVKIVDELVKVMGHNP</u>	

ANIIIEARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNICKNVENQQLQNDRLFLYYIQNGKD
MYTGETLDINNLSQYDIDHIIPQAFIKDNSDLNRVLTRSDKNRGKSDDVPSIEVVHEMKSFWSK
LLSVKLITQRKF DNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEDGDNKRR
IRNVKIITLKSNLVSNFRKEFELYKVREINDYHHAAHDAYLNAVVGNA LLKYPQLEPEFVYGEY
PKYNSYRSRKSA TEKFLFYSNILRFFKKEDIQTNE DGEIAWNKEKH KIILRKVL SYPQVNIVKK
TEEQTGGFSKESI LPKGESDKLIPRKT KNSYWDPKKYGGFDS PVVAYSILVFADVEKGKSKKLR
KVQDMVGITIMEKKRFEKNPVDFLEQRGYRNRLEKIIKLPKYSLFELNKRRLLASAKELQK
GNELVIPQRFTTLLYHSYRIEKDYPEHREYVEKHDEFKELLEYISVFSRKYVLADNNLT KIE
MLFSKNKDAEVSSLAKSFISLLTFTAFGAPAAFNFGENIDRKRYTSVTECLNATLIHQ SITGL
YETRIDLSKLGEDGKRPAATKKAGQAKKKGS YPYDVPDYA PYDVPDYA PYDVPDYA (SEQ
ID NO: 20)

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

MPAAKRVKLD TSEKGPSTGDPTLRRRIESWEFDVFYDPRELKETCLLYEIKWGMSRKIWRSSG
KNTTNHVEVNFIKKFTSERFHSSISCSITWFLSWPCWEC SQAIREFLSQHPGVTLVVIYVARL
FWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMMLYALELH
CIIISLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHPSVTWRLKSGGSSGGSS
GSETPGTSESATPESGGSSGGS PKKKRKV GGKPYSIGLAIGTNSVGAVVTDDYKVPACKMKV
LGNTDKQS IKKNLLGALLFD SGETA EATRLKRTARR YTRRKRNRLRYLQEIFTGEMNKVDENFF
QR LDDSFLVDEDKRGEHPIFGNIAAEV KYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMI
KFRGHFLYEGDLKAENTDVQALFKDFVEEYDKTIEESH LSEITVDALSILTEKVSKSSRLENLI
AHYPTEKKNTLFGNLIALS LD LHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLF
ASAKNLYDAILLSGILT VDDNSTKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDK
NKKGYASYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQE
MHAILRRQGEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRF AWA EYKADEKITPWNFDDIL
DKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQE
IFDHVFKENRKVTDKLLNYLNKEFEEFRIVNLTGLDKENKAFNSSLGTYHDLRKILDKSFLDD
KANEKTIEDIIQTTLFEDREMIRQRLQKYS DIFTKAQLKKLERRHYTGWGRLSYKLINGIRNK
ENKKTILDYLI DDGYANRNFMQLIN DDAL SFKEE IARAQI IDVDDIANVVHDLPGSPA IKKG
LQSVKIVDELVKVMGHNPANIIIEARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNICKNVEN
QQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSDLNRVLTRSDKNRGKS
DDVPSIEVVHEMKSFWSKL LSVKLITQRKF DNLTKAERGGLTEEDKAGFIKRQLVETRQITKH
AQILDERFNTEDGDNKRRIRNVKIITLKSNLVSNFRKEFELYKVREINDYHHAAHDAYLNAVVG

ALLLKYPQLEPEFVYGEYPKNSYRSRKSATEKFLFYSNILRFFKKEDIQTNEEDGEIAWNKEKH
 IKILRKVLSYPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKTNSYWDPKKYGGFDSPVVA
 YSILVFADVEKGKSKKLKVQDMVGITIMEKKRFEKNPVDFLEQRGYRNVRLEKIIKLPKYSLF
 ELENKRRRLLASAKELQKGNELVIQRFTTLLYHSYRIEKDYEPHREYVEKHDEFKELLEYI
 SVFSRKYVLADNNLTIEMLFSKNKDAEVSSLAKSFISLLTFTAFGAPAAFNFFGENIDRKRYT
 SVTECLNATLIHQSITGLYETRIDLSKLGEDGKRPAATKKAGQAKKKGSSGGGGSGGS TNLS
DIEKETGKQLVIQESILMLPEEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWA
LVIQDSNGENKIKMLSGGSGGSGGSTNLSDIEKETGKQLVIQESILMLPEEEVIGNKPESD
ILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLYDVPDYAYPYDVPDYAYP
 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).

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDPTAHAEIMALRQGGLVMQNYRLYDATLYVTTFEPVCVMCAGAMIHSR1GRVVFGVRNAKTGAAGSLMDVLHHPGMNRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESATPESSGPKKKRKVGAKNKDIRYSIGLAIGTNSVGWAVMDEHYELLKKGNHHMWGSRLFDAAEPAATTRASRSIRRYNKRERIRLLRDLLGDMVMEVDPTFFFIRLLNVSFLDEEDKQKNLGNDYKDNYNLFIEKDFNDKTYDKYPTIYHLRKECENKEADPRLIYLALHHIVKYRGNFLKEGQSFAKVYEDIEEKLDNTLKKFMSLNDLDNLFVDNDINSMITVLSKIYQRSKKADDLKIMNPTKEERAAYKEFTKALVGLKFNVSKMILAQEVKKDDKDIELDFSNVDYDSTVDGLQAEGEYIEFIMLHSINSWVELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPDKYNEVREDNPKLHNYLGYIKYPKNTPVEEFYEYIKRLLAKVDTGEAREILERIDLEKFMLKQNSRTNGSIPYQMQKDEMIQIIDNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFEDKQKERILPWNYDQIVDIDATAEGFIERMQNTGTYFPDKPVMAKNSLTVSKFEVLNELNKIRINGKLIPVETKKELLSDLFMMKNKTITDKKLKDWLVTHQYYDTNEELKIEGYQKDLQFSTSLAPWIDFTKIFGEINASNYQLIEKIIYDISIFEDKKKILKRRLKKVYQLDDLLVDKILKLNYTGWSRLSEKLLTGISKNSKETILSILENSNMNLMEIINDESLGFKQIIESNKKDIEGPFRYDEVKKLAGSPAIKRGIWQALLVVQEITKFMKHEPSHIYYIEFAREEQEKVRTESRIAKLQKIYKDLNQTKEDQLVYESIKKEDAKKKIDTDALYYYLQMGKSMYSGKPLDIKLSTYHIDHILPRSLIRTDFNEKDKRFINRQLVETRQIIKNVAINDHYTNTNVVTVRAELSHQFRERYKIYKNRDINDLHHADYIACILGQFIHQNFGNMDVNMIYGQYKNYKDVEHNNYGFILNSMNHIFNDDNSVIWDPSYIGKIKSCFCYKDVYVTKKLQNDAKLFDLTILPSDKNSENGVTKAKIPVNKYRKDVNKYGGFSGDAPIMLAIEADKGKHVRQVIAFPLRLKNY

DEERIKFIEKEKNLKNVKILTEVKKNQLILINHQYFFITGTNELVNATQLLSAKNTKNLFNLV
 DANKHNKLESIDDANFNEVIQELICKLQEPIYSRYNSIGKEFEDSYEKINAVTKQDKLYIEYL
 IAIMSAKATQGYIKPELAREIGTNGKNKGRIKSFTIDLNNKTTFISTSVTGLFSKKYKLGKRPA
TKKAGQAKKKGS YPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 6)

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

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRIVGEGWNRAIGLHDP
TAHAEIMALRQGGLVMQNYRLYDATLYVTFEPVCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLM
DVLHHPGMNHRVEITEGILADECALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESAT
PESSGPKKKRKVGSIINFQRRLMETQASNQLISSHLKGYPIKDYFVGLAIGTSSVGWAVTNKA
 YELLKFRSHKMWSRLFDEGESAVARRGFRSMRRLERKRLKLLEELFADAMAQVDPTFFMR
 LRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHHI
 LKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGKTKSDKAKAI
 EHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANVLGLNASLIDLFGSVEELEEDLKKLQI
 TGDTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPLTISESKVKAFNKHDDLAILK
 SLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKIVEGLSDSKDKEYIL
 SQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIMLEFRI
 PYVGPNTHHNVDNGGFAWAVRKASGRVTPWNFDDKIDREKSAAFIKNLTNKCTYLLGEDVL
 PKSSLLYSEFMILLNELNNVRIDGKPLEKVKEHLIEAVFKQDHKKMTKNRIEQFLKDNGYISET
 HKHEITGLGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGESKKMLRETLRKKFASCL
 DDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGTPETIIILMRNFSYNLMELLGDKFSFME
 RIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVDEVAHIKKALPARIFVEVTRSNK
 NEKKKKDSRQKRLSDLYAAIKDDVLLNGLNNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGR
 CAYTGEIIELSLLNTDNYDIDHIYPRSLTKDDSFDNVLCKRTANAQSDAYPISEEIQKTQKP
 FWTFLKQQGLISERKYERLTRITPLTADDLSGFIARQLVETNQSVKAATTLLRRLYPGVDVVFV
 KAENVTDFRHDDNNFIKVRSLNHHHHAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAK
 MFNYDVNCTNAKDGKAWDVTSMDTVKKMMDSDVRVTKRLLEQTGALADATIYKATVAGKAKD
 GAYIGMKTSSVFADVKYGGMTKIKNAYSIIIVQYTGKGEVIKEIVPLPIYLTNRNTTDQDLI
 NYVASIIIPQAKDISIIYGKLCINQLVKVNGFYYLGKTN SKFCIDNAIQVIVSNEWIPYLKVL
 EKFNNMRKDNLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLVSKLKMPIYQKMKGNAAE
 LSEKGYGLFKKMSLEEQSIIHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINE
 SITGLYSNEVTIVGKRPAATKKAGQAKKKGS YPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 10)

Amino Acid Sequence of Adenine Deaminase, TadA8.13m fused to the C-terminal of nickase VapCas9 (nVapCas9-ABE8, D38A mutant)

MPKKRKVSIIINFQRRGLMETQASNQLISSHLKGYPPIKDYFVG**L**AIGTSSVGWAVTNKAYELLK
FRSHKMWGSRLFDEGESAVARRGFRSMRRLERRKLRLKLLLELFADAMAQVDPTFFMRLRESK
YHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHHILKYRG
NFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGKTKSDKAKAIEHLD
TYIATDTVDTSSKTQKDQVKEDKKRLKAFANVLGLNASLIDLFGSVEELEDLKKLQITGDTY
DDKRDELAKAWSDEIYIIDDCKSVYDAIILSIKEPLTISESKVAKFNKHKKDDLAILKSLLKS
DRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEYILSQIEL
QILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKMLEFRIPIYYVG
PLNTHHNVDNGGFAWAVRKASGRVTWNFDDKIDREKSAAFIKNLTNKCTYLLGEDVLPKSSL
LYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTKNRIEQFLKDNGYISETHKHEI
TGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGESKKMLRETLRKKFASCLDDEAI
KKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGTPETIIILMRNFSYNLMELLGDKFSFMERIQEI
NAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVDEVAHIKKALPARIFVEVTRSNKNEKKK
KDSRQKRLSDLYAAIKDDVLLNGLNNEIFGELKSSLAKYDDAALRSKKLYLYTQMGRCAUTG
EIIELSLLNTDNYDIDHIYPRSLTKDDSFDNLVLCKRTANAQKSDAYPISEEIQKTQKPFWTFL
KQQGLISERKYERLTRITPLTADDLSGFIARQLVETNQSVKAATTLLRRLYPGVVVFVKAENV
TDFRHDDNNFIKVRSLNHHHHAKDAYLNIVVGNVYHERFTRNRAFFKKNGANRTYNLAKMFNYD
VNCTNAKDGKAWDVKTSMDTVKKMMDSDNRVTKRLLEQTGALADATIYKATVAGKAKDGAYIG
MKTKSVFADVSKYGGMTKIKNAYSIIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVAS
IIIPQAKDISIIYGKLCINQLVKVNGFYYYLGGKTNFKCIDNAIQVIVSNEWIPYLKVLEKFNN
MRKDNKDLKANVVSTRALDNKHTIEVRIVEEKNIEFFDYLVSKLKMPIYQKMKGNKAAELSEKG
YGLFKKMSLEEQSIIHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGL
YSNEVTIV**KRPAATKKAGQAKKKSGSETPGTSESATPESSGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLN**
NNRVIGEGWNRAIGLHDPTAHAEIMALRQGLVMQNYRLYDATLYVTFEP
VMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADEC
ALLCRFFRM PRRVFNAQKKAQSSTDPAAKRVKLDGS** YPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO:**

11)

Amino Acid Sequence of Cytidine Deaminase, ppAPOBEC1 fused to the N-terminal of nickase VapCas9 (CBE-nVapCas9, D38A mutant)

MPAAKRVKLD TSEKG PSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGM SRKIWRSSG
KNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWSPCWEC SQAIREFLSQHPGVTLVIYVARL
FWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMMLYALELH
CII SLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHP SVTWRLKSGGSSGGSS
GSETPGTSESATPESSGGSSGGS **PKKKRKV**GS I INFQRRGLMETQASNQLISSHLKG YPIKDYF
VGLA**IGTSSVGWAVTNKAYELLKFRSHKMWGSRLFDEGE SAVARRGFRSMRRLERKRLKL
EELFADAMAQVDPTFFMRLRESKYHYEDKTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSEL
MKSGTDDIRKLFLAVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAISSI
GQILMEAGTKSDKAKAIEHLVDTYIATDVTDTSSKTQKDQVKEDKKRLKA FANLVGLNASLI
DLFGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPLTIS
ESKVKA FNKH KDDLAILKSLLKSDRSIYNTMFKVDEKGLHN VHYIKQGRTEETSCNREDFYKY
TKKIVEGLSDSKDKEYI LQSIELQILLPLQRIKDNGVI PYQLHLEELKAI LAKCGPKFPFLNEV
ADGFSVAEKLIKMLEFRI PYYVGPLNTHHNVDNGGFAWAVRKASGRVT PWNFDDKIDREKSAAA
FIKNLTNKCTYLLGEDVLPKSSL LYSEFMLNELNNVRIDGKPLEKVVKHEH LIEAVFKQDHKKM
TKNRIEQFLKDNGYI SETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFG
ESKKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGT PETIIILMR
NFSYNLMELLGDKFS FMERIQEINAKLTEGQIVNPHDIIDDLALS PAVKRAVWQALRIVDEVAH
IKKALPARIFVEVTRSNKNEKKKDSRQKRLSDLYAAIKDDVLLNGLNNEIFGELKSSLA KYD
DAALRSKKLYLYYTQMGRCA YTGEIIELSLLNTDNYDIDHIYPRSLTKDDSFDNLVLCKRTANA
QKS DAYPISEEIQKTQKFWTFLKQQGLISERKYERLTRITPLTADDLSGFIARQLVETNQSVK
AATTLLRRLYPGVVVFVKAENVTDFRH DNNFIKVRSLNHHHAKDAYLNIVVGNVYHERFTRN
FRAFFKKNGANRTYNLAKMFNYDVNCTNAKDGKAWDVKTSMDTVKKMMDSDVRVTKRLLEQTG
ALADATIYKATVAGKAKDGAYIGMKT KSSVFADVSKYGGMTKIKNAYSIIIVQYTGKKGEVIKEI
VPLPIYL TNRNTTDQDLINYVASIIIPQAKDISIIYGKLCINQLVKVNGFYYLGGKTN SKFCID
NAIQVIVSNEWIPYLKVLEKFNNMRKDNKDLKANVVSTRALDNKHTIEVRIVEKNIEFFDYL V
SKLKMPIYQKMGNKAAELSEKGYGLFKKMSLEEQS IHLIELLNLLTNQKTTFEVKPLGITAS R
STVGSKISNQDEFKVINESITGLYSNEVTIVG **KRPAATKKAGQAKKKGSSGGGGSGS** TNLS
DIEKETGKQLVIQESTILM PEEVEEVIGNKPESDILVHTAYDESTDENVMLL TD APEYKPWA
LVIQDSNGENKIKMLSGGSGGSGGSTNLS **DIIIEKETGKQLVIQESTILM PEEVEEVIGNKPESD**
ILVHTAYDESTDENVMLL TD APEYKPWA **LVIQDSNGENKIKML** YPYDVPDYAYPYDVPDYAY
(SEQ ID NO: 12)**

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

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRIVIGEGWNRAIGLHDP
TAHAEIMALRQGGGLVMQNYRLYDATLYVTFEPVCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLM
DVLHHPGMNHRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESAT
PESSGPKKKRKVGTKVDYYIGLAIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAAER
RGQRGARRRLDRKKERLSLLQDFFAEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKD
FKDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIFEGQKFDTKSSFENSLN
ELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKKELKSVIGDTKFLKAVSAIMIGSSQKLV
DLFENPEDFDDSAIKSVDFSTTSFDDKYSDYELALGDKIALVNILKEIYDSSILENLLKEADKS
KDGNKYISNAFKVKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNNDNYVSYTKSSISNNKRV
KANKFTDQEAFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNGVIP
YQLKLMELNKILENQSKHHFELNVSDEYGSCDKIASIMEFRIPYYVGPLNPNSKYAWIKKQKD
SEITPWNFKDVVDLDSSREEFIDSЛИГРСТЛКДЕКВЛПКАСЛЛНЕМВЛНЛКЛНДЛПИ
TEEMKKIFDQLFKTRKVKVTLKAVANLLKEFNINGEILLSGTGDFKQGLNSYNDFKAIVGDK
VDSDDYRDKIEEИКЛIVLYGDDKSYLQKKIKAGYGKYFTDSEИККМАГЛNYKDWGRLSKLLT
GLEGANKITGERGSIIHMREYNLNLMELMSASFTFTEEIQKLNPVDDRKLSYEMVDELYLSPS
VKRMLWQSLRIVDEИКНИГТDSKKIFIEMARGKEEVKARKESRKНQLLKFYKDГKKAFISEIG
EERYSYLLSEIEGEEENKFRWDNLYLYTQLGRCMYSLEPIDISELSSKNIYDQDHИYPKSKIY
DDSIENRVLVKDЛNSKKGNSYPIPDEILNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDDE
LVQFISRQIVETRQATKETANLLKICKNSEIVYSAENASRFRQEFDIVKCRAVNDLHHMHDA
YINIIVGVNVYNTKFTKDPMNФVKKQEKARSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIK
RIRKELEGTNYRFTRMNYIESGALFNATLQRKNKGSRPLDKGPKSSIEKYGGYTINKACFAV
LDIKSKNKIERKLMPVEREИYAKQKNDKKLSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLЛKID
GSYYFITGGSDKTLELRSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDII
VNKFKTSVFKKSFLNLФQDDKИENIDFKFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKS
DYGRLLSSKTNNIGNYQEFKIINQSITGLFENEVDLLKLGKRPAATKKAGQAKKKGS YPYDVPD
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)

MPKKKRKVTKVДYYIGLAIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAEERRGQRG
ARRRLDRKKERLSLLQDFFAEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKDFKDKN
FHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIFEGQKFDTKSSFENSLNELKVH
LNDEYGLDLEFDNENLINILTDPKLNKTAKKKELKSVIGDTKFLKAVSAIMIGSSQKLVDLFEN
PEDFDDSAIKSVDFSTTSFDDKYSDYELALGDKIALVNILKEIYDSSILENLLKEADKSKDGNK

YISNAFKVKYKNGQDLKEFKRLVRQYHKSAYFDIFRSEKVNDNYVSYTKSSISNNKRVKANKF
 TDQEAFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNGVI PYQLKL
 MELNKILENQSKHEFLNVSDEYGSVCDKIASIMEFRIPIYYVGPLNPNSKYAWIKKQKDSEITP
 WNFKDVVLDSSREEFIDSЛИГРСТЛКДЕКВЛПКАСЛЛНЕМЕЛНЛКЛНДЛПИЕЕМК
 KKIFDQLFKTRKKVTLKAVANLLKEFNINGEILLSGTDGDFKQGLNSYNDFAIVGDKVDSDD
 YRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGKYFTDSEIKKMAGLNYKDWGRLSKKLLTGLEGA
 NKITGERGSIIHFMREYNLNLMELMSASFTFTEEIQKLPVDDRKLSEMVDELYLSPSVKRML
 WQSLRIVDEIKNIMGTDSKKIFIEMARGKEEVKARKESRKNQLLKFYKDGKKAFISEIGEERYS
 YLLSEIEGEEENKFRWDNLYLYYTQLGRCMYSLEPIDISELSSKNIYDQDHIPKSKIYDDSIE
 NRVLVKKDLNSKKGNSYPIPDEILNKNCYAYWKILYDKGLIGQKKYTRLTRRTGFTDELVQFI
 SRQIVETRQATKETANLLKTICKNSEIVYNSKAENASRFRQEFDIVKCRAVNDLHHMHDAYINII
 VGNVYNTKFTKDPMNFKVKKQEKARSYNLENMFYDVKRGGYTAWIADDEKGTVKNASIKRIRKE
 LEGTNYRFTRMNYIESGALFNATLQRKNKGSRPLDKGPKSSIEKYGGYTNINKACFAVLDIKS
 KNKIERKLMPVEREIYAKQKNDKKSDEIFSKYLKDRFGIEDYRVVYPVVKMRTLLKIDGSYYF
 ITGGSDKTLELRSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFK
 TSVFKKSFLNLQDDKNIENIDFKFKSMDFKEKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGR
 SSKTNIGNYQEFKIINQSITGLFENEVDILKLKRPAATKKAGQAKKKSGSETPGTSESATPE
SSGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIDGEWNRRAIGLHDPTAHAEIMA
LRQGGLVMQNYRLYDATLYVTFEPVCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHPGM
NHRVEITEGILADECALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPDVPDYAYPY

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)

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFYDPRELKETCLLYEIKWGMSRKIWRSSG
KNTTNHVEVNFIKKFTSERFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIYVARL
FHWMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPLWMMLYALELH
CIILSLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHPSVTWRLKSGGSSGGSS
GSETPGTSESATPESSGGSSGGS**PKKKRKV**GTKVKDYYIGLAIGTSSVGWAVTDEAYNVLKFNS
 KKMWGVRLFDDAKTAEERRGQRGARRRLDRKKERLSSLQDFFAEEVAKVDPNFFLRLDNSDLYM
 EDKDQKLKSKYTLFNDKDFKDNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHF
 IFEGQKFDTKSSFENSNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKKELKSVIGDT
 KFLKAVSAIMIGSSQKLVDLFENPEDFDDSAIKSVDSTTSFDDKYSDYELALGDKIALVNILK
 EIYDSSILENLLKEADSKDGNKYISNAFKVKYKNGQDLKEFKRLVRQYHKSAYFDIFRSEKV

NDNYVSYTKSSISNNKRVKANKFTDQEAFYKFAKKHLETIKYKINKVNNGSKADLELIDGMLRDM
 EFKNFMPKIKSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDETYGSVCDKIASIMEFRIPIYY
 VGPLNPNSKYAWIKKQKDSEITPWNFKDVVDLDSSREEFIDSЛИGRCCTLKDEKVLPKASLLYN
 EYMLNELNNLKLNDLPITEEMKKIFDQLFKTRKKVTLKAVANLLKEFNINGEILLSGTGD
 FKQGLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGKYFTDSEIKK
 MAGLNYKDWGRSLSKLLTGLEGANKITGERGSIIHFREYNLNLMELMSASFTFTEEIQKLNPV
 DDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIEMARGKEEVKARKESRKN
 QLLKFYKDGKKAFISEIGEERYSYLLSEIEGEENEKFRWDNLYLYYTQLGRCMYSLEPIDISEL
 SSKNIYDQDHIPKSKIYDDSIENRVLVKDKLNSKKGNSYPIPDEILNKNCYAYWKILYDKGLI
 GQKKYTRLTRRTGFTDELVQFISRQIVETRQATKETANLLKICKNSEIVYSAENASRFRQE
 FDIVKCRAVNDLHHMHDAYINIIVGVNVNTKFTKDPMNFKQQEKARSYNLENMFKYDVKRGGY
 TAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRKNKSRPLDKGPKS
 SIEKYGGYTNINKACFAVLDIKSKNKIERKLMPVEREIYAKQNDKLSDEIFSKYLDRFGIE
 DYRVVYPVVKMRTLLKIDGSYYFITGGSDKTLELRSLAQQLILPKKNEWAIKQIDKSSENDYLT
 ERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLQDDKNIENIDFKFKSMDFKECKTLLMLVK
 AIRASGVRQDLKSIDLKSDYGRLLSKTNNIGNYQEFKIINQSITGLFENEVDLLKLGKRPAATK
 KAGQAKKKKGSSGGSGGGS **TNLSDIIIEKETGKQLVIQESIILMPEEEVIGNKPESDILVH**
TAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSAGGSTNLSDIIIEKETGKQ
LVIQESIILMPEEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGEN
KIKMLYPDVDPDYAYPYDVDPDYAY (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)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDPT
AHAETMALRQGGLVMQNYRLYDATLYVTFEPVCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDV
LHHPGMNHRVEITEGILADECAALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESATPES
SGPKKKRKVGKEYHIGLAIGTSSIGWAVTDQFKLMRIKGKTAIGVRLFEEGKTAERRTFRTTR
 RRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEDPAKNQAFIGKLLFPDLLKKNERGY
 PTLIKMRDELPEQRAHYPVTNIYKLREAMINEDRQFDLREVYLVAVHHIVKYRGHFLNNASVDKF
 KVGRIDFDKSFNVLNEAYEELQNCGEGSFTIEPSKVEKIGQLLDTKMRKLDQKAVAKLLEVKVA
 DKEETKRNKQIATAMSKLVLGYKADFATVAMANGNEWKIDLSSETSEDEIEKFREELSDAQNDIL
 TEITSLFSQIMLNEIVPNGMSISESMMDRYWTHERQLAEVKEYLATQPASARKEFDQVYNKYIGQ
 APKEKGFDLEKGLKKILSKKENWEIDEALKAGDFLPKQRTSANGVIPHQMHQQELDRIIEKQAK
 YYPWLATENPATGERDRHQAKYELDQLVSFRIPYYVGPLVTPEVQKATSGAKFAWAKRKEDEIT

PWNLWDKIDRAESAEEAFIKRMTVKDTYLLNEDVLPANSLLYQKYNVLNELNNVRVNGRRLSFGIK
QDIYTELFKKKKTVKAGDVASLMAKTRGVNKPSVEGLSDPKFNSNLATYLDLKSIVGDKVDDN
RYQMDLENIIEWRSVFEDGEIFADKLTEVEWLTDQRSALVKRYKGWGRLSKKLLTGIVDENGQ
RIIDLMWNTDQNFMQIVNQPVFKEQIDQLNQKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQV
VRVVEDIVKAVGNAPKSISIEFARNEGNKGEITRSRRTQLQKLFEDQAHELVKDTSLTEELEKAP
DLSDRYYFYFTQGGKDMYTGDPINFDEISTKYDIDHILPQSFKVDDSLDNRVLVSAENNKKSDR
VPAKLYAAKMKPYWNQLLQGLITQRKFENLTMDVDQTICKYRSLGFGVKRQLVETRQVIKLTANIL
GSMYQEAGTDIETRAGLTQLREEFDLPLKVREVNDYHHAVDAYLTTAGQYLNRRYPKLRFFV
YGEYMKFKHGSDLKLRNFNFFHELMEGDKSQGKVVVDQQTGEPLITTRDEVADYFDWVINLKVMLIS
NETYEETGKYFDASHESSSLYLKNQNKKSKLVVPLKNKLQPEYYGAYTGITQGYMVLKLLDKKG
GFGVYRIPRYAADIILNKCHDEVAYRNKIAEIISSDPRAPKSFEVVPVRLKGTFLVDGEEKFILS
SYRYKVNATQLILPVSDIKLIQDNFKALKKLNVEMQTKKLIEIYDNILRQVDKYYKLYDINKFRA
KLHDGRSKFVELDDFGQDASKEKVIIKILRGLHFGSDLQNLKEIGFGTPLGQFQVSEAGIRLSN
TAFIIIFKSPTGLFNRKLYLKNLGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVP
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)

MPKKKRKVGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAERRTFRTTR
RRLKRRKWRHLHYLDEIFAPHLQEVDENFLRRLQSNIHPPEDPAKNQAFIGKLLFPDILKKNERG
YPTLIKMRDEL PVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAHVHIVKYRGHFLNNASVD
KFKVGRIDFDKSFNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLDQKAVAKLLEV
KVADKEETKRNKQIATAMSKVLGYKADFATVAMANGNEWKIDLSSETSEDEIEKFREELSDAQ
NDILTEITSLSQIMLNEIVPNGMSISESMMDRYWTHERQLAEVKEYLATQPASARKEFDQVYN
KYIGQAPKEKGFDLEKGLKKILSKKENWEIDELLKAGDFLPKQRTSANGVIPHQMHQQELDRI
IEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPYYVGPLVTPEVQKATSGAKFAWAKR
KEDGEITPWNLWDKIDRAESAEEAFIKRMTVKDTYLLNEDVLPANSLLYQKYNVLNELNNVRVNG
RRLSVGIKQDIYTELFKKKKTVKAGDVASLMAKTRGVNKPSVEGLSDPKFNSNLATYLDLKS
IVGDKVDDNRYQMDLENIIEWRSVFEDGEIFADKLTEVEWLTDQRSALVKRYKGWGRLSKKL
LTGIVDENGQRIIDL MWNTDQNFMQIVNQPVFKEQIDQLNQKAITNDGMTLRERVESVLDDAYT
SPQNKKAIWQVVRVVEDIVKAVGNAPKSISIEFARNEGNKGEITRSRRTQLQKLFEDQAHELVK
DTSLTEELEKAPDLSDRYYFYFTQGGKDMYTGDPINFDEISTKYDIDHILPQSFKVDDSLDNRV
LVSRAENNKKSDRVPACKLYAAKMKPYWNQLLQGLITQRKFENLTMDVDQTICKYRSLGFGVKRQL

