
Strand-selective base editing of human mitochondrial DNA using mitoBEs

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Supplementary Figure 1. Test the linker sequences between TALE and nickase.

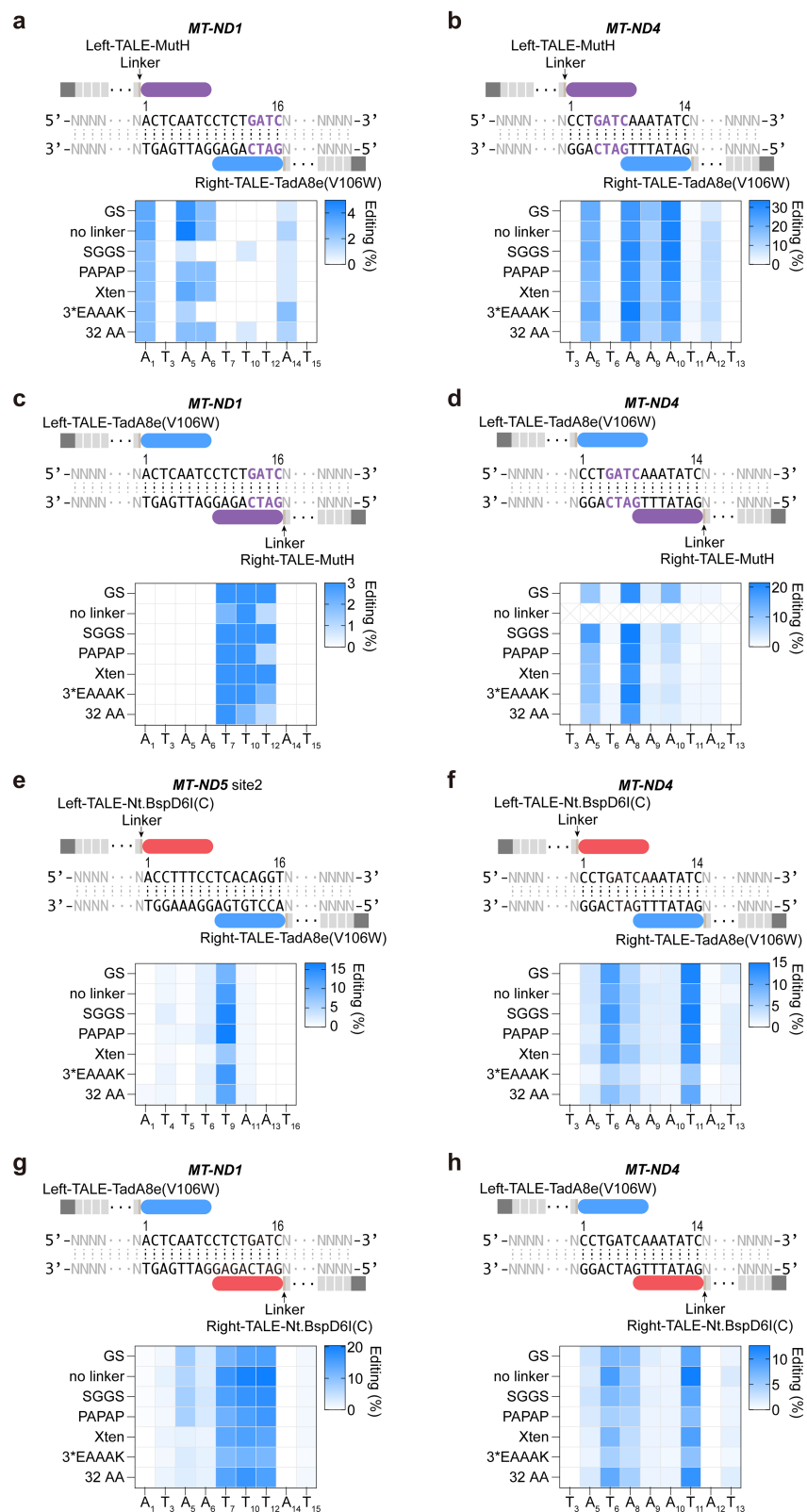
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Supplementary Figure 2. Mitochondrial genome editing specificity of DdCBEs.

