

US011316812B2

(12) United States Patent

Hsu et al.

(10) Patent No.: US 11,316,812 B2

(45) **Date of Patent:** *Apr. 26, 2022

(54) RNA TARGETING METHODS AND COMPOSITIONS

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(*) Notice: Subject to any disclaimer, the term of this

patent is extended or adjusted under 35

U.S.C. 154(b) by 0 days.

This patent is subject to a terminal dis-

claimer.

(21) Appl. No.: 17/352,564

(22) Filed: Jun. 21, 2021

(65) Prior Publication Data

US 2021/0344627 A1 Nov. 4, 2021

Related U.S. Application Data

- (60) Continuation of application No. 16/257,493, filed on Jan. 25, 2019, now Pat. No. 1,122,857, which is a division of application No. 15/937,699, filed on Mar. 27, 2018, now Pat. No. 10,476,825.
- (60) Provisional application No. 62/639,178, filed on Mar. 6, 2018, provisional application No. 62/572,963, filed on Oct. 16, 2017, provisional application No. 62/548,846, filed on Aug. 22, 2017.

(51)	Int. Cl.	
	C12N 15/85	(2006.01)
	H04L 51/00	(2022.01)
	C12N 15/10	(2006.01)
	C12N 15/113	(2010.01)
	C12N 15/82	(2006.01)
	C12N 9/22	(2006.01)
	H04L 9/08	(2006.01)
	H04L 9/30	(2006.01)
	H04L 45/745	(2022.01)
	H04L 51/08	(2022.01)
	H04L 51/18	(2022.01)

(52) U.S. Cl.

(58) Field of Classification Search

See application file for complete search history.

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Primary Examiner — Celine X Qian (74) Attorney, Agent, or Firm — Klarquist Sparkman, LLP

(57) ABSTRACT

Provided herein are CRISPR/Cas methods and compositions for targeting RNA molecules, which can be used to detect, edit, or modify a target RNA.

30 Claims, 87 Drawing Sheets

Specification includes a Sequence Listing.

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FIG. 1A

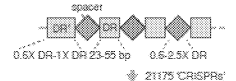
CRISPR system identification

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Obtain bacterial and archaeal genome assemblies and scaffolds from NCS: WGS database



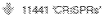
Identify interspaced repeat sequence locations

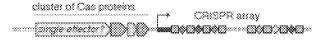


identify putative proteins in 20kb window around CRISPRs



Eliminate candidate arrays with proteins associated with known CRISPA subtypes





Extract proteins >750 as length and ±5 proteins from CRISPR array

CRISPR effector identification

8977 proteins proximal to CRISPR array



all v. all BLASTp single-linkage hierarchical clustering



cluster expansion: SLASTp each cluster against all proteins (-nr database)



sort clusters by member protein proximity to CRISPA array (>70% co-occurrence)



candidate novel single effector CRISPA proteins



expand protein family via this strict metagenome contigs, custom sequencing of genomic DNA tragments



Type Vi Cast 3d family

FIG. 18

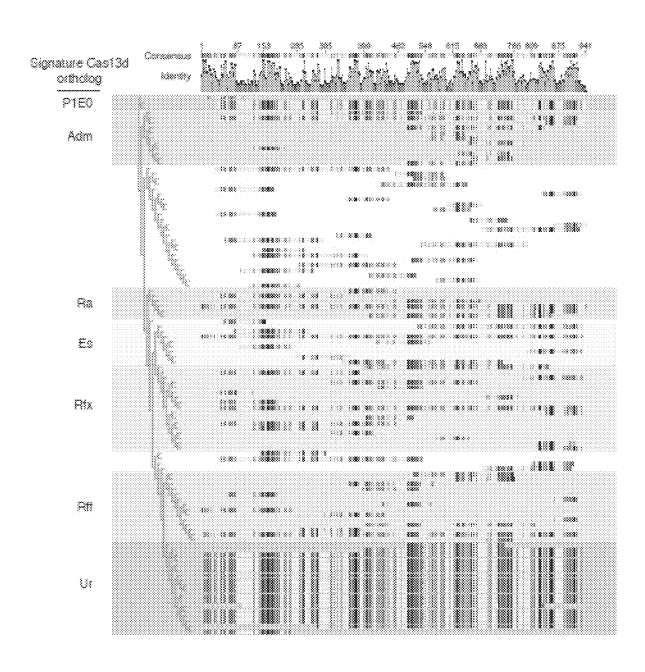


FIG. 2A

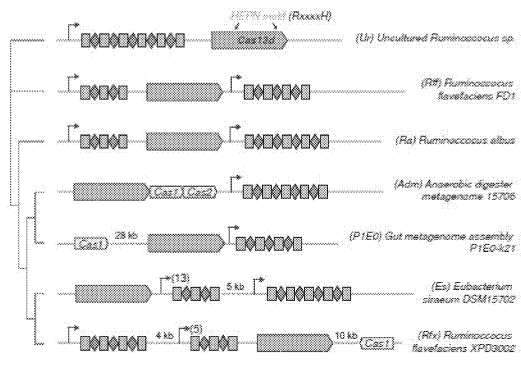
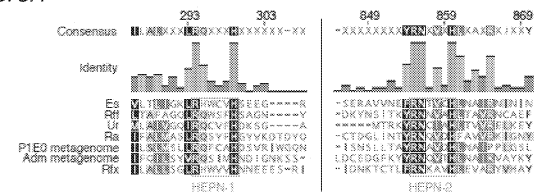
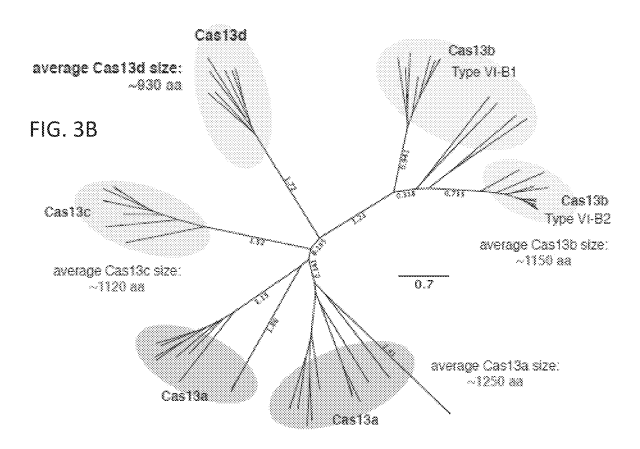
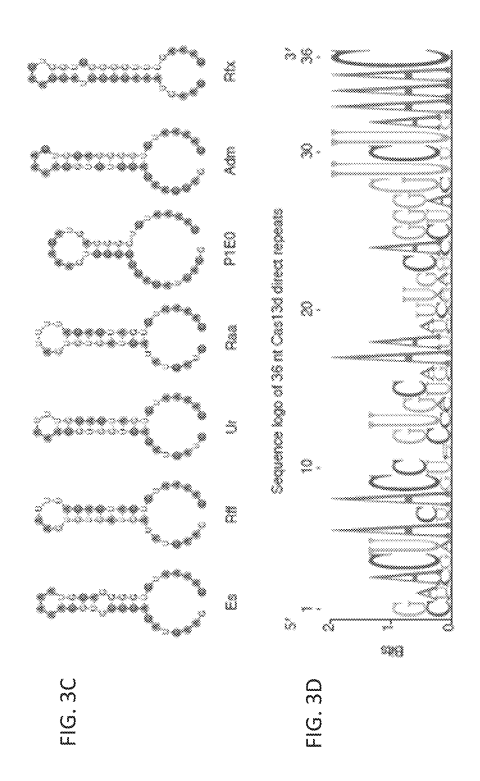


FIG. 2B FIG. 2C EsCas13d 2.884 EDTA undeaved array Uncultured Auminoccocus sp processed guide Cas13d A C COUGAUCAC-5 variable spaces 30nt direct repeat dCas13d



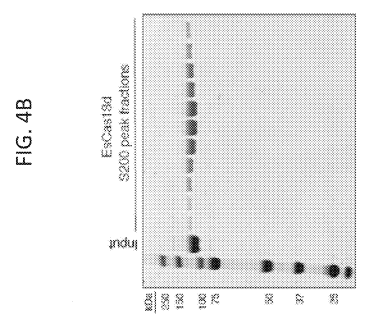


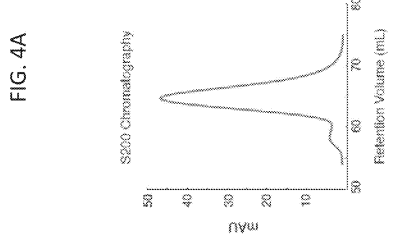


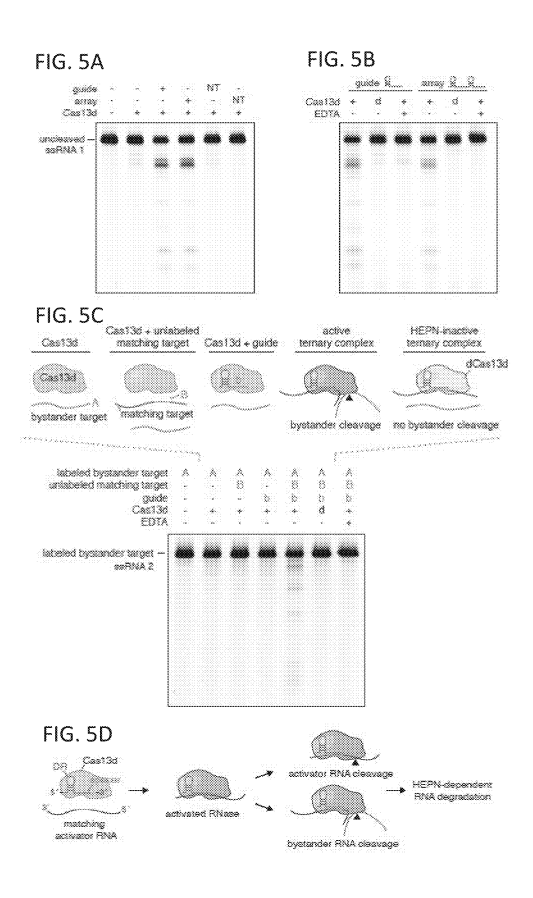


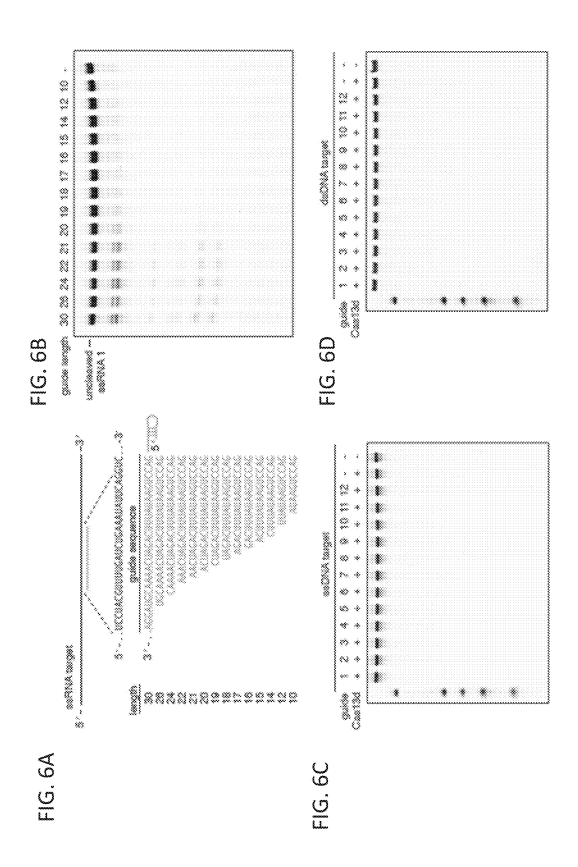
b6taeOb-a3 EsCas13d \$|<u>8 8</u> 8 8

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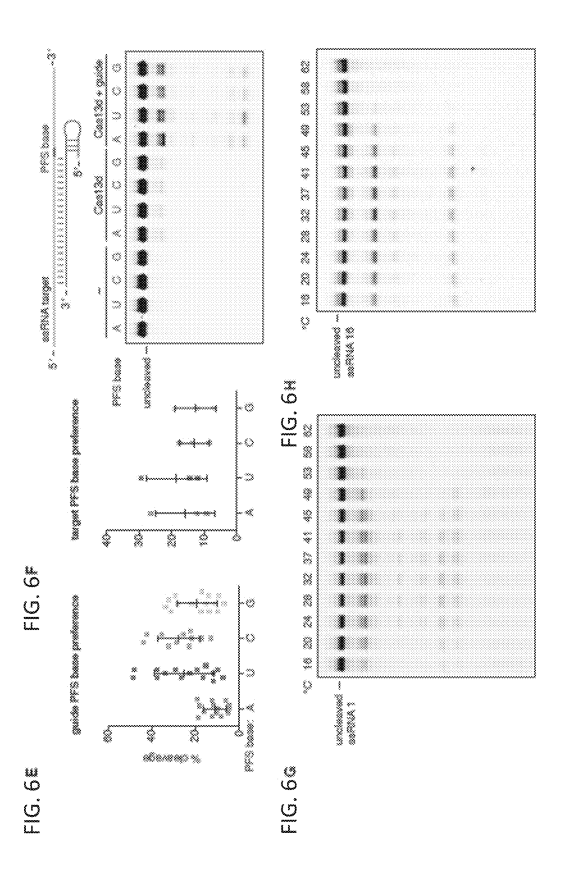


FIG. 7A

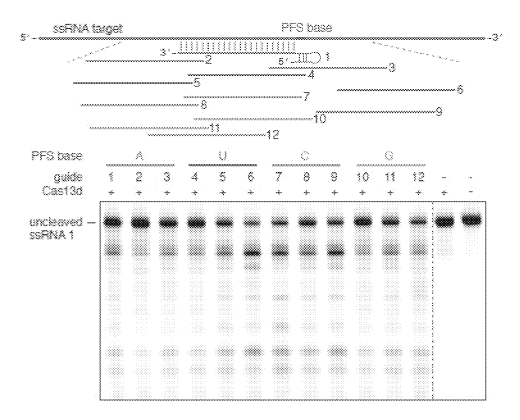


FIG. 7B

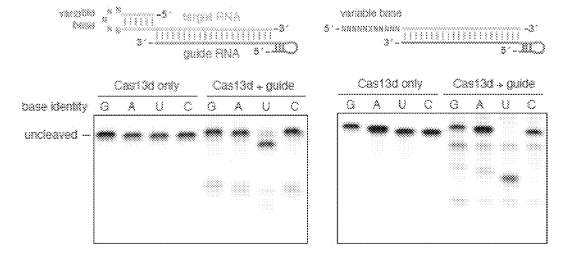
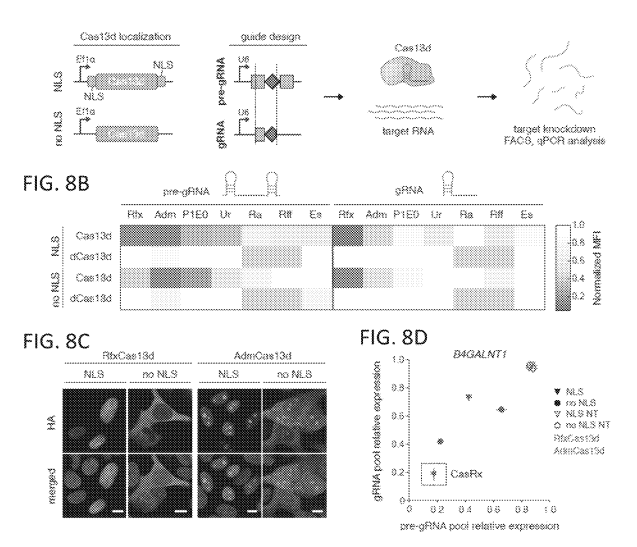
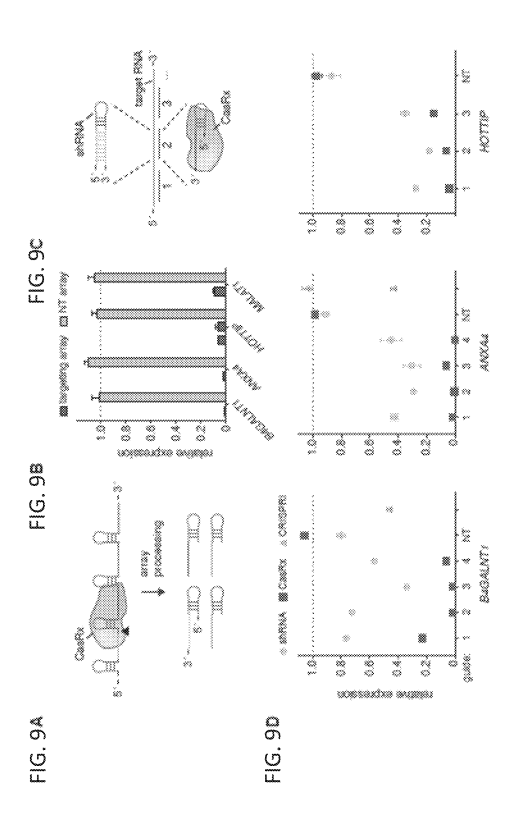
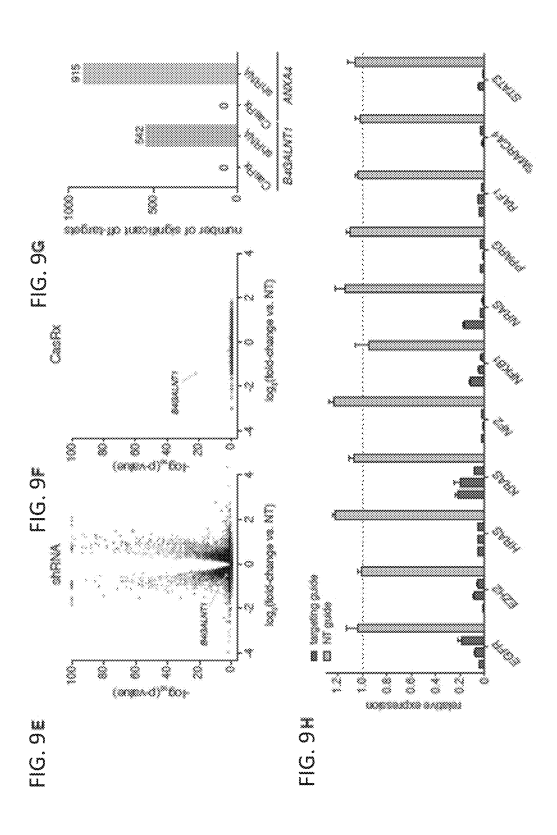
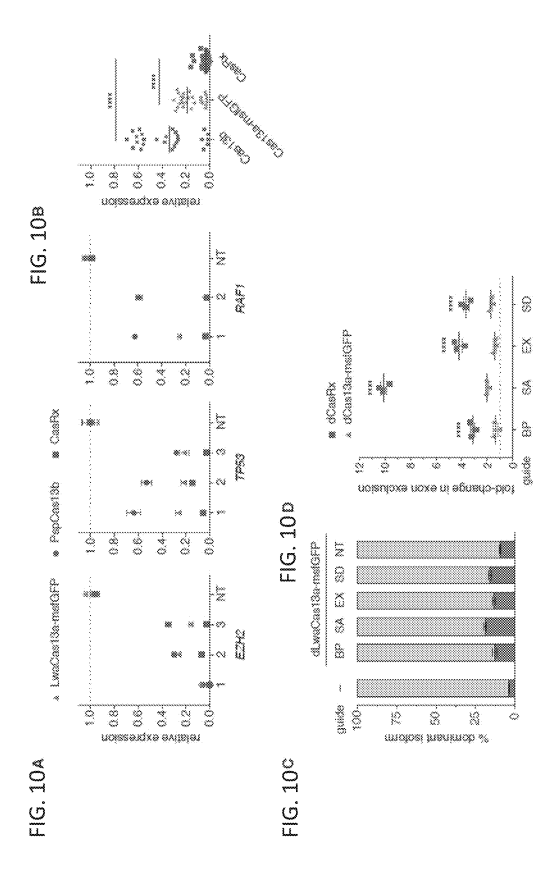


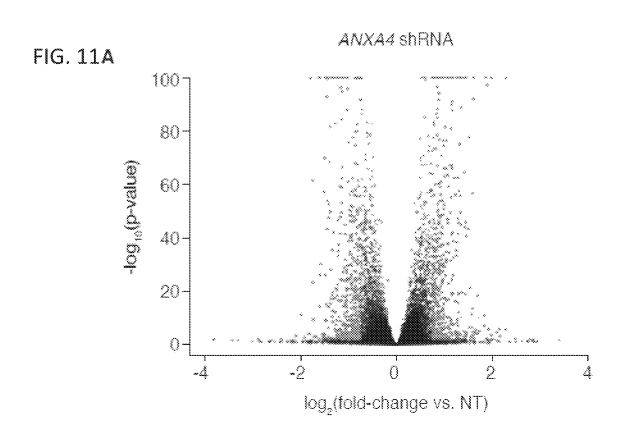
FIG. 8A











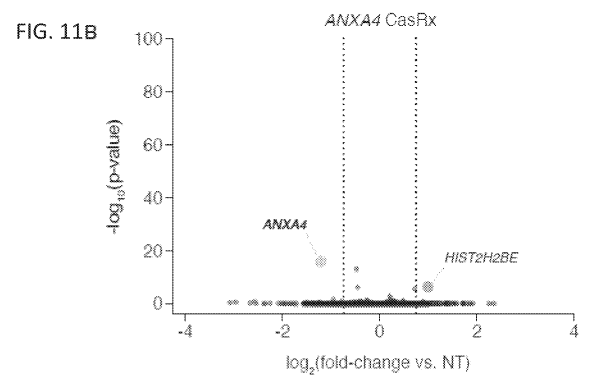


FIG. 12A

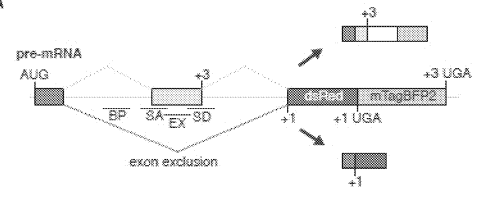


FIG. 128

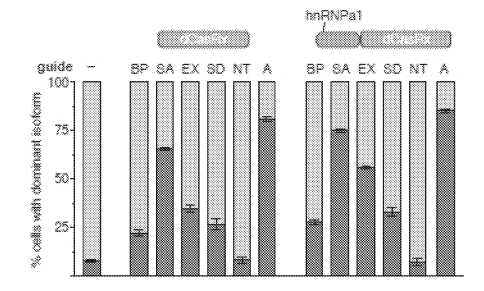


FIG. 12 c

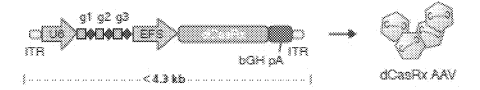
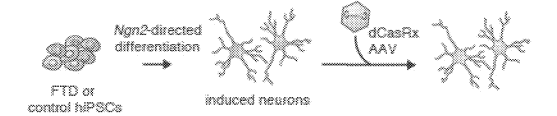


FIG. 12D



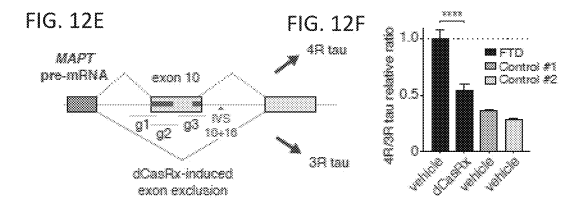


FIG. 13

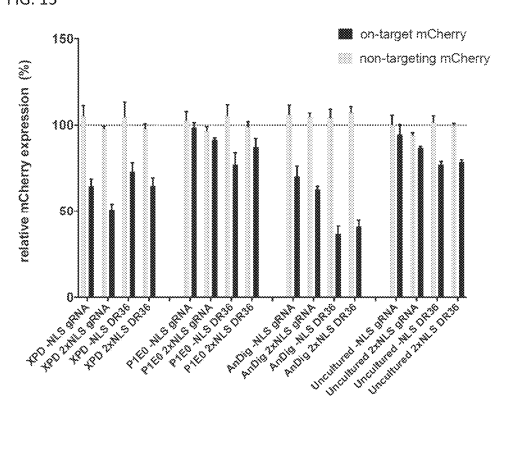
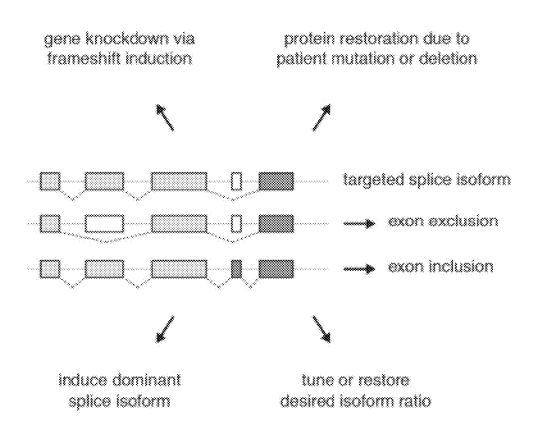


FIG. 14



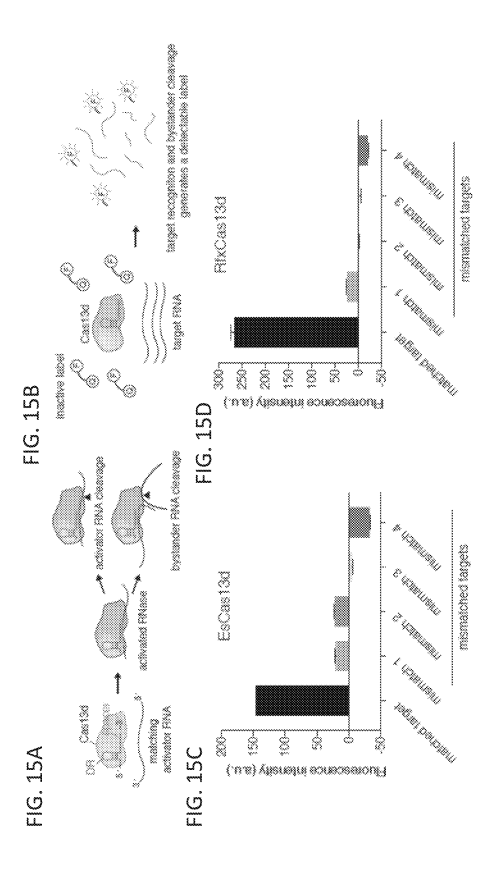
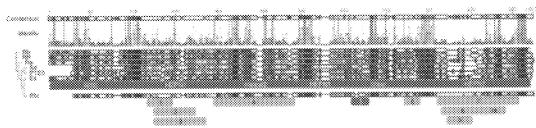
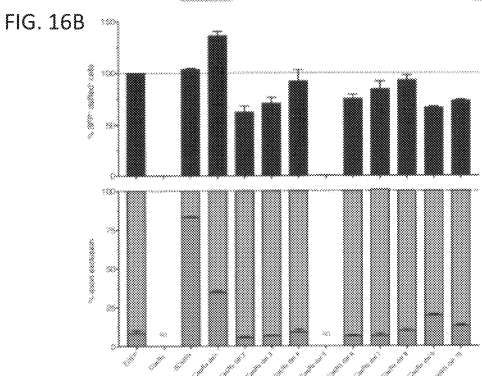


FIG. 16A





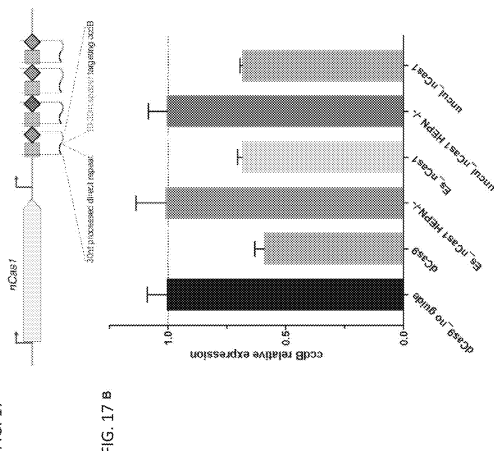


FIG. 174

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CLUSTAL O(1.2.4) multiple sequence a	alignment	
037emb OIZA01000315.1	MAKKKRNIAKERKQNHRESLMK	2.2
emb OCTW011587266.1		0
k8711092736		٥
BMZ-11B_GL0037915		0
BMZ-11B_GL0037771		0
BMZ-11B_GL0069617		٥
Ga0099364_10024192		0
530373_GL0023589		0
emb ODAI011611274.1	KL	20
EMG_10003641		0
Cask_PlE0_metageno		0
emb OIZX01000427.1	KQLREEMQQQRKQ	23
160582958_gene49834	AKNSVIFKLIQAQENKEAAR	20
gi 1198542314 gb NFIR01000008.1	KE	20
emb OGNF01009141.1		0
Ga0129306_1000735		0
MH0288_GL0082219	AKKMTAKERREQEKOKRAE	ত্য শৌ
PIG-022_GL0026351		0
PIG-028_GL0185479		0
250cwins_35838_GL0110300	CO	22
PIG-014_GL0226364		0
250twins_36050_GL0158985	QE	23