VETRQVIKLTANILGSMYQEAGTDIIETRAGLTQLREFDLPKVREVNDYHHAVDAYLTTFAG
 QYLNRRYPKLRSFFVYGEYMKFKHGSDLKLRNFNFFHELMEGDKSQGVVDQQTGELITTRDEV
 ADYFDWVINLKVMLISNETYEETGKYFDASHESSSLYLNQNKKSKLVVPLKNKLQPEYYGAYT
 GITQGYMVILKLLDKGGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKSFEVVVP
 RVLKGTFLVDGEEKFILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLNVEMQTKKLIEIYDN
 ILRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEVIIKILRGLHFGSDLQNLKEIG
 FGTTPLGQFQVSEAGIRLSNTAFTIFKSPTGLFNRKLYLKNLKRPAATKKAGQAKKKSGSETP
GTSESATPESSGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHD
PTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSL
MDVLHHPGMNHRVEITEGILADECALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPY
 DVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 89)

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

MPAAKRVKLDTSEKG PSTGDPTLRRRIESWEFDV FYDPRELKETCLLYEIKWGMSRKIWRSSG
KNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWPCWECSQAIREFLSQHPGVTLVIYVARL
FWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMMLYALELH
CIIISLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHPSVTWRLKSGGSSGGSS
GSETPGTSESATPESSGGSSGGSPKKKRKVGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTA
 IGVRLFEEGKTAERRTFRTT RRLKRRKWRLHYLDEIFAPHLQEVDENFLRRLKQSNIH PEDP
 AKNQAFIGKLLPDLLKKNERGYPTLI KMRDEL PVEQRAHYPVTNIYKLREAMINEDRQFDLRE
 VYLAVH HIVKYRGHFLNNASVDKF KVGRIDFDKS FNVLNEAYEELQNGEGSFTIEPSKVEKIGQ
 LLLDTKMRKLD RQKAVAKLLEV KVADKEETKR NKQI ATAMS KLV LGYKADFATVAMANGNEWKI
 DLSSETSEDEIEKFREELS DAQNDILTEITS LFSQIMLNEIVPNGMSI SESMMDRYWTHE RQLA
 EVKEYLATQPASARKEFDQVYNKYIGQAPKEKGFDLEKGLKKILSKKENWEIDE LLKAGDFLP
 KQRTSANGVIPHQMHQQELDR II EKQAKYY PWLATENPATGERDRHQAKYELDQLVSFRIPYYV
 GPLVTPEVQKATSGAKFAWAKR KEDGE IT PWNLWDKIDRAESA EAFIKRMTVKDTYLLNEDVLP
 ANSLLYQKYNVLN E LNNVRVNGRRLSVGIKQDIYTEL FK KKKTVKAGDV ASLVMAKTRGVNKPS
 VEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENII EWR SVFEDGE IFADKLTEVEWL T
 DEQR SALV KKRYKG WGR LS KLLT GIVDENGQRIIDL MWNTDQNF MQIVNQPVFKEQIDQLNQK
 AITNDGMTL RERVESV LDDAYTSPQNKAIWQVVRVVEDIVKAVGNAPKSISIEFARNEG NKGE
 ITRSR RTQLQKL FEDQ AHEL VKD TS LTEE LEKA PDSL DRY YFY FT QGGKDMYT GDP INF D EIST
 KYDIDHILPQS FVK DDSLD NRVL VSRA ENN KKS DRVPAK LYAA KMKP YWNQ LLKQ GLIT QRK FE
 NL TM DV DQTI KYRSLGFV KRQL VETRQVI KLTANILGSMYQEAGTDIIETRAGLTQLREEFDL

PKVREVNDYHHAVDAYLTTFAGQYLNRRYPKLRSFFVYGEYMKFKHGSDLKLRNFNFFHELMEG
DKSQGVVDQQTGEPLITRDEVADYFDWVINLKVMLISNETYEETGKYFDASHESSSLYLNQN
KKSKLVVPLKNKLQPEYYGAYTGITQGYMVLKLLDKGGFGVYRIPRYAADILNKCHDEVAYR
NKIAEIISSDPRAPKSFEVVVPRVLKGTFLVGEEKFILSSYRYKVNATQLILPVSDIKLIQDN
FKALKKLNVEMQTKKLIEIYDNILRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEK
VIKILRGLHFSDLQNLKEIGFTTPLGQFQVSEAGIRLSNTAFIIFKSPTGLFNRKLYLKNL
GKRPAATKKAGQAKKKGSSGGSGGS TNLSDIIIEKETGKQLVIQESTILMLPEEEVIGNK
PESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSGGSGGSTNLSDI
IEKETGKQLVIQESTILMLPEEEVIGNK PESDILVHTAYDESTDENVMLLTSDAPEYKPWALV
IQDSNGENKIKML YPYDVPDYAYPYDVPDYAY (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)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDP
TAHAEIMALRQGLVMQNYRLYDATLYVTFEPVCVMCAGAMIHSRIGRUVFGVRNAKTGAAGSLM
DVLHHPGMNHRVEITEGILADECALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPGTSESAT
PESSGPKKKRKVGEKKTNYTIGLAIGTDSVGWAVVKDDLELVKKRMKVLGNTEINYIKKNILWGS
LLFESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFRLNESFLVPEDKA
FSKNPIFGTLGEDKTYKTYPTIYHLRQHLADSEEKADVRЛИYLALAHMIKYRGHFLIEGKLDTEH
IAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDGRANKQS
PMMAFITLIVGKKAKFKAANLEEEISLNLTDDSYDENLEILLNTIGSDFADLFDAQRVYNAV
ELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKKDLKG
YAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVHLAEMEA
ILDRQENYYPWLKENREKIISLLTFRIPYYVGPLADGQSEFAWLERKSDEKIKPWNFSDVVDLDR
SAEKFIEQLIGRDTLPDEYVLPKKSLIYQKYMVFNELTKIAYLDERQKRMNLSSVEKKEIFET
LFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMMDDPDNE
EEIVKILTVFEDRKMRKQLEKYKERLSPEQIKELAKKHYTGWGRSLSKLLVGIRDKETQKT
DYLVEDDNHSGGRQHNRNLMQLINDDRSLFKKTIaelQMIDPSADLYAQVQEIA
GSPAIKKGI
LLGLKIVDEIIRVMGEK PENIVIEMARENQTTARGKALSKRREAKIKEGLAALGSSILKENLPG
NADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHIIPQSFTVDNSLDNLVLTNS
SQNRGN
KKDDVPSLEVNRQLAYWRSLKDAGLMTQRKFDNLTKAMRGGLTDKDRERFIQRQLVETRQITK
NVAKLLDMRLNDKDEAGNKIRETNIVLLKSAMASEFRKMFRPLYKVRELNDYHHAHDAYLNA
AINLLALYPYMADDFVYGEFRYKKPQAEKATYEKLQRQWNLIKRFGEKQLFTPDHEDCWNK
ERD

IKTIKKVVMGYRQVNVVKAEERTGMLFKETINGKTNKGSRIPIKKDLDPSKYGGYIEEKMAYYA
VISYEDKKKKPGKTIVGISIMDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIYPDGRRRYI
TGATKTTKGKVELQKANQIAMEQDLVNFIYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHY
TRRFLDAETNINRLRRIYEEEKKDPVDIEALVASFIELLKLTSAAGAPADFIFMGEAISRRRYN
SMTGLFDGQVIYQSLTGLYETRMRFEDKRPAATKKAGQAKKKGS YPYDVPDYAYPYDVPDYA
YPYDVPDYA (SEQ ID NO: 91)

Amino Acid Sequence of Adenine Deaminase, TadA8.13m-nickase fused to the C-terminal of nickase PmaCas9 (nPmaCas9-ABE, D12A mutant)

MPKKKRKVEKKNTYTI~~G~~LAIGTDSVGWAVVKDDLELVKKRMKVLGNTE~~N~~YIKKNLWGSSLFES
GQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFFRLNESFLVPEDKAFSKNPIF
GTLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIAINE
NLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLDIGRANKQS PMMAF
ITLIVGKKAKFKAAFNLEEEISLNLTDDSYDENLEILLNTIGSDFADLFDH~~A~~QRVYN~~V~~NAVELAGI
LSGDVKNT~~H~~AKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKKDLGYAGYI
DGNMS QDS FYKFVKDQLKEVPGSEKFLDS IEKEDFLRKQRSFYNGVIPNQVHAE~~M~~AILDRQE
NYYPWLKENREKI ISLLTFRIPYYVGPLADGQSEFAWLERKSDEKIKPWNFS DVVDLDRSAEKF
IEQLIGRDTYLPDEYVLPKKSLIYQKYMVFNELT~~K~~IAYLDERQKRMNLSSVEKKEIFETLFKKR
SKVTEQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMMDPDNEDLMEEIVK
ILTVFEDRKMRKQLEKYKERLSPEQIKE~~L~~AKKHYTGWR~~L~~SKLLVGIRD~~K~~E~~T~~QK~~T~~ILDYLV~~E~~
DDNHSGGRQH~~L~~NRNLMQLINDR~~L~~SFK~~K~~TI~~A~~ELQ~~M~~IDPSADLYAQVQE~~I~~AGSPAIKKGILLGLK
IVDEIIRVMGEK~~P~~ENIVIEMARENQTTARGKALS~~R~~REAKIKEGLAALGSSLLKENLPGNADLS
QRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHIIPQSFTVDNSLDNLVLTNSSQRGNKKDDV
PSLEVVRQLAYWRS~~L~~KDAGLMTQRKFDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKL
LDMRLNDKKDEAGNKIRETNIVLLKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAAIAINLL
ALYPYMADD~~F~~VYGEFRYKKPQA~~E~~KATYEKL~~R~~QWNLI~~K~~R~~F~~GEKQLFTP DHEDCWNKERDIKT~~I~~K
KVMGYRQVNVVKAEERTGMLFKETINGKTNKGSRIPIKKDLDPSKYGGYIEEKMAYYAVISYE
DKKKKPGKTIVGISIMDKKEFEYDSISYLGKLGFSNPVVQIILKNYSLIYPDGRRRYITGATK
TTKGKVELQKANQIAMEQDLVNFIYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFL
DAETNINRLRRIYEEEKKDPVDIEALVASFIELLKLTSAAGAPADFIFMGEAISRRRYNSMTGL
FDGQVIYQSLTGLYETRMRFEDKRPAATKKAGQAKKKGSSETPGTSESATPESSGSEVEFSHE
YWMRHALTLAKRARDEREVPVGAVLVLNNR~~V~~IGEGWNRAIGLHDPTAHAEIMALRQGGLVMQNY
RLYDATLYVT~~F~~EP~~C~~VMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGIL

ADECAALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGS YPYDVPDYAYPYDVPDYAYPYDVA
PDY (SEQ ID NO: 92)

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

MPAAKRVKLDTSEKGPGTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMRSRKIWRSSG
KNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWSPCWEC SQAIREFLSQHPGVTLVIYVARL
FWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPLWMMLYALELH
CIILSLPPCLKISRRWQNHLAFFRLHQNCHYQTIPPHILLATGLIHPSVTWRLKSGGSSGGSS
GSETPGTSESATPESSGGSSGGSPKKKRKVGEKKTNYTIGLAIGTDSVGWAVVKDDLELVKKRM
KVLGNTEINYIKKNLWGSLLFESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDEN
FFFRLNESFLVPEDKAFSKNPIFGTLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAH
MIKYRGHFLIEGKLDTEHIAINENLEQFFESYNALFSEEPIELKEELIAIENILREKNSRTVK
EKRITSFLKDGRANKQSPMMAFITLIVGKKAKFKAAFNL EEEISLNLTDDSYDENLEILLNTI
GSDFADLF DHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQ
YDMTFVAPKDAQKKDLGYAGYIDGNMSQDSFYKFVQDQLKEVPGSEKFLDSIEKEDFLRKQRS
FYNGVIPNQVHLAEMEAILD RQENYYPWLKENREKIISLLTFRIPIYYVGPLADGQSEFAWLERK
SDEKIKPWNFS DVVDLDRSAEKFIEQLIGRDTYL PDEYVLPKKSLIYQKYMVFNELTKIAYLDE
RQKRMNLSSVEKKEIFETLFKKRSKVTEKQLVKKFENYLQIDNPTIFGIEDAFNADYSTYVELA
KVPGMKSMMDDPDNE DLMEEIVKILTVFEDRKMRKQLEKYKERLSPEQIKE LAKKHYTGWGRL
SKKLLVGIRDKETQKTILDYLV EDDNHSGGRQHLNRNLMQLINDDR LSFKKTIAELQMIDPSAD
LYAQVQEIA GSPA IKKGILLGLKIVDEII RVMG EKPENIVI EMAREN QTTARGKAL SKRREAKI
KEGLAALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHII PQSFTV
DNSLDNLVLTNSSQNRGNKKDDVPSLEV VNRQLAYWRS LKDAGLMTQRKFDNLTKAMRGGLTDK
DRERFIQRQLVETRQITK NVAKL LD MRLND KKDEAGNKIRETNIVLLKSAMASEFRKMFR LYKV
RELNDYHHAHDAYLNAIAINLLALYPYMADD FVYGEFRYKKPQAEKATYEKL RQWNLI KRGF
EKQLFTP DHEDCNKERDIKTIKKVMGYRQNVVKKAEERTGMLFKETINGKTNKGSRIPIKKD
LDPSKYGGYIEEKMAYYAVISYEDKKKKPGKTIVGISIMDKKEFEYDSISYLGKLGFSNPVVQI
ILKNYSLIAYPDGRRYITGATTTKGKVELQKANQIAMEQDLVNFIYHLKNYDEISHPESYAF
VQSHTDYFDRLFDSIEHYTRRF LDAETNINRLRRIYEEKKDPVDIEALVASFIELLKLT SAG
APADFI FMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMRFEDGKRPAATKKAGQAKKKGS
SGGSGGSGGSTNLSDIIIEKETGKQLVIQESIMLPEEEVIGNKPESDILVHTAYDESTDEN
MLLTSDAPEYKPWALVIQDSNGENKIKMLSGGS GGSGSTNLSDIIIEKETGKQLVIQESIMLP

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)

MPAAKRVKLDTNLSDIIEKETGKQLVIQESTILMLPEEVEEVIGNKPESDILVHTAYDESTDEN
MLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSGGGSTNLSSDIIEKETGKQLVIQESTILMLP
EEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGS
SGGSPKKKRKVEKKNTYTIGLAIGTDSVGAVVKDDLELVKKRMKVLGNTENYIKKNLWGSLL
FESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFRFLNESFLVPEDKA
FSKNPIFGTLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIA
INENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDGRANKQS
PMMAFITLIVGKKAKFKAAFNL
EEEISLNLTDDSYDENLEILLNTIGSDFADLF
DHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAP
KDAQKKDLKGYA
GYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVH
LAEMEAILD
RQE
NYPW
LKENRE
KIISLLTFR
IYPL
ADGQSE
FAWLRKS
DEKIK
PWNF
SDVV
DLRSA
EKF
IEQLIGRD
TYLP
DEYVLP
PKSLIY
QKYM
VFNELTK
IAYL
DERQK
RMNLSS
VEK
KEIFETLF
KKRS
KVTE
QVKFF
ENYL
QIDN
PTIF
GIEDAF
NADY
STY
VELAK
VPGM
KSMM
DDPD
NEDLM
EE
IVK
ILT
TVF
EDR
KMRR
KQLE
KYKER
LSPE
QIK
ELAK
KH
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IRD
KET
QKT
ILD
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LVED
DH
NNH
SGGR
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NLM
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DDRL
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QNG
KDI
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FDRL
SQY
DED
HII
PQS
FTV
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LDN
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NKK
DDV
PS
LEV
VN
RQL
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WRS
LKD
AGL
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KNV
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KRPAATKKAGQAKKK
GSSGGSSGGSSGSETPGTSESAT
PESSGGSSGGS
TSEKG
PSTGD
PTLRR
RRIES
WEFDV
FYDPRE
LRKET
CLLYE
I
KWGM
SRK
IWR
SS
GKNT
TTNH
VEVN
FIKK
FTSERR
FHSS
ISCS
ITWFL
SWSPC
WECSQAIREF
FLSQHPGV
TLVIY
VAR
LF
WHMD
QRNR
QGLR
DVL
VNSG
VTIQ
IMRA
SEYY
H
CWRNF
VNYP
PGDEAH
WPQY
PPLW
MM
LYALEL

HCIIILSLPPCLKISRRWQNHLAFFRLHLQNCYQTIPPHILLATGLIHPSVTWRYPYDVPDYAY
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

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

MPKKKRKVGMGKPYSIGLDIGTNSVGWAVVTDDYKVPACKMKVLGNTDKQSIIKKNLLGALLFDS
GETAEATRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDDFLVDEDKRGEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQA
LFKDFVEEYDKTIEEHLSEITVDALSILTEKVSKSSRLENLIAHYPTEKKNTLFGNLIALSLD
LHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAILLSGILTVDNN
STKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVQDEFYKY

LKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHYPFLKENQDK
IEKILTFRIPYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLP
EEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTDKLLNYL
NKEFEFRIVNLTGLDENKAFNSSLGTYHDLRKILDKSFLDDKANEKTIEDIIQTTLFEDRE
MIRQRLQKYSIFTKAQLKKLERHYTGWGRLSYKLINGIRNKENKTIIDYLIDDGYANRNF
QLINDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPAIKKGILQSVKIVDELVKVMGHNPANI
IEMARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNINKVENQQLQNDRLFLYYIQNGKDMYT
GETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKSFWSKLLS
VKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRN
VKIITLKSNLVSNFRKEFELYKREINDYHHADAYLNAVGNALLKYPQLEPEFVYGEYPKY
NSYRSRKSATEKFLFYSNILRFFKKEDIQTNEGEIAWNKEHKIKILRKVLSYPQVNIVKKTEE
QTGGFSKESILPKGESDKLIPRKTKNSYWDPKKYGGFMQPVVAYSILVFADVEKGKSKKLRKVQ
DMVGITIMEKKRFEKNPVDFLEQRGYRNVRLEKIIKLPKYSLELENKRRLLASAKFLQKGNE
LVIPIQRFTTLLYHSYRIEKDYPEHREYVEKHDEFKELLEYISVFSRKYVLADNNLTKEMLF
SKNKDAEVSSLAKSFISLLTFTAFGAPRAFNFFGENIARKEYRSVTECLNATLIHQSITGLYET
RIDLSKLGEDGEGADKRTADGSEFESPKKRKV (SEQ ID NO: 95)