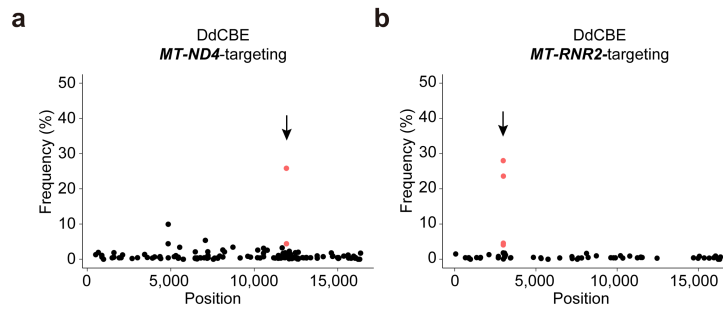
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Supplementary Sequences. The sequence of TALE-Nickases, TALE-Deaminases and monomeric mitoBEs sequence.

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Supplementary Fig. 1 Test the linker sequences between TALE and nickase. a-d, The editing efficiencies of mitoABE^{MutH} with different linkers. **e-h,** The editing efficiencies of mitoABE^{Nt.BspD6I(C)} with different linkers. The mean values from n = 3 biologically independent replicates are shown.



Supplementary Fig. 2 Mitochondrial genome editing specificity of DdCBEs. *MT-ND4*-targeting DdCBE (Left TALE-DddA-G1397-N with Left TALE-DddA-G1397-C) (**a**), *MT-RNR2*-targeting DdCBE (Left TALE-DddA-G1397-N with Left TALE-DddA-G1397-C) (**b**). For **a** and **b**, data are presented as mean values of $n = 3$ independent biological replicates. The arrow points to the targeted editing site and the red dots represent the editing efficiency of cytosines in the editing window.

Supplementary Sequences.

The sequence of TALE-Nickases, TALE-Deaminases and monomeric mitoBEs sequence.

The TALE repeat array sequences are shown in Supplementary Table. 2

TALE-Effector sequence:

MTS-Tag-TALE N-terminal non-repeat-TALE Repeat array-TALE C-terminal non-repeat-2aa
linker-Nickase/Deaminase

Non-Targeting TALE-Effector sequence:

MTS-Tag-TALE N-terminal non-repeat-TALE C-terminal non-repeat-2aa linker-Nickase/Deaminase

MTS:

>SOD2 MTS

MALSRAVCGTSRQLAPVLGYLGSRQKHSLPD

>COX8 MTS

MASVLTPLLLRGLTGSAARRLPVPRAK

Tag:

>3×HA

YPYDVPDYAGYPYDVPDYAGYPYDVPDYA

>3×FLAG

IHSLDYKDHDGDYKDHDIDYKDDDDKMDIA

> TALE N-terminal non-repeat

MDIADLRTLGYSSQQQEKIKPKVRSSTVAQHHEALVGHGFTHAHIVALSQHPAALGTVAVK
YQDMIAALPEATHEAIVGVGKQWSGARALEALLTVAGELRGPPLQLDTGQLLKIARGGVT
AVEAVHAWRNALTGAPLN

>TALE Repeat array (e.g., 20mer)

LTPEQVVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQALETVQRLLPVL
CQAHGLTPEQVVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQALETVQ
RLLPVLCQAHGLTPEQVVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQ
ALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS
??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQ
VVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQALETVQRLLPVLCQAH
GLTPEQVVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQALETVQRLLPV
LCQAHGLTPEQVVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQALETV
QRLLPVLCQAHGLTPEQVVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS??GGK
QALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQALETVQRLLPVLCQAHGLTPEQVVAIA
S??GGKQALETVQRLLPVLCQAHGLTPEQVVAIAS??GGKQALE

> TALE N-terminal non-repeat

SIVAQLSRPDPALAALTNDHLVALACLGGRPALDAVKKGLG

>2aa linker

GS

>SGGS linker

SGGS

>PAPAP linker

PAPAP

>Xten linker

SGSETPGTSESATPES

>3*EAAAK linker

AEAAAKEAAAKEAAAKEAAAKA

>32AA linker

GSGSSGGSSGSETPGTSESATPESSGGSSGGS

Nickase sequences

>MutH

MSQPRPLLSPPETEEQLLAQAQQLSGYTLGELAALVGLVTPENLKRDKGWIGVLLEIWL
GAS
AGSKPEQDFAALGVELKTIPVDSLGRPLETTFVCVAPLTGNSGVTWETSHVRHKLKRVLWIP
VEGERSIPLAQRRVGSPLLWSPNEEDRQLREDWEELMDMIVLGQVERITARHGEYLQIRPK
AANAKALTEAIGARGERILTLPRGFYLKKNFTSALLARHFLIQ