FIG. 18A

U.	S.	Pa	ate	ent			Apr	. 26	5, 20)22		Sl	1eet	24	of S	87			US	S 1	1,3	16,8	812	B2
20	20	20	20	20	0	0	0	0	9	0	0	0	0	0	0	31	4.9	0	20	23	2.0	0	20	
	KQ	KQKREEKERINKQ	KORREERERONKO	KQRREERERONKQ													MLKKCSVCGKRFETNYQGGKPVCRNCRSNKSKPVNPKSSRKPESIDKAI		KA				TE	FIG. 18B
emb[OJMM01002900.1]	emb OGPN01002610.1	emb OHCP01000044.1	PIG-046_GL0077813	pigchimera	emb CIEN01002196,1	O2.UC290_GL0096317	CasR_Anaerobic_dig	emb OGDF01008514.1	emb OGZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD_	Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	ODAI_chimera	31009_GL0034153	DLF014_GL0011914_	CasR_Ralbus	Ga0187910_10040531	tpg DJXD01060002.1	CasR_Fsiraeum	

PIG-018_GL0023397	AAKKLSFRELREOREREROA	20	U.
PIG-025_GL0099734		2.0	S.
Ga0129317_1008067_chimera		0	Pa
EYZ-362B_GL0088915	KQ	20	ate
uncultured_Ru_sp	QL	22	ent
RflavgD-1		20	
tpg DBXI01000091.1		2.0	
			Apr. 26,
037emb OIZA01000315.1	KADSNAEKEKAKKPVVENK	64	, 20
emb OCTW011587266.1	MKQNDRENNNKIKKSAAK	18	22
k87_11092736		σ ₀	
BMZ-11B_GL0037915	MAEHSKKTAAK		Sl
BMZ-11B_GL0037771		10	heet
BMZ-11B_GL0069617		10	25
Ga0099364_10024192	MANTATEQUE	10	of
530373_GL0023589	MKIEMKNGEVKISAKKENCSIAK	23	87
emb ODAI011611274.1	KYQKAQAEBAAAAQQTAAGABSEENPCFDVVKDTKRKALNPLHVELEAPSAKKSSVK	44	
EMG_10003641	KVTKETTIEKQSIKKHKQKSKKTAIK	30	
Cask_PlE0_metageno	MEREVKKPFKKSLAK	15	US
emb OIZX01000427.1	AIQKQQEQRQEKAAAARETAAPEQPAAAFVPKRQRKSLAK	61	S 1:
160582958gene49834	KKAKDIAEQARIAKRNGVVKKEENRINRIQIEIQTQKKSNTQNAYHLKSLAK	را دی	1,31
gi 1198542314 gb NFIR01000008.1	BHKKQEKLRKEQEELRKKQEKQREDQKELEKIKKEEGGEGEKKKSGAK	68	16,8
emb OGNF01009141.1	MADIDKKKSSAK	1.2	812
	FIG. 18C		B2

U.	S.	Pa	ate	nt			Apr	. 26	5, 20	22		Si	1eet	26	of S	87			US	S 13	1,31	16,8	312	B2
26	54	S)	17	55 50	© H	0.9	67	63	58	82	\$ P	(T)	න ස්	0.5	22	ଷ୍ଟ	2 e	10	30	3.0	0	14	06	
QREQQTVTDESERKKKFLKSGAK	VVERQEANRKKAQAAATQDATAPVHHKNAKSLAK	MAESYKIKSKKTRIK	MSESDKIKIKSKKTRIK	KARQAESQRDKIKNMNVEKMKNINTNDIKHTKAK	HIKPEKKQIKQATLIK	KAQKYASEPSPLQSDTAGVECSQKKTVVSHIASSKTTAK	KDMMYTKSTDAVSVPTLKAAPTKAEMSQDTAEASTLITPGTLKTKAK	KWAKQDTPVVPRSKTEEKPVVAASDDKLLKTTQVKKVQTKSKAK	KWAKNDSVIIVPETKERIKTGELQDNNRKRSRQKSQAK	KWAKKQDDATAVFECEANIKPADSKDEDCANISIKRKTQAK	KWAKKQADATAVFECEADIKPADSKDEDCTNIYIKREKKKTQAK.	MERQKRKMKSKSK	MERQKRKKSCAK		MIETKPKREDIAKIPAAKSKK	KSQENEALKKQKLABEAAQKRREELEKKNLAQWEETSAEGRRSRVK			MIEKKSYAK	MIEKKKSFAK	MIEKKKSYAK	MSTKKRFRYSVAAK	resainvevitdethkeakioknefktfdoftke-loetokvngstkkehitknkhtnvk.	FIG. 18D
Ga0129306_1000735	MH0288GL0082219	PIG-022_GL0026351	PIG-028_GL0185479	250twins35838GL0110300	PIG-014_GL0226364	250twins_36050_GL0158985	emb OJMM01002900.1	emb OGPN01002610.1	emb OHCP01060044.1	FIG-046_GL0077813	pig_chimera	emb OlENO1002196.11	02.UC29-0GL0096317	Cask_Anderobic_dig	emb OGDF01008514.1	emb 0GZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD_	Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	

EILIEKLISDOKK KIEKFKIKYAEQETVDKI ZAAVNGVANGTANG		MGSRGVIIMTERRKSRAK TEEIAQSDKIMRTESILVQHTKKSLAK TEEIAQSDKIMRTESILVQHTKKSLAK	ତ ଓ ଓ ଓ ଓ ଓ ଓ ଓ	U.S. Pat
CasR_Ralbus Ga0187910_10040531 tpg DJXD01000002.1 CasR_Esiraeum	AAVNSVNDTPERTEEAN	VSSAFVERRDEQSKRIFAKVSSAFVERREDEQSKRIFAKWKRQKSKRIVSK	6 H 5 G 6 G 6 G 6 G 6 G 6 G 6 G 6 G 6 G 6 G	tent A
FIG-015_GL0099734 FIG-025_GL0099734 Ga0129317_1008067_chimera EYZ-3628_GL0088915	FARQVQEREABKARKER——————————————————————————————————	-QVSVSDIVGAAAVQNENGKKSMAK AELKRFKLDQETRGKLPKNERKSLAK MKKQKKSLVK	2	pr. 26, 2022
uncultured_Ru_sp RflavFD-1 tpg DEYI01000091.1 037emb OIZA01000315.1 emb OCTW011587266.1	KAAEINNNAAPAI		N N N N N N N N N N N N N N N N N N N	Sheet 27 of 87
k87_11092736 BMZ-11B_GL0037915 BMZ-11B_GL0069617 Ga0099364_10024192 530373_GL0023589	-06	KAEIAVRSADPPEETLPTESD	ν ν ν ν ν φ 4 φ ν ν Ο 4	US 11,316,812 B2

AMGLKTVLSFD---D--

RMGLKSTLVFD--N-AMGLKSTLVMG--D-AMGLKSTLVFD---D-

250twins_36050_GL0158985

emb|OGPN01002610,1| emb|OHCP01000044.1|

PIG-046_GL0077813

emb|OJMM01002900.1|

250twins_35838_GL0110300

PIG-014_GL0226364

PIG-028_GL0185479

PIG-022_GL0026351 MH0288_GL0082219

AMGLKTVLGFD---N--

AMGLKIVLGFD--N-MAGVKSVFVIG---D-

AMGLKAVLSFD---N-

RAGLKSSFVN----EN--RAGINSSFVS---EG--KLGLKSTIIAD--K----

AAGLKSTFTLP--TG-

AAGLKSVFVLS--EG-

MSGLKSALVIN--NH-

smb|ODAI011611274,1|

AAGLKSTFVISPQEK-

AAGLKSNFILDPORR-AAGVKSVFAI---GM-

ALGLKSTFILDRDEQ-

gil1198542314|gb|NFIR01000008.1|

emb!OGNF01009141,1

3a0129306_1000735

160582958_gene49834 emb:OIZX01000427.1|

Cask_PlE0_metageno

EMG_10003641

AAGLKSTFAVN---G----

AVGVKSVFVVG--D-

AAGLKSVFFDQ--K--

MAGVKSTFVID---S--

02.UC29-0_GL0096317

emb|OIEN01002196,1

pig_chimera

emb|OGDF01008514,1|

emb|OGZC01000639.1

CasR_Anaerobic_dig

tpg:DJXD01000002,1

Cask E. siraeum

3a0187910_10040531

Cask R. albus

DLF014_GL0011914_

31009_GL0034153

ODAI_chimera

emb|OCVV012889144.1|

Sa0187910_10006931

Ga0224415_10007274 Ga0187911_10069260 3a0187910_10015336

R. flav XPD

FIG. 18G

tpg|DBYI01000001.1

EYZ-362B_GL0088915

uncultured_Ru_sp

R._flav._FD-1

PIG-025_GL0099734 PIG-018_GL0023397

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Ø. 0 0 0	ထ	.හ න	7.9	73	8,2	83	4	101	06	135	149	147	83	80	133	28	80 4	131	7.6	ණ ල ස	140	150	145	
쮼	HISRS	PGEGLIEASVAN	EEGK	and also and the first test that has and and and and and test test test and and and and and and test test	AGE		SS		LADRIVIADD	VRFPNVIADD	VLNHS-TQKMEEIQTE	DDCEPVKVNN	GLAADSLVDN	TAGSSDD	DVQKEGSSVPEVLTDD	GUSATION	KNGIVSDD	DLSEFAN	TTQTAMFEN	CHINVNN	DEINYPN	GIFKKDEKDVKATICN	GTIKKEEKGVKSTICN	
	ATLSTHAKFAKAGRDGREFKCGDVD	LGGFNVTGVDENTVDAERGI	GKRGFEVRKIDENVDLYGDL	EKSFTVNKIDDEIDVQRS	RNITANKIVSEGIQISGII	NIFSATPISERIVDITGET	RDPELFSAKPLETGYRIQRFNA	NKNKKFDANNDSKVVVIKGISN	GEKQQFQVQRQDESRFRLQNSR	PVQQFQVEPASAAKYRLKNSR	SSPEQYSAEFQNKQFKIKGNIK	NKKAALSVEVCGKSFNISKKEN	EPEVFSVIPCDKKFELQPAKR	DNKEAFSAALQNKRFEVFGR	EGDPAFSVIPMPKQFEIQGRAI	NKPRFSVEFGNKEYIINDK	NKPRFSVEFGNKE?IINDK	YTPRMFRSEINPGEIVISKGD	QDEKMFRIDVEKQQIYIEREFPDENNKII	EHERMFSADVGEKNIELSKND	ERPRMFNTSINAKKVDLSKDN	ENARMFDSSVDEQNVNIEKRMTIEEKQND	VNSKMFESSVNKRDINIEKRITIERPQQD	FIG. 18H
emb OCTW011587266,1	k8/_11092/36	BMZ-11B_GL0037915	BMZ-11B_GL0037771	BMZ-11B_GL0069617	Ga0099364_10024192	530373_GL0023589	emb ODAI011611274.1	EMG_10003641	Cask_P1E0_metageno	emb OIZX01000427.1	160582958_gene49834	gi 1198542314 gb NFIR01000008.1	emb OGNF01009141.1	Ga0129306_1000735	MH0288_GL0082219	PIG-022_GL0026351	PIG-028_GL0185479	250twins_35838_GL0110300	PIG-014_GL0226364	250twins_36050_GL0158985	emb 0JMM01002900.1	emb OGPN01002610.1	emb OHCP01000044,1	

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149	151	ტ თ	83	106	න භ	144	φ. σ.	ති ග	98	85	დ აე	හ හ	155	154	96	136	132	127	60 63	<u>क</u>	332	1.54	ന യ	
GTIKKDEKDVKSTICN	GIIKKDEKDVKSTICN	NNIRAKVNN	NKEHVAINN	SLPLSKRORIRAERIKRAREENR	GIKSHISATADN	GISNGRIVLPEISVDN	EKFRHPKGYDIISNN	KKFKHPKRYDVIADN	KKFGAPKGYDVVANN	AKFSHPKGYAVVANN	VKFSHPNGYDVVANN	HHAGVHALVDQ	KRSNVKGLADN	RIKAEGGPVEFDG	TKRGINTINKKKQYAN	SRRCKNIESAN	GRIKDKTARPKD	RMQDMNLTADH		RRGETVSAVIDN	HRGQSVCATADN	KRIACLNTQADN	PFPKQAEAKN	
EDVRMFDSNVDECSINLEKRMTVEERQKD	EDVRMFDSNVDECSINLEKRMTVEERQKD	N-PAAYDAVYGTDSIRVKKTN	0-EQAFDAQYGKEKIFVTNAD	PPAFDLELKEKTFYISGKNNINTSRENPLASASLPLSKRQRIRAERIKRAREENR	V-KPRFSVEKPATSYSSSF	E-EETFTAKLKFAQTEPTVATSI	D-KEAFAASIANKNVGYKIKND	E-KESFSAEIFKKNIGCKIE-N	K-KEAFVAEITDKNAGYKII-N	E-GEAFSAEMADKNAGYKIG-N	E-GEAFSAEMTDNNAGYKIG-N	E-NPAFDAVINDKKYALTG	ETENQKKYVVTN	GEAFALAKEPVNHKLVLKSG	D-KPVYDVVGVEKETGIINVVG	T-KPAYEIVDIEKNGDLIKVQ	GAFEVAERDESTLILES	PQAALKNVHAPNKQKIHFIG	SSAGLKNVEVS-GKKIKFQG	DPAYQLNVVTMNGYSVTG	ANPAYEITDISSVNYSVEG	E-KENFTVDTDGITDTIVPVQS	SDSEKHTLKINSITDTELRLSG	FIG. 181
PIG-046_GL0077813	pig_chimera	emb OIEN01002196.1	O2.UC29-0_GL0096317	Cask_Anaerobic_dig	emb OGDF01008514.1	emb OGZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD_	Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	ODAI_chimera	31009_GL0034153	DLF014_GL0011914_	Cask_Ralbus	Ga0187910_10040531	tpg DJXD01000002.1	Cask_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	

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8 8 8 E 8 8 8 8 8 8 8 8	136 136 136 127		1 1 1 1 1 8 6 8 1 1 1 2 8 8 1 1 1 1 1 1 1 1 1 1 1 1 1
IS-EHSELHVLGADDLAVSFSGTRRMKDVSELKADNSSKDNSNIQLGGVNEVNITFSS	PNKHIE-TDNMHIQALYNIED TDKLIBDKNSGMDDLGFKRELELEVFGQCFDDSIHIQIAHAVFD EYESLISNPAESFREDYLGLKGTLERKFFGDEY-FKDNLRIQIIYSILD PVNDLIDESRKEYGQDYLMLKDQLEKVFFGQTF-PGDNLRIQIIYNILD TIEALLINPSEKVGEDYLKLKGTLERRFFGREF-PHDNIRIQLIYNILD	SLLGTLENPAENDPVDYLGLKSTLEKEFFGQEF-FNDSIRIQIIHNILD GHATFLNNPAEHVGTDYLKEKETLEMEFFGKSF-PGDSVRIQIIHQILDKKTAQVVLPACSDNQLHAKDTIEQMYFGKTYTDNIHIQIAYNIMDLAYRPAGVRPDQIGAKAALEKRYFGKTYTDNIHIQIAYNIMD PLFDQSPTAIQPNRISGNDMIGIRRMLERKYFVHNEENKEFQDNIRIQIAYCILD PLHRA-ETPRRQPLGAGMDQLRRKAILEQKYFGRTFDDNIHIQLIYNILD PLKRRKDGGFVPGMDALRRKNVLEQRYFGADNIHIQMIYSIILD	SGNKKDDDLIHCRKNLEEMYFG
EYZ-362B_GL0088915 uncultured_Ru_sp RflavFD-1 tpg:DBYI01000091.1;	037emb 01ZA01000315.1 emb ccrw011587266.1 k87_11092736 BMZ-11B_GL0037915 BMZ-11B_GL0037771	BMZ-11B_GL0069617 Ga0099364_10024192 530373_GL0023589 emb ODAI011611274.1 EMG_10003641 CasR_P1E0_metageno emb OIZX01000427.1	gi 1198542314 gb NFIR01000008,1 emb OGNF01009141.1 Ga0129306_1000735 MH0288_GL0082219 PIG-022_GL0026351

DATE
KTY-NKDNLBVQIAYNIFDRTF-PNENLRVQIAYNIFDRTF-PNDNLRVQIAYNIFDRTF-PNDNLRVQIAYNIFDRTF-PNDNLRVQIAYNIFDKTF-DDNIRIQLISKILDKTF-SDNIRIQLISKILDKTF-SDNIRIQLISKILD
KPSNDNIHIQVVSKIMD 136KEYSDNIKIQIIYNILD 148KTF-SDDNIHIQLIYNILD 133KSFNDDLHIRLIYNILD 133SAD-GIDNVCIQVIHNILD 127SAD-GIDNVCIQVIHNILD 128STUGNNTICIQIIHNILD 128STUGNNTICIQIIHNILD 128
KSSGGDNNLCIQIIANIID 133SAD-GTDNVCIQVIANILD 127
SAD-GTDNVCIQVIHNILD 129SSVSGNDNICIQVIHNILD 128STDGNNTICIQIIHNILD 128STDGNNTICIQIIHNILD 128DTFADNIHVQIAYNILD 122RTFNDNIHIQIIHNILD 198
SSIDGNDICIQVIANILD 128SSIDGNNICIQIIANILD 128
SSTUGNNTICIQIIHNILD 128DTFADNIHVQIAYNILD 122RTFNDNIHIQIIHNILD 198ENFNDNIHIQLIYNILD 138ENFNDNIHIQLIYNILD 178
RTFNDNIHIQIIANNID 198 SNKVLQDNIRIQIAHNIMD 198ENFNDNIHIQLIXNILD 138
GNKVLQDNIRIQIAHNIMD 198 ENFNDNIRIQLIYNILD 138 RTFNDNIBIQLIYNILD 178
ENFNDNIRIQLIYNILD 138 RTFNDNIBIQLIYNILD 178
RIFNDNIHIQLIYNILD 178

Patent	Cask_Ralbus	PRHITVDTQGKFKEDMLGIRSVLEKKIFGKTFDDNIHVQLAYNILD 178	
PLEKK	0_10040531	LHSHDGERAVGADLLCAKDKLEQLYFGRTFNDNIHIQLIYQILD	
PLREWGRKK.—DEPEOSUPTOMICLARTLERUYEG.——KHF.—DDNIHIQLINNID 179 PLHED	11000002.1	LFKPQPGMDLLCLKDKLEMHYFGKTFDDNIHIQLIYQILD 12	m
PLHEDGSSSVPTDMICLAKPTLERVYFGKTFDDM.HIGLINNILD 179 PLHESG	siraeum	LRRFNGRKKDEPEQSVPIDMLCLKPTLEKKFFGKEFDDNIHIQLIYNILD 18	S
### PLYRK	GL0023397	LHHDGGSSVPTDMLCLKPTLERVYFGKTFDDNLHTQLIYNILD	σı.
PTHK	GL0099734	LYRK	ത
PLHSGK-NNLYPENGTDILGLEGETEZEYFF-DDNVHIQLIYNILD	7_1008067_chimera	THKKDNEQKNTRQDMLGLKSTLEKFYFGSTFDDNIHIQIIHNIQD	o,
PTHRS	GL0088915	LHSGKNNLVPKNGTDILGLKGILEQRYYGRNFDDNVHIQIIYNILD	
	ced_Ru_sp	THRSGESSPVRGDMLGLKSELEKRFFGKTFDDNIHIQLIYNILD	
PLBSGKDKPNKSACQDMLGLKSELEKRYFGKIFDDNLHIQIIHNILD		LYGGKDNPEKPYGRDMLGLKDKLEERYFGCIFNDNLAIQIIYNILD	
EKILAKHITNIYTVNSEDRNYNQSGNDTIGFDLNYRV 174	101000091,18	LHSGKDKPNKSAGQDMLGLKSELEKRYFGKIFDDNLH1Q11HN1LD	20
IEKILAKHITNIIYTVWSFDRNYNQSGNDTIGFDLNYRV			S
IQKSLAAVIPNVLYTLNNLDRSYST	nb orza01000315.1	EKILAKHITNIIYTVNSFDRNYNQSGNDTIGFDLNYRV	
1 100 10	WÓ11587266.1	QKSLAAVIPNVLYTLNNLDRSYSTDNTSDKKDIIGNTLNYQH	
TYKILGMNVADILYALGNNQDTE	92736	BDTAGTDGTC-D	
IXKILGMNVADILYALGNMQDTE	_GL0037915	LIDDDWTSDVIGLALGK	
1 IQKILGIYITDIIYCINNLRDETHL	GL0037771	YKILGMNVADILYALGNMQDIE	23
1 MKKIFGVYSNNITYSINNLLEE	GL0069617	SLSKVEDCLGQSLSK	90
MKKIFGVYSNNITYSINNLLEBREGSDFLGVFYTEN 164 IEKLLAVYISNIIYAVNNYTGVSAMKDSKGRPVDLLGDYGILGEE 233 IEKILMPHINNICFEINNMLRL-EGYQEDSFMGS	54_10024192	QKLLGIYITDIIYCINNLRDETHLDHESDIVGLSMSN 16	ထ
IEKILAVYISNIIYAVNNVTGVSAMKDSKGRPVDLLGDYGILGEE 233 IEKILMPHINNICFEINNMLRL-EGYQEDSFMGSFNLYK 194 IHKMLAVPANHIVHTLNLLGGYGEIDFVGMLPAGL 174 FIG. 18L	31.0023589	KKIFGVYSNNITYSINNLLEEFYTEN	
IEKILMPHINNICFEINNMLRL-EGYQEDSFMGSFNLYK 194 IHKMLAVFANHIVHTINLLGGYGEIDFVGMLPAGL 174 FIG. 181	1011611274.1	EKLLAVYISNIIYAVNNVTGVSAMKDSKGRPVDLLGDYGILGEE 23	8
IHKMLAVFANHIVBTLNLLGGYGETDFVGMLPAGL 174)3641		ধা
	10_metageno	GEIDFVGMLPAGL 17	\$P
		FIG. 18L	B2

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216	237	227	164	177	215	7.62	164	222	176	218	215	228	224	228	230	174	174	188	175	227	175	163	89 9 11	
LAAHV	-NSKDDFFGCDSHESVAYLYDELKA	HNAST	LNLK	GYLILKN	GGLTLKN		MSIVN	slyyoa	KQIGP	TLYAYK		SLYAFA	SISIM	TLYMLK	TLYMLK	KNTQ	KART	SESSII	YLYIGN	NFSTRN	QIXNSSY	GGEKPIY	GKFSTVY	
KDRDFIGM	NSKDDFFGCDS	GEEEFDLIGS	GMEYNDYIGT	DPETDDFIGS	GTEGIDFIGI	XDI3GGX	TDEYDDFIGY	QSENSGRDCDVIG	BDKNDLLG	RDERMYDDLIG	AKKDIIG	SDIIG	SINDVIG	SDNDVIG	8INDNIG	IDRPIDIFGD	TONPIDXMSN	IDGKIDVIG	YDREQDYLG	DDFLS	BIIDIIGI	EDIIGM	KDI1GF	
IHKILAAASGHIVHLLNIVNGS	IEKLLTPYINNIIYTLNBLMRD	IEKILAVQIMNIVFILNNLLRW-S	IEKILSVHVNDIVYSVNNILSRGE	INKILAVHANNIVYTLNNLDRE-A	IKKVLAIHANNIVYTLÖNĻSRR-E	LEKILAVHSMNIVYTFDNLRRK-E	LEKILAVHSNNIVYTFDNLRRK-E	IKĶILSVYINNIIYMFYNLARSEEYDIFYNSQSENSGRDCDVIG-	IKKIICPYIMNVVNIFYNLGRGKLK	IKKILGTYVNNIIYIFYNLNRAGTG	IKKIIGAYINNIIYIFYNLGREEYD	IQKILGTYVNNIIYSFYNLRRDG-K	VQKILGTFVNNIIYSFYNLSRDEVQ	IQKIFGTYINNIIYSFYNLSRDEFQ	IQKILGTYINNIIYSFYNLSRDESQ	IEKTFSVVIGNIVYAINNLSLEQS	TEKTFSFVMGNIVXAINNLÖPEQN	INKIISPYINDIVYSMNNLARNDEY	IRKILSTYANNVVFTINSMRRLDE	IEKILAEYTTNAVFAIDNVSGCS	IEKILSEYIPNVVYAFNNIAGFKDE	IEKILAEYITNAVYSFDNIAGFG	IEKILAETITNAAXAVNNIAGLD	FIG. 18M
emb OIZX01000427.1	160582958gene49834	gi 1198542314 gb NFIR01000008.1	emb OGNF01009141,1	Ga01293061000735	MH0288_G10082219	PIG-022_GL0026351	PIG-028_GL0185479	250twins_35838_GL0110300	PIG-014_GL0226364	250twins_36050_GL0158985	emb OJMM01002900.1	emb OGPN01002610,1	emb OHCP01000044.1	FIG-046_GL0077813	pigchimera	emb OIEN01002196.1	O2.UC290_GL0096317	Cask_Anaerobic_dig	emb OGDF01008514.1	emb oGZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	

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164	164	163	242	236	176	216	214	213	164	224	217	237	167	219	21.7	220	21.8		287	200	හ ස	194	
GKFSTVY	KKFTSIK	NFSAGT	GGISTSK	YRGLDK	YASTIN	YLSARA	YMSIRN	GRMRATI	GGWRINL	KRITDE	NMTIDY	ALSFQQ	YMGTSR	YMNTMN	YLSTWN	XMNTIN	YMNTEN		-EAKPHLGY	-AAKDRFSY-	-KALPYMGF-	-SMSDALGF-	
KDIIGF	NDVIGG	T-QADELDSIG	NSAEKPIDMIGA	NEGDDF11G			NDNDMG	DELSDDFVGM	EELTEDFIGA	IENSDFFM	DESSDIFQ	SERGDFIG	VEFSDMIG	DIEHEDFIG	SESHDDFIG	KYLNSDYIG			PNKI SAWKKRENFINF YN	KRGEFTEYYN	EKMQKLLS	GRGKEIIE	
IEKILAEYITNAAYAVNNISGLD	IEKILABYITNAVYATNNIIDPD	ITKILIVYANNVVYALNNLVHADDD	IKKILAVHTNNIVYALDNIHERGRE	IEKILAVHINNIVYSVNNLFRVQG	IEKILSVYVNNIVYALGNLERDNS	IEKILSVYINNAVYALGNLERKEN	VEKIMAQYVSDIVYMLHNIDKTER	IQKILALHANNIIFALDNLLHKKN	IEKILAVHVNNIVFTLDNVLAFQK	IEKILAVYSTNAIYALNNMSADEN	IEKILAIYSTNAVYALNNMIADED	1QKILIVYSVNTVYTLNNLFGKEI	IAKILAAHSNNAGYALDNMLAYQG	IEKNLĄIYVTNVVAĄLDHMCDETL	IEKILAVYVTNIVYALNNMLGVKG	IEKILAVHSANITTALDHMVDEDDE	IEKIIAVYATNITAAIDAMVDDDNE	*	PYLEYGGGKDSNGKP	SYESFNVE			FIG. 18N
Rflav_XPD_	Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	ODAI_chimera	31009_G10034153	DLF014_GL0011914_	Cask_Ralbus	Ga0187910_10040531	tpg:DJXD01000002.1	Cask_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	EYZ-3628_GLC088915	uncultured_Ru_sp	RflavFD-1	tpg DBXI01000091.1		037emb OIZA01000315.1	emb ocTW011587266.1	k87_11092736	BMZ-11B_GL0037915	

BMZ-11B_GL0037771	RMKPYMGF 1	U. 62.1
BMZ-118_GL0069617	SITPYLGE-	S.
Ga0099364_10024192	TOWNTYTOWN TOWN TOWN TOWN TOWN TOWN TOWN TOWN	184 P 3
530373_GL0023589	SYQAMNEADAIIKSKLLTKNSKYLNLDMNNVWF-NK-EVSELSSFRKYV-	ate
emb ODAI011611274.1	GLIKRLQRIPECQADERAKALQAFLCSERLSY-	ent
EMG_10003641	FYDAFIATTDDTSKQVRY-	225.
CasR_P1E0_metageno	PYDKLRVVKKKKRPQLAYVDIKADIAAYAKRPQLAY-	2.07
emb ofZX01000427.1	LYNELNEEAKSPRLIY-	Apı 5₹0
160582958_gene49834	GYSDRLKTKPNLSKNIDRIWNNFCNYMN-SDSGNTEARLAY-	r. 26
gi 1198542314 gb NF1R01000008.11	TYERFRGRNKKQSQMRY-	5 , 2 0
emb OGNF01009141.1	SFETYKNNLVKIPQLAY-	261
Ga0129306_1000735	TFEIYCDFAALQNFRLAY-	214
MH0288_G10082219	SYAAYCEPRIFFDEGKEKNIIENIEKSHGEFLEYMKNPRLAY-	257
PIG-022_GL0026351	SYEEFWNPEKLKNPRLGKDQSVYDNINRSRETFLEYRKNPRLGY-	h ee f
PIG-028_GL0185479	SYEEFISPEKLKNFRLGY-	37
250twins_35838_GL0110300	SYRNQDANRFEDIRAYYTY-	254
PIG-014_GL0226364	KSEEMVEDLLKNSSAYFVY-	87
250twins_36050_Gto158985	PMEAQQTYLLKNTSAYYVY-2	253
emb OJMM01002900.1	2 STATESTITAL SKILNNISAYFTY-	233
emb OGPN01002610,1	DFDNQLKDKNTEAYFSY-	326 256
emb OHCP01000044.1	DYDROKETGIEAYYAY-2	5 1
PIG-046_GL0077813	DFDGQKETRIEAYYSY-	1,3
pig_chimera	DFDGQKETRTEAYYSY-	3.6.4
emb OIEN01002196.1	TMLPRCEXL	හා ග
	FIG. 180	B2