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

MPKKKRKVGMGKPYSIGLDIGTNSVGWAVTDDYKVPACKMKVLGNTDKQS IKKNLLGALLFDS
GETAEATRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFVDEDKRGEHHP
GNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQA
LFKDFVEEYDKTIEESHLEITVDALSIYTEKVSKSSRLENLIAHYPTEKKNTLFGNLIALSLD
LHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAILLSGILTVDNN
STKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEFYKY
LKGILLKINGSGDFLDKIDREDFLRKQRTFDNGIIPHQIHLQEMHAILRRQGEHYPFLKENQDK
IEKILTFRIPYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLP
EEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTDKLLNYL
NKEFEFRIVNLTGLDENKAFNSSLGTYHDLRKILDKSFLDDKANEKTIEDIIQTTLFEDRE
MIRQRLQKYSIFTKAQLKKLERLHYTGWGRLSYKLINGIRNKENKTIIDYLIDDGYANRNF
QLINDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPAIKKGILQSVKIVDELVKVMGHNPANI
IEMARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNINKVENQQLQNDRLFLYYIQNGKDMYT

```
GETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKSFWSKLLS  
VKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRN  
VKIITLKSNLVSNFRKEFELYKvreINDYHHAHDAYLNAVGNALLLKYPQLEPEFVYGEYPKY  
NSYRSRKSATEKFLFYSNILRFFKKEDIQTNEDGEIAWNKEKHIKILRKVLSYPQVNIVKKTEE  
QTGGFSKESILPKGESDKLIPRKTNSYWDPKKYGGFMQPVVAYSILVFADVEKGKSKKLRKVQ  
DMVGITIMEKKRFEKNPVDFLEQRGYRNVRLEKIIKLPKYSLELENKRRLLASAKFLQKGNE  
LVIPIQRFTTLLYHSYRIEKDYPEHREYVEHKDEFKELLEYISVFSRKYVLADNNLTKEMLF  
SKNKDAEVSSLAKSFISLLTFTAGAPRAFNFFGENIARKEYRSVTECLNATLIHQSITGLYET  
RIDLSKLGEDGEGADKRTADGSEFESPKKRKV (SEQ ID NO: 96)
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NLS (bold italics)

Linker (bold underlined)

5 ScoCas9-NGC variants were used to target a genomic locus that was randomly 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 variants recognized an exemplary NGC 3' PAM sequence, AGC. Cells were harvested 72 hours post-transfection and total DNA was extracted.

15 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 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)

MSEVEFSHEYWMRHALTLAKRARDEREVPGAVLVLNNRVIGEGWNRAIGLHDPTAHAEIMALR
QGGLVMQNYRLYDATLYSTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNH
RVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDSGGSSGGSSGSETPGTSESATPESS
GGSSGGSGKPYSIGLAIGTNSVGWAVVTDDYKVPACKMKVLGNTDKQSIKKNLLGALLFDGET
AEATRLKRTARRRYTRRKNRRLYIQLQEIFTGEMNKVDENFFQRLDDFLVDEDKRGEHHPIFGNI
AAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFK
DFVEEYDKTIEESHlseitvdalsiltekvsksrrlenliahyptekntlfgnlialslldhp
NFKTNFQLSEDAKLQFSKDTYEDLEGFLGEVGDEYADLFASAKNLYDAILSGILTVDNSTK
APLSASMVKRYEEHQDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEFYKYLKG
ILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHYPFLKENQDKIEK
ILTFRIPYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLPEEK
VLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTDKLLNYLNKE
FEEFRIVNLTGDKENKAFNSSLGTYHDLRKILDKSFLDDKANEKTIEDIIQTTLFEDREMIR
QRLOQKYSDFITKAQLKKLERRHYTGWGRLSYKLINGIRNKENKTIIDYLIDDGYANRNFMQLI
NDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPAICKGILQSVKIVDELVKVMGHNPANIIIE
MARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNIKNVENQQLQNDRLFYYIQNGKDMYTGET
LDINNLSQYDIDHIIPQAFIKDNSLDNRVILTRSDFKNRGKSDDVPSIEVVHEMKSFWSKLLSVKL
ITQRKF DNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEDGDNKRRIRNVKI
ITLKSNLVSNFRKEFELYKREINDYHHAHDAYLNAVGNALLKYPQLEPEFVYGEYPKNSY
RSRKSATEKFLFYSNILRFFKKEDIQTNEDEGIAWNKEKHIKILRKVLSPQVNIVKKTEEQTG
GFSKESILPKGESDKLIPRKTNSYWDPKKYGGFMQPVVAYSILVFADVEKGSKKLRKVQDMV
GITIMEKKRFEKNPVDFLEQRGYRNRLEKIIKLPKYSLELENKRRLLASAKFLQKGNELVI
PQRFTTLLYHSYRIEKDYPEHREYVEKHDEFKELLEYISVFSRKYVLADNNLTKIEMLFSKN
KDAEVSSLAKSFISLLTFTAFGAPRAFNFFGENIARKEYRSVTECLNATLIHQSTITGLYETRID
LSKLGEDGEEGADKRTADGSEFESPKKKRKV (SEQ ID NO: 98)

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

MSEVEFSHEYWMRHALTLAKRARDEREVPGAVLVLNNRVIGEGWNRAIGLHDPTAHAEIMALR

*QGGLVMQNYRLYDATLYSTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNH
RVEITEGILADECALLCRFFRMPRRVFNAQKKAQSSTDSGGSSGGSSGSETPGTSESATPESS*
GGSSGGS*GKPYSIGLAIGTNSVGWAVTDDYKVPACKMKVLGNTDKQS IKKNLLGALLFDGET
AEATRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRDDSFLVDEDKRGEHHPIFGNI
AAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFK
DFVEEYDKTIEESHLEITVDALSILTEKVSKSSRLENLIAHYPTEKKNTLFGNLIALSDLHP
NFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAILLSGILTVDNSTK
APLSASMVKRYEEHQDKLKKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEFYKYLKG
ILLKINGSGDFLDKIDREDFLRKQRTFDNGIIPHQIHLQEMHAILRRQGEHYPFLKENQDKIEK
ILTFRIPYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLPEEK
VLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVTDKLLNYLNKE
FEEFRIVNLTGLDENKAFNSSLGTYHDLRKILDKSFLDDKANEKTIEDIIQTTLFEDREMIR
QRLQKYSDFTKAQLKKLERLHYTGWRSLSYKLINGIRNKENKKTILDYLIDDGYANRNFMLI
NDDALSFKEEIARAQIIDVDDIANVVHDLPSPAICKGILQSVKIVDELVKVMGHNPANIIIE
MARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNICKVENQQLQNDRLFLYYIQNGKDMYTGET
LDINNLSQYDIDHIIIPQAFIKDNSLDNRVLTRSDKNRGKSDDVPSIEVVHEMKSFWSKLLSVKL
ITQRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEDGDNKRRIRNVKI
ITLKSNLVSNFRKEFELYKREINDYHHADAYLNAVGNALLKYPQLEPEFVYGEYPKYN SY
RSRKSATEKFLFYSNILRFFKKEDIQTNEEDGEIAWNKEKHICILRKVLSPQVNIVKKTEEQTG
GFSKESILPKGESDKLIPRKTKNSYWDPKKYGGFMQPVVAYSILVFADVEKGKSKLRKVQDMV
GITIMEKKRFEKNPVDFLEQRGYNRVLEKIIKLPKYSLELENKRRLLASAKFLQKGNELVI
PQRFTTLLYHSYRIEKDYPEHREYVEKHDEFKELLEYISVFSRKYVLADNNLTKIEMLSKN
KDAEVSSLAKSFISLLTFTAFGAPRAFNFFGENIARKEYRSVTECLNATLIHQSITGLYETRID
LSKLGEDGEEGADKRTADGSEFESPKKRKV (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 above Description, but rather is as set forth in the following claims.

10

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

MGKPYSIGLDIGTNSVGAVVTDDYKVPACKMKVLGNTDKQSIKKNLLGALLFDGETAEAT
RLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSFVDEDKRGEHHPIFGNIAA

10 EVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFK
DFVEEYDKTIEESHLSEITVDALSILTEKVSKSSRLENLIAHYPTEKKNTLFGNLIALSLDL
HPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAILLSGILTVD
NSTKAPLSASMVKRYEEHQDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEF
YKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHYPLK

15 ENQDKIEKILTFRIPYYVGPLARKGSRFAWEYKADEKITPWNFDDILDKEKSAEKFITRMT
LNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKVYVNEQGEAKFFDTNMKQEIFDHVFKENRKV
TKDKLLNYLNKEFEEFRIVNLTGLDENKAFCNSSLGTYHDLRKILDKSFLDDKANEKTIEDI
IQTTLFEDREMIRQRLQKYSIFTKAQLKKLERRHYTGWGRLSYKLINGIRNKENKKTILD
YLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPAICKGILQSVKI

20 VDELVKVMGHNPANIIEEEARENQTTDKGRRNSQQRLLQDSILKNLDNPVNICKNVENQQLQ
NDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRSDKNRGKSDD
VPSIEVVHEMKSFWSKLLSVKLITQRKF DNLTKAERGGLTEEDKAGFIKRQLVETRQITKH
AQILDERFNTEFDGNKRRIRNVKIIITLKSNLVSNFRKEFELYKVREINDYHHAHDAYLNAV
GNALLLKYPQLEPEFVYGEYPKYN SYRSRKSATEKFLFYSNILRFFKKEDIQTNEDGEIAWN

25 KEKHKILRKVLSPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKTKNSYWDPKKYGGF
DSPVVAYSILVFADVEKGSKKLRKVQDMVGITIMEKKRFEKNPVDFLEQRGYRNRLEKII
KLPKYSILFELENKRRLLASAKELQKGNELVIPQRFTLLYHSYRIEKDYPEHREYVEKHK
DEFKELLEYISVFSRKYVLADNNLTKIEMLFSKNKDAEVSSLAKSFISLITFTA FGAPA AFN
FFGENIDRKRYTSVTECLNATLIHQ SITGLYETRIDLSKLGED (**SEQ ID NO: 1**).

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

MAKNKDIRYSIGLDIGTNSVGWAVMDEHYELLKKGNHHMWGSRLFDAAEPAATRASRSIRR
RYNKRERIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDQKQNLGNDYKDNYNLFIEKDF
NDKTYYDKYPTIYHLRKELCENKEADPRLIYLALHHIVKYRGNFLKEGQSFAKVYEDIEEK
LDNTLKKFMSLNLDNLFVDNDINSMITVLSKIYQRSKKADDLKIMNPTKEERAAYKEFTK
5 ALVGLKFNVSKMILAQEVKKDDKDIELDFSNDYDSTVDGLQAELEYIEFIEMHSINSWV
ELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPDKYNEVFREDNPKLHNYLGYIKY
PKNTPVEEFYEYIKRLLAKVDTGEAREILERIDLEKFMLQNSRTNGSIPYQMOKDEMIIQII
DNQSVYYPQLKENREKLISILEFRIPYYFGPLNTHSEFAWIKKFEDKQKERILPWNYDQIVD
IDATAEGFIERMQNTGTYPDKPVMAKNSLTISKFEVNLNELNKIRINGKLIPVETKSELLSD
10 LFMKNKTITDKKLKDWLVTHQYYDTNEELKIEGYQKDLQFSTSAPWIDFTKIFGEINASNY
QLIEKIIYDISIFEDKKILKRRLKVVYQLDDLLVDKILKLNWTGWSRLSEKLLTGKSKNSK
ETILSILENSNMNLMEIINDES LGFKQIEESNKKDIEGPFRYDEVKLAGSPAIKRGIWQA
LLVVQEITKFMKHEPSHIYIEFAREEQEKVRTESRIAKLQKIYKDLNLQTKEDQLVYESLKK
EDAOKKIDTDALYLYYLQMGKSMYSGKPLDIDKLSTYHIDHILPRSLIKDDSLDNRVLVLPK
15 ENEWKLDSETVPFEIRNKMMGFWQKLHENGMSNKKFFSLIRTDFNEKDKRFINRQLVETR
QIIKNVAVIINDHYTNVVTVRAELSHQFRERYKIYKNRDLNLDHHADAYIACILGQFIH
QNFGNMDVNMIYGQYKKNYKKDVQEHNNYGFILNSMNHIHFNDDNSVIWDPSYIGKIKSCFC
YKDVTVKKLEQNDAKLFDLTILPSDKNSENGVTAKIPVNKYRDKVNKYGGFSGDAPIMLA
IEADKGKKHVRQVIAFPLRLKNYNDEERIKFIEKEKNLKNVKILTEVKKNQLILINHQYFFI
20 TGTNELVNATQLKLSAKNTKNLFNLVDANKHNLESIDANFNEVIQELICKLQEPIYSRYN
SIGKEFEDSYEKINAUTKQDKLYIEYLIAIMSAKATQGYIKPELAREIGTNGKNKGRIKSF
TIDLNKTTFISTSVTGLFSKKYKL (SEQ ID NO: 4).

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

25 MSIINFQRRLMETQASNQLISSHLKGYPKDYFVGLDIGTSSVGWAVTNKAYELLKFRSHK
MWGSRLFDEGESAVARRGFRSMRRLERKRLRLKLLLEELFADAMAQVDPTFFMRLRESKYHY
EDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHHILKYRGN
FLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGKTKSDKAKAIEHLV
DTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEEDLKKLQITG
30 DTYDDKRDELAKAWSDEIYIIDCKSVYDAIILLSIKEPLTISESKVKA FNKHKKDDLAILK
SLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEY
ILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKML
EFRIPYYVGPLNTHHNVNDNGFAWAVRKASGRVT PWNFDDKIDREKSAAAFIKNLTKCTYL

LGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTKNRIEQFLK
DNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGESKKMLRE
TLRKKFASCLDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGTPETIIILMRNFSYNL
MELLGDKFSFMERIQUEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVDEVAHIKKA

5 LPARI FVEVTRSNKNEKKKDSRQKRLSDLYAAIKDDVLLNGLNNEIFGELKSSLAKYDDA
ALRSKKLYLYTQMGRCAVTGEIIELSLLNTDNYDIDHIYPRSLTKDDSFDNLVLCKRTANA
QKS DAYPISEEIQKTQKPFWTFLKQQGLISERKYERLTRITPLTADDLSGFIARQLVETNQS
VKAATTLLRRLYPGVDVVFVKAENVTDFRHDDNNFIKVRSLNHHHAKDAYLNIVVGNVYHER
FTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKGKAADVKTSMDTVKKMMDSDNDVRTKR
10 LLEQTGALADATIYKATVAGKAKDGAYIGMKTSSVFADVKYGGMTKIKNAYSIIIVQYTGK
KGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISIIYGKLCINQLVKVNNGFYYLG
GKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNNMRKDNLKANVVSTRALDNKHTIEVRIV
EEKNIEFFDYLVSKLKMPIYQKMGNKAAELSEKGYGLFKKMSLEEQSIHЛИELLNLTNQK
TTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIV (SEQ ID NO: 8).

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

MTVKDYYIGLDIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAEERRGQRGARRRL
DRKKERLSLLQDFFAEEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKDFKDKNFHK
KYPTIHLLMDLIEDDSKKDIRLVLYACHYLLKNRGHFIFEGQKFDTKSSFENSNELKVHL
20 NDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQKLVDLFE
NPEDFDDSAIKSVDFSTTSFDDKYSDYELALGDKIALVNILKEIYDSSILENLLKEADSKD
GNKYISNAFKVKKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNDNYSYTKSSISNNKRV
KANKFTDQEAFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNGV
IPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRIPYYVGPLNPNSKYAWIK
25 KQKDSEITPWNFKDVVDLDSREEFIDSILIGRCTYLKDEKVLPKASLLYNEYMVNLNELNNLK
LNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTDGDFKQGLNSYND
FKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGKYFTDSEIKKMAGLNYKD
WGRLSKKLLTGLEGANKITGERGSIIFMREYNLNLMELMSASFTFTEIQLNPVDDRKLS
YEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSKKIFIEMARGKEEVKARKESRKNQLLK
30 FYKDGKKAFISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYQQLGRCMYSLEPIDISELSS
KNIYDQDHYPKSKIYDDSIENRVLVKKDLNSKKGNSYPIPDEILNKNCYAYWKILYDKGLI
GQKKYTRLTRRTGFTDELVQFISRQIVETRQATKETANLLKICKNSEIVYASKENASRFR
QEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKDPMNFVKQEKARSYNLENMFKYDVK

RGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRKNKGSRPLK
DKGPKSIEKYGGYNINKACFAVLDIJKSKNKKIERKLMPVEREYAKQKNDKLSDEIFSKY
LKDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITGGSDKTLELRSALQLILPKKNEWAIKQID
KSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLFQDDKIENIDFKFKSMDF
5 KEKCKTLLMLVKAIRASGVRQDLKSIDLKSDYGRLOSSKTNNIGNYQEFKIINQSITGLFENE
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

MKEYHIGLDIGTSSIGWAVTD SQFKLMRIKGKTAIGVRLFE EGKTAERRTFRTT RRRLKRR
10 KWRLHYLDEIFAPHLQEVDENFLRRLKQSNIHPEDPAKNQAFIGKLLFPDLLKKNERGYPTL
IKMRDEL PVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLA VHHIVKYRGHFLNNASVDKF
KVGRIDFDKS FNVLNEAYEEELQN GEGSFTIEPSKVEKIGQLLLDTKMRKLDRQKAVAKLLEV
KVADKEETKRNKQIATAMSKLVLYKADFATVAMANGNEWKIDLSSETSEDEIEKFREELSD
AQNDILTEITS LFSQIMLNEIVPNGMSISESMMDRYW THERQLAEVKEYLATQPASARKEFD
15 QVYNKYIGQAPKEKGFDLEKGLKKILSKKENWKEIDELLKAGDFLPKQRTSANGVI PHQMHQ
QELDRIIEKQAKYYPWLATE NPATGERDRHQAKYELDQLVSFRIPYYVGPLVTPEVQKATSG
AKFAWAKRKEDGEITPWNLWDKIDRAESA EAFIKRMTVKDTYLLNEDVLPANSLLYQKYNVL
NELNNVRVNGRRLSVG IKQDIYTEL FK KKKTVKAGDV ASLVMAKTRGVN KPSVEGLSDPKKF
NSNLATYLDLKSI VGD KVDDNRYQMDLEN IIEWRSV FEDGEI FADKL TEVEWL TDE QRSALV
20 KKRYKGWGR LS KKL LTGIVDENGQ RIIDL MWNTD QNF MQIVN QPV FKEQ IDQLN QKAIT NDG
MTLRERVESV LDDAYTSPQNK KAI WQV VRVVEDIVKAVGNAPKSISIEFARNEG NKGEITRS
RRTQLQKL FEDQA HELV KDT SLTEE LEKAP DLS DR YYFY FTQGGKDMY TGDP INF DEIST KY
DIDHILPQS FVK DDS LDNRV LVSRAENN KKSDRV PAKLYAAKM KPYWNQ LLQGLIT QRK F
NLTMDVDQT IKYRSLGFV KRQLVETRQVI KLTAN ILGSMYQEAGTDIIETRAGLTKQLREEF
25 DLPKVREVNDYH AHDAYLTT FAGQ YLN RRY PKLRS FFVYGE YM KFKHGSDLKLRN FNFFHE
LMEGDKS QGKVVDQQT GELIT TRDEVADY FDW VINV LKVML ISNET YEET GK YFDASHES SSL
YLKNQNKSKLVVPLKNKLQPEYYGAYTG ITQGYM VILKLLDKGGFGVYRIPRYAADI LNK
CHDEVAYRN KIAE IISSD PRAPKS FEVV VPRV LKGTF LVDGEEK FILSSYRYKVNATQLI LP
VSDIKLIQDNF KALKKL NVEM QT KKLIEI YDNILRQVDKYYKLYDINKFRAKLHDGRSKFVE
30 LDDFGQDASKEK VI IKILRGLHFGSDLQNLKEIGFGTTPLGQFQVSEAGIRLSNTAFIIFKS
PTGLFNRKLYLK NL (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

MEKKTNYTIGLDIGTDSVGWAVVKDDLELVKKRMKVLGNTETNYIKKNLWGSLLFESGQTAK
DRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFRLNESFLVPEDKAFSKNPIFGTL

5 GEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIAINEN
LEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKDGRANKQS PMMA

FITLIVGKKAKFKAAFNLEEEISLNLTDDSYDENLEILLNTIGSDFADLFDAQRVYNAVEL
AGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKKDLKG

YAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVH LAEME
10 AILDRLQENYYPWLKENREKIISLLTFRIPIYYVGPLADGQSEFAWLERKSDEKIKPWNFS DVV

DLDRSAEKFIEQLIGRDTYLPDEYVLPKSLIYQKYMVFNELTKIAYLDERQKRMNLSSVEK
KEIFETLFKKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYSTYVELAKVPGMKSMM

DPDNEDLMEEIVKILTVDERKMRKQLEKYKERLSPEQIKELAKKHTGWGRLSKKLLVGI
RDKETQKTILDYLVEDDNHSGGRQHNRNLMQLINDDRLSFKKTIABELQMIDPSADLYAQVQ

15 EIAGSPAIIKGILLGLKIVDEIIRVMGEKPENIVIEMARENQTTARGKALS KRREAKIKEGL
AALGSSLLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLSQYDEDHII PQSFTVDN

SLDNLVLTNSSQNRGNKDDVPSLEVNRQLAYWRSI KDA GLMTQRKFDNLTKAMRGGLTDK
DRERFIQRQLVETRQITKNAKLLDMRLNDKDEAGNKIRETNIVLLKSAMASEFRKMFR LY

KVRELNDYHHAHDAYLNAAIAINLLALYPYMADDFVYGEFRYKKKPQAEKATYEKL RQWNLI
20 KRFGEKQLFTP DHEDCWNKERDIKIKKVMGYRQVN VVKKAEERTGMLFKETINGKTNKGS R

IPIKKDLDPSKYGGYIEEKMAYYAVISYEDKKKPGKTIVGISIMDKKEFEYDSISYLGKLG
FSNPVVQIILKNYSLIAYPDGRRYITGATKTTKGKVELQKANQIAMEQDLVNFIYHLKNYD

EISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIYEEEKKDPVDIEALVA
SFIELLK LTSAGAPADFIGMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMRFED (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

MPKKKRKVGGKPYSIGLDIGTNSVGWAVVTDDYKVPACKMKVLGNTDKQS IKKNLLGALLFD
SGETAETRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRGEHH
5 PIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAEN
TDVQALFKDFVEEYDKTIEESHLSITVDALSILTEKVSKSSRLENLIAHYPTEKKNTLFGN
LIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAIL
SGILTVDNSTKAPLSASMVKRYEEHQKDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYIE
10 SGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRRQ
GEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRFAWAEYKADEKITPWNFDDILDKEKSA
EKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKVYVNEQGEAKFFDTNMKQEIFDH
VFKENRKVTDKDLLNYLNKEFEEFRIVNLTLGDKENKAFNSSLGTYHDLRKILDKSFLDDKA
NEKTIEDIIQTTLFEDREMIRQRLQKSDIFTKAQLKKLERRHYTGWGRRLSYKLINGIRNK
ENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVVHDLPGSPAICK
15 GILQSVKIVDELVKVMGHNPNANIEMARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNIK
NVENQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSDLNRVLTRSD
KNRGKSSDVPSIEVVHEMKSFWSKLLSVKLITQRKF DNLTKAERGGLTEEDKAGFIKRQLVE
TRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKSNLVSFRKEFELYKVREINDYHHAH
DAYLNAVVGNNALLKYPQLEPEFVYGEYPKNSYRSRKSATEKFLFYSNILRFFKKEDIQTN
20 EDGEIAWNKEKHIKILRKVLSYPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKTNSYW
DPKKYGGFSPVVAYSILVFADVEKGKSKKLKVQDMVGITIMEKKRFEKNPVDFLEQRGYR
NVRLEKIIKLPKYSLELENKRRLLASAKELQKGNELVIPQRFTTLLYHSYRIEKDYEP
REYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMLFSKNKDAEVSSLAKSFISLLFTA
FGAPAAFNFFGENIDRKRYTSVTECLNATLIHQSTITGLYETRIDLSKLGEDGKRPAATKKAG
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

MPKKKRKVGMGKPYSIGLDIGTNSVGWAVVTDDYKVPACKMKVLGNTDKQS IKKNLLGALLF
DSGETAEATRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFLVDEDKRGEHH
30 HPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAE
NTDVQALFKDFVEEYDKTIEESHLSITVDALSILTEKVSKSSRLENLIAHYPTEKKNTLFG
NLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYDAIL

LSGILTVDNSTKAPLSASMVKRYEEHQDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYI
ESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAILRR
QGEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRAWAHEYKADEKITPWNFDDILDKEKS
AEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFD
5 HVFKENRKVTDKLLNYLNKEFEEFRIVNLTGLDENKAFNSSLGTYHDLRKILDKSFLDDK
ANEKTIEDIIQTTLFEDREMIRQLQKYSIFTKAQLKLERRHYTGWGRLSYKLINGIRN
KENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVHDLPSPAIK
KGILQSVKIVDELVKVMGHNPANIIEMARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNI
KNVENQQLQNDRFLFLYIYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRS
10 DKNRGKSDDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLV
ETRQITKHVAQILDERFNTFEDGNKRRIRNVKIITLKSNLVSFRKEFELYKVREINDYHHA
HDAYLNAVVGNAALLKYPQLEPEFVYGEYPKYNSYRSRKSATEKFLFYSNILRFFFEDIQT
NEDGEIAWNKEKHIKILRKVLSYPQVNIVKTEEQTGGFSKESILPKGESDKLIPRKTNSY
WDPKKYGGFMQPVVAYSILVFADVEKGKSKLRKVQDMVGITIMEKKRFEKNPVDFLEQRGY
15 RNRVLEKIIKLPKYSLFELNKRRLLASAKFLQKGNELVIPQRFTTLLYHSYRIEKDYEPE
HREYVEHKDEFKELLEYISVFSRKYVLADNNLTKIEMLFSKNKDAEVSSLAKSFISLLTFT
AFGAPRAFNFFGENIARKEYRSVTECLNATLIHQSITGLYETRIDLSKLGEDGEGADKRTAD
GSEFESPKKRKV (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

MPKKKRKVGMGKPYSIGLDIGTNSVGAWVTDDYKVPACKMKVLGNTDKQS~~IKKNLLGALLF~~
DSGETAEATRLKRTARRRYTRRKNRRLRYLQEIFTGEMNKVDENFFQRLDDSFVDEDKRGEH
HPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDLKAE
NTDVQALFKDFVVEYDKTIEESHLSEITVDALSILTEKVKSSRLENLIAHYPTEKKNTLFG
25 NLIALS~~LDLHPNF~~KTNFQLSED~~AKLQFS~~KDTYEEDLEGFLGEVGDEYADLFASAKNLYDAIL
LSGILTVDNSTKAPLSASMVKRYEEHQDLKKLKDFIKVNAPDQYNAIFKDKNKKGYASYI
ESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGIIPHQIHLQEMHAILRR
QGEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRAWAHEYKADEKITPWNFDDILDKEKS
AEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQEIFD
30 HVFKENRKVTDKLLNYLNKEFEFRIVNLTGLDENKAFNSSLGTYHDLRKILDKSFLDDK
ANEKTIEDIIQTTLFEDREMIRQLQKYSIFTKAQLKLERRHYTGWGRLSYKLINGIRN
KENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVHDLPSPAIK
KGILQSVKIVDELVKVMGHNPANIIEMARENQTTDKGRRNSQQLKLLQDSLKNLDNPVNI

KNVENQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNRVLTRS
DKNRGKSDDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKRQLV
ETRQITKHVAQILDERFNTEDGDNKRRIRNVKIITLKSNLVSFRKEFELYKVREINDYHHA
HDAYLNADVGNALLKYPQLEPEFVYGEYPKNSYRSRKSATEKFLFYSNILRFFKKEDIQT
5 NEDGEIAWNKEKHIKILRKVLSPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKTKN SY
WDPKKYGGFMQPVVAYSILVFADVEKGKSKLRKVQDMVGITIMEKKRFEKNPVDFLEQRGY
RNVRLEKIIKLPKYSIFELENKRRLLASAKFLQKGNELVIPQRFTTLLYHSYRIEKDYPE
HREYVEKHDEFKELLEYISVFSRKYVLADNNLTKEMLFSKNKDAEVSSLAKSFISLLTFT
AFGAPRAFNFFGENIARKEYRSVTECLNATLIHQSITGLYETRIDLSKLGEDGEGADKRTAD
10 GSEFESPKKRKV (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

MPKKKRKVGAKNKDIRYSIGLDIGTNSVGWAVMDEHYELLKKGNHHMWGSRLF DAAEPAATR
RASRSIRRYNKRRERIRILLRDLLGDMVMVEDPTFFIRLLNVSFLDEEDKQKNLGNDYKDNY
15 NLFIEKDFNDKTYDKYPTIYHLRKELCENKEKADPRLIYLALHHIVKYRGNFLKEGQSFAK
VYEDIEEKLDNTLKKFMSLNDLDNLFVDNDINSMITVLSKIYQRSKKADDLLKIMNPTKEER
AAYKEFTKALVGLKFNVSKMILAQEVKDDKIELDFSNDYDSTVDGLQAELEYIEFIEM
LHSINSWVELQDILGNNSTISAAMVERYEEHKNDLRLKKVIREELPDKYNEVFREDNPKLH
NYLGYIKYPKNTPVEEFYEYIKRLLAKVDTGEAREILERIDLEKFMLKQNSRTNGSIPYQM Q
20 KDEM IQIIDNQS VYY PQLKENREKLISILEFRIPYYFGPLNT HSEFAWIKKFEDKQKERILP