>Nt.BspD6I(C)

RQLEEVIDLLEVYHEKKNVIEEKIKARFIANKNTVFEWLTWNGFIILGNALEYKNNFVIDEEL
QPVTHAAGNQPDMEIYEDFIVLGEVTTSTKGATQFKMESEPVTRHYLNKKKELEKQGVEKE
LYCLFIAPEINKNTFEEFMKYNIVQNTRIPLSLKQFNMLLMVQKKLIEKGRRLSSYDIKNLM
VSLYRTTIECERKYTQIKAGLEETLNNWVVDKEVRF

>Fok1-Fok1(D450A)

FKQLVKSELEKKSELRHKLKYVPHEYIELIEIARNSTQDRILEMKVMEFFMKVYGYRGKHL
GGSRKPDGAIYTVGSPIDYGVIVDTKAYSGGYNLPIGQADEMQRYVEENQTRNKHINPNEW
WKVYPSSVTEFKFLFVSGHFKGNYKAQLTRLNHITNCNGAVLSVEELLIGGEMIKAGTLTLE
EVRRKFNNGEINFSSGGGGSGGGGSGGGGSGGGGSSGGGGSGGGGSQLVKSELE
EKKSELRHKLKYVPHEYIELIEIARNSTQDRILEMKVMEFFMKVYGYRGKHLGGSRKPA
GAI
YTVGSPIDYGVIVDTKAYSGGYNLPIGQADEMQRYVEENQTRNKHINPNEWWKVYPSSVTE
FKFLFVSGHFKGNYKAQLTRLNHITNCNGAVLSVEELLIGGEMIKAGTLTLEEVR
RKFNNGE
INF

>Nb.BsaI(C, N441D/R442G)

CRSHADRGRWEKNLRSYTTDRRAFEYWVDGDWVAADKLMGLIRTNEQIKKETCLNDNHP
GPCSADHIGPISLGFVHRPEFQLLCNSCNSAKNNRMTFSDVQHLINAENNGEEVASWYCKHI
WDLRKHDVKNNENALRLSKILRDNRHTAMFILSELLKDNHYLFLSTFLGLQYAERSVSFSNI
KIENHIITGQISEQPRDTKYTEEQKARRMRIGFEALKSYIEKEDGNALLVINDKIIDKINEIKNIL
QDIPDEYKLLNEKISEQFNSEEVSDELLRDLVTHLPTKESEPANFKLARKYLQEIMEIVGDEL
SKMWEDERYVRQTFADLD

>Nt.BsaI(C, R236D)

CRSHADRGRWEKNLRSYTTDRDAFEYWVDGDWVAADKLMGLIRTNEQIKKETCLNDNHP
GPCSADHIGPISLGFVHRPEFQLLCNSCNSAKNNRMTFSDVQHLINAENNGEEVASWYCKHI
WDLRKHDVKNNENALRLSKILRDNRHTAMFILSELLKDNHYLFLSTFLGLQYAERSVSFSNI
KIENHIITGQISEQPRDTKYTEEQKARRMRIGFEALKSYIEKENRNALLVINDKIIDKINEIKNIL
QDIPDEYKLLNEKISEQFNSEEVSDELLRDLVTHLPTKESEPANFKLARKYLQEIMEIVGDEL
SKMWEDERYVRQTFADLD

>Nb.BsmBI (C, R438D)

KDPGRHDDNMRLYNHDRRAFMWWSEGDWALADALYNKAGAGKCADPDCQKEVEKISPD
HVGPISCGFKQIPFFKPLCASCNSAKNRRFSYQDVKELLYENYTGDSVASWQVRALWDNC
KHLVKNDDDSKLLSNLMRSLQDYLYRLSLYKLFSNGFAHLLSYFLTPEYAHYKITFEGLNTST
LEYERYYYKTFKKTKSTSSLAARIVRIAFEELEIYNSKDINEDKLIKFDTSWEKDFENIISYATK
NLSLDEEASKWNKVLTDKNLSSTEKDKKISSLLEDKNYEVYKKQFYILKDLLVEHFNKIGE
QIAKDYMK

>Nt.BsmAI(C, R221D)

CRASQDKGRSKENLKSYSYTKDRDAYEYWSDGNIHAANQFMGSPFFNNISADHIGPISLGFVH
DPRYLQPMMSGGDNSSKRDRQLQDDIEKIIETEKRTNVYPMSWYSKLIWEYIKKNYSTHKS
LIS