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DENLKRVLRRCMYL	DYSSFMSFNKDASKPYMRY-	SYERLLDIADKYAVDGEDWRNTAAGISNDFEKKQFQTINGFWDLLDMIEFYMCY-	QWDEFQNPEQHREHFGNKDNVICSVKKQQDLFFNFFKNNRIGY-	TYADFSKDKSDK	TYKQFKEPDKNSRLGY-	TFDEFAEPDRHKERFIKDGKLDIKLINQLKNQYDEFDAFLDDIRFGY	TYDEFKDPEHHRAAFNNNDKLINAIKAQYDEFDNFLDNPRLGY-	TFAQFSASDNPRLGY-	SYAKSKSKSKSKKEIHGY-	EYEQYCSEKSDNNERIAY- 2	NYSDFCKNKPE	SYEDFINKNLNGNELGY	GYNKVINKAINNGNRLGY-	TYKIFCDISNSGRLGY-	GYDAFRNSINQRKELLY	DYQTLRGQINRKELLY	TYDDFEKKKESTTNSREKADFDAFEKFIGNYRLAY-	TYDKFRGRSENPRLAY- 2	TYDDYTSDSKNKRSLGY-	TFDNYDFNHKNLPRLGY-	SFDVFMHPSQSTKRLGY- 2	IYDVFIDPDNSLSDDKKANVRKSLSKFNALLKTKRLGY-	TYDVFMDFSKNSLSPKDRKNIDNSRAKFEKLLSTKRLGY-
02.UC29~0_GL0096317	Cask_Anaerobic_dig	emb OGDF01008514.1	emb:0GZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD_	Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	ODAi_chimera	31009_GL0034153	DLF014_GL0011914_	CasR_Ralbus	Ga0187910_10040531	tpg DJXD01000002.1	Cask_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	EYZ-362B_GL0088915	uncultured_Ru_sp	RflavFD-1

B2

FIG. 18P

tpg(DBY161060091.1	TYEVFMEPSKNPRLDSNARKNIENSRERFEYLLDTQRLGY-	U.S.
037emb OIZA01000315.1)	XENIFYDHG	Pate
emb oc1w011587266.1	FPDILCVLEKVNGNDR	ent
k87_11092736	FGSIDVFKVTKKRE	199
BMZ-11B_GL0037915	FGSAFKPIPKKPGDKISGKYAGGKLKER	222
BMZ-11B_GL0037771	FGGIFRTQKKEDRKKDSNLSEEEKEEKKKK	Ap ı 60₹
BMZ-11B_GL0069617	LGAGFKQSERPKHQKNSRPDPRQKĮE	r. 26
Ga0099364_10024192	FGBAFRPVGDDKVKEITLSDEVRKNIEKIIALEEQKRNPSTPRFKQENINLEIENAMGKF	547
530373_GL0023589	YSRSYANKISTAKKICICLQEDFHFFDYATVKKAADGFTLLKRIGRS) 22
emb ODAI011611274.1	FGKEFCLVRNSPKQP	279
EMG_10003641	LGNALYSDSLSNLIKDE	242
Cask_F1E6_metageno	LGAAFYDVIPGKSKR	522
emb OIZX61006427.1	YSAAFYETLIDINGK	39
160582958_gene49834	FGELFYKPKETGDAKS	of 263
gi!1198542314 gb NFIR01000008.1	FGSTFCLFNENEERITS	87
emb OGNF01009141.1	FGSAFYNTPEDISAK	207
Ga0129306_1000735	YGNAFFRKLSKAERLARGREIFDKESPERRQEILGSRGKN	254
MB0288_GL0082219	YGBAFYRRLSNNERSKLGMERYERMSDAQRDAFDRKYPIDFCDG	303
PIG-022_GL0026351	FGKVFYEKVIDPKNK	S 1
PIG-028_GL0185479	FGKVFYEKSTDAKDK	022
250twins_35838_GL0110300	FDGLFSVPKREDDGKI	16,8 04.8
PIG-014_GL0226364	FGDVFKQVKLSKEQQE	812
	FIG. 18Q	B2

YGTLFEKVKAKSKKEQ. FDGVFKQITDRDSNK
GDVFKKSKKGKKDE
DDVFKKNKKPDKNKE
DNVFKKIDKKKKRN
DNVFKKIDKNKKKS
HNILNSDSDNNSKMN
DLGFEYAEPSPYKKS
GKVFIRDVKKSKLST
SEAFFCETTVKDPDS
GKAFFHAESE
WGKAFFTGQGN-
GKAFFEKNDLKHNPN
GKAFFCKEGD-
GQAFFSKEGR-
GKAFFYKDGK-
GDTFAFLDKRI
GNAFFKDEGNK
GAAFAEKVOPT
GNVFYKNKNE-
GDIFFKKEEVSKNGK
GEAFMVNSGNS
GSAFYNGDT
geaf yhene-
FIG. 18R

Cask_Esiraeum	FADAFYVNKKNPRGK	273	U.
PIG-018_GL0023397	YSDAFYBKKGKG	256	S.
PIG-025_GL0099734	YGHIFFKDKK	273	P
Ga0129317_1008067_chimera	FGSAFYSOKG-KD	205	ate
EYZ-3628.GL0088915	FGFAYNPKDR	267	ent
uncultured_Ru_sp	FGLEEPKTKD-NR	268	
RflavFD-1	FGFDYDANGKDKK	2.73	
tpg DBYI01000091.1	LSLEYDKRSKDKR	271	Apr.
037emb OIZA01000315.1	BPISEEKFYNYLNILNFIRNNTFHYKDN	294	. 26, 2022
emb ocrw011587266.1		251	
k87_11092736	ERAAADEHNAKVFRALGAIRQKLAHFKWKERAAAADEHNAKVFRALGAIRQKLAHFKWK	229	Sl
BMZ-118_GL0037915	DAAIAEDVAVLRVLCGLRQISAHFRSEKKGE	253	heet
BMZ-118_GL0037771	QEELLQKDLQHNMDVARCISALRHATAHNKPATAHDKQDE	249	41
BMZ-11B_GL0069617		228	of
Ga0099364_10024192	KSKDAFETAKKKYNRIVADETNAKTLRILGAMRQITAHFKDQ	286	87
530373_GLO023589	NYFFDMLCDSAGNYQCKNAQILLSILGMLRQESFHENRST	298	
emb ODAI011611274.1	DKEEKRQYKLMRVLCLLGELRQFLVHGKKKEKE	312	
EMG_10003641	ILDGKRSKELKKYYQELCLLGMVRQSMIHSNQF	275	US
Cask_P1E0_metageno	DAARGRVKREQDVYAILSLMSLLRQFCAHDSVRIWGQNT	261	S 1.
emb OIZX01000427.1	SERRSNEDIFNILALMTCLRNFSSHHSIAIKVKDYS	289	1,3
160582958_gene49834	DYKTHLSNNQKEEWELKSDKEVYNIFAILCDLRHFCTHGESITFSG	339	16,8
gi 1198542314 gb NFIR01000008.1	ENKKEWKRFEKKCYHLLAVLGMMRQATAHGDSK	304	812
	FIG. 18S		B2

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253	782	344	252	254	305	259	309	287	321	317	320	3,2.2	26.0	253	279	281	87 87	249	247	258	253	236	245
SDAEDFDGT	KSVDDEIRALAPEWVKREERDVYSELVLMSELRQSCFHGQQKN	ELHFREIEKLAPEVVRKSEKEIYYILAMLSELRQCVFHDSAAS	SKKKMRLISEKEMYYIFALLGGLRQFCTHDKNR	hkkkmrlisekemyifallgglrofcthdknra	KESEKERAKDQNFDVLRLLSVGRQLTFHSDKSNNE	-KSEEEKRRIWQELWNKNFDHNKNIVNSLIKFRQMCFHQNYENSDKEVS	RAKEAEIDACTAHNYDVLRLLSLMRQLCMHSVAGTAFKLA	DREIKNSYNALVLKVLYYLRQFCMHGNTYTKRNEESFLS	NNEDYEKNLRHNFNVLRVLSFLRQICTHAYVKCTGGAKNNGDSTKVEAES	GDNSKQYQENLRHNFNILRVLSFLRQICMHAEVHVSDDEGCTRTQNYTDS	DDYEGKRNEILRYNFWVLRVLSFIRQICAHAKVKVSSDNDREKGGVFVDS	DDCKRERNEILRYNFNVLRVLSFLRQICAHAQVKISNEADREKGGGLVDS	YNKVNKGKEEKDNRNNENIEKLKKALEVIKIIRVDSFHGVDGIKGDQKF	EQQIEGERERERERELETIKKALDVISVMRTDSFHGYDMCNSTSAIVKF	GKGEKIEVMYRSDEEIFTIFQILSYVRQSIMHNDIGKGEKIEVMYRSDEEIFTIFQILSYVRQSIMHNDI	GRIVPCLEQRSDGDIYNILRILSIVRQTCMHDNASM	RKIVKKTEKEVYHILTLIGSLRQWITHSTEGGIS		KKKRDKNPYILKYDNECYYIIALLSGLRHWNIHSHAKDDLVS	KYLNKQDNERYHILALLSGLRNWVVANNEVESKID	NYIINYGNECYDILALLSGLRHWVVHNMEEESRIS	RNNRQKDPIECYALLALLCGLRNWVVHNNEEKDLIK	ADADKEKQVYAMLACLGSLRQACSHYRIRYSVNGKNVDADA
emb OGNF01009141,1	Ga0129306_1000735	MH0288_GL0082219	PIG-022_GL0026351	PIG-028_GL0185479	250twins_35838_GL0110300	PIG-014_GL0226364	250twins_36050_GL0158985	emb OJMM01002900.1	emb OGPN01002610.1	emb OHCP01000044.1	PIG-046_GL0077813	pigchimera	emb OIEN01002196.1	O2,UC29-0_GL0096317	Cask_Anaerobic_dig	emb OGDF01008514.1	emb OGZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD_	Ga0224415_10007274	Ga0187911_10069260

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FIG. 18T

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VEVLKSEKEIFDILALLGSLROFCFHYDEAVFENEDDGVIDKEYND 315 -TKILRPEKEIZHITALLGSLROFCFHYDEAVFENEDDGWIDKEZND 296 RRDEKVIZHILSLAASVROFCFHYDZISDGKGRA 306 RRDEKVIZHILGKLRHWCVHSEEGRA 301 KRSKKEIYDILALIATIKGWCHSEE
-TKLRPEKEIXHIFALMASLRQSYFHCXVKDTDYGSRRDEKVIXHILSLAASVRQFCFHNDYISDDGKGRA
RRDEKVIYHILSLAASVRQFCFHNDVISDDGKGFIK 290
######################################
KNYLREDKELYSVITLIGKLRHWCVHSEEGRA
-SKERSEKEIYSMLAMLGKLEHWCVHSESGEA 287
-FEKKSDEEVYNICALMGQIRQCCFHGKQEKYQ 237NQKKRLYYLMSMIGQLRQFAFHXDS
-VSQAYKKRVYHMLAIVGQIRQCVFHDKS
-KSEEIKKRLYHLVAFAGQLRQWSFHSAGGLP 302
-KSEEIKKRLYHLVAFAGQLRQWSFHSVEGLP 302 * *
* *
PIRFFQGATGGRQLWNDVIAPLWKKRIERVRKS-F 274
T
T
1LKKEFSKADW-QTVEDYXAKLVDRINEG-F 327

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332	346	334	294	32.2	374	346	300	888 888	377	286	288	349	293	343	324	35.88	354	357	359	294	284	312	322	
	QEYRKLLGEFYDAQVDKVNK-SF	SSIYTLDSSYDSTMNTAELLGKGDDSSLVALATDARVEARAILDELYKKGVDSINN-SF	ODMRDLLDDGWRRALGGVND-HF	PDMKKMLDTFYTEAFIQLMQ-SF		BARGCRPBARKELDELYKKKIHEMNQ-GF	KLYNKKVHMALTGMKKVLDANFNKKVEHLMN-SF	FGVDGARELLDRLYAEKINDLR-SF	ABAMQVLDGVYSQKVAALNNF	BDAKAVLDKYYVSTVSSLNK-HF	DDAKAVLDKYYVSTVSSLNK-HF	AQDENRRQDIQSLLNILNSTCRSNLEGVNGD-F	FINKEEKASLDEIYTGAIDKVNCD-F	SADLKEILDEAFSGAVNKLNDG-F	ADPQINELIDAVYADGIKTINSD-F	TAPELSKIINEIYKEGIDRINND-F	KMPELKTLIDNIYSKGINAINDE-F	VAPELNEVINSVYSKGIYDINDN-F	VAPELNEVINSVYSKGIDDINDN-F	YNEEIQKTISEPENRKVEEVQQD-F	KGKNLIYRNAFKYKTDTIKKQ-Y	YPARFVGFLSDLLKTKTNDVNRM-F	DRKNGFDELAELLDYLYDEKIDIVNRD-F	
SAWMFNLEKVLT	FAWLYRLDRQLS	NSSIYTLDSSYDSTWNTAELL	TAALYHLQALPTAALYHL	AAGLYNLRRLG	KPFPYNLEKNLF	RAEIYKLGKEFDRL	SVIMYNLDNEELYK	SARIFRLDNDLG	RSWLYNLEKVLP	RNWLYSLDTMLN	RNWLYSLDTMLN	-AYLFDLSKLTRA	STALFNIEQFL	ESALFNIEDVL	DIALYNAKEFFAK	LDALFNITEYFAK	LEALFNISKAFGK	LDALFNISSFFDA	LDALFNISRFFDA	PRSKYNLAVN	PSKKFNLDQ	KSSILAIBK	RTVÆTLGQNSVR	FIG. 18V
530373_GL0023589	emb ODAI011611274.1	EMG10003641	Cask_P1E0_metageno	emb 012X01000427.1	160582958_gene49834	gill198542314(gb NFIR01000008.11	emb OGNF01009141.1	Ga0129306_1000735	MH0288_GL0082219	PiG-022_GL0026351	PIG-028_GL0185479	250twins_35838_Gh0110300	PIG-014_GL0226364	250twins_36050_GL0158985	emb 0JMM01002900.1	emb OGPN01002610.1	emb OHCP01060044.1	PIG-046_GL0077813	pig_chimera	emb OlEN01062196.1	02.UC29-0_GL0096317	Cask_Anaerobic_dig	emb OGDF01008514.1	

FIG. 18W

FIG. 18X

FIG. 18Y

-EDN-KVAISLLIDBURKG —— ANEKDIJDLIKAYERE ARGESIKKIRERMIDE 386 -TKN-AINIXIIKEVFPE —— ANEKDIJDLIKAYERE ARGESIKKIRERMING —— ANSOMKKVAFP ARGESIKKIRAKIRIKKIRIKKIRIKKIRIKKIRIKKIRIKKIR	Ga0129317_1008067_chimera
:	l t
PYDKEKEVAKLSSVKHKLYKIYDFVITHYLNSNK ##############################	
### ##################################	GYCPLP
HSSAPDVKRKVDTFRSKFYAILDFIIYEASVSVANSGOMGKVAFW 380ARTHKQFDSYRGKLYTIIDFIIYEMAAED 1KDKQYDSYRGKIYTILDFILFRALSQW	KNIDL
376	HELGER
The contraction of the contrac	HYPD-
The company of the	YASG-
TAPETKKRYDSYRAKIYGLTDFLLFKHIBNT	FYSE
	FVPELT
TLLLCDEGSTVKLNVHDTMKSKFYKNLDFMIYKYYKYG	OOELS
TLILCDEGSTVKLNVHDTMKSKFYKNLDFMLYKYYRYE	YKH-
	CSEA
VIADKRYDTCRSKLYNLMDFLILRVYRTGR 412	PDAA-
	PDAA-
	PDAA-
EFDKISNKKFDSVRSKLMRLADYIIYDYYNK	MDEDK
421	HDSNK
N 466	HAA
	FAP-

FIG. 18AA

436 426	410 P	ate	ent	411	425	Ap ı	117	613	22	423	Sheet	50	425 Jo	87	424	423	400 US	S 1	1,3	16,8	812	B2
YEAECYKAECYNSERQKLYKLIDFVIYYSFIHYK	PELSGYKEDQYNSVRSKLYKLFDFIIAHYFRKH	SDLKSFKEDKYNSVRAKLYKLFDFIITYYYDHH	KGYADKEYDSVRNKLYQMTDFILYTGYINED	TAYTSNPDDRLYDKEIMSVRNKLYQMIDFIIYLEYRDN	TVTV	EEASDFRDKDYDSVRRKLYKLMDFCIYYLYYS-D	ALRDEKIDSFRSKLYRLIDFCLFRIYLK-D	YGFRFKDKQYDSVRSKMYKIMDFLLFCNYYRND	DGADRIKEQDMDSVRSKLYKLIDFCIFKYYBE-F	DEAEKIRSKDMNSVRSKLYKLFDFCIFYQYFI-D		DKLLLEIVETLRLSKND-DEKENVYKKYAE-KLFKADDVINPIKAISKLFAE	DSFVDDFVAALRASQSE-EEKEKLYAQYSB-RLFADEGLKSALKKAVDMISD	KGAIDNALVKLREAPDE-EAKEKIYNVLAA-SIRNDSLFLRLKSACDKFG	GAFLEESVQKLRSTPSE-EEKEKVYDVLAE-TIWNATKDDVKKVFDIRQQ	QGIIDKTVSSLRLTKDE-EEKDHVYQNAAE-LVWGMVSNCLTPYFNDPKN	-EELETMVADLRESESD-EKKEELYHDYAE-LFWDGVKDLIVPFFAAFDG	-KQLEEWVAVLRETSNE-DAKENLYDEFAR-TAMNTVGDSAKQLIENMQS	EEQIDQFVRKLRASTSN-VEKEALYQQESE-RTWNEIRGQIIDNVIPFID	AERREAFVAKLRAVMTA-EAKQRAYADEAA-EIWNDEGSGIRAAFLEILE	PEKGEKLIEDLRSKIKGKKKED-EDKKQRYABESA-CILKAKRDIIKKDLTEAAN	FIG. 18BB
cask_Ralbus	Ga0187910_10040531	tpg:DJXD01000002.1	Cask_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	EYZ-362B_GL0088915	uncultured_Ru_sp	RflavFD-1	tpg DBY101000091.1		037emb OIZA01000315.1	emb OCTW011587266.1	k87_11092736	BMZ-118_GL0037915	BMZ-11B_GL0037771	BMZ-11B_GL0069617	Ga0099364_10024192	530373_GL0023589	emb ODAIO11611274.1	EMG_10003641	

Sa0224415_10048792_chimera

amb|OGDF01008514.1|

emb|0GZC01000639.1

Cask_Anaerobic_dig

emb|OCVV012889144.1

gill198542314:gb|NFIR01000008.11

emb|OGNF01009141.1

3a0129306_1000735

160582958_gene49834

emb|OIZX01000427.1|

Cask_P1E0_metageno

250twins_35838_GL0110300

PIG-014_GL0226364

PIG-028_GL0185479

MH0288_GL0082219 PIG-022_GL0026351 250twins_36050_GL0158985

emb|OGPN01002610.1|

emb|OHCF01006044.1|

PIG-046_GL0077813

pig_chimera

emb!OJMM01002900,1

emb|OlEN01002196.1| O2.UC29-0_GL0096317

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4 ቁ ቁ ይ ቁ ይ	422 422	491	491	41. Q	484	474	458	403	471	459	473	406	459	467	471	471		513	464	475	
LSEKDIFITSLRGSFSE-EQKDKLYSDEAE-RLWAKLGKLMLETKKFRGQ LSEKDIFVINLRGSFND-DQKDALXYDEAN-RIWRKLENIMHNIKEFRGN	LSEKDIFVISLRGSFNE-DQKDRLYYDEAQ-RLWSKVGKLMLKIKKFRGK PDMVDAFVVELRSLAKDEDAKNAVYEKYAK-AVWNDVKQPIAVMLSYMNG	PDKINDFVEKLRIQNKN-DEKIKLYYDBAV-CLLSELGREIHTMTSCVHN	AKEKDILVSSLRASLND-TKKENLYAGBAK-TIAESIKDRMDTVCRMYNK	KDRIDDIVSKLRGAIYD-YKKDDIYKABAD-WVWKKLNRQIKALIKLLSN	QDEVDEIVNYLRSTLSE-EQKKEKYIKLAG-NIWAEYKTQFNKLKDILLA	PARIEEIVDKLRESVND-EEKESIYSVEAK-YVYESLSKVLDKSLKNSVS	PEKGEEMVDCLRLCMTE-DEKDSHYEGTAK-KLVRELAYDNQEAAEQANG	AFEKEALVSSLRSSLTE-ENKEELYIKTAR-TLASALGADFKKAAADVNA	SDRADDIVNTLRSSL	KSIIDTIVSKLRSSVTD-EAKEAVYREEAA-NLRGRLLDAVALIAKSLNG	KEHADSIVEKLRATLSG-ESKEEIYKAEAN-KIWNLYEDIIMNKIEPALD	SERNENLYSRLRESLTD-ENKDIIYSKEAK-IVWNELRKKFSTILDNVKG	PDRAEKNVRFLRSCINE-QEKEQFYTQEAN-ALKSENPGLVNDFCMQLKD	IAAGESIVRKLRFSMTDDEKEGIYADEAAKLWGKFRNDFENIADHMNG	PELSEKNYDIIRAAVSD-TKKDNLYSDEAA-RLWSIFKEKFLGFCDKIVV	EERSRENVNYLRSTLND-EQKDAFYEEEGK-RLWSENRKKFIYFCDNINK	* * *	KGNKLFKEKIIIKKEYIEDVSIDKNIYDFTKVIFFMTCF	TKSNIFKMKTPLDKALIENIKVNSDASDFCKLIYVFTRF	AEGNRPVFPNELRNNRDIRNVRSEWLEATQDVDAAAFVQLIAFLCNF	PIG. 18DD
Ga0187910_10006931 Rflav_XPD_	Ga0224415_10007274 Ga0187911_10069260	Ga0187910_10015336	ODAL_chimera	31009_GL0034153	DLF014_GL0011914_	CasR_Ralbus	Ga0187910_10040531	tpg:DJXD01000002.1:	CasR_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	EYZ-362B_GL0088915	uncultured_Ru_sp	RflavFD-1	tpg DBXI01000091.1		037emb OIZA01000315.1	emb Ocrw011587266.1	k87_11092736	

U.	S.	Pa	ate	ent			Apı	. 26	5, 20)22		SI	heet	53	of	87			US	S 1	1,3	16,8	812
462	464	438	508	513	534	535	483	517	562	543	489	516	576	474	476	525	471	526	498	547	535	540	542
DIKNAAAQSLODVNLDSVTVEKEFFVVKLLAFLCNF	KYILKYKDAKIPGDFEDWITSKISEDDGEPFVKVLSFLCNF	DFTVFTTEKVPADTSKEPFLSSANTEPLVKLLAFLCNF	YFTKKEKEITRTAQPVLSTSSIAHTSKKITQFSSFAKLLAFLCNF	SIQKQENEDKLIKKIDQSIIDKAVPHFVNSKDLSYFAKVVYVVASF	AVDFGSAVKGIKARSSVAGDKRFAEWLEEVRIRPEGVSCFIKLMYLLTRF	KDLFADLVKSNKNEKQKFKNEYEBLLKPFMIPVKVDYFSELIYLVTRF	GKKLQDKDK-KKSDELGLSRDVLDGVLFRPAQQGSRANADYFCRLMHLSTWF	PENLSRLSGOKRKGELSLDDAMLKECLYEPGPVPEDAAPEEANAEYFCRMIYLATLF	IPSLSKDE	RILSEMDERRNKKVNQESSDTDREEPLDSELAEGITFIKETAHSFSEMIYLLTVF	KIIGNMQPDSTITASMLHNTGKDWHPISENAHYFTKWIYTLTFF	VVKGRTKLINKLKLSADESTLVRNAIDGVRISPRASYFTKLIYLMTLF	VILSNNTPGSIKIKELALSKYDRDLVENAIDSAVDERGRELKITDEASYFSKLIYLLILF	SVKSKYGKLELDKQTINQLKRAVDDISINTKGDYFTKLMYLISLL	LVKSKYGKLELDTQTVNQLKRAVDDISINTKGDYFTKLMYLISLL	QYKTKFKSEFKGGISIENMQQQMILLQTENIDYFSKYVLFLTKF	EKEKKKEFSSKIKIDLRSVQKIEDQNASVFAKMVYLISRF	EKRNKFRSKVALPDVSGAAYMLSSENIDYFVKMLFFVCKF	ESLSKFKGYKDIDESLISRYGITVANTDTLVKILYFLCKF	ELVPVKVEGKNDPQQFTHGKLGKKEIESFCLSDKNTSDIAKVVYFLCNF	EAEGIIIPGKEDPVKFSHGKLDKKEIESFCLTTKNTEDITKVIYFLCKF	BAKGVIVPDKEEAVKFSNGALDKSBIBRFCITSANTDSVAKIIYFLCKF	EAKDVIVEDKKKPVKFSHGKLDKNEIERFCITSANTDSVAKIIVFLCKF
BMZ-11B_GL0037915	BMZ-11B_GL0037771	BMZ-11B_GL0069617	Ga0099364_10024192	530373_GL0023589	emb ODAI011611274.1	EMG_10003641	Cask_P1E0_metageno	emb 012X01006427.1	160582958_gene49834	gi 1198542314 gb NFIR01000008.1	emb OGNF01009141.1	Ga0129306_1000735	MH0288_GL0082219	FIG-022_GL0026351	PIG-028_GL0185479	250twins_35838_GL0110300	FIG-014_GL0226364	250twins_36050_GL0158985	emb OJMM01002900,1	emb OGPN01002610.1	emb OHCP01000044.1	PIG-046_GL0077813	pig_chimera

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FIG. 18EE

U.	S.	Pa	ate	nt			Apr	. 26	5, 20	22		Sł	ıeet	54	of S	87			US	5 13	1,31	16,8	312	B2
483	470	494	200	52.6	467	468	480	475	45.58	464	534	535	462	532	518	503	443	513	201	5135	447	505	508	
YIKNSREKNLSGGSSLPKYSFIEGFTKRSKKINDNDEKNADLFCNMLYYLAQF	SIKLYSTKNNVDDQFMLPFEQVSKDIKGGDAAREKADAFCDMTYYLTQF	GDALKEIKRKNRDRKLPQSVIATVQVNSDANVFSGLIYFLTLF	PAVIGKIKGPAVIGKIKG	QVKEYSSKN-MPIPIQKQIQNILKFAEQVTYFTKLMYLLTMF	LASRIPDRD-GNISEFVESLPRIHRLLPRGQRISNFSKLMYLLIMF	KVKEYKKYKEYKKKEVPRIERILPQGKDISAFSKLMYMLSMF	MIRDYKKSDIPILNRILPESEDVSIFSKIMYALIMF	KTREYKKTRAFILPRILPAGRDVSAFSKLMYALTMF	DTRKYKNMGTPRIRRLIPEGRDISTFSKIMYALTMF	SAIKNIKAFEL	IENTSYEITDKKQKEYYKMQINSLNSADKVSDFSKVIYLLF	GEGIKHITVDRFIAGAGAGKLIDKCSISTECDYFCKFIXLMTLF	RRIFSQDNDITNWNGVEFEKIGEQKLEDMADYFCKLMYFVTLF	NIKEFSDAKEDDVYYEEFKNFKFNEVGKEKLGENAEYFCKLMYLLTLF	GETIKDLQKRYDDETANRIWDISQRSISGNVNCFCKLIYIMTLM	SNITQMQKNEQQGKT-KGMFAIRDEIRVSRKPVSYFSKVIYVMTLL	KNIRDYQKKANDYRISFEDIKIGNTGIGYFSELIYMLILL	DNIKKLSKSNIEIQEDKLRKCFISYADSVSEFTKLIYLLTRF	VNIKKLQQNTLEIFDSKLENCFIKSAEKVSAFTKLNYLLTRF	EGVISGLKKNKSYDNMSIEGIVAEBTANVSYFSKIIYLLAQF	SNIKKLENVKEKFISEDEFDDIKLDIDISYFSKLMYVMCYF	RYSSGSNSTENARKKEDFPVFFHCHDRQKVSNVSLFAKLMYAFCLF	DVIKELGKADMDFDEKILDSEKKNASDLLXFSKMIYMLTYF	FIG. 18FF
emb OIEN01002196,1	02.UC29-0_GL0096317	Cask_Anaerobic_dig	emb!OGDF01008514.1	emb OGZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XFD_	Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	ODAL_chimera	31009_GL0034153	DLF014_GL0011914_	CasR_Ralbus	Ga0187910_10040531	tpg bJXD01000002.1	CasR_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	EYZ-362B_GL0088915	uncultured_Ru_sp	