WNYDQIVDIDATAEGFIERMQNT GTYFPDKPVMAKNSLT VSKEVLN E N KIRINGK LIP V E
TKKELLSDLFMKNKTITDKKLKDWLVT HQYYDTNEELKIEGYQKDLQFSTS LAPWIDFTKIF
GEINASNYQLIEKIIYDISI FEDKKILKRRLLKKVYQL DLLVDKILKLN YT GWSRLSEKLLT
GIKS KNSKETILSILENSNMNLMEIINDES LGFKQIIEESNKKDIEGPFRYDEVKKLAGSPA
25 IKRGIWQALLVVQEITKFMKHEPSHIYIEFAREEQEKVRTESRIAKLQKIYKDLNLQTKEDQ
LVYESLKKEDAKKIDTDALYLYLQMGKSMYSGKPLDIDKLSTYHIDHILPRSLIKDDSLD
NRVLVLPKENEWKLDSETVPFEIRNKMMGFWQKLHENGLMSNKKFFSLIRTFNEKDKKRFI
NRQLVETRQIINKNAVIINDHYTNVVTVRAELSHQFRERYKIYKNRDNLHHAHDAYIA
CILGQFIHQNFGNMDVNMIYGQYKKNYKKDVQEHNNYGFILNSMNHIHFNDDNSVIWDPSYI
30 GKI KSCFCYKDVYVTKKLEQNDAKLFDLTILPSDKNSENGVT KAKIPVNKYRKDV NK YGGF S
GDAPIMLAIEADKGKKHVRQVIAFPLRLKNYND EERIKFIEKEKNLKNV KILTEVKKNQLIL
INHQYFFITGTNELVNATQLKLSAKNTK NL FNLVDANKHNKLESIDDANFNEVIQELICKLQ
EPIYSRYNSIGKEFEDSYEKINA VTQDKLYII EYLIAIMS AKA TQGYIKPELAREIGTNGK

NKGRIKSFTIDLNTTFFISTSVTGLFSKKYKLGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (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 MPKKRKVGSIINFQRRGLMETQASNQLISSHLGYPIKDYFVGLDIGTSSVGWAVTNKAYELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRRLERRKLRLKLEELFADAMAQVDPTFFMRLRESKYHYEDKTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGKTKSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFAVLVGLNASLIDLFGSVEELEED

10 LKKLQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPLTISESKVKAFNKKHDLLAILKSLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLSDSKDKEYIILSQIELQILLPLQRIKDNGVI PYQLHLEELKAILAKCGPKFPFLNEVADGFSVAEKLIKMLEFRIPYYVGPLNTHHNVDNGGFAAWRKASGRVT PWNFDDKIDREKSAAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTK

15 NRIEQFLKDNGYISETHKHEITGLGEIKNDLASYRDMVRILGDGFDRSMAEEIITDITIFGESKKMLRETLRKKFASCLDDEAIKKLTLYRDWGRLSQKLLNGIEGCDKAGDGTPETIIILMRNFSYNLMELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVEDEVAHIKKALPARIFVEVTRSNKNEKKKDSRQKRLSDLYAAIKDDVLLNGLNNEIFGELKSSLAKYDDAALRSKKLYLYTQMGRCAYTGEIIELSLLNTDNYDIDHIYPRSLTKDDSFNLV

20 LCKRTANAQKSDAYPISEEIQKTQKPFWTFLKQQGLISERKYERLTRITPLTADDLSGFIARQLVETNQSVKAATTLLRRLYPGDVVFVKAENVTDFRHDNNFIKVRSLNHHHAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAKMFNYDVNCTNAKDGKAWDVKTSMDTVKKMMDSNDVRVTKRLLEQTGALADATIYKATVAGKAKDGAYIGMKTSSVFADVS KYGGMTKIKNAYSIIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISIYGKLCINQLVKV

25 NGFYYYLGGKTN SKFCIDNAIQVIVSNEWI PYLK VLEKFNNMRKDNKDLKANVVSTRALDNKHTIEVRIIVEEKNIEFFDYLVS KLKMPIYQKMKGNAEELSEKGYGLFKKMSLEEQS IHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIVGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 9).

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

MPKKRKVGTVKDYYIGLDIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAERRGQRGARRRLDRKKERLSLLQDFFAEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKD

FKDKNFHKKYPTIHHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIFEGQKFDTKSSFENS
LNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSS
QKLVDLFENPEDFDDSAIKSVDfsttsFDDKYSDYELALGDKIALVNILKEIYDSSILENLL
KEADKSKDGNKYISNAFKVVKYNKHGQDLKEFKRLVRQYHKSAYFDIFRSEKVNDNYVSYTKS

5 SISNNKRVKANKFTDQEAFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPK
IKSSSDNGVIPYQLKLMELNKILENQSKHEFLNVSDEYGSVCDKIASIMEFRIPYYVGPLNP
NSKYAWIKKQKDSEITPWNFKDVLDSSREEFIDSЛИRCTYLDEKVLPKASLLYNEYMV
LNELNNLKLNDLPITEEMKKKIFDQLFKTRKKVTLKAVANLLKKEFNINGEILLSGTGDFK
QGLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGKYFTDSEIKK
10 MAGLNYKDGRSLSKLLTGLEGANKITGERGSIIFMREYNLNLMSASFTFTEEIQKLN
PVDDRKLSYEMVDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSSKKIFIEMARGKEEVKARKE
SRKNQLLKFYKDGGKAFIGEIGEERYSYLLSEIEGEENKFRWDNLYLYYTQLGRCMYSLEP
IDISELSSKNIYDQDHIPKSKIYDDSIENRVLVKDLNSKKGNSYPIPDEILNKNCYAYWK
ILYDKGLIGQKKYTRLTRRTGFTDELVQFISRQIVETRQATKETANLLKTICKNSEIVYSK
15 AENASRFRQEFDIVKCRAVNDLHHMHDAYINIIVGNVYNTKFTKDPMNFKKQEKARSYNLE
NMFKYDVKGYYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRK
NKGSRPLDKGPKSSIEKYGGYNINKACFAVLDIKSKNKIERKLMPVEREYAKQKNDKKL
SDEIFSKYLDKDRFGIEDYRVVYPVVKMRTLKIDGSYYFITGGSDKTLELRSALQLILPKKN
EWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIIVNKFKTSVFKKSFLNLFQDDKNIEND
20 FFKFMSDFKEKCKTLLMLVKAIRASGVVRQDLKSIDLKSDYGRLLSKTNNIGNYQEFKIINQS
ITGLFENEVDLLKLGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA
(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 MPKKKRKVGKEYHIGLDIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAERRTFR
TTRRLKRRKWRHLHYLDEIFAPHQEVDENFLRRLKQSNIHPEPDPAKNQAFIGKLLFPDLLKK
NERGYPTLIKMRDEL PVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAHHIVKYRGHFL
NNASVDKFKVGRIDFDKSFNVLNEAYEEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLDRK
AVAKLLEVVKVADKEETKRNKQIATAMSKLVLYKADFATVAMANGNEWKIDLSSETSEDEIE
30 KFREELSDAQNDILTEITSLFSQIMLNEIVPNGMSISESMMDRWTHERQLAEVKEYLATQP
ASARKEFDQVYNKYIGQAPKEKGFDEKGLKKILSKKENWKEIDELLKAGDFLPKQRTSANG
VIPHQMHQQELDRIIEKQAKYYPWLENPATGERDRHQAKYELDQLVSFRIPYYVGPLVT
EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAFAFIKRMTVKDTYLLNEDVLPANSI

LYQKYNVLNELNNVRVNGRRLSVGIKQDIYTELFKKKTVKAGDVASLVMAKTRGVNKP SVE
GLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIIIEWRSVFEDGEIFADKLTEVEWLT
DEQRSAVKKRYKGWGRLSKKLLTGIVDENGQRIIDLMWNTDQNFMQIVNQPVFKEQIDQLN
QKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSISIEFARNEG
5 NKGEITRSRRTQLQKLFEDQAHELVKDTSLTEELEKAPDLSDRYYFYFTQGGKDMYTGDPIN
FDEISTKYDIDHILPQSFKDDSLDNRLVSRAENNKSDRVPAKLYAAKMKPWQNOLLQG
LITQRKFENLTMDVDQTICKYRSLGFVKRQLVETRQVIKTANILGSMYQEAGTDIETRAGL
TKQLREEFDLPKVREVNDYHHAVDAYLTTFAGQYLNRRYPKLRSFFVYGEYMKFKHGSDLKL
RNFNFFHELMEGDKSQGKVVWDQQTGELITTRDEVADYFDWVINLKVMLISNETYEETGKYFD
10 ASHESSSLYLKNQNKKSKLVPPLKNKLQPEYYGAYTGITQGYMVLKLLDKGGFGVYRIPR
YAADILNKCHDEVAYRNKIAEIISSDPRAPKSFEVVVPRVLKGTFLVDGEEKFILSSYRYKV
NATQLILPVSDIKLIQDNFKALKKLNVEMQTKKLIEIYDNILRQVDKYYKLYDINKFRAKHL
DGRSKFVELDDFGQDASKEKVIIKILRGLHFGSDLQNLKEIGFGTTPLGQFQVSEAGIRLSN
TAIFIIFKSPTGLFNRKLYLKNLGKRPAATKKAGQAKKKGSYPYDVPDYAYPDVPDYAYPD
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

MPKKKRKVGEKKTNYTIGLDIGTDSVGAWAVVKDDLELVKKRMKVLGNTE NYIKKNLWGSLL
FESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFRLNESFLVPEDKA
20 KNPIFGTLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHFLIEKLD
EHIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITSFLKD
NKQSPMMAFITLIVGKKAKFKAAFNLEEEISLNLTDDSYDENLEILLNTIGSDFADLF
DHAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYDMTF
VAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRK
25 QRSFYNGVIPNQVHLAEAMEAILDQENYYPWLKENREKIISLLTFRIPIYYVGPLADGQSE
FAWLERKSDEKIKPWNFSDVVDLDRSAEKFIEQLIGRDTLPDEYVLPKKSLIYQKYM
FVNELTKIAYLDERQKRMNLSSVEKKEIFETLFKKRSKVTEQLVKFFENYLQIDN
PTIFGIEDAFNADYSTYVELAKVPGMKSMMDDPDNE
30 DLMEETIVKILT
VFEDRKMR
RKQLEKYKERLSPEQIKE
LA
KKHYTG
WGRLSKKLLVG
I
RD
KET
QKT
T
ILD
YLV
EDDN
HSG
GRQ
H
LN
RNL
MQL
IN
DDRL
SF
KKT
IA
EL
Q
MID
PSADLYAQQEIA
GSPAI
KKG
ILL
GLK
IV
DEI
IR
VM
GEK
PEN
IV
EMARE
NQ
TT
ARG
KAL
SKRR
EAK
IKE
GLA
ALG
SS
LL
KEN
LPG
NAD
LSQ
RK
IY
LY
YT
QNG
KDI
Y
L
D
E
PL
F
D
RL
S
QY
D
E
H
II
PQS
FT
VD
NS
LD
NL
VL
TN
S
QN
RG
N
K
DD
V
P
S
LEV
VN
R
Q
LAY
W
R
S
L
K
D
A
G
L
M
T
Q
R
K
F
D
N
L
T
KA
MRGG
LTD
KDR
RER
FI
QR
QL
VET
RQ
IT
K
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K
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E

FRKMFRFLYKVRELNDYHHADAYLAAIAINLLALYPYMADDFVYGEFRYKKPQAEKATYE
KLRQWNLIKRFGEKQLFTPDHEDCNKERDIKTIKKVMGYRQVNWKKAERTGMLFKETIN
GKTNKGSRIPIKKDLDPSKYGGYIEEKMAYYAVISYEDKKKPGKTIVGISIMDKKEFEYDS
ISYLGKLGFSNPVVQIILKNYSLIYPDGRRRYITGATKTTKGKVELQKANQIAMEQDLVNF
5 IYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIYEEKKDP
VDIEALVASFIELLKLTSAAGAPADFIFMGEAISRRRYNSMTGLFDGQVIYQSLTGLYETRMR
FEDGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (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.
- 25 20c. The Cas9 protein of claim 20, wherein the mutation at an amino acid position 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 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 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 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 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)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH
DPTAHAEIMALRQGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA

GSLMDVLHHPGMNRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG
TSESATPESSGPKKRKVGKPYSIGLAIGTNSVGWAVTDDYKVPACKMKVLGNTDKQSICK
KNLLGALLFDSEGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQLDDSFL
VDEDKRGEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHF
5 LYEGDLKAENTDVQALFKDFVEEYDCTIEESHLEITVDAISILTEKVSKSSRLENLIAHYP
TEKKNTLFGNLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDEGLGEVGDEYADLFAS
AKNLYDAILLSGILTVDNSTKAPLSASMVKRYEEHQDLKKLKDFIKVNAPDQYNAIFKDK
NKKGYASYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHL
QEMHAIRRRQGEHYPFLKENQDKIEKILTFRIPYYGPLARKGSRFawaeyKADEKITPWNF
10 DDILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKVYVNEQGEAKFFD
TNMKQEIFDHVFKENRKVTKDKLLNYLNKEFEFRIVNLTGLDENKAFNSSLGTYHDLRKI
LDKSFLDDKANEKTIEDIIQTTLFEDREMIRQLQKYSDFITKAQLKKLERRHYTGWRGS
YKLINGIRNKENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVVH
DLPGSPAICKGILQSVKIVDELVKVMGHNPNANIIEMARENQTTDKGRRNSQQLKLLQDSL
15 KNLDNPVNICKNVENQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNS
LDNRVLTRSDKNRGKSDDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGGLTEEDK
AGFIKRQLVETRQITKHVAQILDERFNTEDGDKRRIRNVKIITLKSNLVSFRKEFELYKV
REINDYHHAHDAYLNAVGNALLKYPQLEPEFVYGEYPKNSYRSRKSATEKFYLFSNILR
FFKKEDIQTNEGEIAWNKEKHICILRKVLSPQVNIVKKTEEQTGGFSKESILPKGESDKL
20 IPRKTKNSYWDPKKYGGFDSPVVAYSILVFADVEKGKSKKLRKVQDMVGITIMEKRFKEKNP
VDFLEQRGYRNVRLEKIIKLPKYSIFELENKRRLLASAKELQKGNELVIPQRFTTLLYHSY
RIEKDYEPHREYVEKHDEFKELLEYISVFSRKYVLADNNLTKIEMLFSKNKDAEVSSLAK
SFISLLTFTAFGAPAAFNFFGENIDRKRYTSVTECLNATLIHQSTITGLYETRIDLSKLGEDG
KRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 20);

25 (b)

MPAAKRVKLDGSEVEFSHEYWMRHALTAKRARDEREVPVGAVLVNNRVIGEGWNRAIGLH
DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAA
GSLMDVLHHPGMNRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG
TSESATPESSGPKKRKVGAKNDIRYSIGLAIGTNSVGWAVMDEHYELLKGNHHMWGSRL
30 FDAAEPAATRRASRSIRRYNKRERIRLLRDLLGDMVMEVDPTFFIRLLNVSFLDEEDKQK
NLGNDYKDNYNLFIEKDFNDKTYDKYPTIYHLRKELENKEADPRLIYLALHHIVKYRGN
FLKEGQSFAKVYEDIEEKLDNTLKKFMSLNLDNLFVDNDINSMITVLSKIYQRSKKADDLL
KIMNPTKEERAAYKEFTKALVGLKFNVSKMILAQEVKDDKDIELDFSNVDYDSTVDGLQAE