GVYRDALKQNMSNFMYLWYILEHCNQDGEHFLEEALLKPNYDYFQYSYTFNELGEIVSIN
PRHFTDRNQYETERYKRIAFESVYDYNEKENRNIKANLIDNEQRMLNKLQCQEISSGVPVEQC
KKLLIELMEVIQKRIISTL

>Nb.BsrDI(C)

IPEELFNWPRTDKVNFKSPQGLIKYDEL CYQLEKAVGSKKAYCLSNNAGAKPQKLESLKEW
INSQKKLFDKAPKLTPPAEFNMKLDAFPVTSNNNNYYVTTSKNILYLFDYWKDLRIAIAETAFP
RLKGKLPTDIDEKPALIYICNSVKPGRLFGDPFTGQLSAFSTIFGKKNIDMPRIVVAYYPHQIY
SQALPKNNKSNKGITLKKELTDFLIFHGGVVVKLNEGKAY

>Nt.CviPII

MYIYMSTPQAKTKYYEQRFVNDFYKELERNKVSLPVTIVLKDNLGIKQVIQNGSGVRVLRD
KANAKSPSKI KSEELGRHVTSKADIALFTEEKNGTKVDVAWISPQSHKDFLGKKITPAQYFD
ASSDVMFKTKIGQPKEIKELKNKMISLSVPLTATKYCWPKYKSGTSLRIWDDVQSTILMNM
AIFGVEFGKAYCRNNANILMVGDP LIEVKDDKTII LTTKENGFSLANGFAEYIPSKDKPIFFTK
PTSGKKTVVDGKTIEGVSVWIIYRSYAGSKNRKIDDV LKNKIELISSSCSVKKKDNFVSIMQS
KKITSPPKSKKITSPPKSKKITSPSKSKKITNFFMKK

>BspQI(C)

NNSFNPVRTK DQLHESAVITREKKILLKEPEILQKIKNRNNGEGLKSIIWKKFDKKCFNCEKE
LTIEEVRLDHTRPLAYLWPIDEHATCLCEKCNNTKHDMFPIDFYQGDEDKLRRRLARITGLDY
ESLVKRDVNEVELARIINNIEDFATNVEARTFRSIRNKVKEVRPDTDLFEILKSKNINLYNELQ
YELLTRKD

>N.AlwI(C)

YHLEELLFENNEKKFAENQKNEWDEILAYMDLLISPKPISIEIADKEISIPSGERPAYFEWVLW
RAFLALNHLIIEPQQCRRFKVDQDFKPIHNAPGGGADVIFEYENFKILGEVTTSGGATQFKME

SEPVTRHYLNKKKELEKQGVEKELYCLFIAPEINKNTFEEFMKYNIVQNTRIPLSLKQFNML
LMVQKKLIEKGRRLLSSYDIKNLMVSLYRTTIECERKYTQIKAGLEETLNNWVVDKEVRF

>I-TevI

KSGIYQIKNTLNNKVYVGSAKDFEKRWKRHFKDLEKGCHSSIKLQRSFNKHGNVFECSILEE
IPYEKDLIERENFWIKELNSKINGYNIADATFGDTCSTHPLKEEIIKKRSETVKAKMLKLGPD
GRKALYSKPGSKNGRWNPEETHKFCKCGVRIQTSAYTCSKCRNRSGENNSFFNHKHSQGPSA
D

Deaminase sequences

>TadA8e(V106W)

SEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVNLRVIGEGWNRAIGLHDPTAHAEIM
ALRQGGLVMQNYRLIDATLYVTFEPCVMCAGAMIHSRIGRVVFGWRNSKRGAAGSLMNV
LNYPGMNHRVEITEGILADECAALLCDFYRMPRQVFNAQKKAQSSIN

>rAPOBEC1-2×UGI

SSETGPVAVDPTLRRRIEPHEFEVFFDPRELKTCCLLYEINWGGRHSIWRHTSQNTNKHVE
VNFIEKFTTERYFCPNTRCSITWFLSWSPCGECSRAITEFLSRYPHVTLFIYIARLYHHADPRN
RQGLRDLISSGVTIQIMTEQESGYCWRNFVNYSNEAHWPYPHLLWVRLYVLELYCIILGL
PPCLNILRRKQPQLTFFTIALQSCHYQRLPPHILWATGLKGGSGGSTNLSDIIEKETGKQLVIQE
SILMLPEEVEEVIGNKPESDILVHTAYDESTDENVMMLLTSDAPEYKPWALVIQDSNGENKIK
MLSGGSGGSGGSTNLSDIIEKETGKQLVIQESILMLPEEVEEVIGNKPESDILVHTAYDESTDE
NVMLLTSDAPEYKPWALVIQDSNGENKIKML