RflavFD-1 tpg DBXI01000091.1	WVIGEHEK	509 509	U.S.
	•		Pate
	LDGKEINDLITNIISKLQVIEDHNNVIKFIFHNKD	548	ent
	LDGKEINILLNSLIKKFODIHSFNTTVKKLSE	499	
	LEGKEINELVTALIKKFEGIQALIDLERNLEGV	508	
	LDGKEINELLSTFVNRLENVQALIDLAKNLGET	495	Apı
	LEGKEINELLTAYIHKFECIQDFLNVISSLGEN	497	:. 26
	WDGKEVNEVLSAYIRKFESIQAFIDVLESKEINEKV	474	5, 20
	WEGKEINELLSAYIHKFENIQEFINLLEKLEGK	541	22
	LDSKEINMFFDALINELENIASFNDVLEQLDM	545	
	LDGKEINELLTGLINKLENIQSFLDVMQQEHA	566	SI
	LSGKEINDLLTQLINKFENIAAFIRMYQNDQG	567	1eet
	MDGKEINTLLTTLISKLENIDSLRSVLESMGL	515	55
	MDGKEINTLLTTLISKFENIAAFLQTMEQLNI	549	of S
	LDGKEINLLCTSLIEKFENIASFNEVLKSPQIG	595	87
gi 1198542314 gb NFIR01000008.1	LDGKEINILLTQLIHCFONISSFMDTMEEENL	57.00	
	MDGKEINDLVTTLINKFUNIASFIEVLKSQSV	521	
	LDGKEINDLLTTLIHAFENIDSFLSVLGSERL	548	US
	LDGKEINDLITTLVNSFDEIDSFCCVLQKAKL	808	S 12
	IDGKEINDLLTTLIHQFENIASFIDVMKKENI	50.6	1,31
	IDGKEINDLLTTLIHQFENIASFIDVMKKENI	508	16,8
	LDGKEINELLCALINKFONIADLDISKQIGT	557	312
	FIG. 18GG		B2

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503	558	530.	579	567	572	574	515	502	554	532	558	499	50.0	512	507	490	496	566	567	494	564	550	ಸು ಪು	
LDGKEINELCTKLLNKFKNINELVETAKRCGE	LDGKEINELLCALINKFDNIADILDAAQCGS	LDGKEINELCCAMINKFDNINDLIKTAAQCGE	LDGKEINELCCAMMNKFDGIGDLIDTAKQCGE	LDGKEINELCCAMMNKLDGISDLIETAKQCGE	LDGKEINEFCCAMMNKLDGINDLMETAEQCGA	LDGKEINELCCAMMNKLDGINDLIETAEQCGA	LDGKEINIFLTSIHNIFQNIDSFLKVMKEKGM	LDGKEINVFLTTLHNLFENIASFLDVMHKQNI	LDGKEINEMVSNLITKFENIDSLLHVDREIYKSDEKDLDLEIEKLALFFKGVVRPNAKTD	QDGKEINELVSSLVNKFANIQSFVDVMRSQGI	LDGKEINDLLTTLINKFDNISSLLKTMEQLEL	LDGKEINDLLTTLINKFENIQGFLDIMPEINV	LDGKEINDLFTTLINKFDNIQCFLKIMPLIGV	LDGKEINELLTTLINKFDNIQSMLKIMPLIGV	LDGKEINDLLTTLINKFDNIQSFLKVMPLIGV	LDGKEINDLLTTLINKFDNIQSFLKVMPLIGV	LDGKEINDLLTTLVNKFDNIHSFNQVLTALGL	LDGKEINDLLTTLINKFDNIASLLSVLEKQSG	IDGREINDLLTTLINKFDNIDSFSVDKAVNNQ	LDGKEINDLLTTLINKFDNIRSFIEVMEEMSL	LDGKEINDLLTTLINKFDNIRSFVELMKEENI	LDGKEINDLLTTLVNKFDMIASFIDVMDELGL	LDGKEINDLLTTLINKFENIVSFEDVLRQINV	FIG. 18HH
PIG-014_GL0226364	250twins_36050_GL0158985	emb OJMM01002900.1	emb OGPN01002610,1	emb OHCP01000044.1	FIG-046_GL0077813	pig_chimera	emb OlEN01002196.1	02.UC29-0_GL0096317	Cask_Anaerobic_dig	emb OGDF01008514.1	emb OGZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD	Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	ODAI_chimera	31009_GL0034153	DLF014_GL0011914_	Cask_Ralbus	Ga0187910_10040531	

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475 533 479 479	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	577 528 526 526 572 572	5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5
HISFIDILKKLNL HRSFLEIMDELGL HRSFLETMOELGL HRSFIDIANQIGL	IACMIKCIKDLGI IIKEFLKIMKSSAVD IIANQIKTAKELGI IIRSFIDTANFLNI	NA	DIPKDGKRVLSGS
LDGKEINDLLTTLINKFDNIISFIDILKKLML. LSGKEINDLVTTLINKFDNIRSFLEIMDELGL- INGKEINDLLTTLINKFDNIRSFLETMDELGL- IDGKEVNDLTTTLINKFDNIRSFIDTANQIGL- LDGKEINDLLTTLINKFDNIGSIIEAATQIGI-	LDGKEINELLSALLNKFDTIACMIKCTRDLGI LDGKEINDLTTLINKFDNIANQIKTAKELGI LDGKEINDLLTTLINKFDNIRSFIDTANFLNI**:* . : :	AVYKDYSDKYAIFR 1INADYVDDYSLFEDSIRFENEFALFNVNFSKNYAVFN VFSAKKGRNFSAFN VFSAKKGRNFSAFN	ETGLSDAFSFFE
tpg:DJXD01000002.1; CasR_Esiraeum PIG-018_GL0023397 PIG-025_GL0099734 Ga0129317_1008067_chimera	EXZ-36ZB_GL0088915 uncultured_Ru_sp RflavFD-1 tpg DBY101000091.1	037emb 01ZA01000315.1 emb 0CTW011587266.1 k87_11092736 BMZ-11B_GL0037915 BMZ-11B_GL0037771 BMZ-11B_GL0069617 Ga0099364_10024192	emb ODAIO11611274.1 EMG_10003641 CasR_P1E0_metageno emb OIZX01000427.1

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602	548	575	635	533	535	584	530	585	557	606	594	599	601	542	529	583	559	585	526	527	539	534	51.7	
KEISRELRIINSF	AEICSELSAMNSF	GVIAQELRAVNSF	AEISQELRAINSF	KNIACELRIVNSF	EIIAHELRTVNSF		AKLAEEMRFVQSV	RRISAQİRIVKNI	NDLSDQIRIVKSI	RKITNDIRVAKSI	ATMSNQIRIVKNI	GTISGQIRVVKSI	ETISDQIRIVKSI	GHVAKKIEIVISL	AYIVKKLETVISV		GRISRELHILKGI	SRLCKEITQLKSF	HEIAGELKLIKGF	EKIADELRLIKSF	EKIADELKLIKSF	AKIADELRLIKSF	EKIADELRLIKSF	
S.H	BG	SQ	r u r r r r R S H r r r r r r r r r r r r r r r r r r		NS	DA		RS	SN	NC	KC		NC	HA	DS	SA	Ś.Ţ	SÖ	KS	K.S.	SN	SO	SN	
LIKLKEDYBIFE	CIHFSERKMEI	ERTEDANYRIFA	DRGFTEGYRFFA	NYEFAEKYKEFN	ECEFTERFAFFK	PVVECADYESLN	TIEFCPEYKMFK	SVWFVDSYRFFE	DIEFVKEYKLFI	EVKFIEEFACLS	DVEFVDQFKCLS	KİREVDNENVLS	KVEFVDKFSVLS	ECKFORDFKMFS	SYDFIEQYVMFY	TGAGEISKSFSIFQ	DSGFTADYAMFA	QTTFKEDYTFFQ	NAKFEPEYVFFN	NVKFTEDYAFFN	NAKESSDYAFFN	NAKEVEEYAFFK	NAKFAEEYSFFW	FIG. 18JJ
gi 1198542314 gb NFIR01000008.1	emb OGNF01009141.1	Ga0129306_1000735	MH0288_GL0082219	PIG-022_GL0026351	FIG-028_GL0185479	250twins_35838_GL0110300	PIG-014_GL0226364	250twins_36050_GL0158985	emb OJMM01002900.1	emb OGPN01002610.1	emb OHCF01000044.1	PIG-046GL0077813	pig_chimera	emb OIEN01062196.1	02.UC29-0_GL0096317	Cask_Angerobic_dig	emb QGDF01008514.1	emb 0GZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD_	Ga0224415_10007274	

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523	609	595	521	591	577	564	505	572	560	574	506	564	569	568	568		620	566	571	558	55.59	544	
-DSGRVVEYLREINSF	SSNLLKEKTINKSENYTCKIVEELREINSF	KAGEICESLRSINSF	-ESKDICLRLKEIHSF	ESKNVCSTLREINSF	-DSKAICLDLQFINSF	-Y-DRCRNISGELRLINSE	-NMT	-GSTKYLAELVELNSF	-DSQKYRDELIELNSF	TSDIVKKELHVIKNL	-RSKDISVELNIIRNF	DSBQRAEEMNLVKSI	DSPELFIVKNI	HSEKYVDELNIVKNI	IICDYAGELNIIKNI		NAPREPLIKDALIALGVSSNDFDERYERYFRIDVD	-NINLSFMYDDALRTLGVSDENLPEVKREYF	-DAKRVLYKSALEILGAPPDEVS	SLKPLFIRFALQILGGARISDEDV	EAKRPLYKAAIRMLCPPEKWEKYIS	GAKRALFRDAVETLGIVENSNWENEDGTIS	
SASYEADYKIFE	-KKVEFVENYSFFN	-NRVEFGDKYDMFN	KCEFMEEYKFFN	ECIFDKKEAFFD	EHSFŢDNYKMFA	DCTEKPEFAFG	EFKFKPEYADFF	DRIFIAEYSFFE	ERTETENEEE	ECNTIDEYQFFN	NIEFIDDEKFFD	TAPESKDFNFFE	-VECELTAGYKLFN	NIAFVKNYDFFW	DVKETKDYDFFN		ARMENKIENAPKEPLLI	SKMDFGLDNINLSFMY	GKMKPDMTBAKRVLY	GKMDAGLUSLKPLFI	GKMKPDLTEAKRPLY	GKMKPDLEGAKRALF	FIG. 18KK
Ga0187911_10069260	Ga0187910_10015336	ODAIchimera	31009_GL0034153	DLF014_GL0011914	Cask_Ralbus	Ga0187910_10040531	tpg DJXD01000002.1	CasR_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	BYZ-362B_GL0088915	uncultured_Ru_sp	RflavFD-1	tpg DBYI01000091.1		037emb OIZA01000315.1	emb OCTW011587266.1	k87_11092736	BMZ-11B_GL0037915	BMZ-11B_GL0037771	BMZ-11B_GL0069617	

FIG. 1811

FIG. 18MM

REC. 11092736 REC. 11 11092736 REC. 11 11092736 REC. 11 11092736 REC. 11 11092736 REC. 11092736 REC. 11092736 REC. 11092736 REC. 11092736 REC. 11092737 REC. 1	037emb 01ZA01000315.1		620	U.
Colored Colo			566	S.
1	7_11092736		592	Pa
1 4.1 AEXIQPSEER	11B_GL0037915		576	ıte
7 4.1 ARYLOFSEER	11B_GL0037771		580	nt
92 ————————————————————————————————————	1-11B_GL0069617	n' yan app yan app pan an an yan a	563	
4.1 AEYLOFCEEEK	0099364_10024192		654	A
# 1 AEXILGESEER. 649 111 AEXILGESEER. 649 111	3373_GL0023589		617	Apr.
No YOMTDYCKIYKDFLSANKTLDCKSKNMQAFVSELKNAALSENYEGAETYELADT 711 11 12 134 14 15 16 17 18 18 19 19 19 19 19 19 19 19		AEYLQF SEEEK	649	26,
11 615 627 642 642 642 642 642 642 642 643 644 6	1,10003641		7111	202
11 11 12 13 14 15 15 15 15 15 15 15	R_P1E0_metageno		582	2
NFTRO1000008.1 667 INFIRO1000008.1 642 1	OIZX01000427.1		615	
NFIR0100008.1	582958 <u></u> gene49834		667	She
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1198542314 gb NF1R01000008.1		642	et 6
1 1 2 2 2 2 2 2 2 3 4 4 4 4 4 5 7 8 7 8 7 8 7 8 7 8 7 8 7 8 7 8 7 8 7	OGNE01009141.1		586	2 of
1 L0110300 L0158985 L01589885 L015898	129306_1000735		617	f 87
0110300	288_GL0082219		677	
0110300	-022_GL0026351		573	
0110300	-028_GL0185479		575	U
0158985	twins_35838_GL0110300		644	S 1
158985 ——————————————————————————————————	-014_GL0226364		598	1,3
FIG. 18NN	twins_36050_GL0158985		655	16,
	OJMM01002900.1		621	812
		FIG. 18NN		B2

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667	655	660	662	585	572	627	611	623	564	565	577	572	555	566	670	939	ଅବସ୍ଥ	65 60 60	620	୧୦୧	545	615	603	
GKRLYIQEYKYFNDMFFNA	GKKVYTKDYNNFGDMFFEG	GKVLRTKEYNNIRKMLCKK	GKYLYSKEYDDFQYMFFKD		······································							.			DWKKRIDDEQQEYDRLLN	EWEW								FIG. 1800
emb OGPN01002610.1	emb OHCP01000044.1	PIG-046_GL0077813	pigchimera	emb OIEN01002196.1	O2.UC29-0_GL0096317	CasR_Anaerobic_dig	emb OGDF01008514.1	emb 0GZC01000639.1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD	Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	ODAI_chimera	31009_GL0034153	DLF014_GL0011914_	Cask_R,_albus	Ga0187910_10040531	tpg DJXD01000002.1	CasR_Esiraeum	PIG-018_GL0023397	

PIG-025_GL0099734 Ga0129317_1008067_chimera	618 S44	U.S.
EYZ-362B_GL0088915	\hat{g}_{0}	
uncultured_Ru_sp	607	
RflavFD-1	611	
tpg DBY1010000091.1		
		Apı
037emb OIZA01000315.1	ADKDHQKVSTFLMNNVINNSRFKY 644	
emb OCTW011587266.1	GKTKNLSAYIRNNVLENRRFKY 588	တ
k87_11092736	KTVNPFRNYIAKNVITSRSFYY 614	
BMZ-11B_GL0037915	RSNTFRNFITNNVIKSRRFVY 597	<i>t</i> -
BMZ-11B_GL0037771	KQVNPFRNFIAGNVIESRRFMY 602	
BMZ-11B_GL0069617	RAIKPFRNFIANNVIESRRFMY 585	ieet
Ga0099364_10024192	WRSLIDTEYLMPETNPFRNFVAKQVIESRRFMY 687	
530373_GL0023589		
emb ODAI011611274.1		
EMG_10003641	NLPAYFSEEDKEKLARYIVHSDGTYKKFLKESFYALEELPNEGFRNFISNNVINSRRFNY 771	er.
CasR_P1E0_metageno		E
emb OIZX01000427.1		
160582958_gene49834	XIYSFSGKKIPNKNFRNFIINNVITSRRFLY 698	
gi 1198542314 gb NFIR01000008.1	TKNKINDKKKKGFVRYIMNNVIKSTRFRY 671	
emb OGNF01009141.1	FDKSASKKEKGFRNFTRNNVVDSMRFKY 614	ঘা:
Ga0129306_1000735		
	FIG. 18PP	B2

MH0288_GL0082219		
PIG-022_GL0026351	KKTRLINARSKADTGLRNFIISNVIKSSRFKY 604	
PIG-028_GL0185479	KKTRLIARSKADMGLRNFIISNVIKSSRFKY 606	Pa 90
250twins_35838_GL0110300	T	
PIG-014_GL0226364	DBMSMNDAEKSFVAKGHQKRNFIISNVVLNKRFLY 631	
250twins_36050_GL0158985	D	erel '
emb OJMM01062900.1	DEFGRVKYNKKGKPVINARRRNFTINNVLSSKWFFY 65	1,
emb OGFN01002610.1	GNHKVRNFIANNVKQSKWFFY 688	_
emb OHCP01000044.1	KNHRVRNFVSNNVIKSKWFSY 676	:. 26
PIG-046_GL0077813	679	
pig_chimera	SARVRNFISNSVIKSKWFSY 682	
emb OIEN01002196.1	RESCATGENAGEDHIENFLVSNVIRSERFNY 617	17
02.UC29-0_GL0096317		
Cask_Anaerobic_dig		neet
emb OGDF01008514.1	YVSSKILIGGADKNLRNFITNNVIKNRRFLY 641	
emb 0GZC01000639.1		S
Ga0224415_16048792_chimera	GINVIKSKRFAY 596	
emb OCVV012889144.1		97
Ga0187910_10006931	FKDSNGKLLHRGKHGMRNFIINNVVNNKRFHY 609	- 60
Rflav_XPU_	SLDENGNKLKKGKAGMRNFTINNVISNKRFHY 604	
Ga0224415_10007274		
Ga0187911_10069260	YNQNGEKIKVRVDIGFRNFIANNVVESSRFHY 598	
Ga0187910_10015336	KHHYFKSGKKLPDIGLRNFIINNVIESRRFNY 702	
ODAI_chimera		
	FIG. 18QQ	B2

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592	662	651	637	576	645	635	650	575	629	634	643	643			889	632	6 5 9	641	645	629	740	169	739	
QKDNKNAKKDFRNFIIKNVIKSNRFIY	BREDKGIKKDFRNFIIKNIIKSNRFMY	LDKEGNKLKGARHDFRNFIANNVIKSSRFKY		PVGADGKFIKGKQGFRNFIASNVIESSRFHY	BANGD		ISNVLKSSRFRY	TDKSGKKLAQSKKGFRNFIINNVVESSRFKY	QYFQKGAKHDFRNFTINNVINSTRFIY	TKEKGKGIHGLRNFITNNVIESSRFVY	KIDPVTGRPLKGKNPLRNFIANNVIENSRFIY	TINPYTGKTEKGKNPFRNFTANNVIENKRFIY	* *		VVKYINPADINGLAKNRYLVKFVLSKIPEEQIDSYYKLFSNEEE	VIKYIHPSDVQKIACNKAIAGFVLNRMPDTQIKRYYDSLINKGA	LVRYAKFTAVRKLMSNFKIVRYVLKRLPEKQVASYYSAIWTQSES	LVRYTKADAVRAVMKNGKIVRYVLARIPETQVDSYYKSVVDSSD	LVRYSKPKAVRAIMQNRSIVNYVLHRLPSEQIKTYSRVFPEDF	LARYAKPKSVRELMKHEEIIRYVLTRLPEKQIDTYFSNISEGDE	LVRYTKPKTVRALMSNRAIVHYVLSRIAD1QDHHMTESQIDRYYQNLPQYNEQ	VMRFMNPKSARRIMQCESLVAFSLKDIPDTQIERYCKSVNIPFDS	VIRXVSFETARKXARQEALVRFALHRVPLLQLRRYYQSCCGPKKD	FIG. 18RR
31009_GL0034153	DLF014_GL0011914_	Cask_Ralbus	Ga0187910_10040531	tpg DJXD01000002.1	CasR_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	EYZ-362B_GL0088915	uncultured_Ru_sp	RflavFD-1	tpg DBXI01000091.1			037emb OlZA01000315.11	emb OCTW011587266.1	k87_11092736	BMZ-11B_GL0037915	BMZ-11B_GL0037771	BMZ-11B_GL0069617	Ga0099364_10024192	530373_GL0023589	emb ODAIO11611274.1	

EMG_10003641	IMRFCNPEKIANIGKNKVLISFALSSLAEKTDMIAKYYRVFCDRID	U.
Cask_F1E0_metageno	LLRYMTPEQARVLAQNEKLIAFVLSTVPDTQLERYCRICGREDI	S.
emb OIZX01000427.1	LIRYSDPAQLHQLASNKKLVRFVLSSIPDTQINRYYETCGQIKL	Pa 869
160582958_gene49834	LIRYGNPEKIRKIAINPSIISFVLKQIPDEQIRRYYPFCIGKRI	ate
gi 1198542314 gb NFIR01000008.1	LVRYADPKKVRAFAANKKVVAFVLKDIPDDQIRAYYNSCFRQNSDSSS	ent
emb OGNF01009141.1	LTRYIDISSVKAFSNNKALVKFAIKDIFQEQILRYYNSCFGASER	659
Ga0129306_1000735	LVRYCEPRAVRDYMSCRPLIRLTLRDMPDTILRRYYEQSVGAAT	969
MH0288_GL0082219	VVRYGNPKKLREVMQNEAVVKFVLNDLPEEQINRYSENILGLDA	Apı
PIG-022_GL0026351	LIRYSNPKKVKAIAENKKLVRFVLDKIFDPQIIRYCNTCGF-KV	·. 26
PIG-028_GL0185479	LIRYSNPKKVKAIAENKKLVSFVLDKIPDTQIIRYCNTCGF-KV	6, 20
250twins_35838_GL0110300	IAKYVKPADCARMASNKKMIEFALRDLPETQIKRYYYTITGNEA	122
PIG-014_GL0226364	VAKYGSPTACNOMMKNOKIVRMALESISEDQLRKYARRVYDNKEA	676
250twins_36050_GL0158985	VVKYNRPSSCRELMKNKEILRFVLRDIPDSQVRRYFKAVQGEEA	S I
emb 0JMM01002900.1	VAKYNRPSECOKFMKSKKLIALVLKDVPETQIARYYQSVTGGRT	heed
emb OGPN01002610.1	VVRYNKPAECQIIMRNKTLVKFTLDDLPDMQIQRYYSSVFGDNN	28L
emb OHCP01000044,1	VVRYNKPAECQALMRNSKLVKFALDELPDSQIEKYYISVFGEKS	of 3
PIG-046_GL0077813	IVRYNQPSECQAIMKNKTLVKFALDELPDLQIQRYFVALYGDED	87
pig_chimera	IVRYNQPSECRAIMKNKTLVKFALDELPDLQIQRYFVALYGDED	726
emb OIEN01002196.1	LSRYSNLAEVKKLAQNPSLVQFVLSRIEPSLICRYYESSQGIS	550
O2.UC29-0_GL0096317	LARYSDMATVSKLSKNKEVVKFVLKRVDPKQIERYYRSVCDVR	U \$
Cask_Anaerobic_dig	IARHMNTHYVKQLANNETLINRFVLNKMGDAKIINRYYESISGNT	S 1 .
emb OGDF01008514.1	TVRYMNPKRAKKLVQNDALVVLALSGIPETQIDRYYKSCIEKR	1, 3]
emb OGZC01000639,1	LIRYGNPQKLHTLSQNETVVRFVLSRIAKNQRVQGMNGKNQIDRYYETCGGTN	1 6,8
Ga0224415_10048792_chimera	LIRYGDPAHLHKIATNKNVVRFVLGRIADMQKKQGQKGKNQIDRYYEVCVGNK	312
	FIG. 18SS	B2

emb OCVV012889144.1	LIRYGDPAHLHEIAKNEAVVRFVLGRIADIQKKQGKDGKNQIDRYYEICVGKD	U.
Ga0187910_10006931	IIRYGDPAHLHBIAKNEVVVRFVLGRIADIQKKQGKGGKNQIDRYYBICIGNG	S.
Rflav_XPD_	LIRYGDPAHLHEIAKNEAVVKFVLGRIADIQKKQGQNGKNQIDRYYETCIGKD	P:
Ga0224415_10007274	LIRYGNPVHLHEIAKNEAVVKFVLGRIADIQKKQGQNGRNQIDRYYETCIGKD6	ate
Ga0187911_10069260	LIRYCHPRKIRNLAGNAALIEYQLRRLPELQILRYYEACTEPIK 64	ent
Ga0187910_10015336	IVRYADPRKIRKCTENNELLKFAFKDVFDSQVDRYYNICVTNK74	ιń
ODAL_chimera	LVRYANTEKIAALARNETIVRFVIKKLPAAQIVRYYIACNRIREN 72	7
31009_GL0034153	LIKYSNPTNVRKFASNTNVIKFVLGTLPDKQLERYYKSCGLSD 63	ın
DLF014_GL0011914_	LVKYSNPTDVRNLASNRNVVKFVLNAIPDIQIDRYYKSCIQHY	c . 26
Cask_Ralbus	LVKYSSADGMIKLKINEKLIGFVLDKLPETQIDRYYESCGLD	5, 2 (ε ₆₉
Ga0187910_10040531	LVRYNNPHKTRMIAQNEAIVRFVLSEIPDEQIRRYYDVCRDPK	089
tpg DJXD01000002.1	LVRYNNPHKTRILVKNPNVVKFVLEGIPEI	6 E
CasRRsiraeum	LVRYGNPKKIRETAKCKPAVRFVLNEIPDAQIERYYEACCPKN 68	රා
PIG-018_GL0023397	LVRYGNPKKIRETAKCPPAVKFVLDGIPDSQIERYYNSGFPED	hee1
PIG-025_GL0099734	LVKYCNPKKIRKIANNEKIVKFVLGRITEIQIERYYYSCNPELK 65	694 89
Ga0129317_1008067_chimera	IVRYSNPQKIRKLANNSVVVGFVLGKLPDAQIBSYFNSCLPN61	r-
EYZ-362B_GL0088915	VIRFCNPKTAGQIIRNENILRFVLQRMPQE61ERYYHACTKN67	87
uncultured_Ru_sp	LIKYANAQKIREVAKNEKVVMFVLGGIPDJQIERYYKSCVEFP	21.5
RflavFD-1	LIKFCNPENVRKIVNNIKVTEFVLKRIPDAQIERYYKSCTDS	685
tpg DBXI01000091.1	VIKFCNPKNVRKLVNNIKVTEFVLKRMPETGIDRYFESCIEG	US
	*	S 11,
037emb OIZA01600315.1		,316,
emb ocrw011587266.11	TDIQAQAKALLDCITGISFDAIKDDKHLHKSKE 66	w)
	FIG. 18TT	B2

k87_11092736	NSNEMVKLIEMIDRLITEIAGFSFAVLKDKKDSIVSASR 698	
BMZ-11B_GL0037915	AGNGKPIKEKIKALAEKLGEFSFDLIAENRDGIIDNAKL 680	
BMZ-11B_GL0037771	KRGFSP 684	ਵਜ
BMZ-11B_GL0069617	SEISRKIDTLIKALTGFSFNSLFINRDKIVENTKL 664	ate
Ga0099364_10024192	QHKNVSLETKIDALADYLCKYTFEKNVLKQKNGIV-LNT 778	
530373_GL0023589	ANPNYAHMKEQLTYKLLQVQFCNFSKVŞNSSHQANPNYAHMKEQLTYKLLQVQFCNFSKVŞNSSHQ	St.
emb ODAI011611274.1		
EMG_10003641	848	Apı
CasR_P1E0_metageno	TGREAQIRYLTAQIMGVRYESFTDVEQRGRG	
emb OIZX01006427.1	AGRAAKVETLIDMIAAIREDQFRDVNQKERG724	
160582958_gene49834	DDVTLMRDELGKMLQSVNFEQFSRVNKQNAK774	
gi 1198542314 gb NFIR01000008.1	NNSNASWDADSNKRDISVSDMRKALTEKITGLNFGDFEEESKKGIR765	\$2
emb OGNF01009141,1	YYNDGMSDKLVEAIGKINLMQFNGVIQQADRNYNLPE 695	۲O
Ga01293061000735	VDRERILDTLADKLLSLRFTDFENVNQRANAE728	
MH0288_GL0082219		. 69
PIG-022_GL0026351		
PIG-028_GL0185479		
250twins_35838_GL0110300	GDAESIKGVIIEQLHAFSIKNTLLSIKNMGEGEY 756	99
PIG-014_GL0226364	FQNQTADEIRENLAKKLKNFNIGAELKTVGNLSEKD 712	2
250twins_36050_GL0158985	YASAEAMRTRLVDALSQFSVTACLDEVGGMTDKEF 770	
emb oJMM01002900.1	QANSEAMRMTLIKLLHEFSIKNVLSDVGTMTASEN 736	
emb OGPN01002610.1	MPAVDEMRKRLIDKINQFSVRGFLDELDEIVLMSDEE 769	
emb OHCP01000044,1	SSSNEEMRRELLKKLCDFSVRGFLDEIVLLSEDEM 755	
PIG-046_GL0077813		7
	FIG. 18UU	B2