LGEYIEIFIEMLHSINSWVELQDILGNNSTISAAMVERYEEHKNDLRVLKKVIREELPKYNE
VFREDNPKLHNYLGYIKYPKNTPEEFYFYIKRLLAKVDTGEAREILERIDLEKFMLKQNSR
TNGSIPYQMOKDEMIQIIDNQSVPYQPLKENREKLISILEFRIPIYYFGPLNTHSEFAWIKKF
EDKQKERILPWNYDQIVDIDATAEGFIERMQNTGTYFPDKPVMAKNSLTFSKFEVLNELNKI
5 RINGKLIPVETKKEPLLSDLFMKNKTITDKKLKDWLVTHQYYDTNEELKIEGYQKDLQFSTSL
APWIDFTKIFGEINASNYQLIEKIYDISIFEDKKILKRLKKVYQLDDLLVDKILKLNYTG
WSRLSEKLLTGIKSNSKETILSILENSNMNLMEIINDES LGFKQIIIESNKKDIEGPFRYD
EVKKLAGSPAICKRGIWQALLVVQEITKFMKHEPSHIYIEFAREEQEKRATESRIAKLQKIYK
DLNLQTKEDQLVYESLKEDAKKKIDTDALYLYLQMGKSMYSGKPLDIDKLSTYHIDHILP
10 RSLIKDDSLDNRVLVLPKENEWKLDSETVPFEIRNKMMGFWQKLHENGLMSNKKFFSLIRTD
FNEKDKKRFINRQLVETRQIINKAVIINDHYTNVVTVRAELSHQFRERYKIYKNRDLND
LHHAHDAYIACILGQFIHQNFGNMDVNMIYGQYKKNYKKDVQEHNNGFILNSMNHIHFNDD
NSVIWDPSYIGKIKSCFCYKDVYVTKLEQNDAKLFDLTILPSDKNSENGVTKAKIPVNKYR
KDVKYGGFSGDAPIMLAIEADKGKKHVRQVIAFPLRLKNYNDERIKFIEKEKNLKNVKIL
15 TEVKKNQLILINHQYFFITGTNELVNATQLLSAKNTKNLFNLVDANKHNKLESIDDANFNE
VIQELICKLQEPIYSRYNSIGKEFEDSYEKINAVTKQDKLYIIEYLIAIMSAKATQGYIKPE
LAREIGTNGKNKGRIKSFTIDLNTTFISTSVTGLFSKKYKLGKRPAATKKAGQAKKKKGSY
PYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 6);

(c)

20 MPAAKRVKLDGSEVEFSHEYWMRHALTAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLH
DPTAHAEIMALRQGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGRUVFGVRNAKTGAA
GSLMDVLHHPGMNHRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG
TSESATPESSGPKKRKVGSIINFQRRLMETQASNQLISSHLKGYPYIKDYFVGLAITSSV
GWAVTNKAYELLKFRSHKMWGSRLFDEGESAVARRGFRSMRRLERKLRKLLEELFADAM
25 AQVDPTFFMRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTD
DIRKLFLAVHHILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAISSIGQIL
MEAGKTKSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDL
FGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPLTIS
ESKVKA FNKHKDDLAILKSLLKSDRSIYNTMFVDEKGLHNYVHYIKQGRTEETSCNREDFY
30 KYTKKIVEGLSDSKDKEYILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPP
LNEVADGFSVAEKLIKMLEFRIPIYYVGPLNTHHNVDNGGFAWA VRKASGRVT PWNFDDKIDR
EKSAAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNENNRIDGKPLEKVVKEHLIEAV
FKQDHKKMTKNRIEQFLKDNGYISETHKHEITGLGEIKNDLASYRDMVRILGDGFDRSMAE

EIITDITIFGESKKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAG
DGTPETIIILMRNFSYNLMEELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKR
AVWQALRIVDEVAHIKKALPARIFVEVTRSNKNEKKKDSRQKRLSDLYAAIKDDVLLNGL
NNEIFGELKSSLAKYDDAALRSKKLYLYYTQMGRCAVTGEIELSLLNTDNYDIDHIYPRSL
5 TKDDSF DNVLCKRTANAQKSDAYPISEEIQKTQKFWTFLKQQGLISERKYERLTRITPLT
ADDLSGFIARQLVETNQSVKAATTLLRRLYPGDVVFVKAENVTDFRHNNFIKVRSLNHHH
HAKDAYLNIVVGNVYHERFTRNFRAFFKNGANRTYNLAKMFNYDVNCTNAKGKAWDVKTS
MDTVKKMMDSDVRVTKRLLEQTGALADATIYKATVAGKAKDGAYIGMKTSSVFADVSKYG
GMTKIKNAYSIIIVQYTGKGEVIKEIVPLPIYLTNRNTTDQDLINYVASIIPQAKDISHIYG
10 KLCINQLVKVNGFYYLGKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNNMRKDNKDLKAN
VVSTRALDNKHTIEVRIVEEKNIEFFDYLVSKLKMPIYQMKGNKAAELSEKGYGLFKKMSL
EEQSIHLIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTI
VGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 10);

(d)

15 MPKKKRKVSIINFQRRGLMETQASNQLISSHLKGYPKDYFVGLAITSSVGWAVTNKAYEL
LKFRSHKMWGSRLFDEGEESAVERGFRSMRRLERKRLRLKLEELFADAMAQVDPTFFMRL
RESKYHYEDKTTGHSSKHILFIDKNYNDQDYFKEYPTVYHLRSELMKSGTDDIRKLFLAVHH
ILKYRGNFLYEGATFDSNASTLDDVIKQALENITFNCFDCNSAIISSIGQILMEAGTKSDKA
KAIHLVDTYIATDTVDTSSKTQKDQVKEDKKRLKAFANLVLGLNASLIDLFGSVEELEEDL
20 KKLQITGDTYDDKRDELAKAWSDEIYIIDDCKSVYDAIILLSIKEPGLTISESKVKAFNKHK
DDLAILKSLLKSDRSIYNTMFKVDEKGLHNYVHYIKQGRTEETSCNREDFYKYTKKIVEGLS
DSKDKEYILSQIELQILLPLQRIKDNGVIPYQLHLEELKAILAKCGPKFPFLNEVADGSVA
EKLIKMLEFRIPIYYVGPLNTHHNVDNGGFAAWRKASGRVT PWNFDDKIDREKSAAAFIKNL
TNKCTYLLGEDVLPKSSLSEFMLLNELNNVRIDGKPLEKVVKEHLIEAVFKQDHKKMTKN
25 RIEQFLKDNGYISETHKHEITGLDGEIKNDLASYRDMVRILGDGFDRSMAEIIDITDITFGE
SKKMLRETLRKKFASCLDDEAIKKLTKLRYRDWGRLSQKLLNGIEGCDKAGDGT PETIIILM
RNFSYNLMEELLGDKFSFMERIQEINAKLTEGQIVNPHDIIDDLALSPAVKRAVWQALRIVDE
VAHIKKALPARIFVEVTRSNKNEKKKDSRQKRLSDLYAAIKDDVLLNGLNNEIFGELKSS
LAKYDDAALRSKKLYLYYTQMGRCAVTGEIELSLLNTDNYDIDHIYPRSLTKDDSF DNVL
30 CKRTANAQKSDAYPISEEIQKTQKFWTFLKQQGLISERKYERLTRITPLTADDLSGFIARQ
LVETNQSVKAATTLLRRLYPGDVVFVKAENVTDFRHNNFIKVRSLNHHHHAKDAYLNIVV
GNVYHERFTRNFRAFFKNGANRTYNLAKMFNYDVNCTNAKGKAWDVKTSMDTVKKMMDSN
DVRVTKRLLEQTGALADATIYKATVAGKAKDGAYIGMKTSSVFADVSKYGGMTKIKNAYSII

IVQYTGKKGEVIKEIVPLPIYLTNRNTTDQDLINYVASTIIPQAKDISIIYGKLCINQLVKVN
GFYYYLGGKTNSKFCIDNAIQVIVSNEWIPYLKVLEKFNNMRKDNKDLKANVVSTRALDNKH
TIEVRIVEEKNIEFFDYLVSKLKMPIYQKMKGNAEELSEKGYGLFKKMSLEEQSITHIELL
NLLTNQKTTFEVKPLGITASRSTVGSKISNQDEFKVINESITGLYSNEVTIVKRPAATKKAG
5 QAKKKKSGSETPGTSESATPESSGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNR
VIGEGWNRAIGLHDPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGR
VVFGVRNAKTGAAGSLMDVLHHPGMNHRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQ
SSTDPAAKRVKLDGSYPYDVPDYAYPYDVPDYAYPYDVPDYA (SEQ ID NO: 11);

(e)

10 MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRIGEGWNRAIGLH
DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPCVMCAGAMIHSRIGR VVFGVRNAKTGAA
GSIMDVLHHPGMNHRVEITEGILADECACALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG
TSESATPESSGPKKRKVGTKVKDYYIGLAIGTSSVGWAVTDEAYNVLFNSKKMWGVRLFD
DAKTAEERRGQRGARRRLDRKKERLSLLQDFFAEEVAKVDPNFFLRLDNSDLYMEDKDQKLK
15 SKYTLFNDKDFKDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYIACHYLLKNRGHFIFEGQK
FDTKS FENS LNELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLK
AVSAIMIGSSQKLVDLFENPEDFDSAIAKSVDFSTTSFDDKSDYELALGDKIALVNILKEI
YDSSILENLLKEADSKDGNKYISNAFKVKKYNKHQDLKEFKRLVRQYHKSAYFDIFRSEKV
NDNYVSYTKSSISNNKRVKANKFTDQEAFYKFAKKHLETIKYKINKVNGSKADLELI DGLMR
20 DMEFKNFMPKIKSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFR
IPYYVGPLNPNSKYAWIKKQKDSEITPNFKDVVDLDSSREEFIDS LIGRCTYLKDEKVLPK
ASLLYNEYMVLNELNNLKLNDLPITEEMKKIFDQLFKTRKKVTLKAVANLLKEFNINGEI
LLSGTDGDFKQGLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGY
KYFTDSEIKKMAGLNYKDWGRSLSKLLTGLEGANKITGERGSIIFMREYNLNLMELMSASF
25 TFTEEIQKLPVDDRKLSYEMDELYLSPSVKRMLWQSLRIVDEIKNIMGTDSSKKIFIEMAR
GKEEVKARKESRKNQLLKFYKDGKKAFISEIGEERYSYLLSEIEGEEENKFRWDNLYLYYTQ
LGRCMYSLEPIDISELSSKNIYDQDHIPKSKIYDDSIENRVLVKDKLNSKGNSYPIPDEI
LNKNCYAYWKILYDKGLIGQKKYTRLRTGFTDELVQFISRQIVETRQATKETANLLKTI
CKNSEIVYNSKAENASRFRQEFDIVKCRAVNDLHHMH DAYINIIVGNVYNTKFTKDPMFVKK
30 QEKA RSYNLENMFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESG
ALFNATLQRKNKGSRPLDKGPKS SIEKYGGYTNINKACFAVLDIKSKNKIERKLMPVEREI
YAKQKNDKKSDEIFS KYLKDREGIEDYRVVYPVVKMRTLLKIDGSYYFITGGS DKTLELRS
ALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNL

FQDDKIEINIDFKFKSMDFKEKCTLLMLVKAIRASGVRQDLKSIDLKSDYGRLLSKTNNIGN
YQEFKIINQSITGLFENEVDLLKLGKRPAATKKAGQAKKKGSYPYDVPDYAYPYDVPDYAY
PYDVPDYA (SEQ ID NO: 16);

(f)

5 MPKKKRKVTKVKDYYIGLAIGTSSVGWAVTDEAYNVLKFNSKKMWGVRLFDDAKTAEERRGQ
RGARRRLDRKKERLSLLQDFFAAEVAKVDPNFFLRLDNSDLYMEDKDQKLKSKYTLFNDKDF
KDKNFHKKYPTIHLLMDLIEDDSKKDIRLVYLACHYLLKNRGHFIFEGQKFDTKSSFENSL
NELKVHLNDEYGLDLEFDNENLINILTDPKLNKTAKKELKSVIGDTKFLKAVSAIMIGSSQ
KLVDLFENPEDFDDSAIKSVDFSTTSFDDKYSYELALGDKIALVNILKEIYDSSILENLLK
10 EADKSKDGNKYISNAFKVKKYNKGQDLKEFKRLVRQYHKSAYFDIFRSEKVNDNYVSYT
ISNNKRVKANKFTDQEAFYKFAKKHLETIKYKINKVNGSKADLELIDGMLRDMEFKNFMPKI
KSSDNGVIPYQLKLMELNKILENQSKHHEFLNVSDEYGSVCDKIASIMEFRIPYYVGPLNPN
SKYAWIKKQKDSEITPWNFKDVVDLDSSREEFIDSIGRCTLKDEKVLPKASLLYNEYMVL
NELNNLKLNDLPITEEMKKIFDQLFKTRKKVTLKAVANLLKEFNINGEILLSGTGDFKQ
15 GLNSYNDFKAIVGDKVDSDDYRDKIEEIIKLIVLYGDDKSYLQKKIKAGYGKYFTDSEIKKM
AGLNYKDWGRSLSKLLTGLEGANKITGERGSIIHFMRREYNLNLMSASFTFTEEIQKLNP
VDDRKLSYEMVDELYLSPVKRMLWQSLRIVDEIKNIMGTDSKKIFIEMARGKEEVKARKES
RKNQLLKFYKDGKKAFISEIGEERYSYLLSEIEGEEENKFRWDNLYLYTQLGRCMYSLEPI
DISELSSKNIYDQDHIFYPKSKIYDDSIENRVLVKDLNSKKGNNSYPIPDEILNKNCYAYWKI
20 LYDKGLIGQKKYTRLTRRTGFTDDELVQFISRQIVETRQATKETANLLKICKNSEIVYSA
ENASRFRQEFDIVKCRAVNDLHHMDAYINIIVGNVYNTKFTKDPMNFVKKQEKARSYNLEN
MFKYDVKRGGYTAWIADDEKGTVKNASIKRIRKELEGTNYRFTRMNYIESGALFNATLQRKN
KGSRPLDKGPKSSIEKYGGYNINKACFAVLDIKSKNKIERKLMPVEREIYAKQKNDKKLS
DEIFSKYLDRFGIEDYRVVYPVVKMRTLLKIDGSYYFITGGSDKTLELRSALQLILPKKNE
25 WAIKQIDKSSENDYLTIERIQDLTEELVYNTFDIIVNKFKTSVFKKSFLNLFQDDKIEINDF
KFKSMDFKEKCTLLMLVKAIRASGVRQDLKSIDLKSDYGRLLSKTNNIGNYQEJKIINQSI
TGLFENEVDLLKLKRPAATKKAGQAKKKSGSETPGTSESATPESSGSEVEFSHEYWMRHAL
TLAKRARDEREVPVGAVLVLNNRVIDEGWNRAIGLHDPTAHAEIMALRQGGLVMQNYRLYDA
TLYVTFEPCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNRVEITEGILADE
30 CAALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPYDVPDYAYPYDVPDYAYPYDVP
DYA (SEQ ID NO: 17);

(g)

MPAAKRVKLDGSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIDEGWNRAIGLH

DPTAHAEIMALRQGGLVMQNYRLYDATLYVTFEPVCVMCAGAMIHSRIGRVVFGVRNAKTGAA
GSLMDVLHHPGMNHRVEITEGILADECALLCRFFRMPRRVFNAQKKAQSSTDGSSGSETPG
TSESATPESSGPKKRKVGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGK
TAAERRTFRTTRRLKRRKWRLHYLDEIFAPHLQEVDENFLRLKQSNIHPEDPAKNQAFIG
5 KLLFPDLLKKNERGYPTLIKMRDEL PVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAHV
HIVKYRGHFLNNASVDKFVKGRIDFDKS FNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLD
TKMRKLDQKAVAKLLEVKVADKEETKRNKQIATAMS KLVLYKADFATVAMANGNEWKIDL
SSETSEDEIEKFREELS DAQNDILTEITS LFSQIMLNEIVPNGMS ISESMMDRYWTHERQLA
EVKEYLATQPASARKEFDQVYNKYIGQAPKEKGFDLEKGLKKILSKKENWKEIDELLKAGDF
10 LPKQRTSANGVIPHMHQQELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVS FRI
PYYVGPLVTPEVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAEEFIKRMTVKDTYLL
NEDVL PANSLLYQKYNVLNELNNVRVNGRRLSVGIKQDIYTEL FKKKKTVKAGDVASLVMAK
TRGVNKPSVEGLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENII EWRSVFEDGEIFA
DKLTEVEWLTDEQRSA LVKKRYKGWGRLSKKLLTGIVDENGQRIIDL MWNT DQN FMQIVNQP
15 VFKEQIDQLNQKAITNDGMTLRERVESVLDDAYTSPQNKAIWQVVRVVEDIVKAVGNAPKS
ISIEFARNEGNGEITRSRRTQLQKL FEDQAHELVKDTS LTEELEKAPDLSDRYYFYFTQGG
KDMYT GDPINFDEI STKYDIDHILPQS FVKDDS LDNRVL VSRAENNKKSDRVPACKYAAKMK
PYWNQLLKQGLITQRKFENLTMVDQQT IKYRSLGFVKRQLVETRQVIKTANILGSMYQEAG
TDIIETRAGLTKQLREEFDLPKVREVNDYH AVDAYLTT FAGQYLNRRYPKLRSFFVYGEYM
20 KFKHGSDLKLRNFNFHELMEGDKSQGKVVDQQT GELITTRDEVADYFDWVINLKVMLISNE
TYEETGKYFDASHESSSLYLNQNKSKLVVPLKNKLQPEYYGAYT GITQGYM VILKLLDKK
GGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKSFEVVVPRVLKGTFLVDGEEK
FILSSYRYKV NATQLI LPVSDIKLIQDNFKALKLN VEMQTKKLIEIYDNILRQVDKYYKLY
DINKFRAKLHDGRSKFVELDDFGQDASKEKVIKI LRGHLFGSDLQNLKEIGFGTTPLGQFQ
25 VSEAGIRLSNTAIFIIFKSPTGLFNRKLYLKNLGKRPAATKKAGQAKKKGSYPYDVPDYAYP
YDVPDYAYPYDVPDYA (SEQ ID NO: 88);

(h)

MPKKKRKVGKEYHIGLAIGTSSIGWAVTDSQFKLMRIKGKTAIGVRLFEEGKTAERRTFRT