Monomeric mitoBE sequence

MTS-Tag-TALE N-terminal non-repeat-TALE Repeat array-TALE C-terminal non-repeat-2aa
linker-Nickase-2aa linker-Deaminase

MTS-Tag-TALE N-terminal non-repeat-TALE Repeat array-TALE C-terminal non-repeat-2aa
linker-Deaminase -2aa linker-Nickase

>MutH-2aa linker-TadA8e(V106W)

MSQPRPLLSPPETEEQLLAQAQQLSGYTLGELAALVGLVTPENLKRDKGWIGVLLLEIWL
GASAGSKPEQDFAALGVELKTIPVDSLGRPLETTFVCVAPLTGNSGVTWETSHVRHKLKRV
LWIPVEGERSIPLAQRRVGSPLLWSPNEEEDRQLREDWHEELMDMIVLGQVERITARHGEY
LQIRPKAANAKALTEAIGARGERILTLPRGFYLKKNFTSALLARHFLIQGSSEVEFSHEYW
MRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMALRQGGLVMQ
NYRLIDATLYVTFEPCVMCAGAMIHSRIGRVVFGWRNSKRGGAAGSLMNVLNYPGMNHR
VEITEGILADECAALLCDFYRMPRQVFNAQKKAQSSIN

> TadA8e(V106W)-2aa linker- MutH

SEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIM
ALRQGGLVMQNYRLIDATLYVTFEPCVMCAGAMIHSRIGRVVFGWRNSKRGGAAGSLMNV
LNYPGMNHRVEITEGILADECAALLCDFYRMPRQVFNAQKKAQSSINGSMSQPRPLLSPPE
TEEQLLAQAQQLSGYTLGELAALVGLVTPENLKRDKGWIGVLLLEIWL
GASAGSKPEQDFAALGVELKTIPVDSLGRPLETTFVCVAPLTGNSGVTWETSHVRHKLKRV
LWIPVEGERSIPLAQRRVGSPLLWSPNEEEDRQLREDWHEELMDMIVLGQVERITARHGEY
LQIRPKAANAKALTEAIGARGERILTLPRGFYLKKNFTSALLARHFLIQ

> Nt.BspD6I(C)-2aa linker-TadA8e(V106W)

RQLEEVIDLLEVYHEKKNVIEEKIKARFIANKNTVFEWLTWNGFIILGNALEYKNNFVIDEEL
QPVTHAAGNQPDMEIYEDFIVLGEVTTSTKGATQFKMESEPVTRHYLNKKKELEKQGV
EKELYCLFIAPEINKNTFEFMKYNIVQNTRIPLSLKQFNMLLMVQKKLIEKGRR
LSSYDIKNLMVSLYRTTIECERKYTQIKAGLEETLNNWVVDKEVRFGSSEVEFSHEYW
MRHALTLAKRARDEREVPVGAVLVLNRRVIGEGWNRAIGLHDPTAHAEIMALRQGGLVMQ
NYRLIDATLYVTFE

PCVMCAGAMIHSRIGRVVFGWRNSKRGAAAGSLMNVLNYPGMNHRVEITEGILADECAALL
CDFYRMPRQVFNAQKKAQSSIN

> TadA8e(V106W)-2aa linker-Nt.BspD6I(C)

SEVEFSHEYWMRHALTLAKRARDEREVPVGAVLVNLRVIGEGWNRAIGLHDPTAHAEIM
ALRQGGLVMQNYRLIDATLYVTFEPCVMCAGAMIHSRIGRVVFGWRNSKRGAAAGSLMNV
LNYPGMNHRVEITEGILADECAALLCDFYRMPRQVFNAQKKAQSSIN~~GS~~RQLEEVIDLLEVY
HEKKNVIEEKIKARFIANKNTVFEWLTWNGFIILGNALEYKNNFVIDEELQPVTHAAGNQPD
MEIIYEDFIVLGEVTTSTKGATQFKMESEPVTRHYLNKKKELEKQGVEKELYCLFIAPEINKNT
FEEFMKYNIVQNTRIPLSLKQFNMLLMVQKKLIEKGRRLLSSYDIKNLMVSLYRTTIECERKY
TQIKAGLEETLNNWVVDKEVRF