ACCOUNTY ACCOUNTY	pig_chimera	LPSYGEMRKILLKKLHDFSIKGFLDEIVLLSDLDM	192	U.
	b olen01002196.1	Ā		S.
	:.UC29-0_GL0096317	NISAEDEIEKLVDIIYNFNIDELEQVNNAEIGIG		Pa
	sR_Anaerobic_dig			ate
	ab oGDF01008514.1	SFNPDLNEKIAALSEMITTLKIDDFEDVKONPEKNA		ent
	ab OGZC01000639.1	***************************************	ro.	•
	10224415_10048792_chimera		<#	
	ub cCVV012889144.1	and that that had had that had had had place of	7	Apı
0.01274	a0187910_10006931	of the day dos the test test has not been the t		r. 26
	flav_xPD_			5, 2(
	a0224415_10007274	per per per ser ser ser ser ser ser ser ser ser s)22
	20187911_10069260	RIARTMDEKIGTLIBLIVKMDFSQFEDVQQMDRVRV	ଚଞ୍ଚ	
INSNVLTSDDMAESLVKLVTGMDFGKIKGVRNDDR	a0187910_10015336	alan sala alan 'ana' alan 'ana sala sala sala alan sala alan sa		SI
	DAIchimera	the total teat that they does does does does not		heet
	1009_GL0034153			70
	F014_GD0011914_	in and the said the two lates for the lates for		of :
Chimera	ask_Ralbus	***		87
1	0187910_10040531		721	
TALCSANKRREKLADMIAEIKFENFSDAGNYQKANVTS	og DJXD01000002.1	IPPISUKSAQIDVLARIISSVDYKIFEDVPQSAKINK	560	
TAICSADKKRRELAEMISNISFDWFKDAGNVQRVNAEDRAYPGREAAIDDLTKIIINWKFUDFKNVKQSANVEDchimera	ask_E,_siraeum	TALCSANKRREKLADMIAEIKFENFSDAGNYQKANV		US
BDAYPGREAAIDDLIKIIINMKFUDFKNVKQSANVED	:G-018_GL0023397	TAICSADKKRRELAEMISNISFDWFKDAGNVQRVNA	(O	S 1.
AVYSTPDKARESLRDMLHNISFNDFADVKQDDRRA	:G-025_GL0099734	RAAYPGREAAIDDLTKIIINWKFUDFKNVKQSANV		1,3
TFSLCTADEKIQCLSNIIMEVRYKKFCNVAQKSRDT	.0129317_1008067_chimera	RVYSTPDKARESLRDMLHNISFNDFADVRQDDRR		16,8
FIG. 18VV	Z-362B_GL0088915			812
		FIG. 18VV		B2

U.S. P	Patent	Apr. 26	5, 2022	Sheet 71	of 87	US 11,316,812 B2
711.718	27.	724 758 740	741.722.838	9 8 2 8 9 0 3	8 0 0 8 8 0 0 8	7 7 8 4 8 4 7 7 7 3 4 2 4 5 5 7 5 9 9 7 7 9 9 9 9 9 9 9 9 9 9 9 9
	DGFFYK	KSPQRSADRERKKAMLTLYYTIVYIFVKQMLHINSLYTIGFFYLERDQRFIYSRAKKEN- ESRAVNLEVERLKKLTTLYMSIAYIAVKSLVKVNARYFIAYSALERDLYFFNEKYGEEFR DEEKKNVEIERLKAVTGLYLTVAFIAIKNLVKANARYYIAYSAFERDSELLKEKLGEEEF	NRRETAEEIERLKAITGLYLSVAYIAIKNIVKANARYYIAFAVFERDKELVKAKDAR DISNKNVKIERLKALTGLYLTVAFVAVKNLVKANARYFIAFSAFERDYGFMKSHDPDV KSAIKNVEIEHLKALTGLYLTVAYIAVKNLVKANARYYIAFSIFERDYALFEKKLGKDIL	AVERERLKALVGLYLSVLYRIVKSIVRINTSYTIAFAAYERDCVILNRKMGKNLY SEQEKKQKYQAIVGLYLIVVYWIVKNLVNVNSRYVMAFHILERDTVLLEGKRLFVGG KMKEKDRSQKLIGLYITLLXEIVKNLVNINSRYNIAFQRCDNDSIMILQGQYDER-	DNPKKERYKALIGLYLTVLYLAVKNMVNCNARYVIAFYCRDRDTALYQKEVCWYDLANTQKERYKAMLGLYQTVLYLAVKNLVNINARYVMAFRCVERDMFLYDGELTDPKGQNPMGEKARLQACVRLYLTVPYLFIKNMVNINARYVLAFHCLERDHALCFNSRKLNKEESDKNIIRLYLTVLYLYLYGKNLIYVNSRYFLAFHCAERDEMLYNGETIDNNK	EKKRANAQKEKYKSIIRLYLTVCYLFFKNLVYVNSRYYSAFYNLEKDRSLFEINGELKFTRNREKQKMMGIISLYLNVAYQIVKNLVYVNARYTMAYHCAERDTELLLNAAGEGNLRNARKARYQGAMSLYLNVIYQIIKNMVYINSRYTLAFAKYEHDASLLLDNWVYDFYDTFDAIDKAQKQGLISLYLTVLYLFIKNMVYVNSRYVLAFHCLERDSQLLKITSGYN FIG. 18WW
uncultured_Ru_sp RflavFD~1 tpd:DBYI01000091.1:	037emb OTZA01000315.1	emblocrw011587266.1 k87_11092736 BMZ-118_GL0037915	BMZ-11B_GL0037771 BMZ-11B_GL0069617 Ga0099364_10024192	530373_GL0023589 emb ODAI011611274.1 EMG_10003641	Cask_PlE0_metageno emb OIXX01000427.1 160582958_gene49834 gi 1198542314 gb NFIR01000008.1	emb cGNF01009141.1 Ga0129306_1000735 MH0288_GL0082219 PIG-022_GL0026351 PIG-028_GL0185479

FIG. 18XX

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777	784	774	790	705.	763	767	768	772		8 0/3	756	808	787	782	772	& & & &	804	881	919	775	811	
ANDRMILKKORYQAIVSLYLTVMYLVIKNLVYVNSRYVMAFHALERDAYLYGIINI	-RISEAEIKRKNQAIIRLYLTVMYIMLKNLVNVNĄRYVIAFHCVERDTKLYAESGLEVG-	-ksrnaeikrkngalirlyltvmylmlknlvnvnsryvigfhcverdaklyklsgltie-	-SSKEAQEKMKYQTIISLYLIVCYHLVKNLVNINARYAMAFHSLERDARLYGIYGSEKDY	"TPEEKVERERYKAIIGLYLIVMYHLVKNLVYVNSRYVMAFHCLERDAMHYDVS"""""	TKQMEKERFRAIIRLYLTAIYLVVKMLVNVNARYVLAFHCLERDADLFGLQIKQTLP	renvakerakaviglylivmylivknlvnvnaryviaihclerdfglykelipel-a	ENMERERFKAVIGLYLIVVYRVVKNLVDVNSRYIMAFHSLERDSQLYNVS	SQEAVEKERFRAIIGLYLTVIYLLVKNLVNVNSRYVMAFACLERDAKLYGVQ	* * * * * * * * * * * * * * * * * * * *	ETLKKKDKKQLYNDDDYLLLPEIFSGSKYRE	KNPSKNSYLNDFRSVTAYFIPSEIMKRIEKNE	LHFIPYELNGRTCQFEYLAILKYYLARDEETLKRKCEICEEIKVG	YKLCIEFKMKKGETGYMNAFALLKYYIDRDEDIWRKEEATHRGRDEN		AQSCLAYERIMDGKSKTKYNEIYALTKYLKKEBENDYHPEPGQPFDKEA	EKYVKPFKYIDKGEEKEGKNNFFALTEYLLDKDNSLRYQWNNDLSDEENK	GL	MKMK	AVQESKLTKKFISNQK	EEDKKSGKQRQVEDYTALTRYFVSQGY	ESVSAFLAVNGKKGVQPQXTLLTQLFIRRDY	FIG. 18YY
Ga0187910_10040531 tpq DJXD01000002.1	CasR_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	EYZ-362E_GL0088915	uncultured_Ru_sp	R.flav.FD-1	tpg:DBXI01000091.1		037emb OIZA01000315.1	emb OCTW011587266.1	k87_11092736	BMZ-11B_GL0037915	BMZ-11B_GL0037771	BMZ-11B_GL0069617	Ga0099364_10024192	530373_GL0023589	emb ODAI011611274.1	EMG10003641	CasR_P1E0_metageno	emb 012X01000427.1	

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859	838	775	830	858	753	755	842	804	864	838	88 86 44	850	852	856	787	781	811	795	824	477	779	790	785	
DDSYNEMANKFOMVRKAKKEQYEKEYKCK	EK	GK	LRRDRSWPARLHLPRRALARRRDRVEVMERDVARGPEAYNRDEWLG	KIVFSDLTRDSLQKGK	NSSYNNLTNYFIDNKY		EKKFDAIINEILLEDDKIRKECDKERAGAKTL	EKKEKGKRFDWLALTQKFLSEDKNYYFEHDLRRKEINKI	KKKSSDYTGEDMLALTRKFVGEDAGLYREWKEKNAEAKDK	LANNEDDKWKKGNYVFALTKHFWDNDEPYFDKYNNALQQIRSI	BMKGGPWDGGAQALAVTRRFLNHDREVFDRYCAREAEIARL	EKAERAWEKGATAFALIRKFLNHDKPTFEQYYTREREISAM	NKQERAWSSGATALALTRKFLNQDKLIFEKHYAREGEISKL	NKQERAWSSGATALALTRRFLNQDKLIFEKHYAREGEISKL	KNKKRSSGGFSFIEMTQFFIDKKLFKVAT	FKDDFDAGKKSKKQIAYRIKYIALTEEYLDKGYIKCAT		YMTRDVFIAKGWI		KLDGFVALTKFCLGDEBRFEDLKAK	AL	NNLESKGFTSVTKLCVGIADDDPVKYKNVBIELKER	KKLEEKGFSSVTKLCAGIDETAPDKRKDVEKEMAER	FIG. 18ZZ
160582958_gene49834	gi 1198542314 gb NFIR01000008.1	emb OGNE01009141.1	Ga0129306_1000735	MH0288_GL0082219	FIG-022_GL0026351	PIG-028_G50185479	250twins_35838_GL0110300	FIG-014_GL0226364	250twins_36050_GL0158985	emb 0JMM01062900.1	emb OGPN01002610.1	emb OHCP01000044,1	PIG-046_GL0077813	pig_chimera	emb OIEN01002196,1	02.UC29-0_GL0096317	Cask_Anaerobic_dig	emb OGDF01008514.1	emb OGZC01000639,1	Ga0224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD_	

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764	764	865	855	761	821	8,0,3	797	740	809	804	814	725	787	791	789	801	803	756	844	796	784	
DVEKEMTER	AEAEYLNLTRAFIEPLDGAXP	GRGYCDLSTVLLFGVDDLQNRNRGSYKYLRDW	NTGDYPADN-ADLRRDIKYCVLIKAHIKQEEVKSRRND	PEQPEKLYKIKPDKPLQPAQYFLFTKYLLQNDFFGE			KGDYRKLTDNLLADENYKKF	TIDENNILL	IKKIIKKIIKKIIKKI	GNIDKNKTLLTATILGITSADRAPSGEIVA	VEKLKKEQAVLVEKLLSDGFETAG	LDNYRDLIRHLISEGDSSCN	DBANDENKP	SSNLKNDYRILSQTLCELCDKSPN		NIGGDYLALTAKLCAEGDDYGKKLSEAKQ			CEKHKKNANPPYEYDQEWIDKKKALNSERKACERRLHFSTH	GKLDYDPDP	H_{Λ}	FIG. 18AAA
Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	ODAIchimera	31009_GL0034153	DLF014_GL0011914_	CasR_Ralbus	Ga0187910_10040531	tpg DJXD01000002.1	Cask_Esiraeum	PIG-018_GL0023397	PIG-025_GL0099734	Ga0129317_1008067_chimera	EYZ-362B_GL0088915	uncultured_Ru_sp	RflavFD-1	tpg DBXI01000091.1	037emb OIZA01000315.1	emb OCTW011587266.1	k87_11092736	BMZ-11B_GL0037915	BMZ-11B_GL0037771	

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Cask_Angerobic_dig		U.
emb OGDF01008514.1		S.
emb 0GZC01000639.1		P:
Ga0224415_10048792_chimera	AQASIKEFR	ate
emb OCVV012889144.1	AKKSLEBLKTTNSKLYENYIKYSDERKAEEAKRQ	ent
Ga0187910_10006931	ALASFKQ	824
Rflav_XPD_	AKESIRESI	81.9
Ga0224415_10007274	AKASFKAKQ	Apı 867
Ga0187911_10069260		:. 26
Ga0187910_10015336	RRSNKDVIETFGDFKGKVSKVVEKKNQGLTNEIYDSLCNVAGTIKTEVQNEIKSILKSNG	5 , 2 0
ODAI_chimera		922
31009_GL0034153		761
DLF014GL0011914		S 3
Cask_Ralbus		eo3
Ga0187910_10040531		. 77
tpg:DJXD01000002.1		of :
Cask_Esiraeum		87 118
PIG-018_GL0023397	DC	808
PIG-025_GL0099734		814
Ga0129317_1008067_chimera		U \$22
EYZ-3628_GL0088915		S 1
uncultured_Ru_sp		1,3
RflavFD-1		1 6,8
tpg DBY101000091.1	NI	812 E08
	PIG. 18CCC	B2

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841	807	889	832	821	814	936	83.8	893	947	801	837	898	864	801	872	884	7.79	781	893	857	916	
ETKNLKLPKEKDRDIMKKYLPNDKDREEYNKFFKQYRN	-GKTSRLRKEDVLLYARYISPDHALKNYKMILNSYRN	QATGWLATQARN	DPTGMLAAQMRN	QATGLLLREARN	HFTGYLGVWARN	SESGYLLTAARN	CINKIYQVYRN	PASTDAAASLIWSYRN	RCNDFIRQYRN	ISNSLLTAYRN	ISDRILLEYRN	MHSYTCRNYRN	SNKWSLKEFRN	SDETAIRTERN	AEPGDAŞLSLLEVYRN	SDNTLIRQYRN	ADEWSIVKFRN	SDEWSIVKFRN	NHIDFAVTSYRN	PDPLKLFIKYRN	NGLAAVIKEYRN	
KNLKLPKEKDRDIM	NKGFLEDFEALWNSCGKTSRLRKEDVLLY	WAQYATKRDENMAKHPQKWYDILASHYDELLAL	RRKHYPKRWLNILKDNLAEMERI	RKTKRHFSKQWREWLNEKIADAKSS	SIRRHFTKKWRDIFRKEIDEAVKI	IRSQRHFSQYWLDIFARQIENAKKT	NDFINETKLSKRVCALVNKNRTL	RIGENKRANRHGLNCVLENRNALGS-D	LNSYSCRYLTHNISQLD		LKRSACEQIQHNMEN	-KQETGTAHTKKIEKLNQQIAYIDKDIKN		INPRASAYLTVNLAN	LVRTLREKRVCDNLHNNYAYLCGAD	LNRRAAKYLQTNLEH	LNPHASRYLKVNISN	LNSHARNYLRVNISN	RFAQIKRRESGCYFKSYHVYDYLSKNSNEFKQ	QHRENDKYYDSCMHFSRHSFEYISKNFAEVQSH	KVLRQNDKMIRKMHFTFHSLNYVQKNLESVQS	FIG. 18DDD
037emb OIZA01000315.1	emb ocTW011587266.1	кв711092736	BMZ-11B_GL0037915	BMZ-11B_GL0037771	BMZ-11B_GL0069617	Ga0099364_10024192	530373_GL0023589	emb ODAI011611274.1	EMG_10003641	Cask_P1E0_metageno	emb OIZX01000427.1	160582958gene49834	g1 1198542314 gb NFIR01000008.1	emb OGNF01009141.1	Ga0129306_1000735	MH0288_GL0082219	PIG-022_GL0026351	PIG-028_GL0185479	250twins_35838_GL0110300	PIG-014_GL0226364	250twins_36050_GL0158985	

emb 0JMM01002900.1	LAYRANDKVVKHTHFNLHSYKYVKHNYEEISK887	U.
emb OGPN01002610.1	PLRKANDKLLKQIHYTNHSYTYIVNNLNSFTDIDYCAKDVGLPAFNDKNDNASILGEMRN 930	
emb OHCP01000044.1	ELRKENDQLLKKTHYSKHAYCYIVDNVNNLTGAVANDNGRGLPCLSEKNDNANLFLEMRN 916	
PIG-046_GL0077813	AMRKVNDQLLKQTHESKHSYCYIVDNVNRLTGGECRIDKRVLPVLNEKNDNAGILSDFRK 918	ate
pigchimera	AMRKVNDQLLKRTHFSKHSYCYIVDNVNRLTGGECRTDKRVLPVLNEKNDNAGILLDFRK 922	
emb OlEM01002196.1	IKKNVLKYNGNPESLNHIPGEYICKNMEGYHENTVRNFRN 829	
02.UC29-0_GL0096317	IKENIKKYGGDFKSIDHIPGETIKKNLEGYCAGSTWTFRN 823	
Cask_Anaerobic_dig		A pı ∞
emb OGDF01008514.1	NPKKFTVKSIKEQYAFLTPYIFTTYRN 822	
emb 0GZC01000639.1	TVYEKABAAKNRALKNVKWNCKTRENLENA	
Ga0224415_10048792_chimera	LNRERVKNARNAYLKNIKNYIMIRLQLRDQTDSSGYLCGEFRD 854	
emb OCVV012889144.1	INRERAKTAMNAHLRNTKWNDIMYGQLKDLADS	ω,
Ga0187910_10006931	INREKAKTALNAALRNIKWNVIIRENIRNT	₩
Rflav_XPD_	INREKAKTALNAYLRNTKWNVIIREDLLRIBOKTCTLFRN 855	Ø.
Ga0224415_10007274	INREKAKTALNAHLRNTKWNVIIREDLLRRDNKACKIFRN 838	. 79 ∞
Ga0187911_10069260	SIRHFRN 791	
Ga0187910_10015336	LDESASSYLSHKLVNKVHSYKYLKQNLDCADNTMINQFRN 965	ιń
ODAI_chimera	TDSKSQQKAKSRRRRWCLLLKEDIDKLNGC	4
31009_GL0034153	YISKKNQYIQKNMKIYSKINKNENVFVRYRN 792	Ο.
DLF014_GL0011914_	KKNKKYGEYTSKKISTYIETNMKNFIGCEQFRTPERYRDTQIDMFVKYRN 871	
Cask_Ralbus	GKLGCNKKVGRYLKNNISCCTDG	m
Ga0187910_10040531	GHFKNKKWRGIAEQNLRNSDVPVIKSFRN 826	
tpg DJXD01000002.1	GHLKNOKGHRKWYVLVKNNLQNSDITAVSSFRN 773	
CasR_Esiraeum	DKSF-ABNAANRYLRNARWYKLILDNLKKSERAVVNEFRN 850	
	FIG. 18EEE	B2

U.S	. Pa	ate	nt			Apr. 20	5, 20)22		SI	heet	80	of 3	87			US	S 1	1,3	16,8	312
80 83 44 44 53 44	761	820	821	826	850		988	851	. Q.	881	872	86 86 86	984	188	936	086	844	881	944		844
DLGR-AETAANRHLRKKRWYVQIFNNLKQADRNVINEFRN NLHLRSKKWNRLMRVNLENYSAAASONFRN	IFGKED	KGYLARNKRLCQCVREDIDNAKHLNGDITDYRN	LETKKNERLRKCVEVDINNADSS	RYLAGNKRLRDCVKQDIDNAKKWFVSDKYNSITKYRN	NQDKVQMPKNYFLARNKRWREAIEQDIDNAKKWFIGEKFNNVKNYRN	*	NIVALKIIA-KLSELTKNIDKDINSYFDIYHYCTQRVMFDXCKKNN	KIAHINVIM-SAGKYTGGIKRMDSYFSVFQHLVQCDILSNPNNKG	DAEHLMPVN-EFDVYIEDLRRYPEGTPKNKDYHIGSYFEIYHYIRQRAYLEEVLAKR	DAMHLNAMF-KIAKFVGDFRKGSSHEMBSYFELFHFLEQKTILDSKEDFV	DVEHLNVLR-AIPDYIQDFRHGEKGETAMNSYFELYHYLMQRLMLKNTELDL	QSAHLNVLTQALPQFIGQFARNSKTGKTARMTSYFQLYHYLLQRLLCEDLNFDV	CALHLINVLI-ALPEFVGEFRKTGDKMTSYFELYHFLLQKIMLAEAGLNL	IIMHLNAVS-AFPQYLEGLKVVHSYFDLYHYTVMMLLHRQNEKE-	AAAHLTAVA-AAQEYVSELR	KVAHLEVVS-NIDEYLSGIKHIESYYALYHYLMQKCLLKNYRIE-	AVDHLNAIP-PLGSLCRDIGRVDSYFALYHYAVQQYLNGRYYRK-	AVAHLNVIA-HLADYSADMREITSYYGLYRYLMQRHLFKRHAWQI	LVAHLNVVS-KLQNYVSELPNDYQITSYFSFYHYCMQLGLMEKVSSK-	SVQALNVIR-DARKYIKYINDNKDVQSYFALYHYLVQRYISERAANRT	TAEHLEALR-NADKYLNDLKQFDSYFELYHYITQRNIKEKCEML-
PIG-018_GL0023397	Ga0129317_1008067_chimera	EYZ-3628_GL0088915	uncaltured_Ru_sp	RflavFD-1	tpg:DBXI01000091.1;		037emb[OIZA61000315.1]	emb OCIW011587266.1	k87_11092736	BMZ-11B_GL0037915	BMZ-11B_GL0037771	BMZ-118_GL0069617	Ga0099364_10024192	530373_GL0023589	emb ODAIO11611274.1	EMG_10003641	Cask_F1E0_metageno	emb OIZX01000427.1	160582958_gene49834	gi 1198542314 gb NFIR01000008.1	emb OGNF01009141.1

B2

FIG. 18FFF

U.	S.	Pa	ate	ent			Apı	:. 26	5, 20)22		Sl	heet	81	of	87			US	S 1	1,3	16,8	312	B2
915	927	823	02 02 03	937	904	958	930	973	959	096	965	873	867	882	866	913	898	006	906	901	880	835	1008	
EAKSWFYVYHFLMQRVLEEFFRNT-	BITSYFGIYHXIMQRDLITRCQKV-		EFKSYFEIYAYTVQKSIKNYYDAES	DVKSYYGVYCXIMQRMLCDELIIKN	-GRAFEKIDIFYDLYCYCLQKMLLSEVD	ADSYYSLYCYCLQMYLSKNFSVG-	EVTSYYSLYCYTLQRLLLDDNNND-	DISSYYAFYCYVLQRRLVGKDPNC-	HITSYYAFFCTVLQRMIIGNNSNE-	GITSYYAFFCYVLQRMPVGNKN-E-	GITSYYAFFCYVLQRMLVGNNLNE-	QIDSYYALYHYCMQMNILQGIEQSG	NVNSYFSLYHYCMQKHILYCARFN	SVHSYFALYHYIMQRALYDSLQAKA	RFKSWFHLYHTVIQHSLIQQYEYDR	AMKRYFDCYHYLLQRELGYILEKSN		EVKSYFRLYHYIMQRRIIDVIENNP	EVNSYFQLYHYINQRIIIDSSGN	EVNSYFQLYHXIMQRIIMNERYE	EVNSYFQFYHYIMQRRIMAERYD	YIGSYYALYHYILQRHLLDKIEEDS	GITSYFQIYHYLMQKALNKBFKKCR	
KAAHLSVLN-KGGRLSGDLK	SVAHLNAVR-NMMLYIGDLR	GTAHLNSVR-NIDICIEGLK	GVAHLNSVR-NIDISIKGLR	NVEHLNVVH-CMTKYFSEVK	NVMHLNIIR-DLNDYLPDFKRC	AVAHLNIIN-RLDEYIGSAR	NVQHLNVMN-SITKYLGDIS	DIAHLNIVH-DMVKYIBELK	KIVHLNVVA-DMVKYINEIK	TIAHLNVVH-KMVDYVAEIK	TIAHLNVVH-KMVDYVDEIK	MVAHLTAVA-RVPLYISEVT	AVAHLALMA-RVSEYASDIK	QVTHLNAIR-VAYKYINEIK	MIAHLAAVT-NAYKYIPQMD	IVAHLWIIR-DADRFITGMG	KVAHLEVAR-HAHEYIGNIK	KAAHLEVAR-YAHMYINDIS	KADHLEVAR-YAYKYINDIS	KAVHLEVAR-YVHAYINDIA	KVAHLEAIR-YAHLYINDIA	ATVHLNVIM-EAHRYTKDIK	NVAHINTIR-NMDGIENVT	FIG. 18GGG
Ga0129306_1000735	MH0288_GL0082219	PIG-022_GL0026351	PIG-028_GL0185479	250twins_35838_6L0110300	PjG014_GL0226364	250twins_36050_GL0158985	emb OJMM01002900.1	emb OGPN01002610.1	emb OHCP01000044.1:	PIG-046_GL0077813	pig_chimera	emb OIEN01002196.1	02.UC29-0_GL0096317	Cask_Anaerobic_dig	emb OGDF01008514.1	emb OGZC01000639.1	GaQ224415_10048792_chimera	emb OCVV012889144.1	Ga0187910_10006931	Rflav_XPD_	Ga0224415_10007274	Ga0187911_10069260	Ga0187910_10015336	

U.S. Pat	ent Apı	2. 26, 2022	Sheet 82 of 87	US 11,316,812 B2
937 836 915	2 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	8 8 8 8 8 9 8 8 8 9 8 8 8 8 8 8 8 8 8 8	8 8 4 22 84 22 24	8 6 6 8 8 1 6 8 8 0 6 6 0 2 4 8 6 0 0 2 2 9
AVAHLNIPR-NAIDYIGGIKRADSYFAIYHYMMQQYLSKRHGLP- NIAHLNSIK-NATKYVNDIGIFTSYFQLYHYIMQCDLKDWVKKEV SIAHLNAVR-KASKYINDIRGFDIYFELYHYIMQRDLRDILEKIA	IGDIQ	AVAHLNPIR-NADVLIEGIENITSYFDIYHYIVQRSVLNRTNKT	NVAHLTAVR-NCAEFIGDITKIDSYFALYHYLIQRQLAKGLDHER NVAHLTAIR-NCAEFIGEITKIDSYFALYHYLIQRQLAGRLDPNH *: *: NVVLA	KEYRDSGSFTDEQLDK-LQKILDDIRARGSYDKNLIKLE QGMRESGSFTDEQLDK-LQKILDDIRARGSYDKNLIKLE S-HWS
ODAL_chimera 31009_GL0034153 DLF014_GL0011914	Ga0187910_10040531 tpg DJXD01000002.1 CasR_Esiraeum PIG-018_GL0023397	PIG-025_GL0099734 Ga0129317_1008067_chimera EYZ-362b_GL0088915 uncultured_Ru_sp	RflavFD-1 tpg DBXI01000091.1 037emb OIZA01000315.1	emblocTw011587266.1 k87_11092736 BMZ-11B_GL0037915 BMZ-11B_GL0069617 Ga0099364_10024192 530373_GL0023589

emb ODAI011611274.1	BLVSKNGPASGK-IAAWANAVDRCHSFCKDWLWLL 976	
EMG_10003641		.S.
Cask_P1E0_metageno	TPR-EQELFAAMAQHRIWCSDLVKAL 869	
emb OIZX01006427.1:	RQPERPTEEEQKLIEQEQKQLAW-EKALFDKTLQYHSYNKDLVKAL 926	
160582958_gene49834	NIPLVESLKNEANDAQSYSAKK-TLEYFDLIEKNRTYCKDFLKAL 988	
gi 1198542314 gb NFIR01000008,1		
emb OGNF01009141.1	REQ-TVKYNNDLLKYHGYSKDFVKAL 869	
Ga0129306_1000735		
MH0288_GL0082219	RMKNPV-VVGYFDKVRTYRTCCKDLVKAL 955	 2 6
FIG-022_GL0026351	SEDRNIEDGKISLCNGVYIHERQKNGKFKITVKINPK-TIBYMESVDKYGSYTKDFVKAL 882	
PIG-028_GL0185479	PBYNESVDKYGSYIKDFVKAL 854	
250twins_35838_GL0110300	QDKPD	భ
PIG-014_GL0226364		Ĺ
250twins_36050_Gt0158985		heet
emb OJMM01002900.1		m
emb OGPW01002610.1	BURF-KAKYAKELINDYGTYNKNLMWML 999	
emb OHCP01000044.1		10
PIG-046_GL0077813	KARYSETAKSYGTYSKDFWWLL 986	Ŷ
pigchimera	KNAI-KEKYSATVKSFGTYSKDFNWLI 991	Ţ
emb OIEW01002196.1	KILRIALENARVHRIYSKDAVKYL 902	
02.UC29-0_GL0096317	KQKQ	
Cask_Anaerobic_dig	KDSRDS-LERKIYSKDLLHVL 910	
emb OGDF01008514.1	DYGRKGAPVVSER-VLQLLEQCREHSNYSRDLLHIL 901	
emb 06%C01000639.1	Q	
	FIG. 18III	B2