TRRRLKRRKWRLHYLDEIFAPHLQEVDENFLRLKQSNIHPEDPAKNQAFIGKLLFPDLLKK
30 NERGYPTLIKMRDEL PVEQRAHYPVTNIYKLREAMINEDRQFDLREVYLAHV HIVKYRGHFL
NNASVDKFVKGRIDFDKS FNVLNEAYEELQNGEGSFTIEPSKVEKIGQLLLDTKMRKLDQK
AVAKLLEVKVADKEETKRNKQIATAMS KLVLYKADFATVAMANGNEWKIDL SSETSEDEIE
KFREELS DAQNDILTEITS LFSQIMLNEIVPNGMS ISESMMDRYWTHERQLA EVKEYLATQP

ASARKEFDQVYNKYIGQAPKEKGFDLEKGLKKILSKKENWEIDELKAGDFLPKQRTSANG
VIPHQMHQQELDRIIEKQAKYYPWLATENPATGERDRHQAKYELDQLVSFRIPIYYGPLVTP
EVQKATSGAKFAWAKRKEDGEITPWNLWDKIDRAESAFAIKRMTVKDTYLLNEDVLPANS
LYQKYNVLNELNNVRVNRRGRLSVGIKQDIYTELFKKKTVKAGDVASLVMAKTRGVNKPSVE

5 GLSDPKKFNSNLATYLDLKSIVGDKVDDNRYQMDLENIEWRSVFEDGEIFADKLTEVEWLT
DEQRSALVKKRYKGWGRLSKKLLTGIVDENGQRIIDLWNTDQNFMQIVNQPVFKEQIDQLN
QKAITNDGMTLRERVESVLDDAYTSPQNKKAIWQVVRVVEDIVKAVGNAPKSISIEFARNEG
NKGEITRSRRTQLQKLFEDQAHELVKDTSLTEELEKAPDLSDRYYFYFTQGGKDMYTGDPIN
FDEISTKYDIDHILPQSFKDSDLNRVLVSRAENNKSDRVPAKLYAAKMKPWNQOLLKQG
10 LITQRKFENLTMDVDQTICKYRSLGFVKRQLVETRQVIKTANILGSMYQEAGTDIETRAGL
TKQLREEFDLPLKVREVNDYHHAVDAYLTTFAQYLNRRYPKLRFFVYGEYMKFKHGSDLKL
RNFNFFHELMEGDKSQGVVDDQQTGELITTDEVADYFDWVINLKVMLISNETYEETGKYFD
ASHESSSLYLNQNKSKLVVPLKNKLQPEYYGAYTGITQGYMVLKLLDKGGFGVYRIPR
YAADIINKCHDEVAYRNKIAEIISDPRAPKSFEVVVPRVLKGTFLVDGEEKFILSSYRYKV
15 NATQLILPVSDIKLIQDNFKALKKLNVEMQTKKLIEIYDNILRQVDKYYKLYDINKFRAKLH
DGRSKFVELDDFGQDASKEKVIIKILRGLHFQSDLQNLKEIGFGTTPLGQFQVSEAGIRLSN
TAFIIIFKSPTGLFNRKLYLKNLKRPAATKKAGQAKKKSGSETPGTSESATPESSGSEVEFS
HEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDPTAHAEIMALRQGGLV
MQNYRLYDATLYVTFEPVCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHPGMNHRVE
20 ITEGILADECALLCRFFRMPRRVFNAQKKAQSSTDPAAKRVKLDGSYPYDVPDYAYPYDVP
DYAYPYDVPDYA (SEQ ID NO: 89); or

(i)

MSEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDPTAHAEIMA
LRQGGLVMQNYRLYDATLYSTFEPVCVMCAGAMIHSRIGRVVFGVRNAKTGAAGSLMDVLHHP
25 GMNHRVEITEGILADECALLCRFFRMPRRVFNAQKKAQSSTDGGSSGGSSGSETPGTSES
ATPESSGGSSGGSGKPYSIGLAIGTNsvgavvtddykvpakmkvlgntdkosiknnllga
LLFDGETAEATRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSLVDEDKR
GEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDL
KAENTDVQALFKDFVEEYDKTIEESHlseitvdalsiltekvsksrrlenliahypteknt
30 LFGNLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYEEDEGLGEVGDEYADLFASAKNLYD
AILLSGILTVDNSTKAPLSASMVKRYEEHQKDLKKLDFIKVNAPDQYNAIFKDKNKGYA
SYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGSIPHQIHLQEMHAI
LRRQGEHYPFLKENQDKIEKILTFRIPYYGPLARKGSRFAWAEYKADEKITPWNFDDILDK

EKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQE
IFDHVFKENRKVTDKLLNYLNKEFEFRIVNLTGLDENKAFCNSSLGTYHDLRKILDKSFL
DDKANEKTIEDIIQTTLFEDREMIRQLQKYSIFTKAQLKKLERRHYTGWRGRLSYKLING
IRNKENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVHDLPGS P
5 AIKKGILQSVKIVDELVKVMGHN PANIIEMARENQTTDKGRNSQQLKLLQDSLKNLDNP
VNICKNENQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNRVL
TRSDKNRGKSDVPSIEVVHEMKS FWSKLLSVKLITQRKF DNLTKAERGGLTEEDKAGFIKR
QLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKSNLVSNFRKEFELYKREINDY
HHAHDAYLNAVGNALLKY PQUEPEFVYGEYPKNSYRSRKSATEKFLFYSNILRFFKKED
10 IQTNEDGEIAWNKEKHICILRKVLSPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKT
NSYWDPKKYGGFMQPVVAYSILVFADVEKGKSKLRKVQDMVGITIMEKKRFEKNPVDFLEQ
RGYRNVRLEKIIKLPKYSIFELENKRRLLASAKFLQKGNELVI PQRFTTLLYHSYRIEKDY
EPEHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMPLFSKNKDAEVSSLAKSFISLL
TFTAFGAPRAFNFFGENIARKEYRSVTECLNATLIHQ SITGLYETRIDLSKLGEDGEADKR
15 TADGSEFESP KKRKV (SEQ ID NO: 98);

(j)

MSEVEFSHEYWMRHALT LAKRARDEREVPVGAVLVLNNRVIGEGWNRAIGLHDPTAHAEIMA
LRQGGLVMQNYRLYDATLYSTFEP CVMCAGAMIHSRIGRVFGVRNAKTGAAGSLMDVLHHP
GMNHRVEITEGILADEC AALLCRFFRMP RRVFNAQKKAQSSTDGGSSGGSSGSETPGTSES
20 ATPESSGGSSGGSGKPYSIGLAIGTNSVGWAVTDDYKVPACKMKV LGNTDKQS IKKNLLGA
LLFDSGETAETRLKRTARRRYTRRKNRLRYLQEIFTGEMNKVDENFFQRLDDSF LVDEDKR
GEHHPIFGNIAAEVKYHDDFPTIYHLRRHLADTSKKADLRLVYLALAHMIKFRGHFLYEGDL
KAENTDVQALFKDFVEEYDKTIEESHLSEITVDALSI LTEKVSKSSRLENLIAHYPTEKKNT
LFGNLIALSLDLHPNFKTNFQLSEDAKLOFSKDTYEEDLEGFLGEVGDEYADLFASAKNLYD
25 AILLSGILTVDNSTKAPLSASMVKRYEEHQDLKKLDFIKVNAPDQYNAIFKDKNKGYA
SYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDREDFLRKQRTFDNGIIPHQIHLQEMHAI
LRRQGEHYPFLKENQDKIEKILTFRIPYYVGPLARKGSRF AWA EYKADEKITPWNFDDILD
EKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTVYNELTKVKYVNEQGEAKFFDTNMKQE
IFDHVFKENRKVTDKLLNYLNKEFEFRIVNLTGLDENKAFCNSSLGTYHDLRKILDKSFL
30 DDKANEKTIEDIIQTTLFEDREMIRQLQKYSIFTKAQLKKLERRHYTGWRGRLSYKLING
IRNKENKKTILDYLIDDGYANRNFMQLINDDALSFKEEIARAQIIDDVDDIANVHDLPGS P
AIKKGILQSVKIVDELVKVMGHN PANIIEMARENQTTDKGRNSQQLKLLQDSLKNLDNP
VNICKNENQQLQNDRLFLYYIQNGKDMYTGETLDINNLSQYDIDHIIPQAFIKDNSLDNRVL

TRSDKNRGKSDDVPSIEVVHEMKSFWSKLLSVKLITQRKFDNLTKAERGGLTEEDKAGFIKR
QLVETRQITKHVAQILDERFNTEFDGNKRRIRNVKIITLKSNLVSNFRKEFELYKVREINDY
HHAHDAYLNAVVGNAALLKYPQLEPEFVYGEYPKNSYRSRKSAATEKFLFYSNILRFFKKED
IQTNEEDGEIAWNKEKHIKILRKVLSPQVNIVKKTEEQTGGFSKESILPKGESDKLIPRKT
5 NSYWDPKKYGGFMQPVVAYSILVFADVEKGKSKLRKVQDMVGITIMEKKRFEKNPVDFLEQ
RGYRNVRLEKIIKLPKYSLFELENKRRLLASAKFLQKGNELVI PQRFTTLLYHSYRIEKDY
EPEHREYVEKHKDEFKELLEYISVFSRKYVLADNNLTKIEMLF SKNKDAEVSSLAKSFISLL
TFTAFCAPRAFNFFGENIARKEYRSVTECLNATLIHQSI TGLYETRIDLSKLGEDGEGADKR
TADGSEFESP KKKRKV (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)

MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR
SGKNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLV
15 YARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMML
YALELHCIIISLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHP SVTWRLKS
GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKRKVGGKPYSIGLAIGTNSVGAVVTDD
YKVPACKMKVLGNTDKQS IKKNLLGALLFDGETAEATRLKRTARRRYTRRKNRLRYLQEIF
20 TGEMNKVDENFFQRLDDSFLVDEDKRGEHHPIFGNIAAEVKYHDDPTIYHLRRHLADTSKK
ADLRLVYLALAHMIKFRGHFLYEGDLKAENTDVQALFKDFVEEYDKTIEESHLS
ILTEKVSKSSRLENLIAHYPTEKKNTLFGNLIALSLDLHPNFKTNFQLSEDAKLQFSKDTYE
EDLEGFLGEVGDEYADLFASAKNLYDAILLSGILTVDNSTKAPLSASMVKRYEEHQDKLKK
LKDFIKVNAPDQYNAIFKDKNKKGYASYIESGVKQDEFYKYLKGILLKINGSGDFLDKIDRE
25 DFLRKQRTFDNGSIPHQIHLQEMHAILRRQGEHYPFLKENQDKIEKILTFRIPYVGPLARK
GSRFAWAEYKADEKITPWNFDDILDKEKSAEKFITRMTLNDLYLPEEKVLPKHSPLYEAFTV
YNELTKVKYVNEQGEAKFFDTNMKQEIFDHVFKENRKVT KDKLLNYLNKEFEEFRIVNL
DKENKAFNSSLGTYHDLRKILDKSFLDDKANEKTIEDIIQTLTFEDREMIRQRLQKYS
30 TKAQLKKLERRHYTGWGRRLSYKLINGIRNKENKKTILDYLIDDGYANRNFMQLIN
EEIARAQIIDDVDDIANVHDLPSPAICKGILQSVKIVDELVKVMGHNPANII
TTDKGRRNSQQRLKLLQDSLKNLDNPVNIKVENQQLQNDRLFLYYIQNGKDMYTGETLDIN
NLSQYDIDHIIPQAFIKDNSLDNRVLTRS DKNRGKSDDVPSIEVVHEMKSFWSKLLSVKLIT
QRKFDNLTKAERGGLTEEDKAGFIKRQLVETRQITKHVAQILDERFNTEFDGNKRRIRNV
ITLKSNLVSNFRKEFELYKVREINDYHHAHDAYLNAVVGNAALLKYPQLEPEFVYGEYPKYN

SYRSRKSA TEKFLFYSNILRFFKKEDIQTNEDGEIAWNKEKH KILRKVLSYPQVNIVKKTE
EQTGGFSKESILPKGESDKLIPRKTKN SYWDPKKYGGFDSPVVAYSILVFADVEKGKS KKL R
KVQDMVGITIMEKKRF EKNPVD FLEQRGYRNVRLEKIIKLPKYSLF ELENKRRLLASAKEL
QKG NELVI PQRFTTLLYHSYRIEKDYEP EHREYVEHKDEFKELLEYISVFSRKYVLADNNL
5 TKIEMLFSKNKDAEVSSLAKSFISLLTFTAFGAPAA FNFFGENIDRKRYTSVTECLNATLIH
QSITGLYETRIDLSKLG EDGKRPAATKKAGQAKKKKGSSGGSGG STNLSDIIEKETGKQ
LVIQESI LMLPEEV EEVIGNKPESDILVHTAYDESTDENVMLLTS DAPEYKPWALVIQDSNG
ENKIKMLSGGSGGSGG STNLSDIIEKETGKQLVIQESI LMLPEEV EEVIGNKPESDILVHTA
YDESTDENVMLLTS DAPEYKPWALVIQDSNGENKIKML YPYDVPDYA PYDVPDYA PYDVP
10 DY A (SEQ ID NO: 21);

(b)

MPAAKRVKLD TSEKGPSTGDPTLRRRIESWEFDVYD PRELRKETCLLYEIKWGMSRKI WRS
SGKNTTNHVEVNFIKKFTSERRHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVY
VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPPLWMML
15 YA LEHL CIILSLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHP SVTWR LKS
GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKRKVGSIINFQRRGLMETQASNQLISSH
LKGYP IKDYFVGLAIGTSSVGWAVTNKAYELLKFRSHKMWSRLFDEGE SAVARRGFRSMRR
RLERRKLRKLLEELFADAMAQVDPTFFMRLRESKYHYEDKTTGHSSKHILFIDKNYNDQDY
FKEYPTVYHLRSELMKSGTDDIRKLFLAVHILKYRGN FLYEGATFDSNASTLDDVIKQALE
20 NITFNCFCDSAIISSIGQILMEAGKTKSDKAKAIEHLVDTYIATDTVDTSSKTQKDQVKEDK
KRLKA FANLVLGLNASLIDLFGSVEELEEDLKKLQITGDTYDDKRDELAKAWSDEIYIIDD C
KSVYDAIILLSIKEPGLTISESKVKAFNKH KDDLAILKSLLKSDRSIYNTMF KVDEKGLHNY
VHYIKQGRTEETSCNREDFYKYTKIVEGLSDSKDKEYIILSQIELQILLPLQRIKDNGVIPY
QLHLEELKAI LAKCGPKFPFLNEVADGF SVAEKL IKMLEFRIPIYYVGPLNTHHNVDNGGFAW
25 AVRKA SGRVT PWNFDDKIDREKSAAFIKNLTNKCTYLLGEDVLPKSSLLYSEFMLLNELNN
VRIDGKPLEKVVK EHLIEAVFKQDHKKMTKNRIEQFLKDNGYISETHKHEITGLDGEIKNDL
ASYRDMVRILGDGFDRSMAEEIITDITIFGESKKMLRETLRKKFASCLDDEAIKKLT KRYR
DWGR LSQKLLNGIEGCDKAGDGT PETIIILMRNFSYNLMELLGDKFSFMERIQEINA KLT E
QIVNP HDI IDDLALS PAVKRAVWQALRIVDEVAHIKKALPARIFVEVTRSNKNEKKKDSRQ
30 KRLSDLYAAIKKDDVLLNGLNNEIFGELKSSLA KYDDAALRSKKLYLYYTQMGR CAYTGEII
ELSLLNTDNYDIDHIYPRSLTKDDSFDNLVLCKRTANAQKSDAYPISEEIQKTQKFWTFLK
QQGLISERKYERLTRITPLTADDLSGFIARQLVETNQSVKAATTLLRRLYPGVDFVKAEN
VTDFRH DNNFIKVRSLNHHHAKDAYLNIVVGNVYHERFTRNFRAFFKKNGANRTYNLAKMF

NYDVNCTNAKDGKAJDVKTSMDTVKKMMDSDVRVTKRLLEQTGALADATIYKATVAGKAKD
GAYIGMKTSSVFADVSKYGGMTKIKNAYSIIIVQYTGKKGEVIKEIVPLPIYLTNRNTTDQD
LINYVASIIPQAKDISIIYGKLCINQLVKVNGFYYYLGGKTNSKFCIDNAIQVIVSNEWIPY
LKVLEKFNNMRKDNLKANVVSTRALDNKHTIEVRIVEKNIEFFDYLVSKLKMPIYQKMK
5 GNKAAEELSEKGYGLFKKMSLEEQSILHIELLNLLTNQKTTFEVKPLGITASRSTVGSKISNQ
DEFKVINESITGLYSNEVTIVGKRPAAKKAGQAKKKGGSGSGSGGSTMNLSDIIIEKETG
KQLVIQESILMLPEEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDS
NGENKIKMLSGGSGGSGGSTMNLSDIIIEKETGKQLVIQESILMLPEEEVIGNKPESDILVH
TAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAYPYDVPDYAY

10 (SEQ ID NO: 12);

(c)

MPAAKRVKLDTSEKGPGSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR
SGKNTTNHVEVNFIKKFTSERRFHSISITWFWSWPCWECSQAIREFLSQHPGVTLV
VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMML
15 YALELHCIILSLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHP
GGSSGGSSGSETPGTSESATPESSGGSSGSPKKRKVGTKVKDYYIGLAIGTSSVG
EAYNVLKFNNSKKMWGVRLFDDAKTAEERRGQRGARRRLDRKKERLSLLQDF
FFLRLDNSDLYMEDKDQKLKSKYTLFNDKDFKDKNFHKKYPTIHLLMD
LIEDDSKKDIRLV
YLACHYLLKNRGHFIFEGQKFDTKSFENSNELKVHLNDEYGLD
LEFDNENLINILTDPKL
20 NKTAKKELKSVIGDTKFLKAVSAIMIGSSQKLVDLFENPEDF
FDDSAIKSVDFSTTSFDDKY
SDYEALGDKIALVNILKEIYDSSILENLLKEADKS
KDGNKYISNAFKYKNGQDLKEFK
RLVRQYHKSAYFDIFRSEKVNDNYSYTKSSISNNKRV
KANKFTDQEAFYKFAKKHLETIKY
KINKVNGSKADLELIDGMLRDMEFKNFMPKIKSSDNG
VIYQLKLMELNKILENQSKHHEFL
NVSDEYGSVC
DKIASIMEFRI
PYVGPLNPNSKYAWIKKQKDSEIT
PWNFKDV
VLDSSREE
25 FIDSLIGRCTYL
DEKVL
PKASLLY
NEY
MVL
NEL
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KLNDL
PITEEM
KKK
IFDQL
FKTRKK
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KAVAN
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QGLNS
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RKELEGTYRFTRMNYIESGALFNATLQRKNKGSRPLDKGPKSSIEKYGGYNINKACFAV
LDIKSKNKIERKLMPVEREIYAKQKNDKKLSDEIFSKYLDRFGIEDYRVVYPVVKMRTLLK
IDGSYYFITGGSDKTLELRSALQLILPKKNEWAIKQIDKSSENDYLTIERIQDLTEELVYNT
FDIIVNKFKTSVFKKSFLNLFQDDKIENIDFKFKSMDFKEKCKTLLMLVKAIRASGVRQDLK
5 SIDLKSDYGRLLSKTNIGNYQEJKIINQSITGLFENEVDLLKLGKRPAATKKAGQAKKKKG
SSGGSGGSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTD
ENVMLLTSDAPEYKPWALVIQDSNGENKIKMLSGGSGGSTNLSDIIEKETGKQLVIQES
ILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENKIKM
LYPYDVPDYAYPYDVPDYAY (SEQ ID NO: 18);

10 (d)

MPAAKRVKLDSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR
SGKNTTNHVEVNFIKKFTSERRHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLV
VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGPDEAHWPQYPPLWM
YALELHCIILSLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHP
5 GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKRKVGKEYHIGLAIGTSSIGWAVTDSQE
KLMRIKGKTAIGVRLFEEGKTAERRTFRTTRRLKRRKWRLHYLDEIFAPHLQEVDENFL
RLKQSNIHPEPDPAKNQAFIGKLLFPDLLKKNERGYPTLIKMRDEL
EAMINEDRQFDLREVYLAHHIVKYRGHF
EGSFTIEPSKVEKIGQLLDTKMRKLDRQKAVAKLLEV
20 YKADFATVAMANGNEWKIDLSSETSEDEIEKFREELS
GMSISESMMDRYW
GERDRHQAKYELDQLVS
RAESAEEAFIKRMTVKDTYLLNEDVLP
25 LFKKKKTVKAGDVASLVM
MDLENIEWRSV
RIIDL
ELEKAPDLS
30 AENNKKSDRVP
TRQVI
EVADY

GAYTGITQGYMVLKLLDKGGFGVYRIPRYAADILNKCHDEVAYRNKIAEIISSDPRAPKS
FEVVVPRVLKGTFLVDGEEKFILSSYRYKVNATQLILPVSDIKLIQDNFKALKKLNVEMQT
KLIIEIYDNILRQVDKYYKLYDINKFRAKLHDGRSKFVELDDFGQDASKEKVIIKILRGLHFG
SDLQNLKEIGFGTTPLGFQVSEAGIRLSNTAIFIIFKSPTGLFNRKLYLKNLGKRPAATKKA
5 GQAKKKKGSSGGSGGSTNLSDIIEKETGKQLVIQESILMLPEEEVIGNKPESDILVH
TAYDESTDENVMLLTSDAPEYKPWALVIQDSNGENIKMLSGGSGGSTNLSDIIEKETG
KQLVIQESILMLPEEEVIGNKPESDILVHTAYDESTDENVMLLTSDAPEYKPWALVIQDS
NGENIKMLYPYDVPDYAYPYDVPDYAY (SEQ ID NO: 90);

(e)

10 MPAAKRVKLDTSEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWR
SGKNTTNHVEVNFIKKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLV
VARLFWHMDQRNRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMML
YALELHCIILSLPPCLKISRRWQNHLAFFRLHLQNCHYQTIPPHILLATGLIHP
GGSSGGSSGSETPGTSESATPESSGGSSGGSPKKRKVGEEKTNYTIGLAIGTDSVG
15 DDLELVKKRMKVLGNTETNYIKKNLWGSLLFESGQTAKDRRLKRVARRRYERRRNRLTELQK
IFAPAIDEVDENFFFRLNESFLVPEDKA
FSKNPIFGTLGEDKTYKTYPTIYHLRQH
LADSE
EKADVRLIYLALAHMIKYRGHFLIEGKLDTEHIA
IENLEQFFESYNALFSEE
PIELKEEL
IAIENILREKNSRTVKEKRITSFLKD
IGRANKQS
PMMAFITLIVGKKAKFKA
AFNLEEEISL
NLTDDSYDENLEILLNTIGSDFADLF
DHAQRVYNA
VELAGILSGDV
KNT
HAKLSAQMVAMYE
20 RHKEQLKEYKSFIKANLPDQYDMTFVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFV
KQDQLKE
VPGSEKFLDSIEKEDFLRKQRSFYNGVIPNQVH
LAEMEA
ILD
RQENYY
PWLK
ENRE
KIISL
TFR
I
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YVG
PLADG
QSE
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YITGATKTTKGKVELQKANQIAMEQDLVNFIFYHLKNYDEISHPESYAFVQSHTDYFDRLFDS
IEHYTRRFLDAETNINRLRRIYEEEKKDPVDIEALVASFIELLKLTSAAGAPADFI FMGEAI
SRRRYNSMTGLFDGQVIYQSLTGLYETRMRFEDGKRPAATKKAGQAKKKGSSGGSGSGGS
TNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSAP
5 EYKPWALVIQDSNGENKIKMLSGGSGGSGSTNLSDIIEKETGKQLVIQESILMLPEEVEEV
IGNKPESDILVHTAYDESTDENVMLLTSAPEYKPWALVIQDSNGENKIKMLYPYDVPDYAY
PYDVPDYAY (SEQ ID NO: 93);

(f)

MPAAKRVKLDTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDE
10 NVMLLTSAPEYKPWALVIQDSNGENKIKMLSGGSGGSGSTNLSDIIEKETGKQLVIQESI
LMLPEEVEEVIGNKPESDILVHTAYDESTDENVMLLTSAPEYKPWALVIQDSNGENKIKML
SGGSGGSGSPKKRKVEKKNTYTIGLAIGTDSVGAVVKDDLELVKKRMKVLGNTEINYIK
KNLWGSSLFESGQTAKDRRLKRVARRRYERRRNRLTELQKIFAPAIDEVDENFFRLNESFL
VPEDKAFSKNPIFGTLGEDKTYKTYPTIYHLRQHLADSEEKADVRLIYLALAHMIKYRGHF
15 LIEGKLDTEHIAINENLEQFFESYNALFSEEPIELRKEELIAIENILREKNSRTVKEKRITS
FLKDGRANKQSPMMAFITLIVGKKAKFKAAFNLEEEISLNLTDDSYDENLEILLNTIGSDF
ADLFDAQRVYNAVELAGILSGDVKNTHAKLSAQMVAMYERHKEQLKEYKSFIKANLPDQYD
MTFVAPKDAQKKDLKGYAGYIDGNMSQDSFYKFVKDQLKEVPGSEKFLDSIEKEDFLRKQRS
FYNGVIPNQVHLAEMEAILEDQENYYPWLKENREKIISLITFRIPYYVGPLADGQSEFAWLE
20 RKSDEKIKPWNFSDVVDLRSAEKFIEQLIGRDTYLPEYVLPKKSLIYQKYMVFNELTKIA
YLDERQKRMNLSSVEKKEIFETLFFKRSKVTEKQLVKFFENYLQIDNPTIFGIEDAFNADYS
TYVELAKVPGMKSMMDPDNEDLMEEIVKILTVFEDRKMRKQLEKYKERLSPEQIKELAKK
HYTGWRSLSKLLVGIRDKETQKTILDYLVEDDNHSGGRQHNRNLMQLINDDRLSFKKTIA
ELQMIDPSADLYAQVQEIAGPSAIKKGILLGLKIVDEIIIRVMGEKPNIVIEMARENQTTAR
25 GKALSKRREAKIKEGLAALGSSLKENLPGNADLSQRKIYLYYTQNGKDIYLDEPLDFDRLS
QYDEDHIIIPQSFTVDNSLDNLVLTNSSQRGNKKDDVPSLEVNRQLAYWRSLKDALMTR
KFDNLTKAMRGGLTDKDRERFIQRQLVETRQITKNVAKLDMRLNDKDEAGNKIRETNIVL
LKSAMASEFRKMFRLYKVRELNDYHHAHDAYLNAIAINLLALYPYMADDFVYGEFRYKKP
QAEKATYEKLRQWNLIKRFGEKQLFTPDHEDCWNKERDIKTIKKVMGYRQNVVKAAERTG
30 MLFKETINGKTNKGSRIPIKKLDPSKYGGYIEEKMAYYAVISYEDKKKPGKTIVGISIMD
KKEFEYDSISYLGKLGFSNPVVQIILKNYSLIAYPDGRRRYITGATKTTKGKVELQKANQIA
MEQDLVNFIFYHLKNYDEISHPESYAFVQSHTDYFDRLFDSIEHYTRRFLDAETNINRLRRIY
EEEKKDPVDIEALVASFIELLKLTSAAGAPADFI FMGEAISRRYNSMTGLFDGQVIYQSLT

GLYETRMRFEDKRPAATKKAGQAKKKGSSGGSSGGSETPGTSESATPESSGGSSGGST
SEKGPSTGDPTLRRRIESWEFDVFYDPRELRKETCLLYEIKWGMSRKIWRSSGKNTTNHVEV
NFIKKFTSERRFHSSISCSITWFLSWSPCWECSQAIREFLSQHPGVTLVIVARLFWHMDQR
NRQGLRDLVNSGVTIQIMRASEYYHCWRNFVNYPGDEAHWPQYPPLWMMLYALELHCILS
5 LPPCLKISRRWQNHLAFLRLHLQNCHYQTIPPHILLATGLIHPSVTWRYPYDVPDYAYPYDV
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

5 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.

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

10 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 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'-

GUUUUAGAGCUGUGCUGUUAAAACAACACAGCAAGGUAAAAUAAGGCUUUGU
CCGUACUCAAGCUUGCAAAAGCGUGCACCGAUUCGGUGCU-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'-

GUUUUAGAGUUGUGUUAUUGAAAAAAUACACAACAGAGUAAAAUAAGCUUA
UGCUUAAAUGCAGCUUUGCUGGUGUCAUUAGAUGACUUUACUAAGGUUGC
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'-

GUUUGAGAGUAGUGUGAAAACAUUACGAGUCAAAUACAAUUAACAA
UGCCUUCGGGCUGCCCACGUAGGGCACCUACUCUCAAUCUUCGGAAUUGAG
20 UU-3' (SEQ ID NO: 13).

46. The method of claim 42, wherein the sgRNA for use with *Ezakiella peruvensis* Cas9

comprises a scaffold comprising a sequence having at least about 80% identity to

5'-

GUUUGAGAGUUAUGUAAUUGAAAAAAUACAUUGACGAGUCAAAUAAAAAUU
25 AUUCAAACCGCCUAAUUAUAGGCCGCAGAUGUUCUGCAUUAUGCUUGCUAU
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 GUCUUGGAUGAGUGUGAAAACACUCAUAGUCAAGAUCAAACGAGUGGUUUUC
CACGAGUUAUUACUUUUGAGGCUCUUAUGGCCAUACAUAAAAAGGAGUCG
GAAUUUCGGCUCCUUUCUU-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' -

GUUUUAGAGCCAUGUAGAAAACAUUGCAAGUUAAAUAAGGCUUUGUCCGU
AAUCAACUUGAAAAAGUGGCGCUGUUUCGGCGCUU-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 comprises a direct repeat sequence and a spacer sequence capable of hybridizing to a target 25 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.

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.

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,

10 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'-

GUUUUAGAGCUGUGCUGUUUAACAAACACAGCAAGUUAAAUAAGGCUUUGU

20 CCGUACUCAAGCUUGCAAAAGCGUGCACCGAUUCGGUGCU-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'-

GUUUUAGAGUUGUGUUAUUGAAAAAAUACACAACACAGAGUUAAAUAAGCUUA

25 UGCUUAAAUGCAGCUUUGCUGGUGUCAUUAGAUGACUUUACUAAGGUUGC
UUCGGCAACCUUUUU-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'-

GUUUGAGAGUAGUGUGAAAACAUUACGAGUCAAAUACAAUUAUUUACAA
UGCCUUCGGGCUGCCGACGUAGGGACCUACUCUCAAUUCUUCGGAAUUGAG
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'-

GUUUGAGAGUAUGUAUUUGAAAAAUACAUGACGAGUCAAAUAAAAAUU
AUUCAAACCGCCUAUUUAUAGGCCGCAGAUGUUCUGCAUUAUGCUUGC
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 GUCUUGGAUGAGUGUGAAAACACUCAUAGUCAAGAUCAAAACGAGUGGUUUUC
CACGAGUUAUUACUUUUGAGGUCUUAUAGGCCAUACAUAAAAGGAGUCG
GAAUUUCCGGCUCCUUUCUU-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'-

GUUUUAGAGCCAUGUAGAAAACAUUGCAAGUUAAAUAAGGCUUUGUCCGU
AAUCAACUUGAAAAAGUGGCGCUGUUUCGGCGCUU-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) 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 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;
 - 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.
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 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.

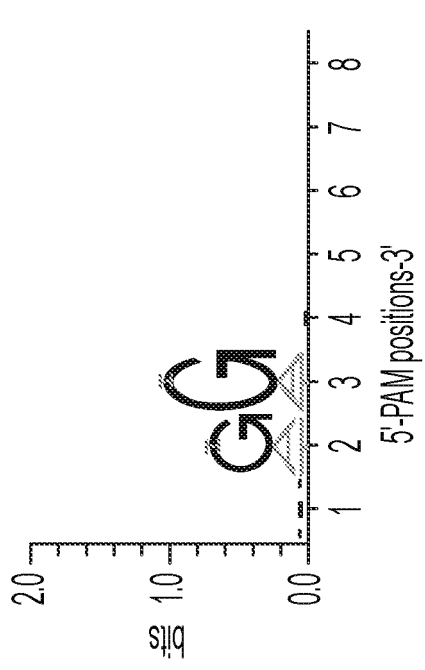


FIG. 1A

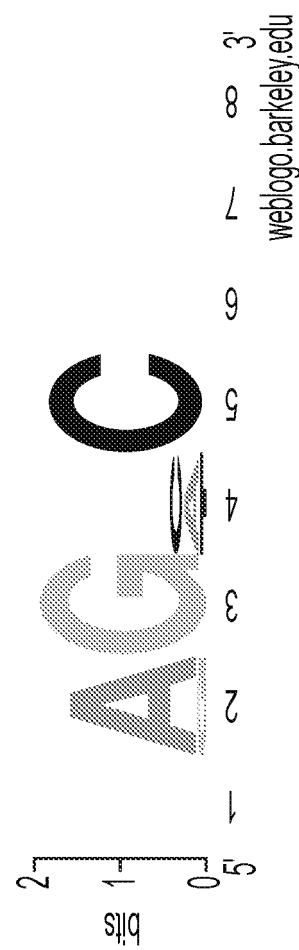


FIG. 1B

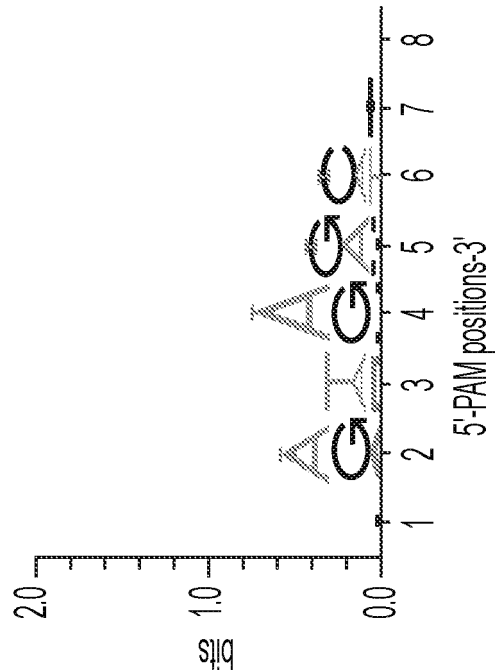


FIG. 1C

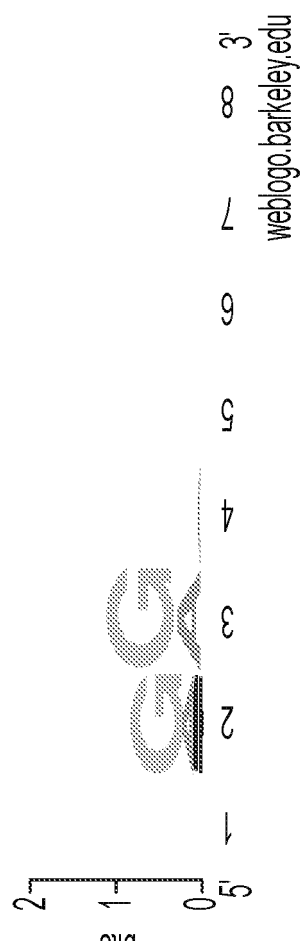


FIG. 1D

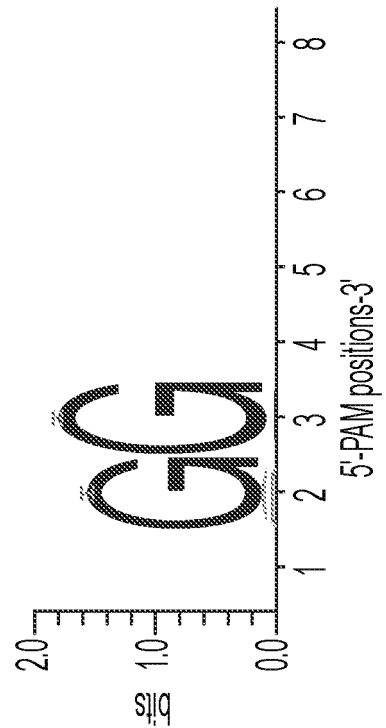


FIG. 1F

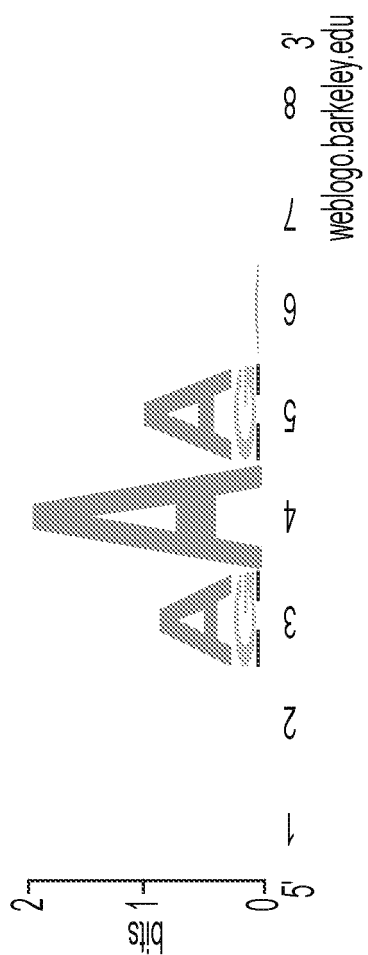


FIG. 1E

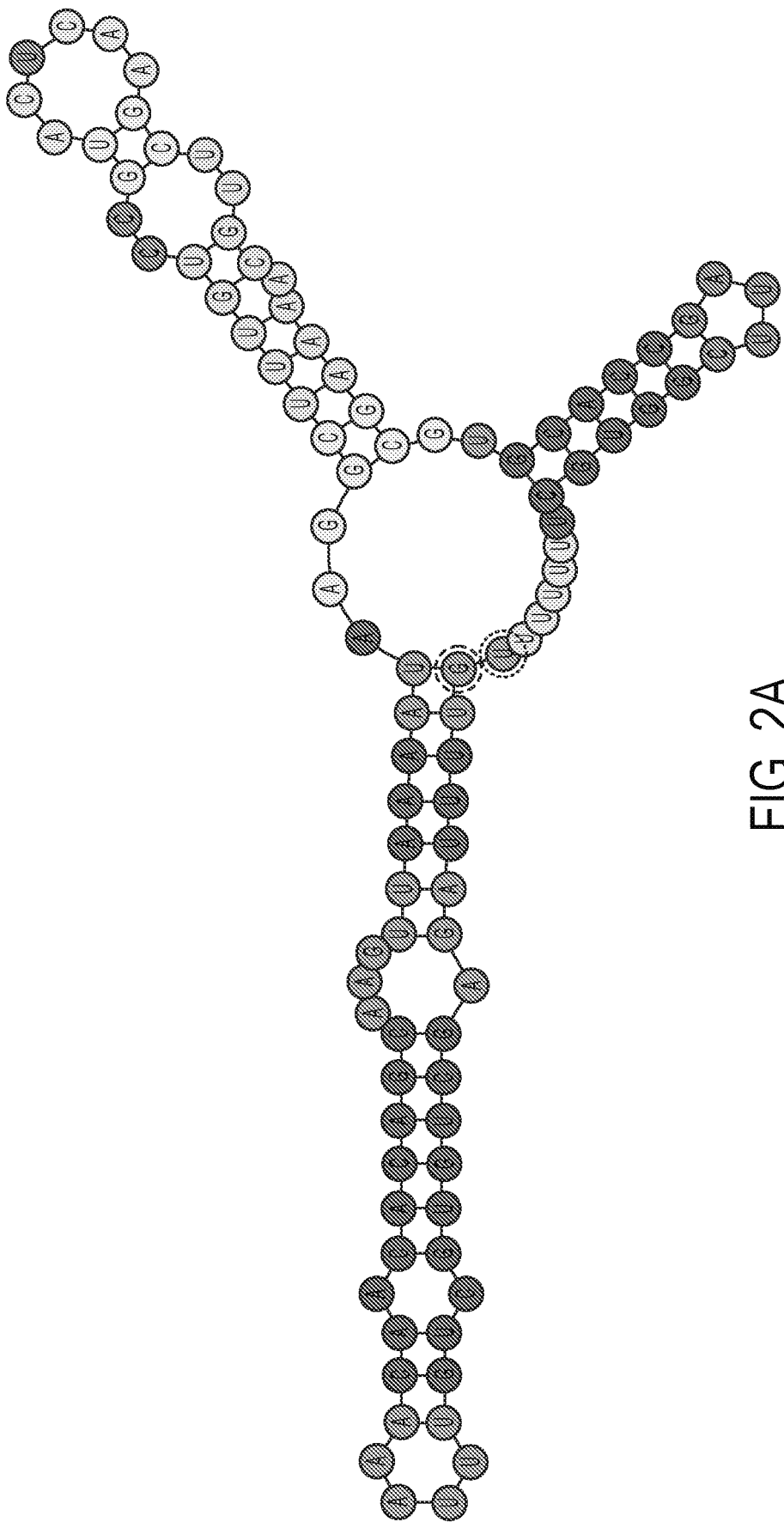


FIG. 2A

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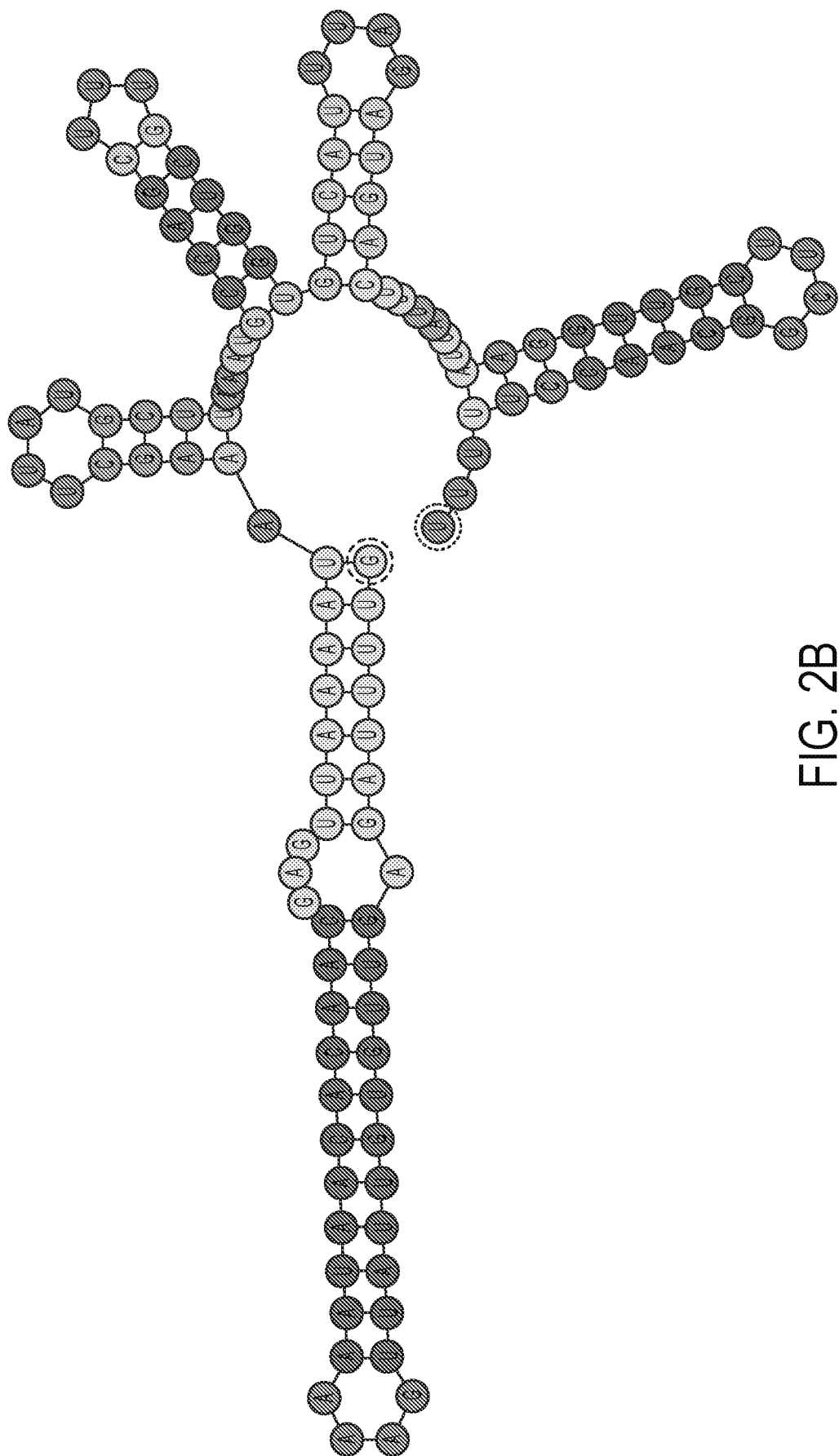


FIG. 2B

SUBSTITUTE SHEET (RULE 26)

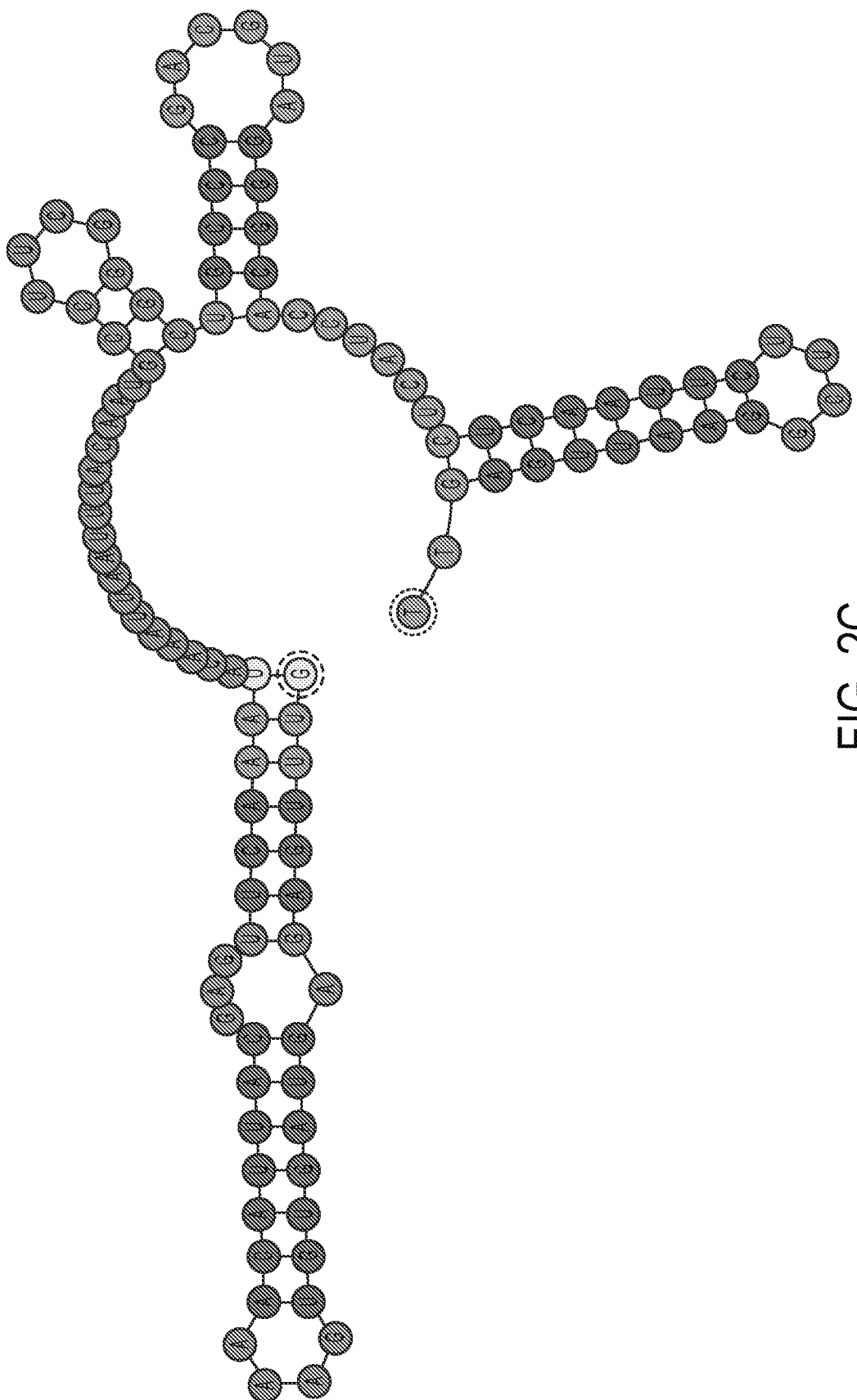


FIG. 2C

SUBSTITUTE SHEET (RULE 26)

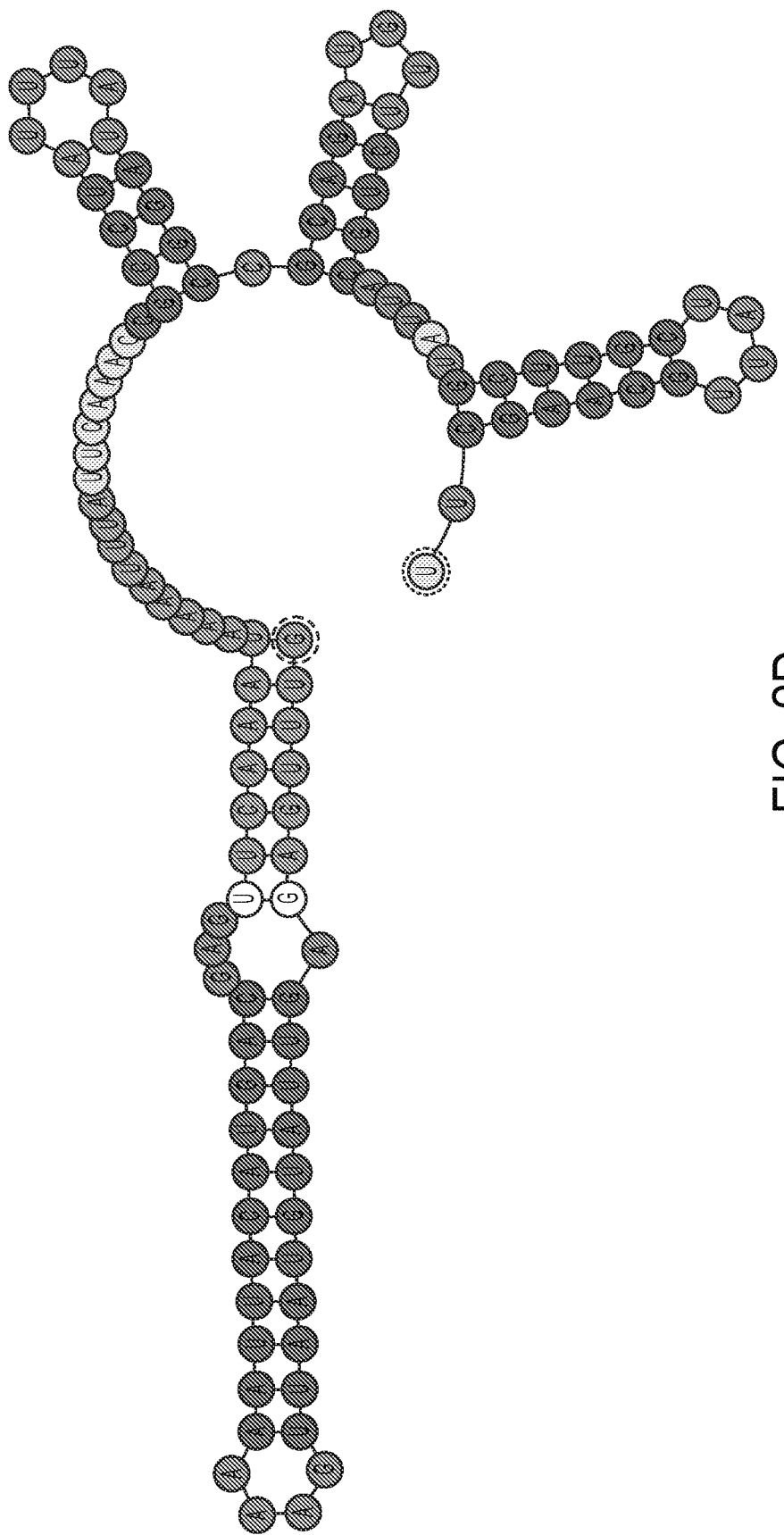


FIG. 2D

SUBSTITUTE SHEET (RULE 26)

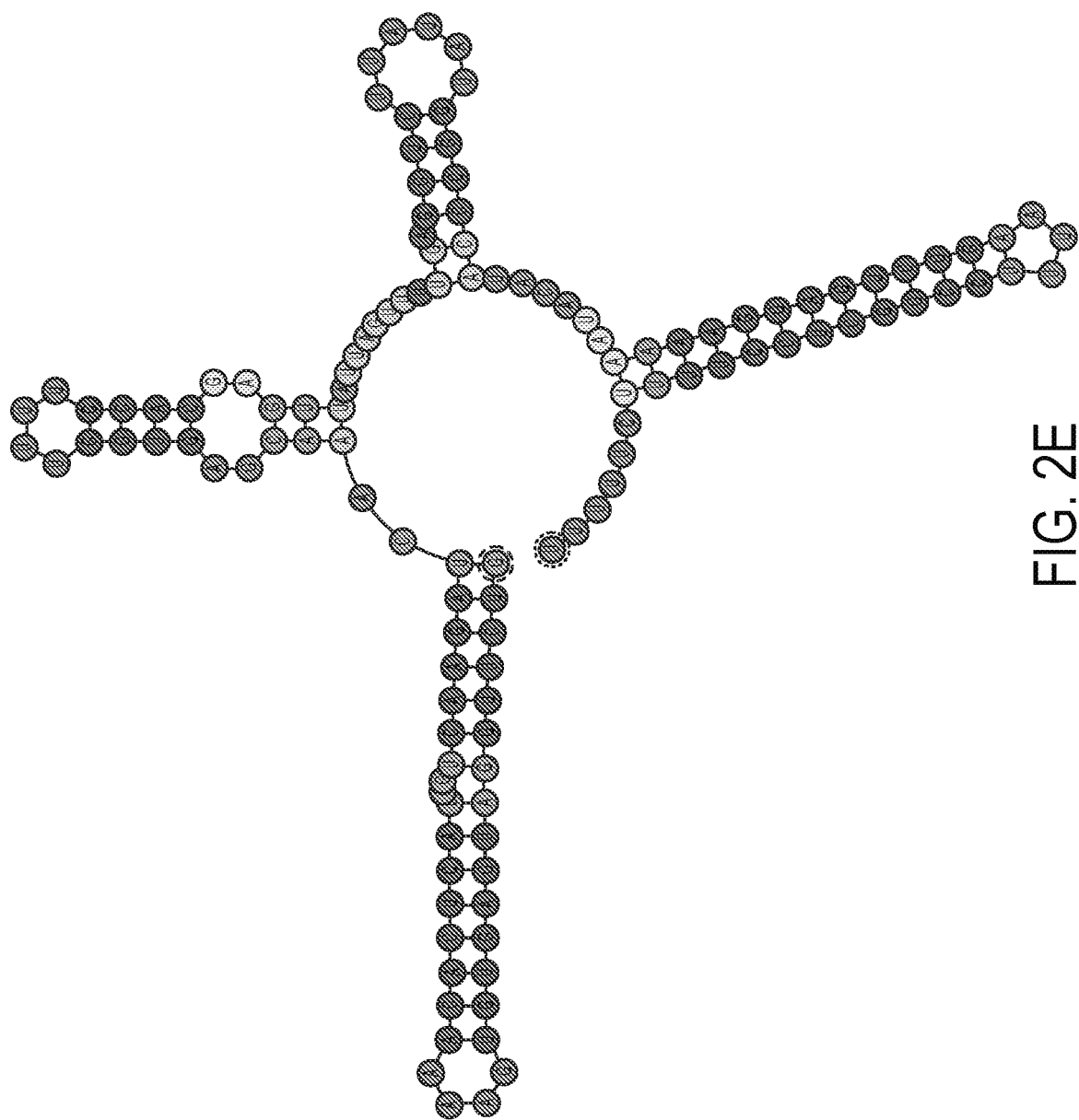


FIG. 2E

SUBSTITUTE SHEET (RULE 26)

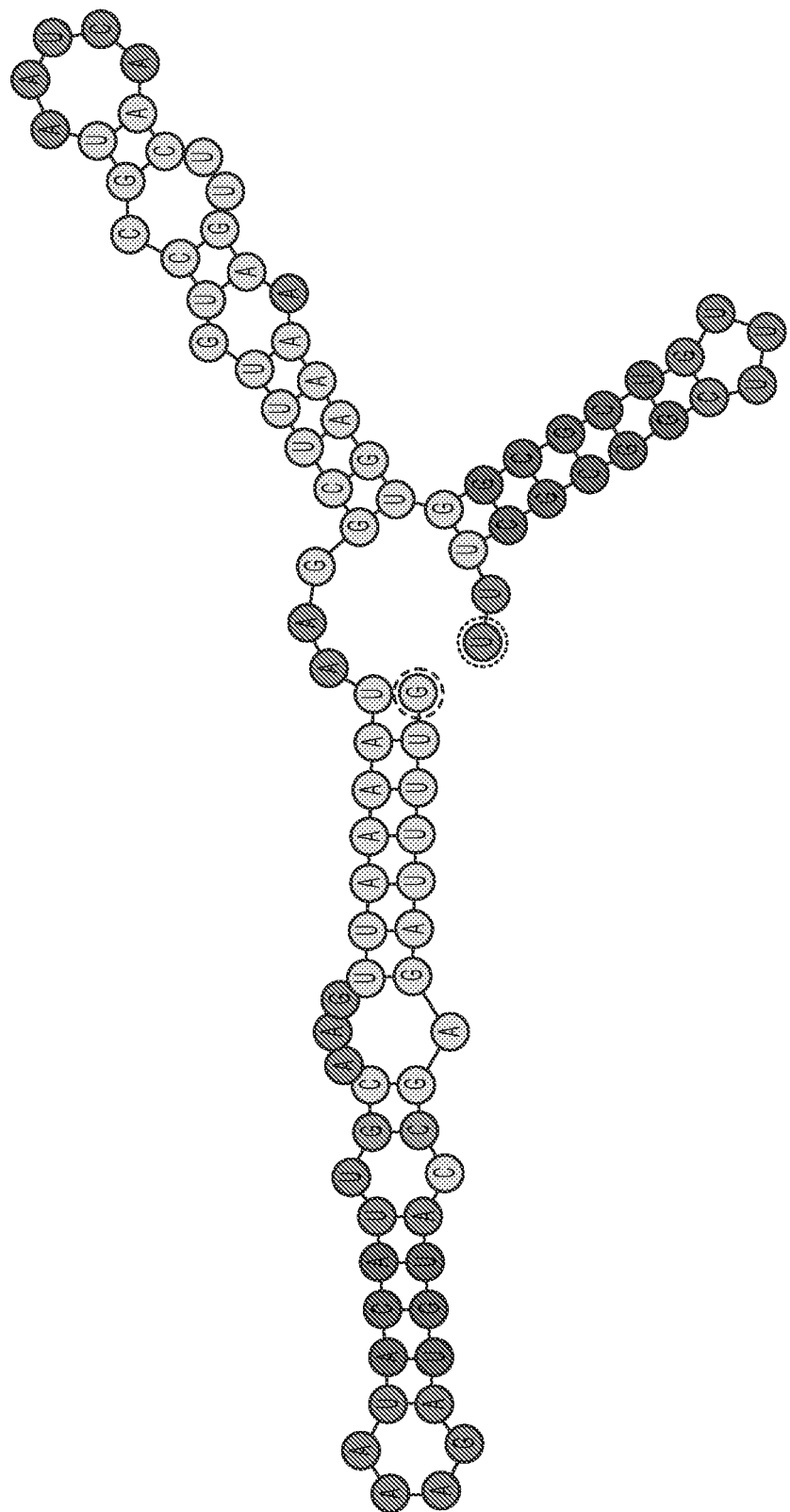


FIG. 2F

SUBSTITUTE SHEET (RULE 26)

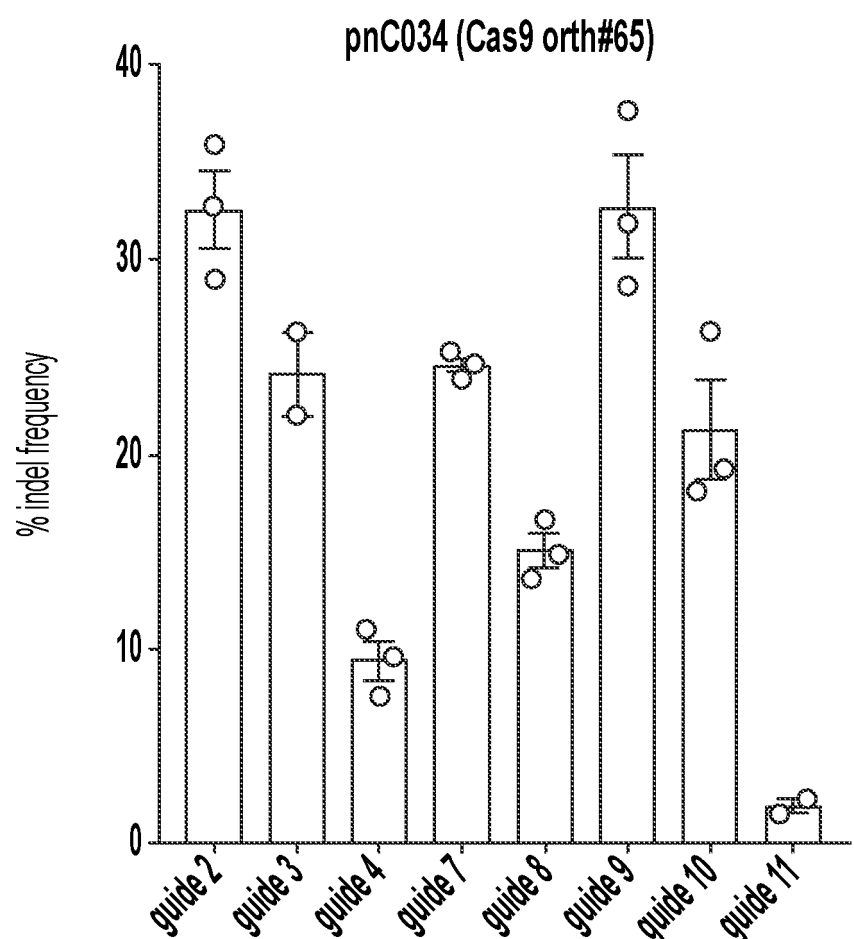


FIG. 3

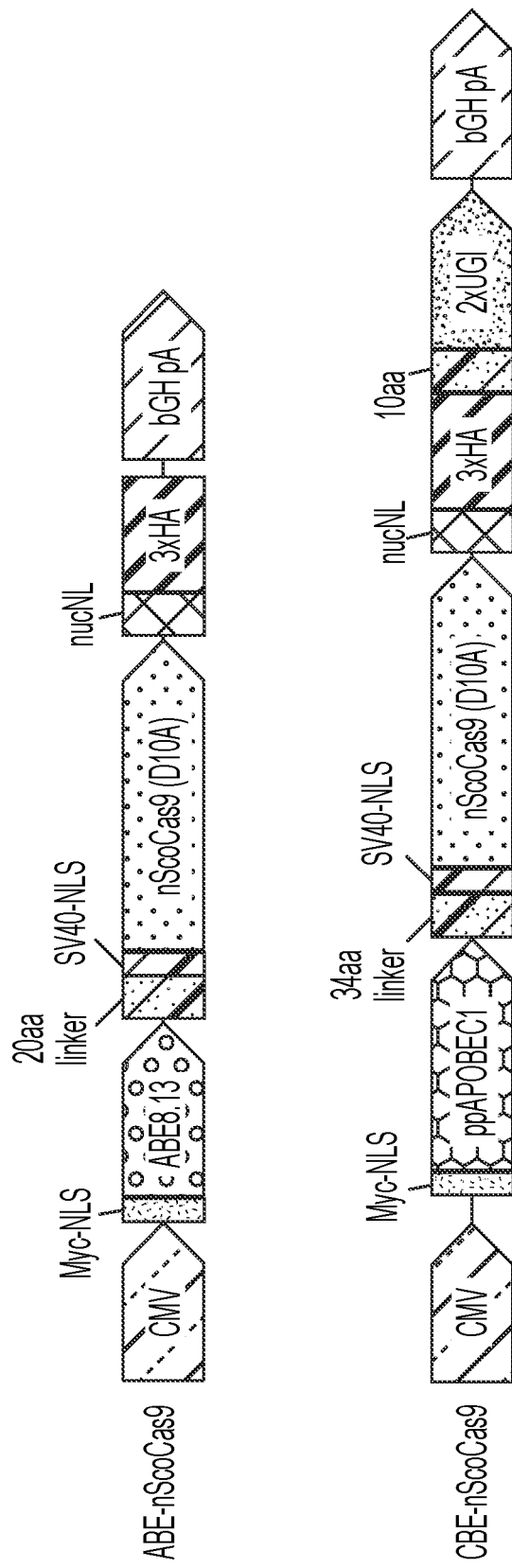
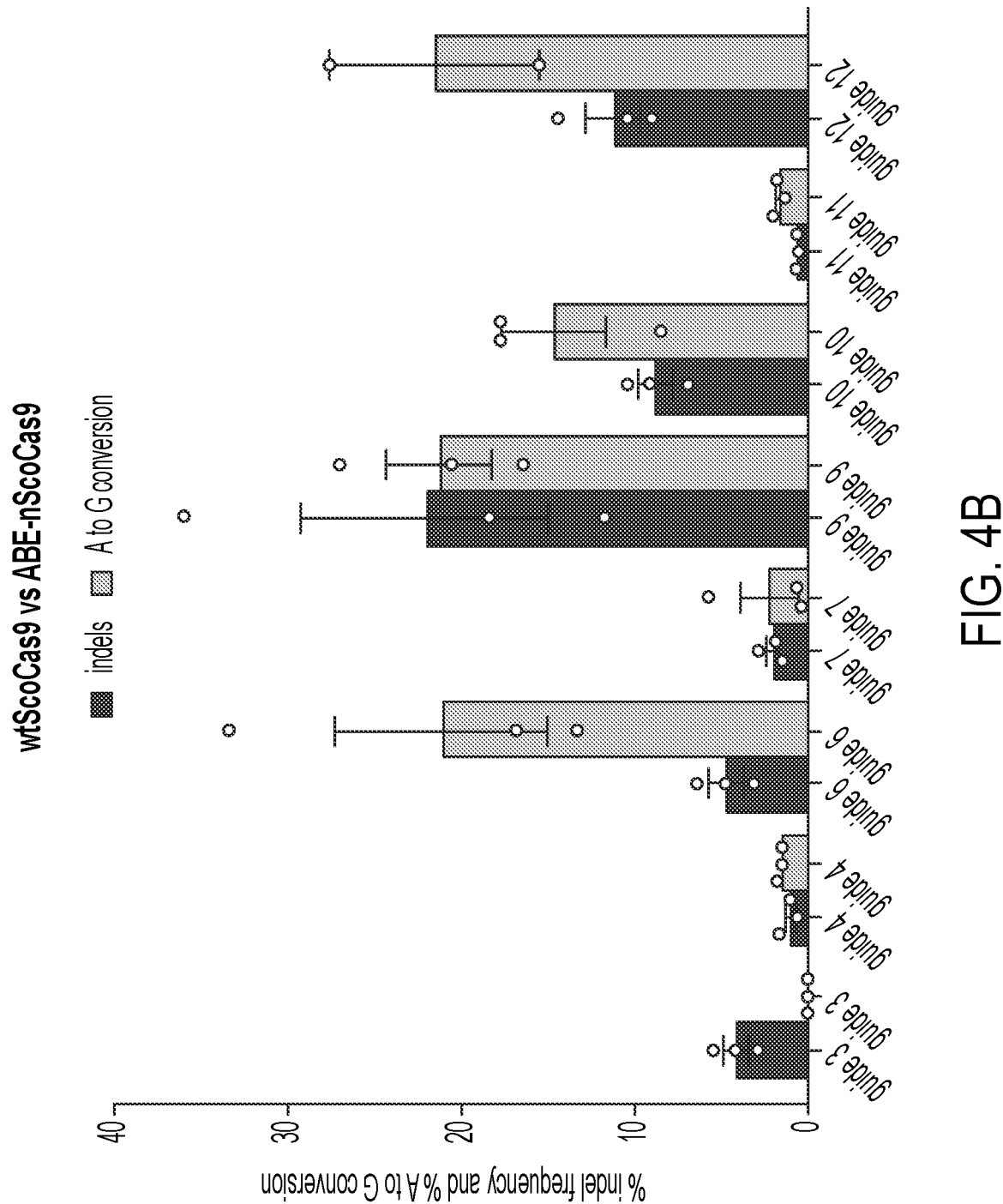


FIG. 4A



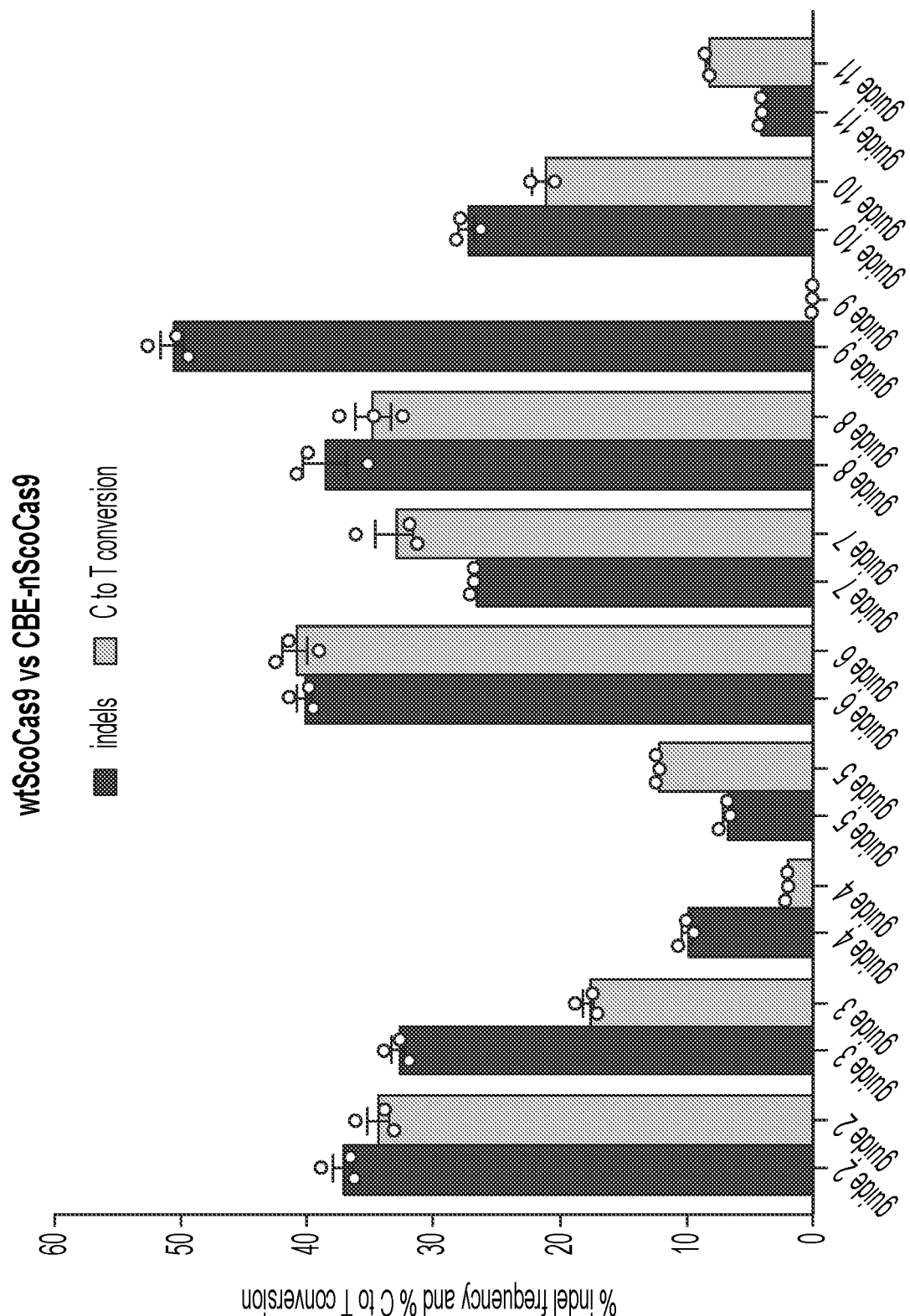


FIG. 4C

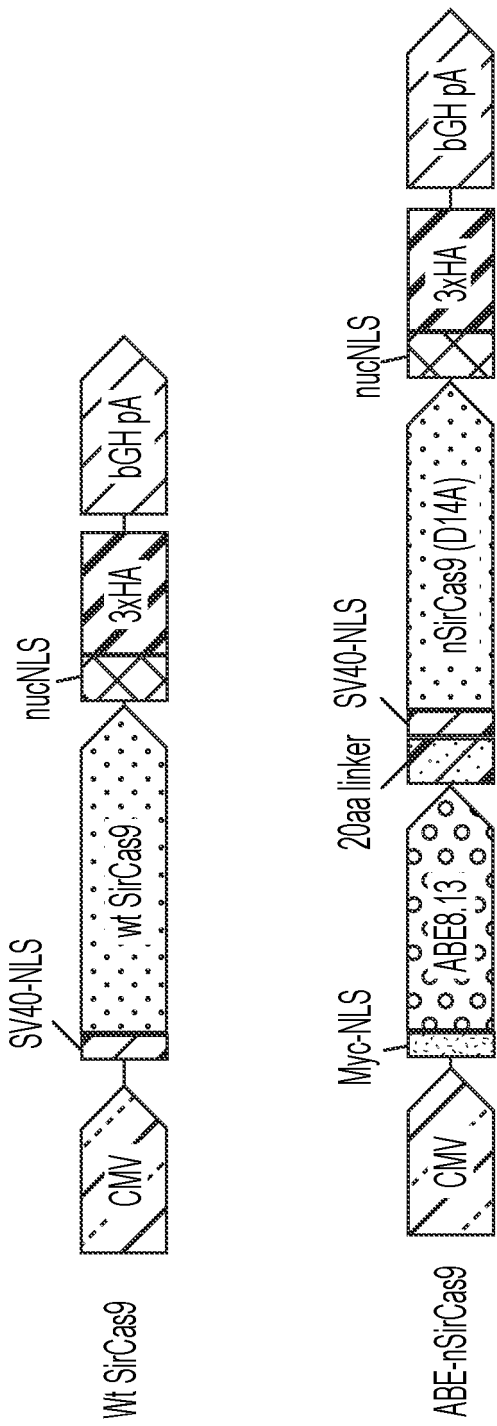
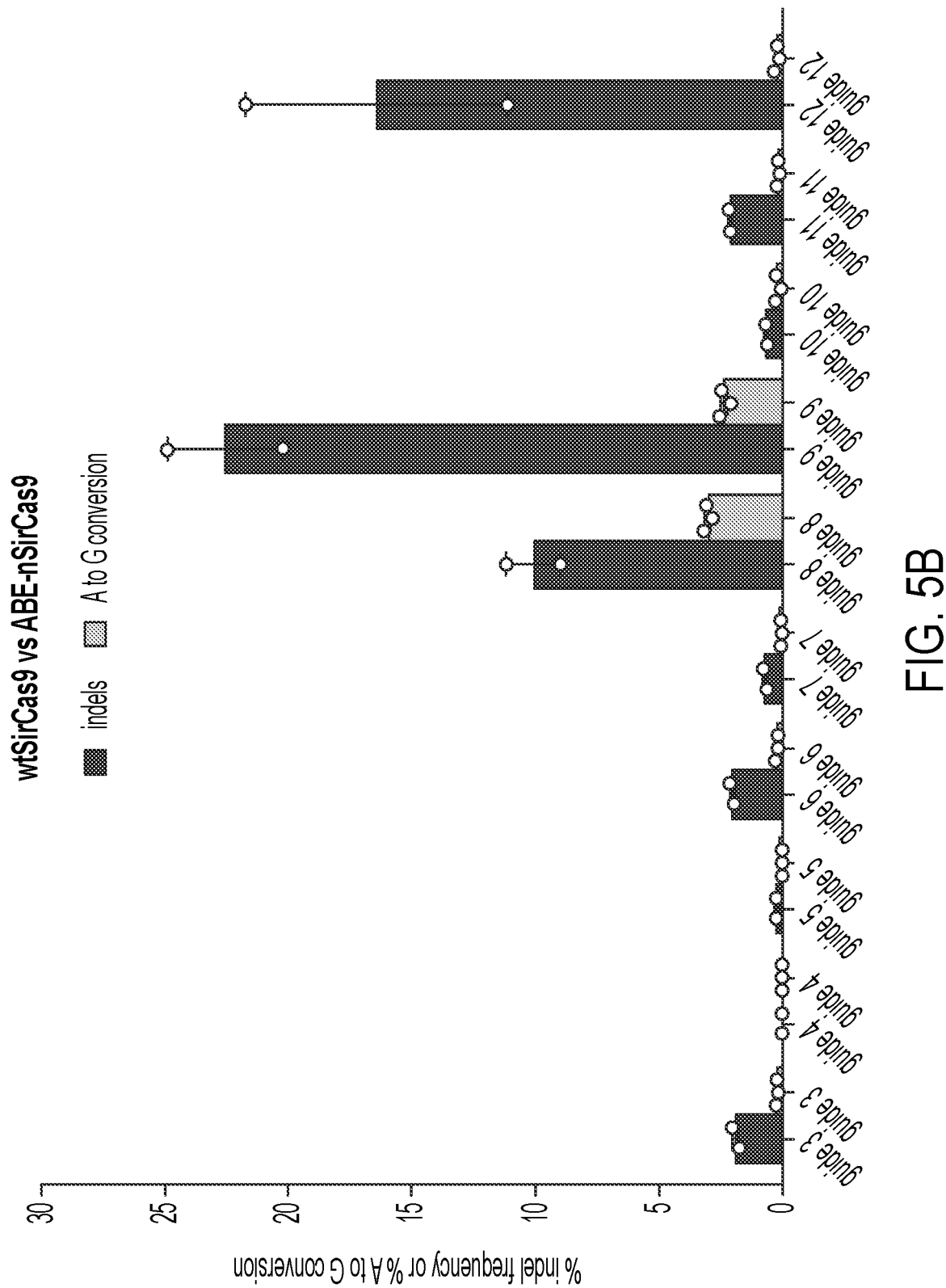


FIG. 5A



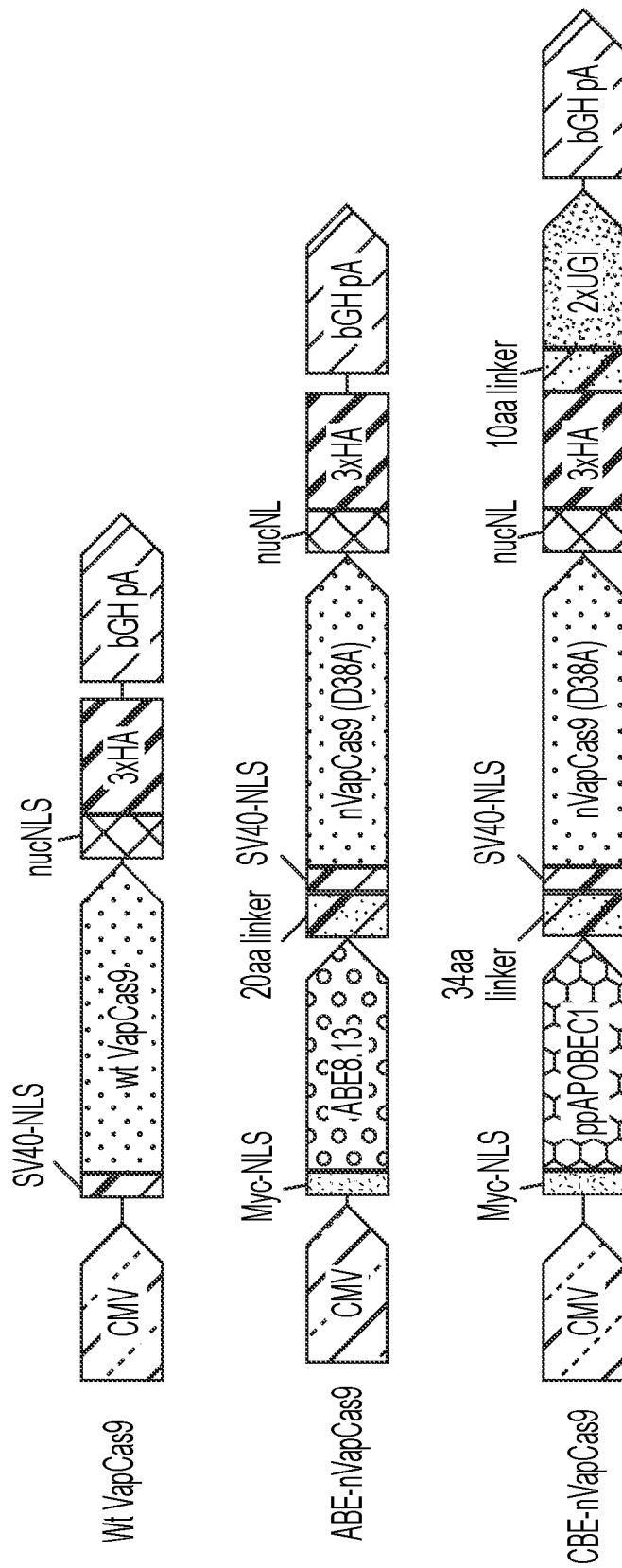
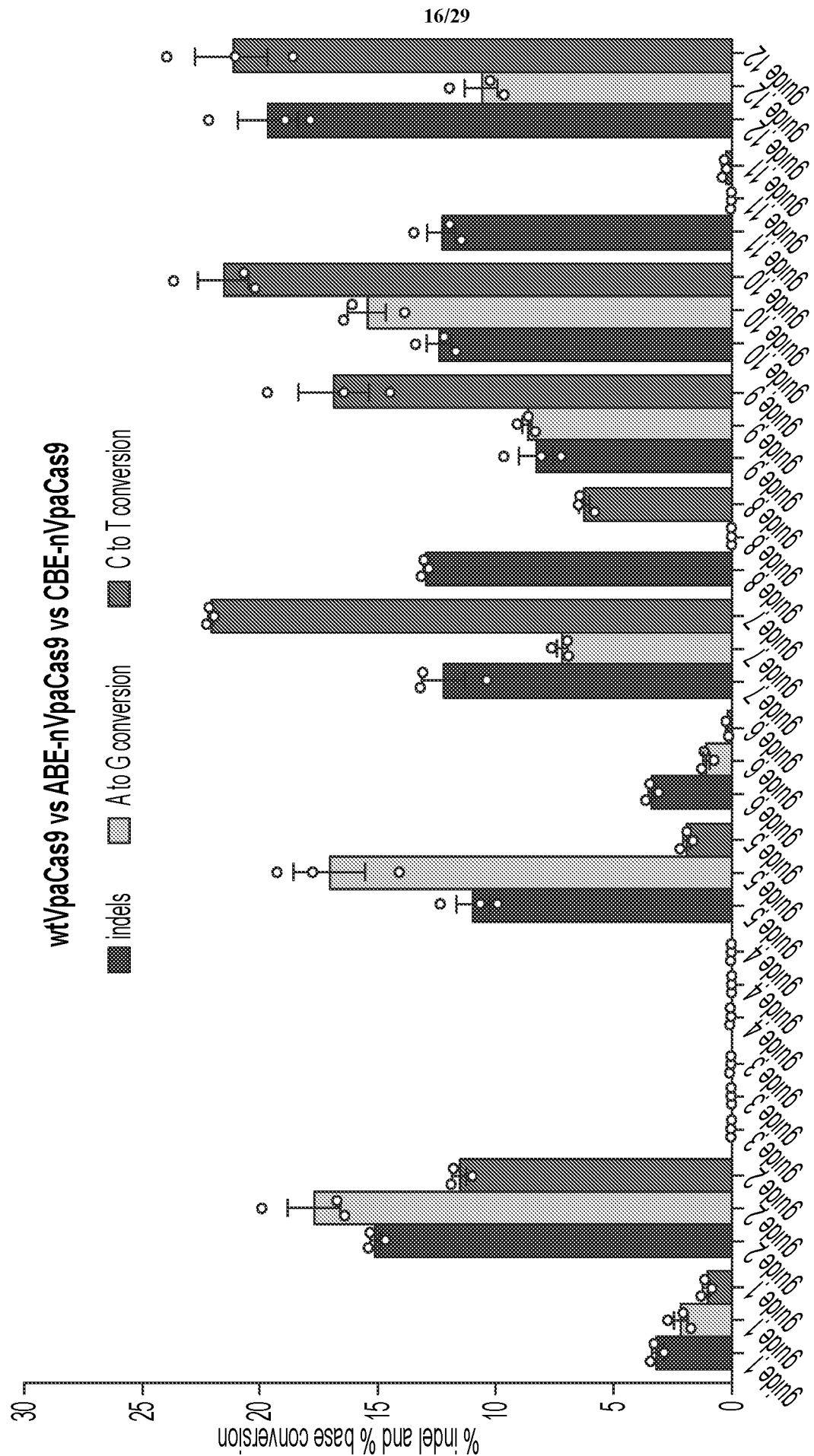


FIG. 6A

**FIG. 6B**

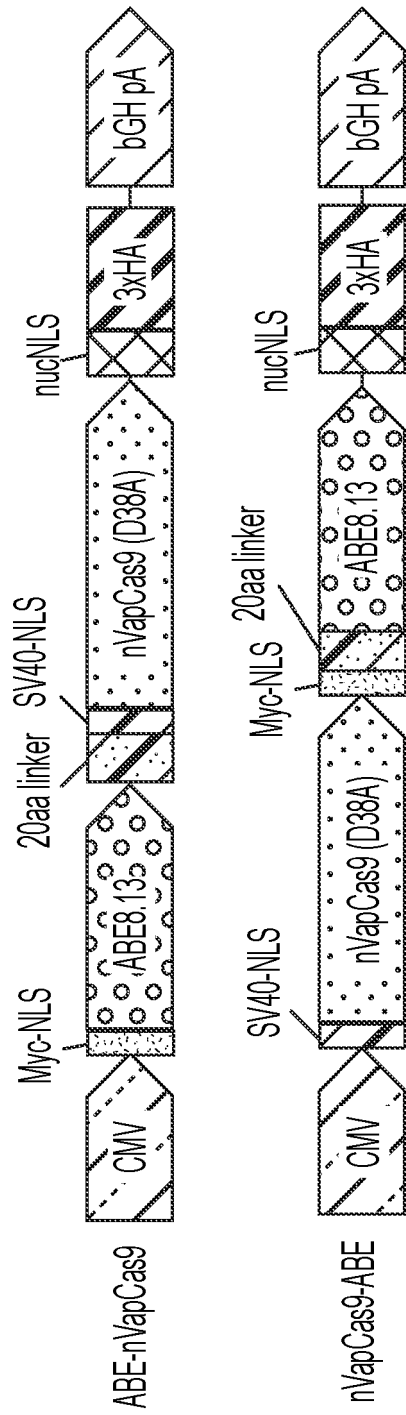


FIG. 7A

ABE-nVapCas9 vs nVapCas9-ABE

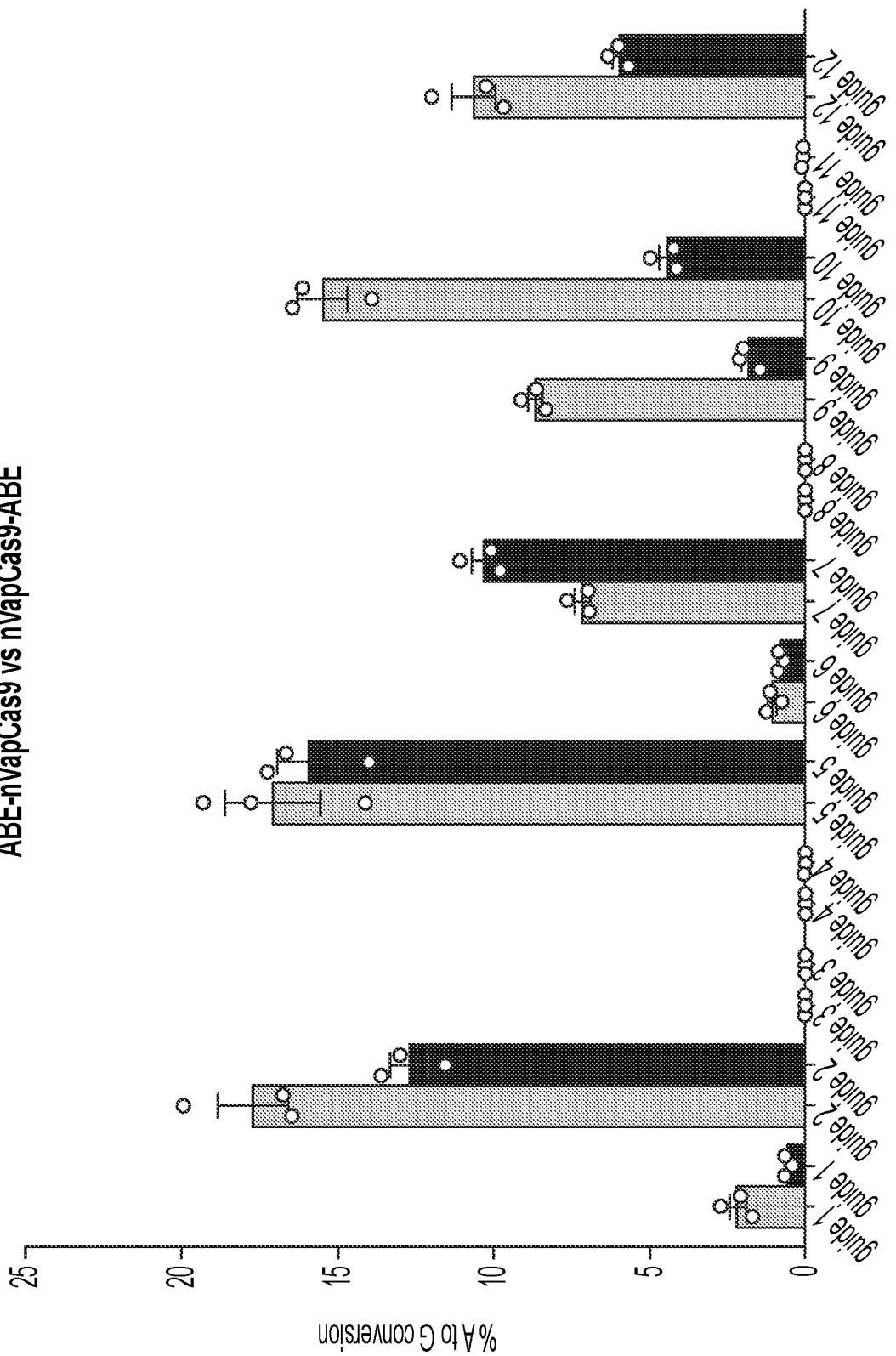


FIG. 7B

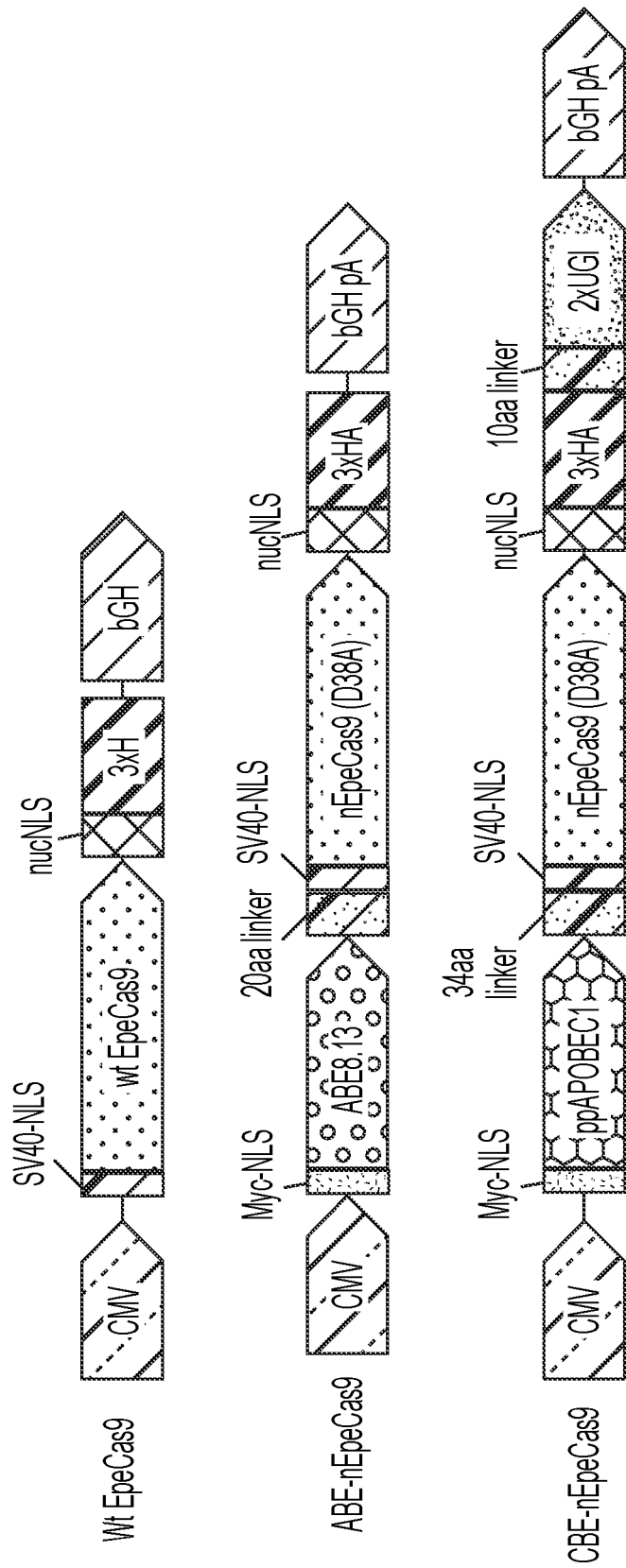


FIG. 8A

wtEpeCas9 ABE-nEpeCas9 CBE-nEpeCas9

■ Indels ■ A to G conversion ■ C to T conversion

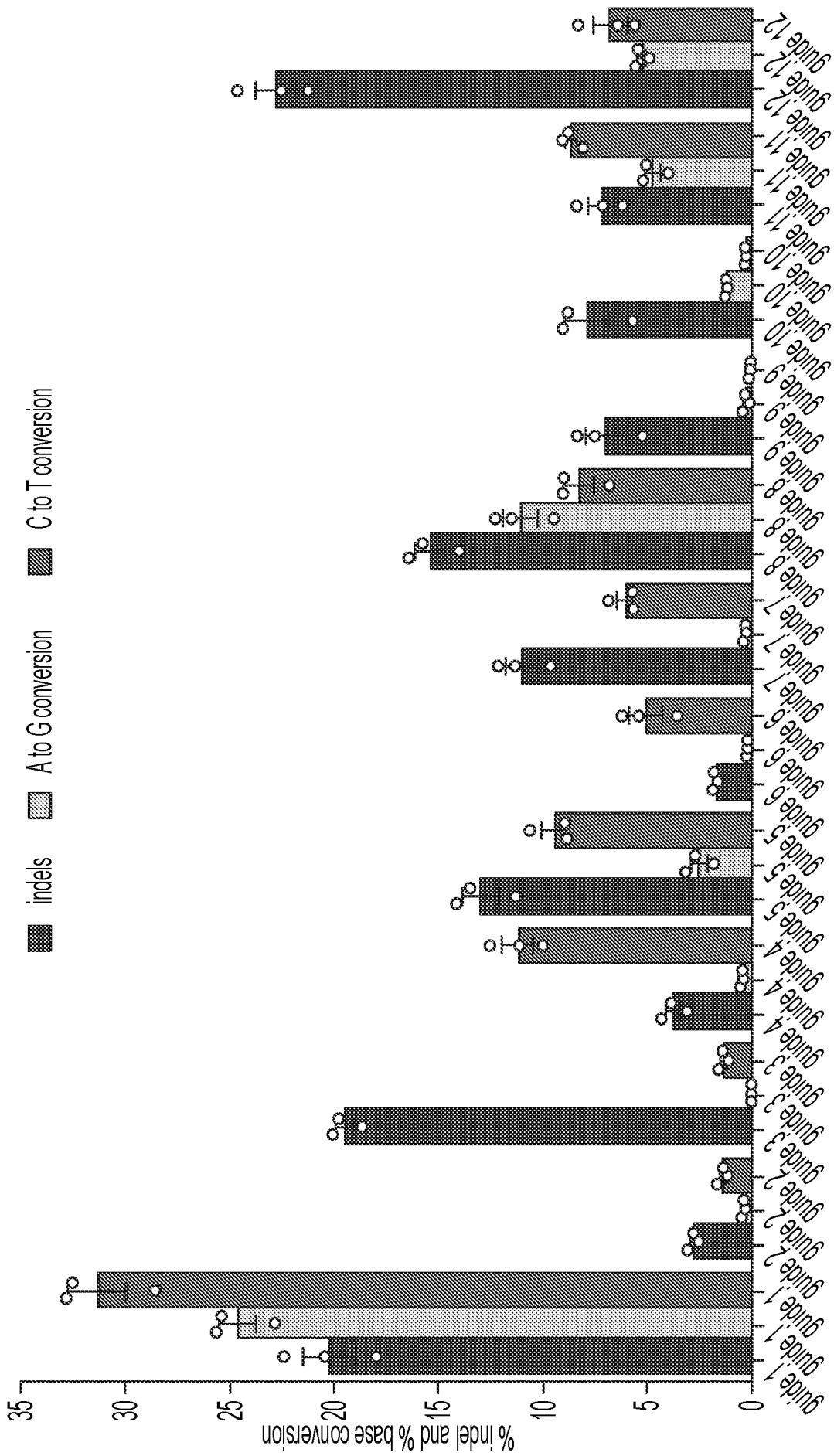


FIG. 8B

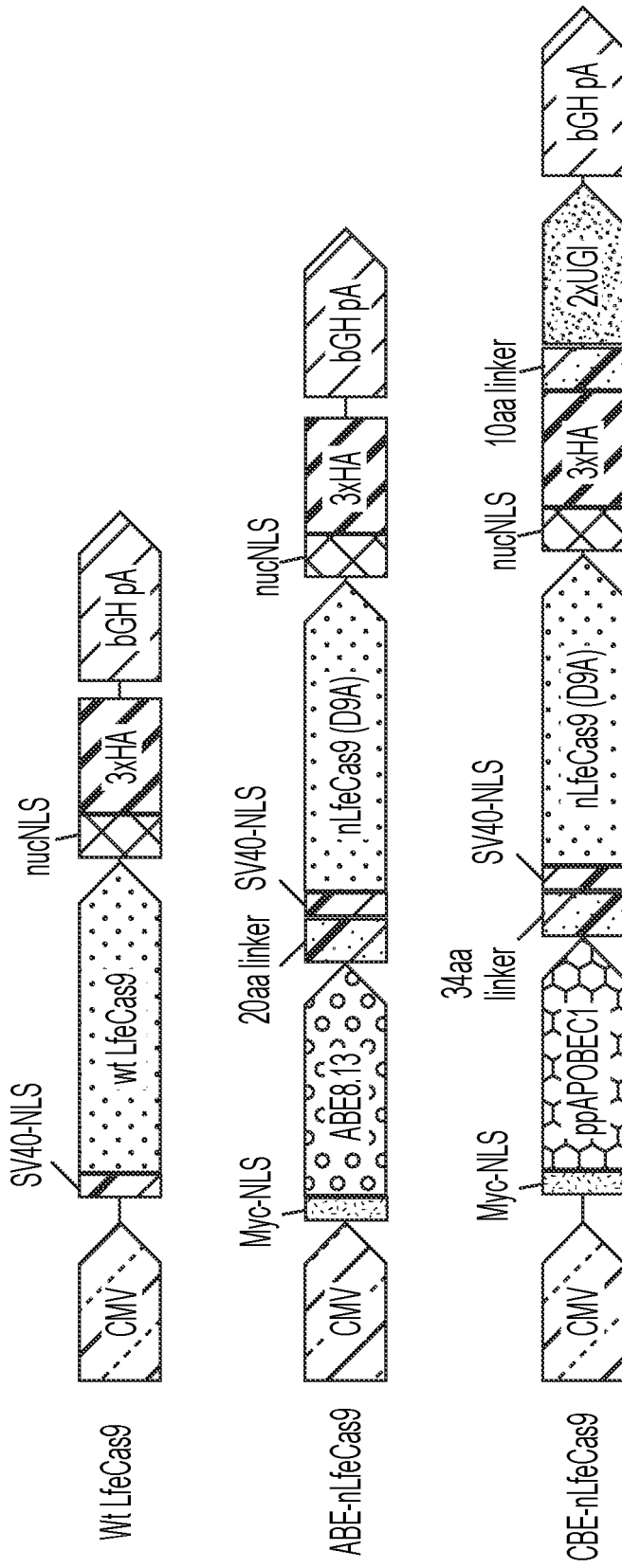
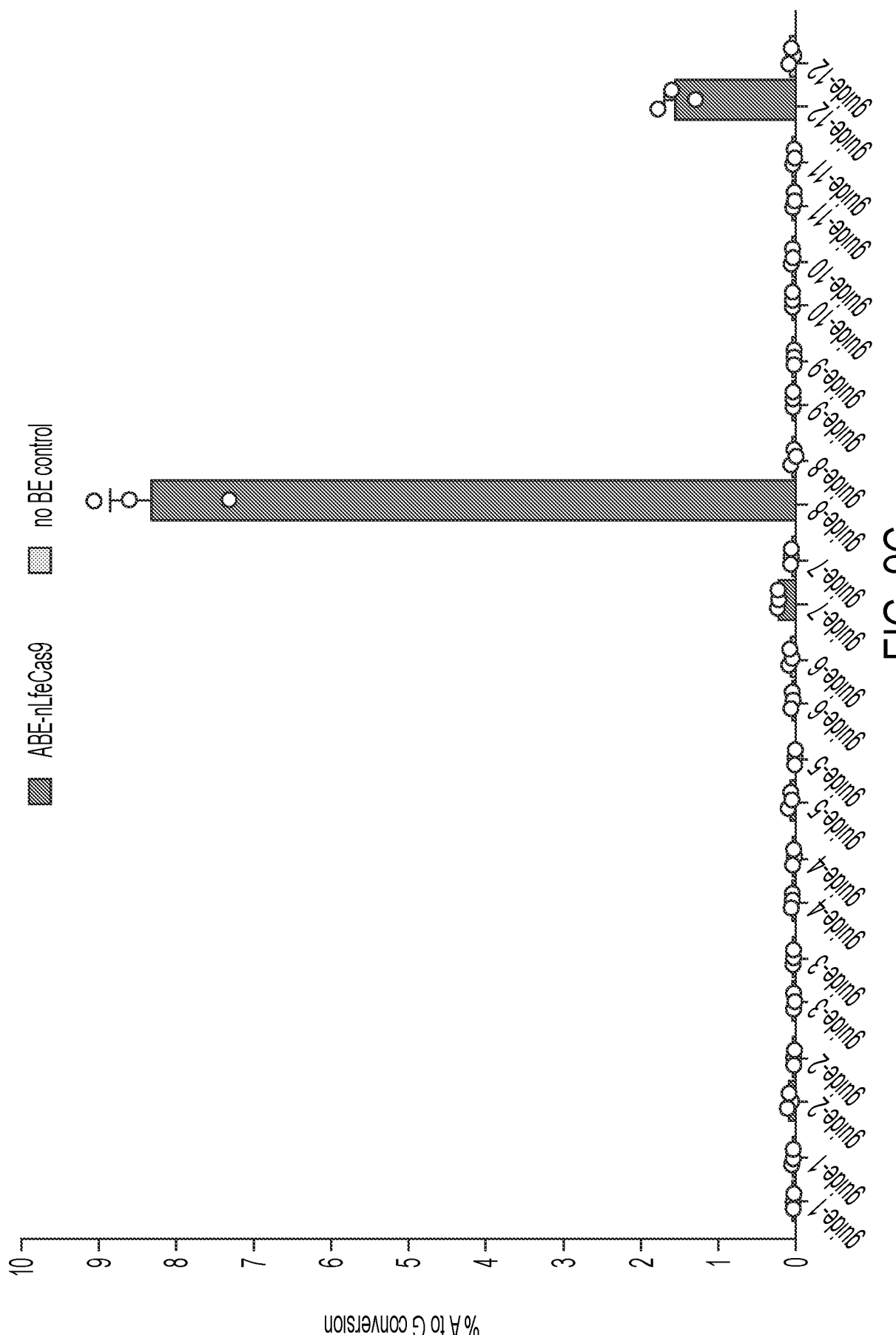


FIG. 9A





SUBSTITUTE SHEET (RULE 26)

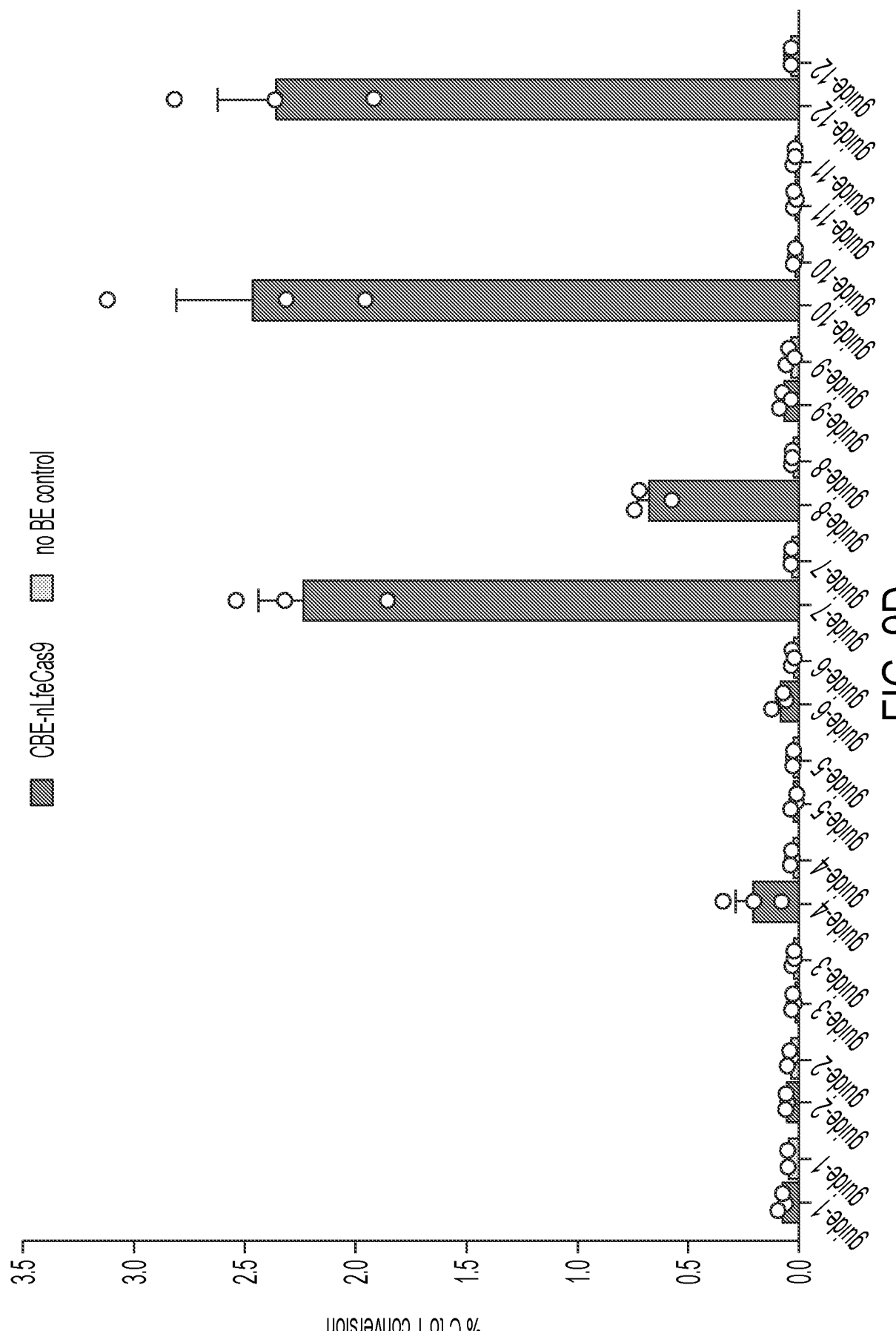


FIG. 9D

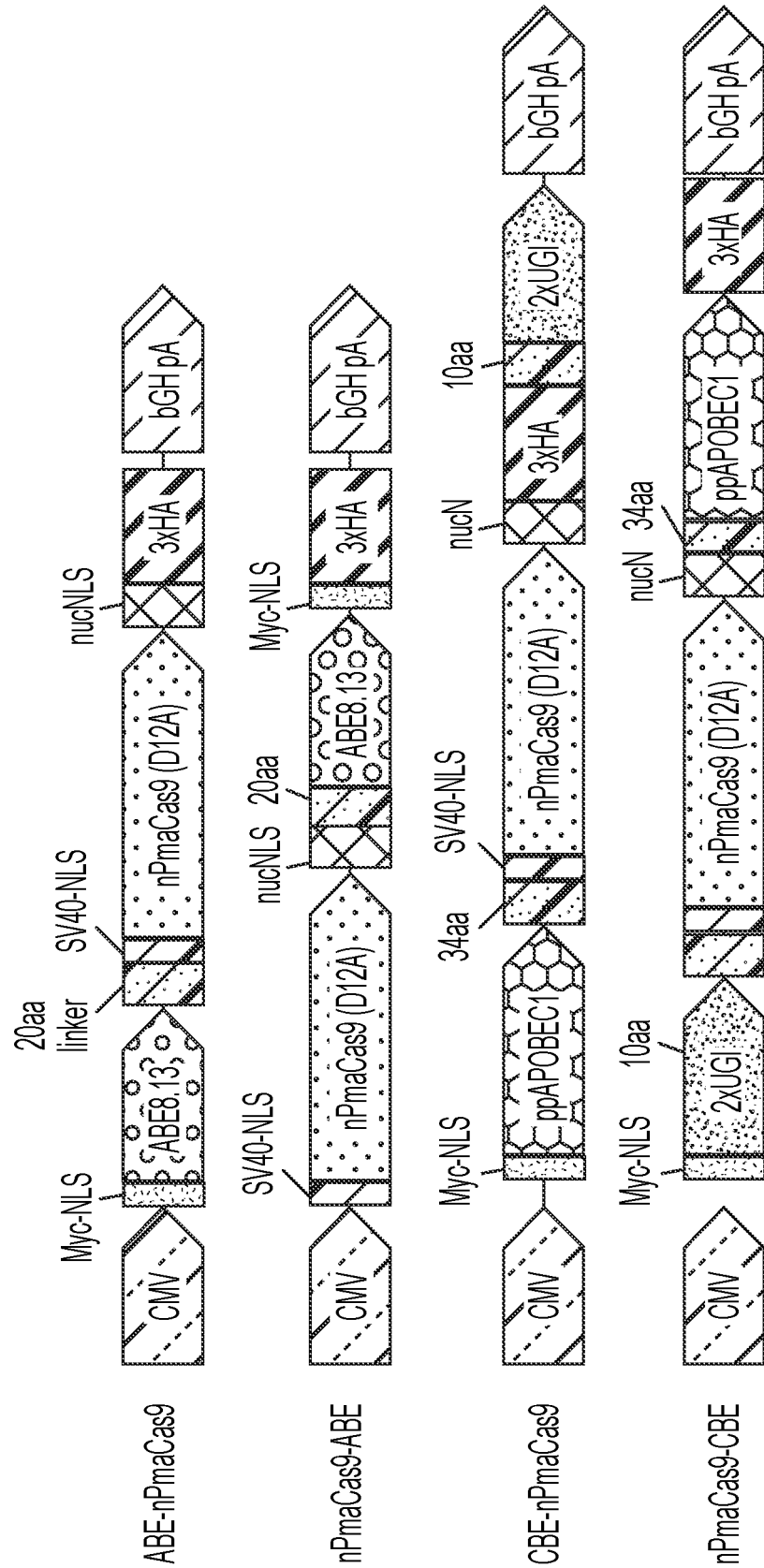


FIG. 10A

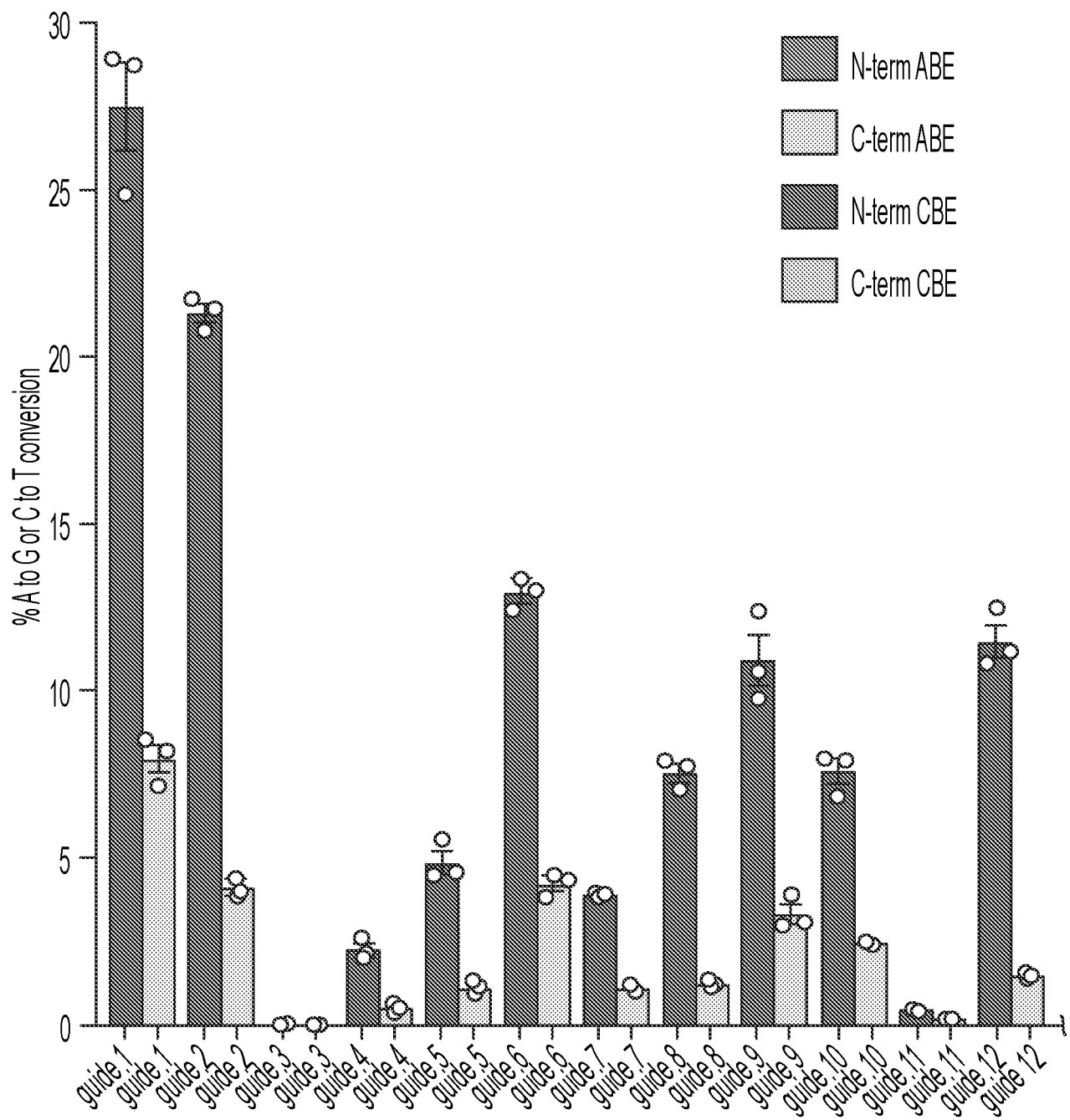
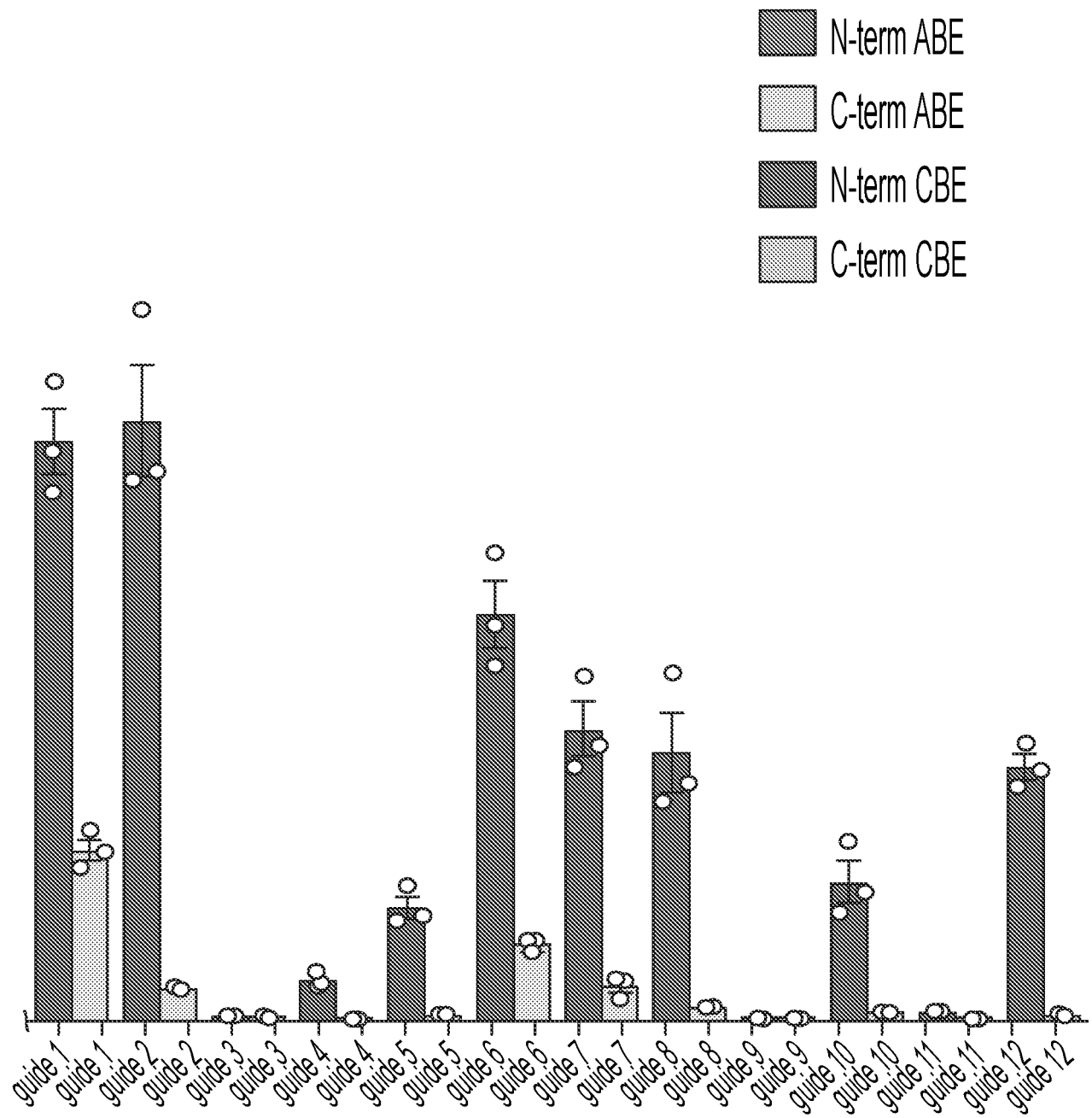


FIG. 10B



**FIG. 10B
CONTINUED**

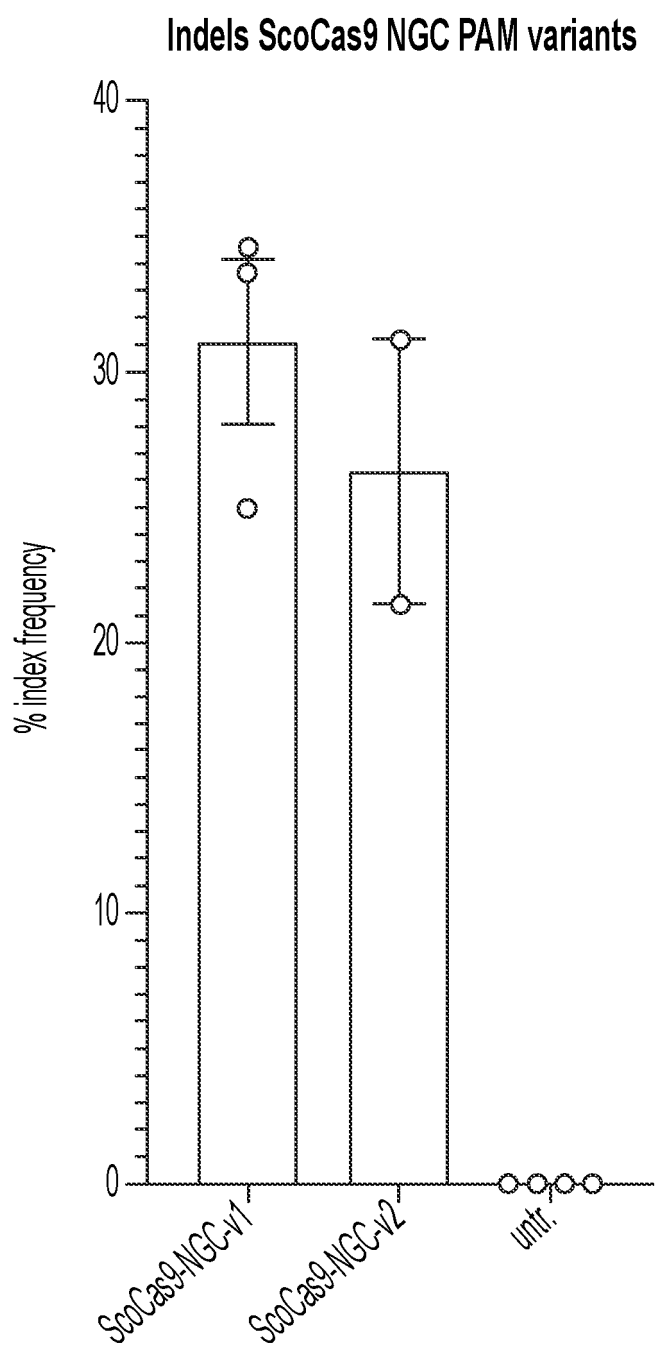
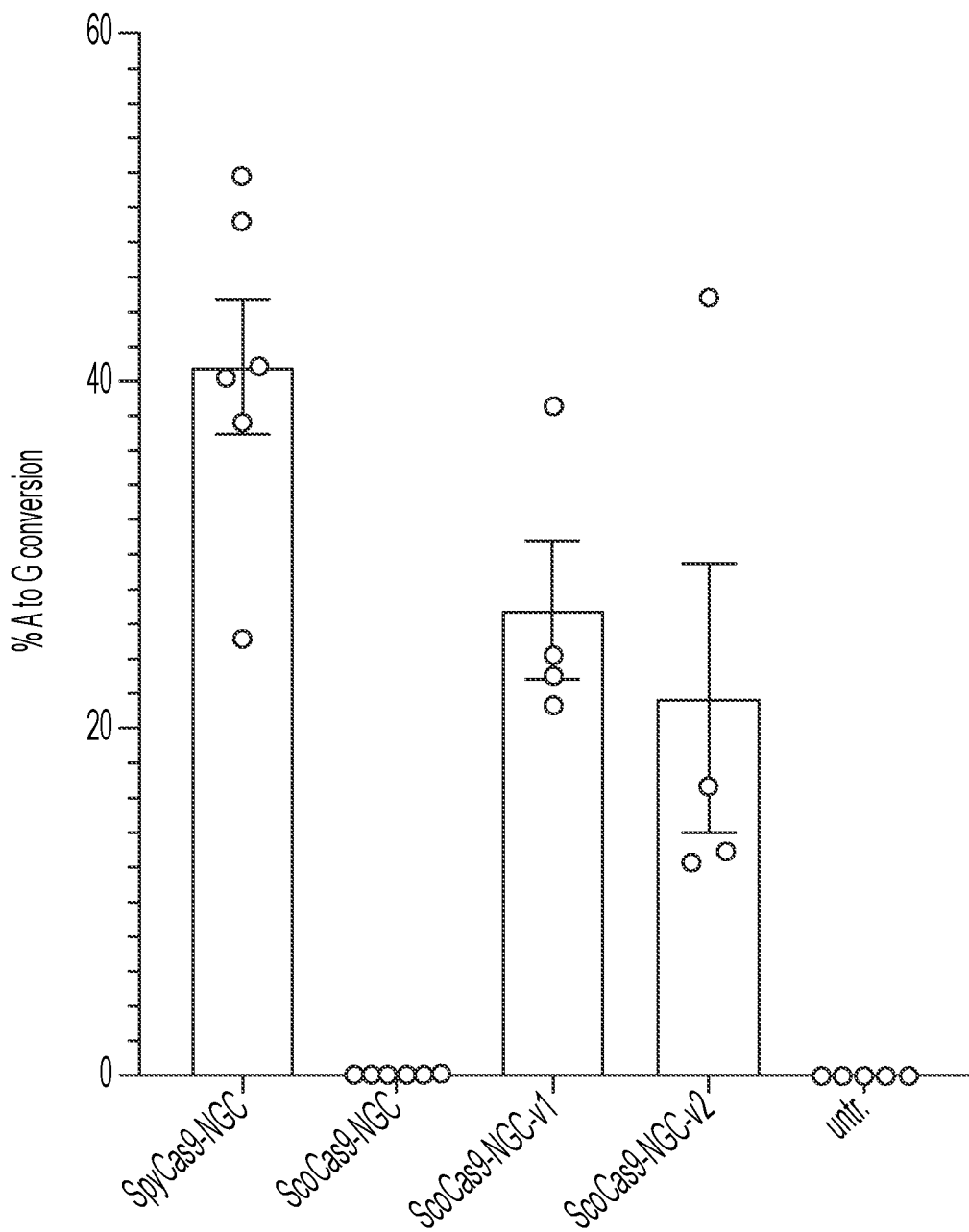


FIG. 11A

ABE ScoCas9 NGC PAM variants**FIG. 11B**