Ga0224415_10048792_chimera	EAM	8 0 0 0 0 0	U.S
		929 9	S.]
	NNANGM-IKKYYESVISDKKYNDRLLKLL	933	Pa
	K	928	ıte
	KKSSGK-VREYFDAVNNEKKYNDRLLKLL	200	nt
	YAEKTYS-EKLWESQISQYGTYSKDFVKAL	864	
	ENAVRKWIFY I TENAEFRYVYWNKKEQQEVEVSFNFRIFGYMENIKNASNIYCKDFVKAL	1,068	
	EGLGCIHGVYCKDLVKAL	955	Apı
	EEKKIEKKDEVKAI	867	:. 26
	EGAAKETEEGQKQDEL-LKTYFDNLDKYGTYVKDFVKAL	953	5, 20
	NNT-ESNYKNSIIKHHTYCKDMVKAL	905)22
	KGLSDQ-TKRYYDALEQYNTYCKDFVKAY	868	
	S-HSCKDFVKAY	844	SI
	VERD-IGDFISKLEEHKTYCKDFVKAY	920	1eet
	FNSF-SAEYISNLEKYHTFSEYFIKAY	913	84
	PK-MKEYEELINKYHIYNKDFVKAL	911	of S
Ga0129317_1008067_chimera	DT-AHKYFEQLTKYKTYVMDMVKAL	828	87
	YRCKYRCK	893	
	QEEKIKYEDDLLKNHGYTKDFVKAL	068	
	SGFBR-NYPQYAPLFKWHIYVKDVVKAL	697	US
	PGF	921	5 11,
			316,
037emb OIZA01000315.1	NLPFAYNVPRFKNLSYKKFFDKQ	945	812
	FIG. 18JJ		B2

U.	S.	Pa	ate	ent			Apr	. 26	5, 20	22		Sì	ıeet	85	of S	87			US	S 1	1,3	16,8	812	B2
911	1034	940	928	923	1054	951	1009	10.57	906	961	1022	1001	904	973	991	924	894	900	90.54 4.20	1020	979	1030	1023	
NIPFGYNVPRYKNLTFEKIYLKSSINE	YLPFAYNLPRYKWLTTEALFDDDSVSGKKRVAEWREREKTREAEREGRRQR	YITLGYSLPRYKIMTVSGLFDPDSEDAKERAKGKRA	FVSLAYNLFRYRNLTKEHHFDDTVLQKIREKESLD	FVPLGYNLPRYKNLTIECLFDEDSVSAKEGKKA	YVSLGYNLPRYKNLTCEPLFDEESATGKERQTRLDEKSKERRQRKGGQK	NSPLAYNAARY INLSCRTKFEEGFGK	NVPFAYNPARYKNLSIANLFDKNEAAPVTEDASEQKEDE	NVPFGYNLPRYKNLSIDELFDRNKLKTGGTIEMKGE	NTPFGYNLARYKNLSIDGLFDREGDRVVREDGERPAE	NAPFGYNLARYKNLSIEPLFSKEAAPAAEIKATHA	NAPFSYNLPRFKNLSTEALFDKNIVYEQADLKKE	NVPFAYNLFRYKNLSIDELFDRNNYLPNKAKKWIPEKKENGEYVMEDCGNKDAGQVENA	CVPFGYNLPRFKNLSIDALFDKNDKREKLKKGFED	NLTFAYNLPRYKNLSIDGRFDKNHPDPSDE	NITFGYNLFRYKNLSIEGLFDKNRPGDKKESALIAE	NIPFGYCLARYKNLSIEFLFDRNTVYNKEDVTEEFSVNSKEQ	MTPFGYCLARYKNLSIEPLFDRNTVYNKKDVKEEFAVNSK	NFPFAYNLARYKNLSNEDLFNAKNNDQKSK	NIIFAYNVPRYKNLSNKELFYRLDSEQ	NIPFAYNLARYKNLSNEKLFYDERAABKADKAENERGE	NIPFAYNLPRYKNLSNERIFYDELQK	NLPFAXNLFRYKNLSSEFLFYDMEYNKKDDE	NLFFAYNLPRYKMLSNEQLFYDBEERRMBKIVGRKNDSR	FIG. 18KKK
emb OCIW011587266.1	k87_11092736	BMZ-11B_GL0037915	BMZ-118_GL0037771	BMZ-11B_GL0069617	Ga0099364_10024192	530373_GL0023589	emb ODAI011611274.1	EMG_10003641	Cask_Pl#0_metageno	emb OIZX01000427.1	160582958_gene49834	gi 1198542314 gb NFIR01000008.1	emb OGNF01009141.1	Ga0129306_1000735	MH0288_GL0082219	FIG-022_GL0026351	PIG-028_GL0185479	250twins_35838_GL0110300	FIG-014_GL0226364	250twins_36050_GL0158985	emb 0JMM01002900.1	emb OGPN01002610.1	emb OACP01000044.1	

PIG-046_GL0077813	nlpfaynlfryknisneqlfydeebrnetegqidrl	U.
pig_chimera	NLPFAYNLPRYKNLSNEQLFYDERERNETEEQIDRL	S.
emb olEN01002196.1	CLPFAYNISRYKALTIKDLFDWTEYSCKKDE	Pe
02.UC29-0_GL0096317	CLPFAYNLFRYKALTVEGLEDWNEYGODDD	ate
CasR_Anaerobic_dig	HSPFGYNTARYKNLSIEALFDKNESRPEVNPLSTND	ent
emb 0GDF01008514,1	NLPFGYNLPRYLNLSSEKYFDANA!	926
emb 0GZC01000639.1	CLPFAYCIFRYKNLSIAELEDRHEPEAEPKEEASSVNNSQFITT	984
Ga0224415_10048792_chimera	CVPFGYCIPRYKNLSMEELFDMNEEKKFKKKRAPENT	Apı 596
emb OCVV012889144.1	CVPFGYCIPRFKNLSIEQMFDMNETDNSDKKKEK	:. 26
Ga0187910_10006931	CVPFGYCIFRFKNLSIEALFDKNEAARYDKIKKKVAVR	5 , 20
Rflav_XFD_	CVPFGYCIPRFKNLSIEALFDRNEAARFDKEKKKVSGNS	196
Ga0224415_10007274	CVPFGYCIPRFKNLSIEALFDMNEAVKFDKEKK	940
Ga0187911_10069260	CCPFGYNLPRFKNLSIEQLFDRNESKEITDAIAPRQ	S l
Ga0187910_10015336	CAPFAYNLPRFKNLSIEELFDMHELSEEPKESMKLTD	1105
ODAL_chimera	NAPFGYNLPRFKNLSVEALFDMNEEPAPEGKERSKNADGCN	966
31009_GL0034153	NVPFAXNYSRFKNLSIDKLEDKNDPRKKKWIVIFNF	of 3
DLF014_GL0011914_	NIPFAYNYFRYKNLSIDELFDKNNTRKTIKIQDKKLEERT	87 866
Cask_Ralbus	NTPFGYDLPRYKNLSIGDLFDRNNYLNKTKESIDANSSIDSQ	944
Ga0187910_10040531	CIPFAYVTPRYKNLTIDGLEDRNRFGEDK	927
tpg DJXD01000002.1	CIPFGYVVFRYKMLTINELFDRNNPNPEFKEEV	US
Cask_Esiraeum	CTFFGYNLVRYKNL11DGLFDKNYPGKDDSDEQK	5 1
PIG-018_GL0023397	CIPFAYNLVRYKNLTIDGLEDRNAPSEK	1,3
PIG-025_GL0089734	CAPFGYNVVRFKALTIYEMFDRNYPEEKRAEBARKSC	1 6,8
Ga0129317_1008067_chimera	CSPFGYNLPRFKNLSIEGKFDMARSK	812
	FIG. 18LLL	B2

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924	913	933	958	
NAPEGYNIPRFKNLIVEQLFDRNADPETITA	NSPFGYNIPRFKNLSIEQLFDRNEYLTEK	NAPFGYNIPRFKNLSIDALFDRNEIKKNDGEKKSDD	NSPFGYNIPRFKDLSIDALFDRNEMKEETDDEKKIQT	** ***

tpg:DBXI01000091.1;

EYZ-362B_GL0088915 uncultured_Ru_sp FIG. 18MMM

RNA TARGETING METHODS AND COMPOSITIONS

CROSS-REFERENCE TO RELATED APPLICATIONS

This application is a continuation of U.S. application Ser. No. 16/257,493, filed Jan. 25, 2019, which is a divisional of U.S. application Ser. No. 15/937,699, filed Mar. 27, 2018, now U.S. Pat. No. 10,476,825, which claims priority to U.S. Provisional Application No. 62/548,846 filed Aug. 22, 2017, U.S. Provisional Application No. 62/572,963 filed Oct. 16, 2017, and U.S. Provisional Application No. 62/639,178, filed Mar. 6, 2018, all of which are herein incorporated by reference in their entireties.

ACKNOWLEDGMENT OF GOVERNMENT SUPPORT

This invention was made with government support under ²⁰ 5 DP5 OD021369-02 and 5 R21 AG056811-02 awarded by The National Institutes of Health. The government has certain rights in the invention.

FIELD

This disclosure relates to a CRISPR/Cas system for modifying (including detecting) RNA, which utilizes novel Cas13d proteins (also referred to a CasR and nCas1) and guide RNAs.

BACKGROUND

Mapping of transcriptome changes in cellular function and disease has been transformed by technological advances 35 over the last two decades, from microarrays (Schena et al., 1995) to next-generation sequencing and single cell studies (Shendure et al., 2017). However, interrogating the function of individual transcript dynamics and establishing causal linkages between observed transcriptional changes and cellular phenotype requires the ability to actively control or modulate desired transcripts.

DNA engineering technologies such as CRISPR-Cas9 (Doudna and Charpentier, 2014; Hsu et al., 2014) enable researchers to dissect the function of specific genetic elements or correct disease-causing mutations. However, simple and scalable tools to study and manipulate RNA lag significantly behind their DNA counterparts. Existing RNA interference technologies, which enable cleavage or inhibition of desired transcripts, have significant off-target effects and remain challenging engineering targets due to their key role in endogenous processes (Birmingham et al., 2006; Jackson et al., 2003). As a result, methods for studying the functional role of RNAs directly have remained limited.

One of the key restrictions in RNA engineering has been 55 the lack of RNA-binding domains that can be easily retargeted and introduced into target cells. The MS2 RNA-binding domain, for example, recognizes an invariant 21-nucleotide (nt) RNA sequence (Peabody, 1993), therefore requiring genomic modification to tag a desired transcript Pumilio homology domains possess modular repeats with each protein module recognizing a separate RNA base, but they can only be targeted to short 8 nt RNA sequences (Cheong and Hall, 2006). While previously characterized type II (Batra et al., 2017; O'Connell et al., 2014) and VI 65 (Abudayyeh et al., 2016; East-Seletsky et al., 2016) CRISPR-Cas systems can be reprogrammed to recognize

2

20-30 nt RNAs, their large size (~1200 amino acids, aa) makes it difficult to package into AAV for primary cell and in vivo delivery.

SUMMARY

This application provides bioinformatic analysis of prokaryotic genomes to identify sequence signatures of CRISPR-Cas repeat arrays and mine previously uncharacterized, compact Cas ribonucleases that can be used for RNA targeting tools. Engineered Type VI-D CRISPR effectors can be used to efficiently knockdown endogenous RNAs in human cells and manipulate alternative splicing, paving the way for RNA targeting applications and further effector domain fusions as part of a transcriptome engineering toolbox.

Provided herein are methods of modifying one or more target RNA molecules, such as a clustered regularly interspaced short palindromic repeats (CRISPR)-CRISPR associated (Cas) system-mediated RNA editing method. Such methods can include contacting one or more target RNA molecules with a non-naturally occurring (e.g., does not naturally occur in the cell or system into which it is introduced) or engineered CRISPR-Cas system. Such a 25 CRISPR-Cas system can include (1) at least one Cas13d protein or at least one Cas13d nucleic acid coding sequence (such as a mRNA or a vector encoding the at least one Cas13d protein) and (2) at least one CRISPR-Cas system guide nucleic acid molecule (such as a guide RNA, gRNA) that hybridizes with the one or more target RNA molecules, or at least one nucleic acid molecule encoding the gRNA. The Cas13d protein forms a complex with the gRNA, and the gRNA directs the complex to the one or more target RNA molecules and modifies (e.g., cuts, detects) the one or more target RNA molecules. In some examples, the one or more target RNA molecules (or a cell containing the one or more target RNA molecules) are contacted with a complex including the at least one Cas13d protein and the at least one gRNA. In some examples, the system includes Mg²⁺. However, in some example, the system does not include Mg²⁺, such as if cleavage of the target RNA is not desired.

In some examples, contacting the one or more target RNA molecules with the non-naturally occurring or engineered CRISPR-Cas system includes introducing into a cell (such as a eukaryotic or prokaryotic cell) containing the one or more target RNA molecules the non-naturally occurring or engineered CRISPR-Cas system, for example using endocytosis, a liposome, a particle, an exosome, a microvesicle, a gene gun, electroporation, a virus, or combinations thereof. In some examples, contacting the one or more target RNA molecules with the non-naturally occurring or engineered CRISPR-Cas system includes contacting a cell-free system (such as a biological or environmental sample, or a cell lysate) containing the one or more target RNA molecules with the non-naturally occurring or engineered CRISPR-Cas system (for example in a diagnostic method to detect a target RNA)

In some examples, the least one Cas13d protein includes one or more HEPN domains, is no more than 150 kD, no more than 140 kD, no more than 130 kD, no more than 120 kD, such as about 90 to 120 kD, about 100 to 120 kD or about 110 kD; includes one or more mutated HEPN domains, and can process the guide RNA, but cannot cleave or cut the one or more target RNA molecules, includes an Cas13d ortholog from a prokaryotic genome or metagenome, gut metagenome, an activated sludge metagenome, an anaerobic digester metagenome, a chicken gut metagenome, and the such as the su

enome, a human gut metagenome, a pig gut metagenome, a bovine gut metagenome, a sheep gut metagenome, a goat gut metagenome, a capybara gut metagenome, a primate gut metagenome, a termite gut metagenome, a fecal metagenome, a genome from the Order Clostridiales, or the 5 Family Ruminococcaceae; includes an Cas13d ortholog from Ruminococcus albus, Eubacterium siraeum, a flavefaciens strain XPD3002, Ruminococcus flavefaciens FD-1, uncultured Eubacterium sp TS28-c4095, uncultured Ruminococcus sp., Ruminococcus bicirculans, or Ruminococcus 10 sp CAG57; includes at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 1534, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 20 from a single array, wherein each gRNA can be different, for 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253, or combinations thereof. In some 25 examples, the least one Cas13d protein has at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 30 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 35 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, includes at least one motif shown in SEQ ID NO: 195, 196 or 197. In some examples, the least one Cas13d protein further includes one or more other agents (e.g., is a fusion protein), such as one or more subcellular localization signals, one or more effector domains, or combinations thereof. 45 In some examples, the least one Cas13d protein that includes one or more HEPN domains, is no more than 1500 aa, no more than 1200 aa, no more than 1100 aa, no more than 1000 aa, such as about 800 to 1500 aa, about 800 to 1250 aa or about 850 to 950 aa.

Also provided are isolated nucleic acid molecules encoding such Cas13d proteins, such as a cDNA, genomic DNA, RNA, or mRNA. Such isolated nucleic acid molecules can be part of a vector (such as a plasmid or viral vector), and can be operably linked to a promoter. In some examples, an 55 isolated nucleic acid molecule encoding a Cas13d protein has at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 124, 125, 126, 127, 128, 139, 140 or 141. In some examples, an 60 isolated nucleic acid molecule encoding at least one Cas13d protein (which can be part of a vector) includes at least one Cas13d protein coding sequence codon optimized for expression in a eukaryotic cell, such as human cell, for example a Cas13d coding sequence having at least 80%, at 65 least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100%

sequence identity to SEQ ID NO: 114, 115, 116, 117, 118, 119, 120, 121, 122, 123, 142, 143, 144, or 145.

In some examples, the gRNA that hybridizes with the one or more target RNA molecules in an Cas13d-mediated manner includes one or more direct repeat (DR) sequences, one or more spacer sequences, such as one or more sequences comprising an array of DR-spacer-DR-spacer. In some examples, the one or more DR sequences have at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 129, 130, 131, 132, 133, 134, 135, 136, 137, 148, 150, 151, 152, 154, 156, 157, 159, 161, 163, 165, 167, 169, 176, 178, 180, 182, 184, 186, 188, 190, 191, 192, 193, 199, 201, 203, 205, 207, 209, 211, 213, 215, 217, 219, 221, 223, 225, 227, 228, 230, 232, 234, 236, 238, 240, 242, 244, 246, 248, 250, 252, or 254. In one example, the gRNA includes additional sequences, such as an aptamer sequence.

In some examples, a plurality of gRNAs are generated example target different RNAs or target multiple regions of a single RNA, or combinations thereof.

Methods of targeting one or more target RNA molecules are provided. In some examples, an entire RNA is targeted. In some examples, a portion of an RNA is targeted. Targeting an RNA molecule can include one or more of cutting or nicking one or more target RNA molecules, activating one or more target RNA molecules, deactivating the one or more target RNA molecules, visualizing or detecting the one or more target RNA molecules, labeling the one or more target RNA molecules, binding the one or more target RNA molecules, editing the one or more target RNA molecules, trafficking the one or more target RNA molecules, and masking the one or more target RNA molecules. In some example, modifying one or more target RNA molecules includes one or more of an RNA base substitution, an RNA base deletion, an RNA base insertion, a break in the target RNA, methylating RNA, and demethylating RNA.

In some examples, such methods are used to treat a 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253, and 40 disease, such as a disease in a human In such examples, the one or more target RNA molecules is associated with the disease

Also provided are isolated proteins, including non-naturally occurring proteins. In some examples, a protein has at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEO ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, or 113. In some examples, an isolated protein is an Cas13d ortholog from a prokaryotic genome or metagenome, gut metagenome, an activated sludge metagenome, an anaerobic digester metagenome, a chicken gut metagenome, a human gut metagenome, a pig gut metagenome, a bovine gut metagenome, a sheep gut metagenome, a goat gut metagenome, a capybara gut metagenome, a primate gut metagenome, a termite gut metagenome, a fecal metagenome, a genome from the Order Clostridiales, or the Family Ruminococcaceae. In some examples, an Cas13d ortholog includes an Cas13d ortholog from Ruminococcus albus, Eubacterium siraeum, a Ruminococcus flavefaciens strain XPD3002, Ruminococcus flavefaciens FD-1, uncultured

Eubacterium sp TS28-c4095, uncultured Ruminococcus sp., Ruminococcus bicirculans, or Ruminococcus sp CAG57. The protein is an Cas13d protein that further includes one or more other agents or domains (e.g., is a fusion protein), such as one or more subcellular localization signals, one or more seffector domains, or combinations thereof.

Also provided are isolated guide RNA (gRNA) molecules. In some examples, an isolated gRNA includes one or more direct repeat (DR) sequences, such as an unprocessed (e.g., about 36 nt) or processed DR (e.g., about 30 nt). In some examples a DR has at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 129, 130, 131, 132, 133, 134, 135, 136, 137,148,150,151,152,154,156,157,159,161,163,165,167, 15169,176,178,180,182,184, 186,188,190,191,192,193,199, 201,203,205,207,209,211,213,215,217,219,221,223, 225, 227, 228, 230, 232, 234, 236, 238, 240, 242, 244, 246, 248, 250, 252, or 254. Such a gRNA can further include one or more spacer sequences specific for (e.g., is complementary 20 to) the target RNA.

Also provided are ribonucleoprotein (RNP) complexes, which include an Cas13d protein provided herein and a gRNA provided herein.

Also provided are recombinant cells that include any ²⁵ Cas13d protein (or nucleic acid molecule encoding Cas13d), any gRNA, any RNP complex, or any vector, provided herein. In one example, the cell is not a bacterial cell. In one example, the cell is a bacterial cell.

Also provided are compositions that include one or more 30 of any Cas13d protein (or nucleic acid molecule encoding Cas13d), any gRNA, any RNP complex, any isolated nucleic acid molecule, any vector, or any cell, provided herein. Such compositions can include a pharmaceutically acceptable carrier.

Also provided are kits. Such kits can include one or more of any Cas13d protein (or nucleic acid molecule encoding Cas13d), any gRNA, any RNP complex, any isolated nucleic acid molecule, any vector, any cell, or any composition provided herein. Such reagents can be combined or in 40 separate containers.

The foregoing and other objects and features of the disclosure will become more apparent from the following detailed description, which proceeds with reference to the accompanying figures.

BRIEF DESCRIPTION OF THE DRAWINGS

FIGS. 1A-1B: Bioinformatic pipeline for the identification of the RNA-targeting class 2 CRISPR system Cas13d. 50 (A) Schematic describing a computational pipeline for CRISPR system identification. A minimal definition for a putative class 2 CRISPR locus was used, requiring only a CRISPR repeat array and a nearby protein >750 aa in length. The initial search was performed on prokaryotic genome 55 assemblies derived from NCBI Genome, and later expanded via TBLASTN of predicted Cas13d proteins against public metagenome sequences without predicted open reading frames. DR, direct repeat. (B) Phylogenetic classification and alignment of full-length Cas13d effectors and metag- 60 enomic fragments. Cas13d effectors and metagenomic Cas13d protein fragments cluster into several distinct branches, which are colored for ease of interpretation. Shading indicates residue conservation using the Blosum62 matrix. Full-length Cas13d effectors used in this study were sampled from distinct branches of the Cas13d family. Alignment of Cas13d proteins and protein fragments was per6

formed using ClustalOmega 1.2.4 and maximum-likelihood tree building was performed with PhyML 3.2.

FIGS. 2A-2C: Type VI CRISPR-Cas13d is a family of single effector CRISPR ribonucleases. (A) Maximum-likelihood phylogenetic tree of Cas13d effectors used herein, with the full Cas13d CRISPR locus depicted along with conserved HEPN RNase domains. Grey rectangles denote CRISPR direct repeats (DRs) and blue diamonds indicate spacer sequences. (B) RNA sequencing of a heterologously expressed Cas13d locus from an uncultured Ruminococcus sp. sample. Mature gRNAs mapping to the CRISPR array indicate a processed 30 nt DR and a variable spacer length from 14-26 nt (SEQ ID NO: 278). Co-fold analysis of direct repeat truncation indicates a strong hairpin structure. (C) Purified E. siraeum Cas13d and catalytically dead Cas13d (dCas13d) protein are each sufficient to process a guide array into its two component gRNAs. Addition of EDTA does not impair gRNA processing. 'd', dCas13d(R295A, H300A, R849A, H854A).

FIGS. 3A-3D. Phylogenetic classification of RNA-targeting class 2 CRISPR effectors and sequence conservation within the Cas13d family. (A) HEPN motif conservation in Cas13d effectors used herein with conserved residues shaded according to Blosum62. Consensus sequence of HEPN1 domain region (SEQ ID NO: 279) and alignment of HEPN1 domain region of seven Cas13d proteins (SEQ ID NOS: 280 to 286). Consensus sequence of HEPN2 domain region (SEQ ID NO: 287) and alignment of HEPN2 domain region of seven Cas13d proteins (SEQ ID NOS: 288 to 294). The RxxxxH HEPN motif is highlighted. (B) Maximumlikelihood tree of type VI CRISPR-Cas families. Average amino acid lengths of Type VI Cas13 superfamily effectors are indicated in red. Alignment of previously described class 2 CRISPR RNA-targeting proteins (Abudayyeh et al., 2017; 35 Cox et al., 2017; East-Seletsky et al., 2017; East-Seletsky et al., 2016; Smargon et al., 2017) and Cas13d effectors was performed using MAFFT 7.38 and maximum-likelihood tree building was performed with PhyML 3.2. Branch labels and scale bar indicate substitutions per site. (C) Predicted Cas13d direct repeat RNA secondary structure. (D) Sequence logo of full length 36 nt Cas13d direct repeats.

FIGS. 4A-4C: Purification of recombinant Cas13d protein. EsCas13d was expressed as N-terminal His-MBP fusion and purified by successive affinity, cation exchange, and size exclusion chromatography. The His-tag was removed by TEV protease cleavage. (A) Chromatogram from Superdex 200 column for EsCas13d. (B) SDS-PAGE gel of size exclusion chromatography fractions for *E. siraeum* Cas13d. (C) SDS-PAGE gel of purified *E. siraeum* Cas13d and dCas13d (R295A, H300A, R849A, H854A mutations of predicted catalytic residues in both HEPN motifs).

FIGS. 5A-5D: Programmable RNA targeting by Cas13d in vitro. (A) *E. siraeum* Cas13d requires a matching guide array or mature gRNA to efficiently cleave complementary ssRNA targets. Denaturing gel depicts cleavage reactions incubated at 37° C. for 1 hour. NT, non-targeting. (B) Substitution with dCas13d or addition of EDTA abrogate Cas13d-mediated RNA targeting with both the guide and array. 'd', dead Cas13d. (C) Denaturing gel depicting guide-target match dependent activation of Cas13d cleavage activity. Scrambled target RNA ('A') is fluorescently labeled, while guide-complementary activator target RNA ('B') is unlabeled. RNA cleavage activity is abolished by the individual removal of guide RNA or complementary target RNA, as well as the addition of EDTA or the catalytic inactivation of Cas13d (indicated as 'd'). (D) A model for

guide and target-dependent activation of Cas13d RNase activity. The ternary Cas13d:gRNA:target RNA complex is capable of cleaving the complementary target RNA or bystander RNAs.

FIGS. 6A-6H. In vitro characterization of Cas13d prop- 5 erties. (A) Schematic showing the length and sequence of gRNA spacer truncations and spacer position relative to the complementary ssRNA target (from top to bottom, SEQ ID NOS: 295 to 309). (B) Denaturing gel depicting EsCas13d cleavage activity of target RNA with different spacer 10 lengths. (C) Denaturing gel depicting EsCas13d cleavage reactions paired with 12 guides from FIG. 3A tiling a complementary ssDNA version of the ssRNA target. (D) Denaturing gel depicting cleavage reactions using EsCas13d paired with the same 12 guides tiling a dsDNA version of the 15 complementary target. (E) Quantification of cleavage efficiency from FIG. 3A. Each PFS base is the average of 3 different spacer sequences tiling a complementary target RNA. Cleavage percentage is determined by the ratio of cleaved band intensity divided by total lane intensity. Mean 20 is depicted ±SD with each data point representing an independent replicate. (F) Cas13d-mediated cleavage of target RNA carrying different PFS bases given an invariant spacer sequence. Quantification of Cas13d cleavage efficiency and a representative denaturing gel depicting EsCas13d cleavage 25 activity are shown. Differences are not significant (one-way ANOVA, P=0.768). Cleavage percentage is determined as above, and mean is depicted ±SD with n=3. (G) and (H) Optimal temperature range for Cas13d activity. Denaturing gels depicting EsCas13d cleavage activity at temperatures 30 ranging from 16-62° C. for two different target RNAs.

FIGS. 7A-7B: Characterization of Cas13d target substrate preference. (A) Cas13d can be generalizably reprogrammed with multiple guides and does not exhibit a protospacer flanking sequence (PFS) requirement. RNA cleavage by 35 EsCas13d and 12 guides tiling the target RNA is shown. Control lanes are from a separate gel run in parallel. (B) Cas13d preferentially cleaves uracil bases in the loop of a hairpin or a linear homopolymer repeat, which is interrupted every 5 nt by a transition mutation (X) to enable synthesis. 40

FIGS. 8A-8D: RNA knockdown activity screen of engineered Cas13d orthologs in human cells. (A) Schematic for mammalian expression constructs encoding for engineered Cas13d effectors and guides. NLS, nuclear localization signal. pre-gRNA, artificial unprocessed guide RNA con- 45 taining a single 30 nt spacer sequence flanked by 2 full length 36 nt DRs. gRNA, predicted mature guide RNA with a single 30 nt processed DR and 22 nt spacer sequence (B) Heatmap of mCherry protein knockdown in a Cas13d ortholog activity screen in human HEK 293FT cells using 50 pools of 4 pre-gRNAs or gRNAs. Normalized MFI, median fluorescent intensity relative to non-targeting condition. Positions in gray were not tested, with n=3. (C) Immunocytochemistry of Cas13d showing localization and expression of engineered constructs. Scale bar, 10 µm. Blue 55 pseudocolor, DAPI staining of nuclei. (D) Comparison of Adm and Rfx Cas13d ortholog constructs for knockdown of endogenous B4GALNT1 mRNA reveals RfxCas13d-NLS (CasRx) to be most effective for both guide RNA architectures. Pools of 4 guides were used for targeting. NT, non- 60 targeting. Values are mean ±SEM with n=3.

FIGS. 9A-9H: CasRx mediates efficient and specific knockdown of diverse human coding and noncoding transcripts. (A) Multiple guide RNAs tiling a target transcript can be expressed as a single array and processed by 65 RfxCas13d-NLS (CasRx) into individual gRNAs within the same cell. (B) Arrays of 4 guides each mediate target

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knockdown by CasRx in 293FT cells via transient transfection. Knockdown relative to GFP vehicle control was determined by qPCR, with n=3. (C) Schematic of CasRx target sequences and spacer position-matched shRNAs. (D) Relative target RNA knockdown by individual position-matched shRNAs and CasRx gRNAs. NT, non-targeting. CRISPRi, dCas9-mediated CRISPR interference for transcriptional repression (n=3) (E) Volcano plot of differential transcript levels between B4GALNT1 targeting and non-targeting (NT) shRNAs as determined by RNA sequencing (n=3). 542 non-specific transcript changes were identified. (F) Volcano plot of differential transcript levels between B4GALNT1targeting CasRx and non-targeting (NT) guide. Targeting guide position is matched to the shRNA shown in (E). B4GALNT1 was the only transcript exhibiting a significant change, with n=3. (G) Summary of significant off-target transcript perturbations by matched shRNAs and CasRx guides. (H) CasRx targeting of 11 endogenous transcripts, each with 3 guides and a non-targeting (NT) guide in 293FT cells. Transcript levels are relative to GFP vehicle control. mean ±SEM with n=3.

FIGS. 10A-10D. Comparison of engineered Cas13 superfamily effectors for targeted knockdown and splicing. (A) Relative target RNA knockdown by individual positionmatched gRNAs for CasRx, NLS-LwaCas13a-msfGFP (Abudayyeh et al., 2017) and PspCas13b-NES (Cox et al., 2017) in HEK 293FT cells. NT, non-targeting. Values are mean ±SEM with n=3. (B) Comparison of Cas13 median knockdown efficiencies. n=3 per guide RNA. **** indicates P<0.0001 according to Friedman's test. (C) Exon exclusion by catalytically inactive NLS-dCas13a-msfGFP on the bichromatic splicing reporter. Guides are position-matched to those reported in FIG. 6B for CasRx. Values are mean ±SEM with n=3. (D) Comparison of splicing modulation by NLS-dCas13a-msfGFP and CasRx. Fold change in targeted exon exclusion relative to non-targeting guide is shown. **** indicates P<0.0001 according to two-way ANOVA.

FIGS. 11A-11B. RNA sequencing from CasRx and shRNA targeting of ANXA4 in human cells. (A) Volcano plots of differential transcript levels between ANXA4 targeting and non-targeting (NT) shRNAs as determined by RNA sequencing (n=3). 915 non-specific transcript changes were identified. (B) Volcano plot of differential transcript levels for an ANXA4 targeting CasRx array used in FIG. 9B containing a guide position matched to the shRNA shown in (A) and a non-targeting (NT) array. ANXA4 was the only transcript exhibiting significant downregulation with n=3. HIST2HBE was the only transcript identified to exhibit significant upregulation. H2B is a dimer partner of H2AX (Du et al., 2006) which has been shown to interact with ANXA4 (Yang et al., 2010).

FIGS. 12A-12F: AAV delivery of catalytically inactive dCasRx splice effectors to manipulate alternative splicing. (A) Schematic of bichromatic exon skipping reporter. +1 and +3, reading frame. BP, intronic branch point-targeting guide. SA, splice acceptor site-overlapping guide. EX, exonic guide. SD, splice donor site-overlapping guide. AUG, start codon. UGA, stop codon. Inclusion of the second exon leads to an out-of-frame (+3), non-fluorescent translation of dsRed followed by in-frame mTagBFP2. Exclusion of the targeted exon leads to an in frame translation of dsRed (+1) followed by a stop codon. (B) Induced exon exclusion by dCasRx and an N-terminal hnRNPa1-dCasRx fusion protein targeted to pre-mRNA. The Gly-rich C-terminal domain of hnRNPa1 is used as the effector domain Exon skipping efficiency is depicted as a relative percentage of cells carrying primarily the dsRed or BFP isoform, determined through flow cytom-

etry. NLS, nuclear localization signal. 'A', CRISPR array carrying all 4 guides. Values are mean ±SEM with n=3. (C) AAV design carrying dCasRx and a three-guide array with total transgene size<4.3 kb, including AAV inverted terminal repeats (ITRs). (D) Schematic of frontotemporal dementia 5 (FTD) disease modeling. Neurons are generated via Neurogenin-2 (Ngn2) directed differentiation of patient-derived and control iPSCs followed by transduction with dCasRx or vehicle control AAV (EFS-mTagBFP2). (E) FTD is associated with SNPs in a putative intronic splice enhancer fol- 10 lowing exon 10 of the MAPT transcript encoding for tau. Alternative splicing of MAPT exon 10 results in 4R tau (by inclusion) and 3R tau (by exclusion). SNPs in the intronic splice enhancer including the indicated IVS 10 +16 mutation result in increased exon inclusion and higher levels of 4R 15 tau. To facilitate reduction of 4R tau levels, gRNAs contained in a dCasRx array were targeted to the exon 10 splice acceptor (g1) as well as two putative exonic splice enhancers indicated in purple (g2, g3). (F) Relative 4R/3R tau transcript ratios in differentiated neurons were assayed via qPCR 20 at 14 days following transduction with AAV. FTD, frontotemporal dementia cells carrying IVS 10+16. Values are mean ±S.D. with n=3. **** indicates P<0.0001.

FIG. 13 is a bar graph showing RNA targeting in human cells using the disclosed methods.

FIG. 14 is a schematic drawing showing how the disclosed Cas13d and DRs can be used to achieve alternative splicing.

FIGS. 15A-15D are a series of panels showing: (A) Cas13d is converted into an active RNase complex upon 30 binding a target matching the spacer sequence of the guide RNA. It is capable of cleaving gRNA-complementary target RNA or non-complementary bystander RNAs. (B) Cas13d target-dependent RNase activity can be converted into a detectable signal, for example through cleavage of a labeled 35 detector RNA that is cleaved only in the presence of a target matching the spacer of the Cas13d guide RNA. In this example, the detector RNA contains a fluorophore, 'F', and a quencher 'Q', that abolishes fluorescence. Only upon bystander RNA cleavage is the fluorophore liberated from 40 Eubacterium siraeum containing a mutated HEPN site. the quencher and fluorescence is generated. (C) Cas13d from E. siraeum produces a visible signal only in the presence of a perfectly matched target and not in the presence of different mismatched targets. (D) Cas13d from R. flavefaciens strain XPD3002 produces a visible signal only in the 45 presence of a perfectly matched target and not in the presence of different mismatched targets.

FIGS. 16A-16B: (A) Alignment of 7 orthologs shows regions with high (green bars) and low (red bars) conservation. Regions selected for deletion are marked 1-10. (B) 50 knock-down (top) and splicing (bottom) activity of fulllength CasRx and CasRx deletion variants. Deletion variant 5 is shown to retain full activity, demonstrating the feasibility of deleting areas of low conservation while retaining

FIGS. 17A and 17B show targeting of ccdB in bacterial cells. (A) construct introduced into bacterial cells expressing ccdB, and (B) relative expression of ccdB under various

FIGS. 18A-18MMM show an alignment of 53 different 60 Cas13d proteins as follows: SEQ ID NOs: 310 (037_-_ emblOIZA01000315.1), 183 (emblOCTW011587266.1), 189 (k87_11092736), 220 (BMZ-11B_GL0037915), 218 (BMZ-11B_GL0037771), 222 (BMZ-11B_GL0069617), 229 (Ga0099364_10024192), 216 (530373_GL0023589), 65 177 (emblODAI011611274.1), 200 (EMG_10003641), 139 (CasR_P1E0_metageno), 179 (emblOIZX01000427.1), 208

(160582958_gene49834), 166 (gil1 198542314lgblN-FIR01000008.1), 185 (emblOGNF01009141.1), 202 (Ga0129306_1000735), 239 (MH0288_GL0082219), 311 (PIG-022_GL0026351), 249 (PIG-028_GL0185479), 210 (250twins 35838 GL0110300), 014_GL0226364), 212 (250twins_36050_GL0158985), 175 (emblOJMM01002900.1), 164 (emblOGPN01002610.1), 160 (emblOHCP01000044.1), 312 (PIG-046_GL0077813), 313 (pig_chimera), 187 (emblOIEN01002196.1), 241 (02.UC29-0_GL0096317), 140 (CasR_Anaerobic_dig), 162 (emblOGDF01008514.1), 155 (emblOGZC01000639.1), (Ga0224415_10048792_chimera), blOCVV012889144.1), 231 (Ga0187910_10006931), 92 (R._flav_XPD_), 198 (Ga0224415_10007274), (Ga0187911_10069260), 233 (Ga0187910_10015336), 253 (ODAI chimera), 214 (31009 GL0034153), (DLF014_GL0011914_), 127 (CasR_R._albus), (Ga0187910_10040531), 153 (tpglDJXD01000002.1), 125 (CasR_E._siraeum), 245 (PIG-018_GL0023397), 247 (PIG-025_GL0099734), 204 (Ga0129317_1008067_chimera), 226 (EYZ-362B_GL0088915), 3 (uncultured_Ru_sp), 126 (R._flav._FD-1), and 149 (tpglDBYI01000091.1), from top to bottom.

SEQUENCE LISTING

The nucleic and amino acid sequences listed in the accompanying sequence listing are shown using standard letter abbreviations for nucleotide bases, and three letter code for amino acids, as defined in 37 C.F.R. 1.822. Only one strand of each nucleic acid sequence is shown, but the complementary strand is understood as included by any reference to the displayed strand. The Sequence Listing is submitted as an ASCII text file, created on Jun. 18, 2021, 924 KB, which is incorporated by reference herein.

SEQ ID NO: 1 is an exemplary Cas13d sequence from Eubacterium siraeum containing a HEPN site.

SEQ ID NO: 2 is an exemplary Cas13d sequence from

SEQ ID NO: 3 is an exemplary Cas13d sequence from uncultured Ruminococcus sp. containing a HEPN site.

SEQ ID NO: 4 is an exemplary Cas13d sequence from uncultured Ruminococcus sp. containing a mutated HEPN

SEQ ID NO: 5 is an exemplary Cas13d sequence from Gut_metagenome_contig2791000549.

SEQ ID NO: 6 is an exemplary Cas13d sequence from Gut_metagenome_contig855000317.

SEQ ID NO: 7 is an exemplary Cas13d sequence from Gut_metagenome_contig3389000027.

SEQ ID NO: 8 is an exemplary Cas13d sequence from Gut_metagenome_contig8061000170.

SEQ ID NO: 9 is an exemplary Cas13d sequence from 55 Gut_metagenome_contig1509000299.

SEQ ID NO: 10 is an exemplary Cas13d sequence from Gut_metagenome_contig9549000591.

SEQ ID NO: 11 is an exemplary Cas13d sequence from $Gut_metagenome_contig 71000500.$

SEQ ID NO: 12 is an exemplary Cas13d sequence from human gut metagenome.

SEQ ID NO: 13 is an exemplary Cas13d sequence from Gut_metagenome_contig3915000357.

SEQ ID NO: 14 is an exemplary Cas13d sequence from Gut_metagenome_contig4719000173.

SEQ ID NO: 15 is an exemplary Cas13d sequence from Gut_metagenome_contig6929000468.

SEQ ID NO: 16 is an exemplary Cas13d sequence from Gut_metagenome_contig7367000486.

SEQ ID NO: 17 is an exemplary Cas13d sequence from Gut_metagenome_contig7930000403.

SEQ ID NO: 18 is an exemplary Cas13d sequence from ⁵ Gut_metagenome_contig993000527.

SEQ ID NO: 19 is an exemplary Cas13d sequence from Gut_metagenome_contig6552000639.

SEQ ID NO: 20 is an exemplary Cas13d sequence from Gut_metagenome_contig11932000246.

SEQ ID NO: 21 is an exemplary Cas13d sequence from Gut_metagenome_contig12963000286.

SEQ ID NO: 22 is an exemplary Cas13d sequence from Gut_metagenome_contig2952000470.

SEQ ID NO: 23 is an exemplary Cas13d sequence from Gut_metagenome_contig451000394.

SEQ ID NO: 24 is an exemplary Cas13d sequence from *Eubacterium_siraeum_*DSM_15702.

SEQ ID NO: 25 is an exemplary Cas13d sequence from 20 gut_metagenome_P19E0k2120140920,c369000003.

SEQ ID NO: 26 is an exemplary Cas13d sequence from Gut_metagenome_contig7593000362.

SEQ ID NO: 27 is an exemplary Cas13d sequence from Gut metagenome contig12619000055.

SEQ ID NO: 28 is an exemplary Cas13d sequence from Gut_metagenome_contig1405000151.

SEQ ID NO: 29 is an exemplary Cas13d sequence from Chicken_gut_metagenome_c298474.

SEQ ID NO: 30 is an exemplary Cas13d sequence from 30 Gut metagenome contig1516000227.

SEQ ID NO: 31 is an exemplary Cas13d sequence from Gut_metagenome_contig1838000319.

SEQ ID NO: 32 is an exemplary Cas13d sequence from Gut_metagenome_contig13123000268.

SEQ ID NO: 33 is an exemplary Cas13d sequence from Gut metagenome contig5294000434.

SEQ ID NO: 34 is an exemplary Cas13d sequence from Gut_metagenome_contig6415000192.

SEQ ID NO: 35 is an exemplary Cas13d sequence from 40 Gut_metagenome_contig6144000300.

SEQ ID NO: 36 is an exemplary Cas13d sequence from Gut_metagenome_contig9118000041.

SEQ ID NO: 37 is an exemplary Cas13d sequence from Activated_sludge_metagenome_transcript_124486.

SEQ ID NO: 38 is an exemplary Cas13d sequence from Gut_metagenome_contig1322000437.

SEQ ID NO: 39 is an exemplary Cas13d sequence from Gut_metagenome_contig4582000531.

SEQ ID NO: 40 is an exemplary Cas13d sequence from 50 Gut_metagenome_contig9190000283.

SEQ ID NO: 41 is an exemplary Cas13d sequence from Gut_metagenome_contig1709000510.

SEQ ID NO: 42 is an exemplary Cas13d sequence from M24_(LSQX01212483_Anaerobic_digester_metagenome) with a HEPN domain

SEQ ID NO: 43 is an exemplary Cas13d sequence from Gut_metagenome_contig3833000494.

SEQ ID NO: 44 is an exemplary Cas13d sequence from Activated_sludge_metagenome_transcript_117355.

SEQ ID NO: 45 is an exemplary Cas13d sequence from Gut_metagenome_contig11061000330.

SEQ ID NO: 46 is an exemplary Cas13d sequence from Gut_metagenome_contig338000322 from sheep gut metagenome

SEQ ID NO: 47 is an exemplary Cas13d sequence from human gut metagenome.

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SEQ ID NO: 48 is an exemplary Cas13d sequence from Gut_metagenome_contig9530000097.

SEQ ID NO: 49 is an exemplary Cas13d sequence from Gut_metagenome_contig1750000258.

SEQ ID NO: 50 is an exemplary Cas13d sequence from Gut metagenome contig5377000274.

SEQ ID NO: 51 is an exemplary Cas13d sequence from gut_metagenome_P19E0k2120140920_c248000089.

SEQ ID NO: 52 is an exemplary Cas13d sequence from Gut_metagenome_contig11400000031.

SEQ ID NO: 53 is an exemplary Cas13d sequence from Gut_metagenome_contig7940000191.

SEQ ID NO: 54 is an exemplary Cas13d sequence from Gut_metagenome_contig6049000251.

SEQ ID NO: 55 is an exemplary Cas13d sequence from Gut_metagenome_contig1137000500.

SEQ ID NO: 56 is an exemplary Cas13d sequence from Gut_metagenome_contig9368000105.

SEQ ID NO: 57 is an exemplary Cas13d sequence from Gut_metagenome_contig546000275.

SEQ ID NO: 58 is an exemplary Cas13d sequence from Gut_metagenome_contig7216000573.

SEQ ID NO: 59 is an exemplary Cas13d sequence from 25 Gut metagenome contig4806000409.

SEQ ID NO: 60 is an exemplary Cas13d sequence from Gut_metagenome_contig10762000480.

SEQ ID NO: 61 is an exemplary Cas13d sequence from Gut_metagenome_contig4114000374.

SEQ ID NO: 62 is an exemplary Cas13d sequence from *Ruminococcus_flavefaciens_*FD1.

SEQ ID NO: 63 is an exemplary Cas13d sequence from Gut_metagenome_contig7093000170.

SEQ ID NO: 64 is an exemplary Cas13d sequence from 35 Gut_metagenome_contig1113000384.

SEQ ID NO: 65 is an exemplary Cas13d sequence from Gut_metagenome_contig6403000259.

SEQ ID NO: 66 is an exemplary Cas13d sequence from Gut_metagenome_contig6193000124.

SEQ ID NO: 67 is an exemplary Cas13d sequence from Gut_metagenome_contig721000619.

SEQ ID NO: 68 is an exemplary Cas13d sequence from Gut_metagenome_contig1666000270.

SEQ ID NO: 69 is an exemplary Cas13d sequence from Gut_metagenome_contig2002000411.

SEQ ID NO: 70 is an exemplary Cas13d sequence from *Ruminococcus albus*.

SEQ ID NO: 71 is an exemplary Cas13d sequence from Gut_metagenome_contig13552000311.

SEQ ID NO: 72 is an exemplary Cas13d sequence from Gut_metagenome_contig10037000527.

SEQ ID NO: 73 is an exemplary Cas13d sequence from Gut_metagenome_contig23 8000329.

SEQ ID NO: 74 is an exemplary Cas13d sequence from 55 Gut_metagenome_contig2643000492.

SEQ ID NO: 75 is an exemplary Cas13d sequence from Gut metagenome contig874000057.

SEQ ID NO: 76 is an exemplary Cas13d sequence from Gut_metagenome_contig4781000489.

SEQ ID NO: 77 is an exemplary Cas13d sequence from $Gut_metagenome_contig12144000352$.

SEQ ID NO: 78 is an exemplary Cas13d sequence from Gut_metagenome_contig5590000448.

SEQ ID NO: 79 is an exemplary Cas13d sequence from Gut_metagenome_contig9269000031.

SEQ ID NO: 80 is an exemplary Cas13d sequence from Gut_metagenome_contig8537000520.

SEQ ID NO: 81 is an exemplary Cas13d sequence from Gut metagenome contig1845000130.

SEQ ID NO: 82 is an exemplary Cas13d sequence from gut_metagenome_P13E0k2120140920_c3000072.

SEQ ID NO: 83 is an exemplary Cas13d sequence from 5 gut_metagenome_P1E0k2120140920_c1000078.

SEQ ID NO: 84 is an exemplary Cas13d sequence from Gut_metagenome_contig12990000099.

SEQ ID NO: 85 is an exemplary Cas13d sequence from Gut_metagenome_contig525000349.

SEQ ID NO: 86 is an exemplary Cas13d sequence from Gut_metagenome_contig7229000302.

SEQ ID NO: 87 is an exemplary Cas13d sequence from Gut_metagenome_contig3227000343.

SEQ ID NO: 88 is an exemplary Cas13d sequence from 15 Gut_metagenome_contig7030000469.

SEQ ID NO: 89 is an exemplary Cas13d sequence from Gut_metagenome_contig5149000068.

SEQ ID NO: 90 is an exemplary Cas13d sequence from Gut_metagenome_contig400200045.

SEQ ID NO: 91 is an exemplary Cas13d sequence from Gut_metagenome_contig10420000446.

SEQ ID NO: 92 is an exemplary Cas13d sequence from new_flavefaciens,strain_XPD3002.

SEQ ID NO: 93 is an exemplary Cas13d sequence from 25 M26_Gut_metagenome_contig698000307.

SEQ ID NO: 94 is an exemplary Cas13d sequence from M36_Uncultured_*Eubacterium*_sp_TS28_c40956.

SEQ ID NO: 95 is an exemplary Cas13d sequence from M12_gut_metagenome_P25C0k2120140920_c134000066. SEQ ID NO: 96 is an exemplary Cas13d sequence from

human gut metagenome. SEQ ID NO: 97 is an exemplary Cas13d sequence from M10_gut_metagenome_P25C90k2120140920,_c28000041.

SEQ ID NO: 98 is an exemplary Cas13d sequence from 35 M11_gut_metagenome_P25C7k2120140920_c4078000105. SEQ ID NO: 99 is an exemplary Cas13d sequence from gut metagenome P25C0k2120140920 c32000045.

SEQ ID NO: 100 is an exemplary Cas13d sequence from M13_gut_metagenome_P23C7k2120140920_c3000067.

SEQ ID NO: 101 is an exemplary Cas13d sequence from M5_gut_metagenome_P18E90k2120140920.

SEQ ID NO: 102 is an exemplary Cas13d sequence from M21_gut_metagenome_P18E0k2120140920.

SEQ ID NO: 103 is an exemplary Cas13d sequence from 45 M7_gut_metagenome_P38C7k2120140920_c4841000003.

SEQ ID NO: 104 is an exemplary Cas13d sequence from *Ruminococcus_bicirculans*.

SEQ ID NO: 105 is an exemplary Cas13d sequence.

SEQ ID NO: 106 is an exemplary Cas13d consensus 50 sequence.

SEQ ID NO: 107 is an exemplary Cas13d sequence from M18_gut_metagenome_P22E0k2120140920_c3395000078.

SEQ ID NO: 108 is an exemplary Cas13d sequence from M17_gut_metagenome_P22E90k2120140920_c114.

SEQ ID NO: 109 is an exemplary Cas13d sequence from *Ruminococcus* sp_CAG57.

SEQ ID NO: 110 is an exemplary Cas13d sequence from gut_metagenome_Pl 1E90k2120140920_c43000123.

SEQ ID NO: 111 is an exemplary Cas13d sequence from 60 M6_gut_metagenome_P13E90k2120140920_c7000009.

SEQ ID NO: 112 is an exemplary Cas13d sequence from M19_gut_metagenome_P 17E90k2120140920.

SEQ ID NO: 113 is an exemplary Cas13d sequence from gut_metagenome_P17E0k2120140920,_c87000043.

SEQ ID NO: 114 is an exemplary human codon optimized *Eubacterium siraeum* Cas13d nucleic acid sequence.

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SEQ ID NO: 115 is an exemplary human codon optimized *Eubacterium siraeum* Cas13d nucleic acid sequence with a mutant HEPN domain.

SEQ ID NO: 116 is an exemplary human codon-optimized *Eubacterium siraeum* Cas13d nucleic acid sequence with N-terminal NLS.

SEQ ID NO: 117 is an exemplary human codon-optimized *Eubacterium siraeum* Cas13d nucleic acid sequence with N- and C-terminal NLS tags.

SEQ ID NO: 118 is an exemplary human codon-optimized uncultured *Ruminococcus* sp. Cas13d nucleic acid sequence.

SEQ ID NO: 119 is an exemplary human codon-optimized uncultured *Ruminococcus* sp. Cas13d nucleic acid sequence with a mutant HEPN domain

SEQ ID NO: 120 is an exemplary human codon-optimized uncultured *Ruminococcus* sp. Cas13d nucleic acid sequence with N-terminal NLS.

SEQ ID NO: 121 is an exemplary human codon-optimized uncultured *Ruminococcus* sp. Cas13d nucleic acid sequence with N- and C-terminal NLS tags.

SEQ ID NO: 122 is an exemplary human codon-optimized uncultured *Ruminococcus flavefaciens* FD1 Cas13d nucleic acid sequence.

SEQ ID NO: 123 is an exemplary human codon-optimized uncultured *Ruminococcus flavefaciens* FD1 Cas13d nucleic acid sequence with mutated HEPN domain.

SEQ ID NO: 124 is an exemplary Cas13d nucleic acid sequence from *Ruminococcus bicirculans*.

SEQ ID NO: 125 is an exemplary Cas13d nucleic acid sequence from *Eubacterium siraeum*.

SEQ ID NO: 126 is an exemplary Cas13d nucleic acid sequence from *Ruminococcus flavefaciens* FD1.

SEQ ID NO: 127 is an exemplary Cas13d nucleic acid sequence from *Ruminococcus albus*.

SEQ ID NO: 128 is an exemplary Cas13d nucleic acid sequence from *Ruminococcus flavefaciens* XPD.

SEQ ID NO: 129 is an exemplary consensus DR nucleic 40 acid sequence for *E. siraeum* Cas13d.

SEQ ID NO: 130 is an exemplary consensus DR nucleic acid sequence for Rum. Sp. Cas13d.

SEQ ID NO: 131 is an exemplary consensus DR nucleic acid sequence for *Rum. flavefaciens* strain XPD3002 Cas13d and CasRx.

SEQ ID NOS: 132-137 are exemplary consensus DR nucleic acid sequences.

SEQ ID NO: 138 is an exemplary 50% consensus sequence for seven full-length Cas13d orthologues.

SEQ ID NO: 139 is an exemplary Cas13d nucleic acid sequence from Gut metagenome P1E0.

SEQ ID NO: 140 is an exemplary Cas13d nucleic acid sequence from Anaerobic digester.

SEQ ID NO: 141 is an exemplary Cas13d nucleic acid sequence from *Ruminococcus* sp. CAG:57.

SEQ ID NO: 142 is an exemplary human codon-optimized uncultured Gut metagenome P1E0 Cas13d nucleic acid sequence.

SEQ ID NO: 143 is an exemplary human codon-optimized Anaerobic Digester Cas13d nucleic acid sequence.

SEQ ID NO: 144 is an exemplary human codon-optimized *Ruminococcus flavefaciens* XPD Cas13d nucleic acid sequence.

SEQ ID NO: 145 is an exemplary human codon-opti-65 mized *Ruminococcus albus* Cas13d nucleic acid sequence.

SEQ ID NO: 146 is an exemplary processing of the *Ruminococcus* sp. CAG:57 CRISPR array.

SEQ ID NO: 147 is an exemplary Cas13d protein sequence from contig emblOBVH01003037.1, human gut metagenome sequence (also found in WGS contigs emblOBXZ01000094.11 and emblOBJF01000033.1.

SEQ ID NO: 148 is an exemplary consensus DR nucleic 5 acid sequence (goes with SEQ ID NO: 147).

SEQ ID NO: 149 is an exemplary Cas13d protein sequence from contig tpglDBYI01000091.11 (Uncultivated Ruminococcus flavefaciens UBA1190 assembled from bovine gut metagenome).

SEQ ID NOS: 150-152 are exemplary consensus DR nucleic acid sequences (goes with SEQ ID NO: 149).

SEQ ID NO: 153 is an exemplary Cas13d protein sequence from contig tpglDJXD01000002.11 (uncultivated Ruminococcus assembly, UBA7013, from sheep gut meta- 15 genome).

SEQ ID NO: 154 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 153).

SEQ ID NO: 155 is an exemplary Cas13d protein sequence from contig OGZCO1000639.1 (human gut meta- 20 genome assembly).

SEQ ID NOS: 156-177 are exemplary consensus DR nucleic acid sequences (goes with SEQ ID NO: 155).

SEQ ID NO: 158 is an exemplary Cas13d protein sequence from contig emblOHBM01000764.1 (human gut 25 metagenome assembly).

SEQ ID NO: 159 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 158).

SEQ ID NO: 160 is an exemplary Cas13d protein sequence from contig emblOHCP01000044.1 (human gut 30 metagenome assembly).

SEQ ID NO: 161 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 160).

SEQ ID NO: 162 is an exemplary Cas13d protein sequence from contig emblOGDF01008514.11 (human gut 35 acid sequence (goes with SEQ ID NO: 200). metagenome assembly).

SEQ ID NO: 163 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 162).

SEQ ID NO: 164 is an exemplary Cas13d protein sequence from contig emblOGPN01002610.1 (human gut 40 metagenome assembly).

SEQ ID NO: 165 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 164).

SEQ ID NO: 166 is an exemplary Cas13d protein sequence from contig NFIR01000008.1 (Euhacterium sp. 45 An3, from chicken gut metagenome),

SEQ ID NO: 167 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 166).

SEQ ID NO: 168 is an exemplary Cas13d protein sequence from contig NFLV01000009.1 (Eubacterium sp, 50 An 11, from chicken gut metagenome).

SEQ ID NO: 169 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 168).

SEQ ID NOS: 171-174 are an exemplary Cas13d motif

SEQ ID NO: 175 is an exemplary Cas13d protein sequence from contig OJMM01002900 human gut metagenome sequence.

SEQ ID NO: 176 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 175).

SEQ ID NO: 177 is an exemplary Cas13d protein sequence from contig ODAI011611274.1 gut metagenome sequence.

SEQ ID NO: 178 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 177).

SEQ ID NO: 179 is an exemplary Cas13d protein sequence from contig OIZX01000427.1.

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SEQ ID NO: 180 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 179).

SEQ ID NO: 181 is an exemplary Cas13d protein sequence from contig emblOCVV012889144.1.

SEO ID NO: 182 is an exemplary consensus DR nucleic acid sequence (goes with SEO ID NO: 181).

SEO ID NO: 183 is an exemplary Cas13d protein sequence from contig OCTW011587266.1

SEQ ID NO: 184 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 183).

SEQ ID NO: 185 is an exemplary Cas13d protein sequence from contig emblOGNF01009141.1.

SEQ ID NO: 186 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 185).

SEQ ID NO: 187 is an exemplary Cas13d protein sequence from contig emblOIEN01002196.1.

SEQ ID NO: 188 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 187).

SEQ ID NO: 189 is an exemplary Cas13d protein sequence from contig e-k87_11092736.

SEQ ID NO: 190-193 are exemplary consensus DR nucleic acid sequences (goes with SEQ ID NO: 189).

SEQ ID NO: 194 is an exemplary Cas13d sequence from Gut metagenome contig6893000291.

SEQ ID NO: 195-197 are exemplary Cas13d motif sequences.

SEQ ID NO: 198 is an exemplary Cas13d protein sequence from Ga0224415 10007274.

SEQ ID NO: 199 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 198).

SEQ ID NO: 200 is an exemplary Cas13d protein sequence from EMG_10003641.

SEQ ID NO: 201 is an exemplary consensus DR nucleic

SEQ ID NO: 202 is an exemplary Cas13d protein sequence from Ga0129306_1000735.

SEQ ID NO: 203 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 202).

SEQ ID NO: 204 is an exemplary Cas13d protein sequence from Ga0129317_1008067.

SEQ ID NO: 205 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 204).

SEQ ID NO: 206 is an exemplary Cas13d protein sequence from Ga0224415_10048792.

SEQ ID NO: 207 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 206).

SEQ ID NO: 208 is an exemplary Cas13d protein sequence from 160582958_gene49834.

SEQ ID NO: 209 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 208).

SEQ ID NO: 210 is an exemplary Cas13d protein sequence from 250twins_35838_GL0110300.

SEQ ID NO: 211 is an exemplary consensus DR nucleic 55 acid sequence (goes with SEQ ID NO: 210).

SEQ ID NO: 212 is an exemplary Cas13d protein sequence from 250twins_36050_GL0158985.

SEQ ID NO: 213 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 212).

SEQ ID NO: 214 is an exemplary Cas13d protein sequence from 31009 GL0034153.

SEQ ID NO: 215 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 214).

SEQ ID NO: 216 is an exemplary Cas13d protein 65 sequence from 530373_GL0023589.

SEQ ID NO: 217 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 216).

SEQ ID NO: 218 is an exemplary Cas13d protein sequence from BMZ-11B_GL0037771.

SEQ ID NO: 219 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 218).

SEQ ID NO: 220 is an exemplary Cas13d protein 5 sequence from BMZ-11B GL0037915.

SEO ID NO: 221 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 220).

SEQ ID NO: 222 is an exemplary Cas13d protein sequence from BMZ-11B_GL0069617.

SEQ ID NO: 223 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 222).

SEQ ID NO: 224 is an exemplary Cas13d protein sequence from -DLF014_GL0011914.

SEQ ID NO: 225 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 224).

SEQ ID NO: 226 is an exemplary Cas13d protein sequence from EYZ-362B_GL0088915.

SEQ ID NO: 227-228 are exemplary consensus DR 20 sp. nCas1 array targeting ccdB. nucleic acid sequences (goes with SEQ ID NO: 226).

SEQ ID NO: 229 is an exemplary Cas13d protein sequence from Ga0099364_10024192.

SEQ ID NO: 230 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 229).

SEQ ID NO: 231 is an exemplary Cas13d protein sequence from Ga0187910_10006931.

SEQ ID NO: 232 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 231).

SEQ ID NO: 233 is an exemplary Cas13d protein 30 sequence from Ga0187910_10015336.

SEQ ID NO: 234 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 233).

SEQ ID NO: 235 is an exemplary Cas13d protein sequence from Ga0187910_10040531.

SEQ ID NO: 236 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 235).

SEQ ID NO: 237 is an exemplary Cas13d protein sequence from Ga0187911_10069260.

SEQ ID NO: 238 is an exemplary consensus DR nucleic 40 acid sequence (goes with SEQ ID NO: 237).

SEQ ID NO: 239 is an exemplary Cas13d protein sequence from MH0288_GL0082219.

SEQ ID NO: 240 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 239).

SEQ ID NO: 241 is an exemplary Cas13d protein sequence from 02.UC29-0 GL0096317.

SEQ ID NO: 242 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 241).

sequence from PIG-014_GL0226364.

SEQ ID NO: 244 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 243).

SEQ ID NO: 245 is an exemplary Cas13d protein sequence from PIG-018_GL0023397.

SEQ ID NO: 246 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 245).

SEQ ID NO: 247 is an exemplary Cas13d protein sequence from PIG-025_GL0099734.

SEQ ID NO: 248 is an exemplary consensus DR nucleic 60 acid sequence (goes with SEQ ID NO: 247).

SEQ ID NO: 249 is an exemplary Cas13d protein sequence from PIG-028_GL0185479.

SEQ ID NO: 250 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 249).

SEQ ID NO: 251 is an exemplary Cas13d protein sequence from -Ga0224422_10645759.

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SEQ ID NO: 252 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 251).

SEQ ID NO: 253 is an exemplary Cas13d protein sequence from ODAI chimera.

SEQ ID NO: 254 is an exemplary consensus DR nucleic acid sequence (goes with SEQ ID NO: 253).

SEQ ID NO: 255 is an HEPN motif.

SEQ ID NOs: 256 and 257 are exemplary Cas13d nuclear localization signal amino acid and nucleic acid sequences, 10 respectively.

SEQ ID NOs: 258 and 260 are exemplary SV40 large T antigen nuclear localization signal amino acid and nucleic acid sequences, respectively.

SEQ ID NO: 259 is a dCas9 target sequence.

SEQ ID NO: 261 is an artificial Eubacterium siraeum nCas1 array targeting ccdB.

SEQ ID NO: 262 is a full 36 nt direct repeat.

SEQ ID NOs: 263-266 are spacer sequences.

SEQ ID NO: 267 is an artificial uncultured Ruminoccus

SEQ ID NO: 268 is a full 36 nt direct repeat.

SEQ ID NOs: 269-272 are spacer sequences.

SEQ ID NO: 273 is a ccdB target RNA sequence.

SEQ ID NOs: 274-277 are spacer sequences.

SEQ ID NO: 278 is a gRNA sequence.

SEQ ID NO: 279 is a consensus sequence of HEPN1 domain region.

SEQ ID NO: 280-286 are HEPN1 domain regions from seven Cas13d proteins.

SEQ ID NO: 287 is a consensus sequence of HEPN2 domain region.

SEQ ID NO: 288-294 are HEPN2 domain regions from seven Cas13d proteins.

SEQ ID NO: 295 is an exemplary RNA target sequence. SEQ ID NO: 296-309 are exemplary gRNA sequences with various truncations.

SEQ ID NO: 310 is an exemplary Cas13d protein sequence from 037_-emblOIZA01000315.1.

SEQ ID NO: 311 is an exemplary Cas13d protein sequence from PIG-022 GL0026351.

SEQ ID NO: 312 is an exemplary Cas13d protein sequence from PIG-046_GL0077813.

SEQ ID NO: 313 is an exemplary Cas13d protein sequence from pig_chimera.

DETAILED DESCRIPTION

Unless otherwise noted, technical terms are used according to conventional usage. Definitions of common terms in SEQ ID NO: 243 is an exemplary Cas13d protein 50 molecular biology can be found in Benjamin Lewin, Genes VII, published by Oxford University Press, 1999; Kendrew et al. (eds.), The Encyclopedia of Molecular Biology, published by Blackwell Science Ltd., 1994; and Robert A. Meyers (ed.), Molecular Biology and Biotechnology: a Comprehensive Desk Reference, published by VCH Publishers, Inc., 1995; and other similar references.

> As used herein, the singular forms "a," "an," and "the," refer to both the singular as well as plural, unless the context clearly indicates otherwise. As used herein, the term "comprises" means "includes." Thus, "comprising a nucleic acid molecule" means "including a nucleic acid molecule" without excluding other elements. It is further to be understood that any and all base sizes given for nucleic acids are approximate, and are provided for descriptive purposes, unless otherwise indicated. Although many methods and materials similar or equivalent to those described herein can be used, particular suitable methods and materials are

described below. In case of conflict, the present specification, including explanations of terms, will control. In addition, the materials, methods, and examples are illustrative only and not intended to be limiting. All references, including patent applications and patents, are herein incorporated 5 by reference in their entireties.

In order to facilitate review of the various embodiments of the disclosure, the following explanations of specific terms are provided:

I. Terms

Administration: To provide or give a subject an agent, such as an Cas13d protein (or Cas13d coding sequence) or guide molecule (or coding sequence) disclosed herein, by any effective route. Exemplary routes of administration include, but are not limited to, injection (such as subcuta- 15 neous, intramuscular, intradermal, intraperitoneal, intratumoral, and intravenous), transdermal, intranasal, and inhalation routes.

Cas 13d (also referred to as CasR, for CRISPR-associated RNase, and Cas13d): An RNA-guided RNA endonuclease 20 Motif 2: enzyme that can cut or bind RNA. Cas13d proteins include one or two HEPN domains (e.g., see SEQ ID NOS: 1-3, 42, 62, 70, 82, 83, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 25 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, and 253). Native HEPN domains include the sequence RXXXXH (SEQ ID NO: 255). Cas13d proteins that include 30 mutated HEPN domain(s), and thus cannot cut RNA, but can process guide RNA, are also encompassed by this disclosure (e.g., see SEQ ID NOS: 2 and 4). An alignment of native Cas13d proteins is shown in FIGS. 18A-18MMM. In addition, Cas13d proteins specifically recognize direct repeat 35 sequences of gRNA having a particular secondary structure (e.g see FIG. 3C). In one example, Cas13d proteins recognize and/or bind a DR having (1) a loop of about 4 to 8 nt, (2) a stem of 4 to 12 nt, stem formed of complementary due to a nt mismatch in the stem, and (3) a bulge or overhang formed of unpaired nt, which can be about 10 to 14 nt (e.g., 5 to 7 to on either side).

In one example, the full length (non-truncated) Cas13d protein is between 870-1080 amino acids long. In one 45 example, the Cas13d protein is derived from a genome sequence of a bacterium from the Order Clostridiales or a metagenomic sequence. In one example, the corresponding DR sequence of an Cas13d protein is located at the 5' end of the spacer sequence in the molecule that includes the Cas13d 50 gRNA. In one example, the DR sequence in the Cas13d gRNA is truncated at the 5' end relative to the DR sequence in the unprocessed Cas13d guide array transcript. In one example, the DR sequence in the Cas13d gRNA is truncated by 5-7 nt at the 5' end by the Cas13d protein. In one example, 55 tary" as used herein refers to a degree of complementarity the Cas13d protein can cut a target RNA flanked at the 3' end of the spacer-target duplex by any of a A, U, G or C ribonucleotide and flanked at the 5' end by any of a A, U, G or C ribonucleotide.

In one example, an Cas13d protein has at least 80%, at 60 least 85%, at least 90%, at least 91%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 65 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74,

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75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253

In one example, an Cas13d protein contains two HEPN Rnase domains which contain a RXXXXH amino acid motif (SEQ ID NO: 255; where X indicates any amino acid). In addition, an Cas13d protein can include one or more of the following amino acid motifs written in the common Prosite format:

```
Motif 1:
                                            (SEO ID NO: 195)
L-x(5)-[FWY]-x(3)-K-[NQS]-[ILM]-[ILMV]-x(2)-N-
x(2) - [FWY] - x(2) - [AG] - x(4) - [DE] - x - D
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(SEQ ID NO: 196) [FWY]-[ILV]-x(2)-[NQS]-[ILV]-x(2)-[DNST]-x(2)-F-x-Y-x(2)-[HKR]-[FHY]

Motif 3: (SEQ ID NO: 197) Y-[CDNSV]-x(2)-R-[FWY]-x-[ADNT]-[LM]-[ST]-x(4)-[FWY]

Thus, in some examples, an Cas13d protein having at least 80%, at least 85%, at least 90%, at least 91%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, nucleotides, which can include a small (e.g., 1 or 2 bp) bulge 40 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253, includes the motif of SEQ ID NO: 195, 196 or 197.

Complementarity: The ability of a nucleic acid to form hydrogen bond(s) with another nucleic acid sequence by either traditional Watson-Crick base pairing or other nontraditional types. A percent complementarity indicates the percentage of residues in a nucleic acid molecule which can form hydrogen bonds (e.g., Watson-Crick base pairing) with a second nucleic acid sequence (e.g., 5, 6, 7, 8, 9, 10 out of 10 being 50%, 60%, 70%, 80%, 90%, and 100% complementary). "Perfectly complementary" means that all the contiguous residues of a nucleic acid sequence will hydrogen bond with the same number of contiguous residues in a second nucleic acid sequence. "Substantially complementhat is at least 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 97%, 98%, 99%, or 100% over a region of 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 30, 35, 40, 45, 50, or more nucleotides, or refers to two nucleic acids that hybridize under stringent conditions.

CRISPRs (Clustered Regularly InterSpaced Repeats): The CRISPR RNA array is a defining feature of CRISPR systems. The term "CRISPR" refers to the architecture of the array which includes constant direct repeats (DRs) interspaced with the variable spacers. In some examples, a CRISPR array includes at least a DR-spacer-DR-spacer (see FIG. 1A). This feature was used in the disclosed computa-

tional pipeline to identify the novel Cas13d protein family (FIG. 1A). In bacteria, the array is transcribed as one single transcript (containing multiple crRNA units), which is then processed by the Cas13d protein and other RNases into individual crRNAs. CRISPRs are found in approximately 5 40% of sequenced bacteria genomes and 90% of sequenced archaea. CRISPRs are often associated with cas genes that code for proteins related to CRISPRs (such as the Cas13d proteins provided herein). The disclosed CRISPR/Cas system can be used for RNA targeting, for example to detect a 10 target RNA, modify a target RNA at any desired location, or cut the target RNA at any desired location.

Downregulated or knocked down: When used in reference to the expression of a molecule, such as a target RNA, refers to any process which results in a decrease in production of 15 the target RNA, but in some examples not complete elimination of the target RNA product or target RNA function. In one example, downregulation or knock down does not result in complete elimination of detectable target RNA expression or target RNA activity. In some examples, the target RNA is 20 a coding RNA. In some examples, the target RNA is non-coding RNA. Specific examples of RNA molecules that can be targeted for downregulation include mRNA, miRNA, rRNA, tRNA, nuclear RNA, lincRNA, circular RNA, and structural RNA. In some examples, downregulation or 25 knock down of a target RNA includes processes that decrease translation of the target RNA and thus can decrease the presence of corresponding proteins. The disclosed CRISPR/Cas system can be used to downregulate any target

Downregulation or knock down includes any detectable decrease in the target RNA. In certain examples, detectable target RNA in a cell or cell free system decreases by at least 10%, at least 20%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 35 90%, at least 95%, at least 96%, at least 97%, at least 98%, or at least 99% (such as a decrease of 40% to 90%, 40% to 80% or 50% to 95%) as compared to a control (such an amount of target RNA detected in a corresponding normal cell or sample). In one example, a control is a relative 40 amount of expression in a normal cell (e.g., a non-recombinant cell that does not include Cas13d or guide RNA).

Effective amount: The amount of an agent (such as the CRISPR/Cas agents provided herein) that is sufficient to effect beneficial or desired results.

A therapeutically effective amount may vary depending upon one or more of: the subject and disease condition being treated, the weight and age of the subject, the severity of the disease condition, the manner of administration and the like, which can readily be determined by one of ordinary skill in 50 the art. The beneficial therapeutic effect can include enablement of diagnostic determinations; amelioration of a disease, symptom, disorder, or pathological condition; reducing or preventing the onset of a disease, symptom, disorder or condition; and generally counteracting a disease, symptom, 55 disorder or pathological condition. In one embodiment, an "effective amount" is an amount sufficient to reduce symptoms of a disease, for example by at least 10%, at least 20%, at least 50%, at least 70%, or at least 90% (as compared to no administration of the therapeutic agent).

The term also applies to a dose that will allow for expression of an Cas13d and/or gRNA herein, and that allows for targeting (e.g., detection or modification) of a target RNA.

Increase or Decrease: A statistically significant positive or 65 negative change, respectively, in quantity from a control value. An increase is a positive change, such as an increase

at least 50%, at least 100%, at least 200%, at least 300%, at least 400% or at least 500% as compared to the control value. A decrease is a negative change, such as a decrease of at least 20%, at least 25%, at least 50%, at least 75%, at least 80%, at least 90%, at least 95%, at least 98%, at least 99%, or at least 100% decrease as compared to a control value. In some examples the decrease is less than 100%, such as a decrease of no more than 90%, no more than 95% or no more than 99%.

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Isolated: An "isolated" biological component (such as an Cas13d protein or nucleic acid, gRNA, or cell containing such) has been substantially separated, produced apart from, or purified away from other biological components in the cell or tissue of an organism in which the component occurs, such as other cells, chromosomal and extrachromosomal DNA and RNA, and proteins. Nucleic acids and proteins that have been "isolated" include nucleic acids and proteins purified by standard purification methods. The term also embraces nucleic acids and proteins prepared by recombinant expression in a host cell as well as chemically synthesized nucleic acids and proteins. Isolated Cas13d proteins or nucleic acids, or cells containing such, in some examples are at least 50% pure, such as at least 75%, at least 80%, at least 90%, at least 95%, at least 98%, or at least 100% pure.

Label: A compound or composition that is conjugated directly or indirectly to another molecule (such as a nucleic acid molecule) to facilitate detection of that molecule. Specific, non-limiting examples of labels include fluorescent and fluorogenic moieties, chromogenic moieties, haptens, affinity tags, and radioactive isotopes. The label can be directly detectable (e.g., optically detectable) or indirectly detectable (for example, via interaction with one or more additional molecules that are in turn detectable).

Modulate: A change in the content of RNA. Modulation can include, but is not limited to, RNA activation (e.g., upregulation), RNA repression (e.g., downregulation), ribonucleotide deletion, ribonucleotide insertion, ribonucleotide chemical modification, ribonucleotide covalent or non-covalent linkage, and/or ribonucleotide substitution.

Non-naturally occurring or engineered: Terms used herein as interchangeably and indicate the involvement of the hand of man The terms, when referring to nucleic acid molecules or polypeptides indicate that the nucleic acid molecule or the polypeptide is at least substantially free from at least one other component with which they are naturally associated in nature and as found in nature. In addition, the terms can indicate that the nucleic acid molecules or polypeptides have a sequence not found in nature.

Operably linked: A first nucleic acid sequence is operably linked with a second nucleic acid sequence when the first nucleic acid sequence is placed in a functional relationship with the second nucleic acid sequence. For instance, a promoter is operably linked to a coding sequence (such as a coding sequence of an Cas13d protein) if the promoter affects the transcription or expression of the coding sequence. Generally, operably linked DNA sequences are contiguous and, where necessary to join two protein-coding regions, in the same reading frame.

Pharmaceutically acceptable carriers: The pharmaceutically acceptable carriers useful in this invention are conventional. *Remington's Pharmaceutical Sciences*, by E. W. Martin, Mack Publishing Co., Easton, Pa., 15th Edition (1975), describes compositions and formulations suitable for pharmaceutical delivery of an Cas13d protein or nucleic acid molecule (or other molecules needed for modifying RNA using the disclosed CRISPR/Cas system with the disclosed Cas13d proteins).

In general, the nature of the carrier will depend on the particular mode of administration being employed. For instance, parenteral formulations usually comprise injectable fluids that include pharmaceutically and physiologically acceptable fluids such as water, physiological saline, balanced salt solutions, aqueous dextrose, glycerol or the like as a vehicle. In addition to biologically-neutral carriers, pharmaceutical compositions to be administered can contain minor amounts of non-toxic auxiliary substances, such as wetting or emulsifying agents, preservatives, and pH buffering agents and the like, for example sodium acetate or sorbitan monolaurate.

Polypeptide, peptide and protein: Refer to polymers of amino acids of any length. The polymer may be linear or branched, it may comprise modified amino acids, and it may be interrupted by non-amino acids. The terms also encompass an amino acid polymer that has been modified; for example, disulfide bond formation, glycosylation, lipidation, acetylation, phosphorylation, or any other manipulation, such as conjugation with a labeling component. As used herein the term "amino acid" includes natural and/or unnatural or synthetic amino acids, including glycine and both the D or L optical isomers, and amino acid analogs and peptidomimetics.

Promoter: An array of nucleic acid control sequences which direct transcription of a nucleic acid. A promoter includes necessary nucleic acid sequences near the start site of transcription. A promoter also optionally includes distal enhancer or repressor elements. A "constitutive promoter" is a promoter that is continuously active and is not subject to regulation by external signals or molecules. In contrast, the activity of an "inducible promoter" is regulated by an external signal or molecule (for example, a transcription factor).

Recombinant or host cell: A cell that has been genetically altered, or is capable of being genetically altered by introduction of an exogenous polynucleotide, such as a recombinant plasmid or vector. Typically, a host cell is a cell in which a vector can be propagated and its nucleic acid 40 expressed. Such cells can be eukaryotic or prokaryotic. The term also includes any progeny of the subject host cell. It is understood that all progeny may not be identical to the parental cell since there may be mutations that occur during replication. However, such progeny are included when the 45 term "host cell" is used.

Regulatory element: A phrase that includes promoters. enhancers, internal ribosomal entry sites (IRES), and other expression control elements (e.g., transcription termination signals, such as polyadenylation signals and poly-U 50 sequences). Such regulatory elements are described, for example, in Goeddel, GENE EXPRESSION TECHNOL-OGY: METHODS IN ENZYMOLOGY 185, Academic Press, San Diego, Calif. (1990) which is hereby incorporated by reference in its entirety. Regulatory elements include 55 those that direct constitutive expression of a nucleotide sequence in many types of host cells and those that direct expression of the nucleotide sequence only in certain host cells (e.g., tissue-specific regulatory sequences). A tissuespecific promoter may direct expression primarily in a 60 desired tissue of interest, such as muscle, neuron, bone, skin, blood, specific organs (e.g., liver, pancreas), or particular cell types (e.g., lymphocytes). Regulatory elements may also direct expression in a temporal-dependent manner, such as in a cell-cycle dependent or developmental stage-dependent 65 manner, which may or may not also be tissue or cell-type specific.

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In some embodiments, a vector provided herein includes a pol III promoter (e.g., U6 and H1 promoters), a pol II promoter (e.g., the retroviral Rous sarcoma virus (RSV) LTR promoter (optionally with the RSV enhancer), the cytomegalovirus (CMV) promoter (optionally with the CMV enhancer), the SV40 promoter, the dihydrofolate reductase promoter, the β -actin promoter, the phosphoglycerol kinase (PGK) promoter, and the EFIa promoter), or both.

Also encompassed by the term "regulatory element" are enhancer elements, such as WPRE; CMV enhancers; the R-U5' segment in LTR of HTLV-I; SV40 enhancer; and the intron sequence between exons 2 and 3 of rabbit β -globin.

RNA Editing: A type of genetic engineering in which a RNA molecule (or ribonucleotides of the RNA) is inserted, deleted or replaced in the genome of an organism using engineered nucleases (such as the Cas13d proteins provided herein), which create site-specific strand breaks at desired locations in the RNA. The induced breaks are repaired resulting in targeted mutations or repairs. The CRISPR/Cas methods disclosed herein, such as those that use an Cas13d, can be used to edit the sequence of one or more target RNAs, such as one associated with cancer (e.g., breast cancer, colon cancer, melanoma), infectious disease (such as HIV, hepatitis, HPV, and West Nile virus), or neurodegenerative disorder (e.g., Huntington's disease or ALS). For example, RNA editing can be used to treat a disease or viral infection.

RNA insertion site: A site of the RNA that is targeted for, or has undergone, insertion of an exogenous polynucleotide. The disclosed methods include use of a disclosed Cas13d protein, which can be used to target a RNA for manipulation at a RNA insertion site.

Sequence identity/similarity: The similarity between amino acid (or nucleotide) sequences is expressed in terms of the similarity between the sequences, otherwise referred to as sequence identity. Sequence identity is frequently measured in terms of percentage identity (or similarity or homology); the higher the percentage, the more similar the two sequences are.

Methods of alignment of sequences for comparison are well known in the art. Various programs and alignment algorithms are described in: Smith and Waterman, Adv. Appl. Math. 2:482, 1981; Needleman and Wunsch, J. Mol. Biol. 48:443, 1970; Pearson and Lipman, Proc. Natl. Acad. Sci. U.S.A. 85:2444, 1988; Higgins and Sharp, Gene 73:237, 1988; Higgins and Sharp, CABIOS 5:151, 1989; Corpet et al., Nucleic Acids Research 16:10881, 1988; and Pearson and Lipman, Proc. Natl. Acad. Sci. U.S.A. 85:2444, 1988. Altschul et al., Nature Genet. 6:119, 1994, presents a detailed consideration of sequence alignment methods and homology calculations.

The NCBI Basic Local Alignment Search Tool (BLAST) (Altschul et al., *J. Mol. Biol.* 215:403, 1990) is available from several sources, including the National Center for Biotechnology Information (NCBI, Bethesda, MD) and on the internet, for use in connection with the sequence analysis programs blastp, blastn, blastx, tblastn and tblastx. A description of how to determine sequence identity using this program is available on the NCBI website on the internet.

Variants of protein and nucleic acid sequences known in the art and disclosed herein are typically characterized by possession of at least about 80%, at least 90%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% sequence identity counted over the full length alignment with the amino acid sequence using the NCBI Blast 2.0, gapped blastp set to default parameters. For comparisons of amino acid sequences of greater than about 30 amino acids,

the Blast 2 sequences function is employed using the default BLOSUM62 matrix set to default parameters, (gap existence cost of 11, and a per residue gap cost of 1). When aligning short peptides (fewer than around 30 amino acids), the alignment should be performed using the Blast 2 sequences function, employing the PAM30 matrix set to default parameters (open gap 9, extension gap 1 penalties). Proteins with even greater similarity to the reference sequences will show increasing percentage identities when assessed by this method, such as at least 95%, at least 98%, or at least 99% sequence identity. When less than the entire sequence is being compared for sequence identity, homologs and variants will typically possess at least 80% sequence identity over short windows of 10-20 amino acids, and may possess sequence identities of at least 85% or at least 90% or at least 95% depending on their similarity to the reference sequence. Methods for determining sequence identity over such short windows are available at the NCBI website on the internet. One of skill in the art will appreciate that these sequence 20 identity ranges are provided for guidance only; it is entirely possible that strongly significant homologs could be obtained that fall outside of the ranges provided.

Thus, in one example, an Cas13d protein has at least 80%, at least 85%, at least 90%, at least 91%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98% or at least 99% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 19, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253.

Subject: A vertebrate, such as a mammal, for example a 40 human. Mammals include, but are not limited to, murines, simians, humans, farm animals, sport animals, and pets. In one embodiment, the subject is a non-human mammalian subject, such as a monkey or other non-human primate, mouse, rat, rabbit, pig, goat, sheep, dog, cat, horse, or cow. 45 In some examples, the subject has a disorder (e.g., viral infection) or genetic disease that can be treated using methods provided herein. In some examples, the subject has a disorder (e.g., viral infection) or genetic disease that can be diagnosed using methods provided herein. In some 50 examples, the subject is a laboratory animal/organism, such as a zebrafish, *Xenopus*, *C. elegans*, *Drosophila*, mouse, rabbit, or rat. Tissues, cells and their progeny of a biological entity obtained in vivo or cultured in vitro are also encompassed.

Therapeutic agent: Refers to one or more molecules or compounds that confer some beneficial effect upon administration to a subject. The beneficial therapeutic effect can include enablement of diagnostic determinations; amelioration of a disease, symptom, disorder, or pathological condition; reducing or preventing the onset of a disease, symptom, disorder or condition; and generally counteracting a disease, symptom, disorder or pathological condition.

Transduced, Transformed and Transfected: A virus or vector "transduces" a cell when it transfers nucleic acid 65 molecules into a cell. A cell is "transformed" or "transfected" by a nucleic acid transduced into the cell when the

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nucleic acid becomes stably replicated by the cell, either by incorporation of the nucleic acid into the cellular genome, or by episomal replication.

These terms encompasse all techniques by which a nucleic acid molecule can be introduced into such a cell, including transfection with viral vectors, transformation with plasmid vectors, and introduction of naked DNA by electroporation, lipofection, particle gun acceleration and other methods in the art. In some example the method is a chemical method (e.g., calcium-phosphate transfection), physical method (e.g., electroporation, microinjection, particle bombardment), fusion (e.g., liposomes), receptor-mediated endocytosis (e.g., DNA-protein complexes, viral envelope/capsid-DNA complexes) and biological infection by viruses such as recombinant viruses (Wolff, J. A., ed, Gene Therapeutics, Birkhauser, Boston, USA, 1994). Methods for the introduction of nucleic acid molecules into cells are known (e.g., see U.S. Pat. No. 6,110,743). These methods can be used to transduce a cell with the disclosed agents to manipulate its genome.

Transgene: An exogenous gene.

Treating, Treatment, and Therapy: Any success or indicia of success in the attenuation or amelioration of an injury, pathology or condition, including any objective or subjective parameter such as abatement, remission, diminishing of symptoms or making the condition more tolerable to the patient, slowing in the rate of degeneration or decline, making the final point of degeneration less debilitating, improving a subject's physical or mental well-being, or prolonging the length of survival. The treatment may be assessed by objective or subjective parameters; including the results of a physical examination, blood and other clinical tests, and the like. For prophylactic benefit, the disclosed compositions may be administered to a subject at risk of developing a particular disease, condition, or symptom, or to a subject reporting one or more of the physiological symptoms of a disease, even though the disease, condition, or symptom may not have yet been manifested.

Upregulated: When used in reference to the expression of a molecule, such as a target RNA, refers to any process which results in an increase in production of the target RNA. In one example, includes direct upregulation, for example if the target RNA participates in a feedback loop with its own transcription. In one example, includes indirect upregulation, such as by knockdown of an inhibitory miRNA that leads to the activation of a target of that miRNA.

In some examples, the target RNA is a coding RNA. In some examples, the target RNA is non-coding RNA. Specific examples of RNA molecules that can be targeted for upregulation include mRNA, miRNA, rRNA, tRNA, nuclear RNA, and structural RNA. In some examples, upregulation or activation of a target RNA includes processes that increase translation of the target RNA and thus can increase the presence of corresponding proteins. The disclosed CRISPR/Cas system can be used to upregulate any target RNA of interest.

Upregulation includes any detectable increase in target RNA. In certain examples, detectable target RNA expression in a cell or cell free system (such as a cell expressing an Cas13d protein and gRNA disclosed herein) increases by at least 20%, at least 30%, at least 40%, at least 50%, at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, at least 95%, at least 100%, at least 200%, at least 400%, or at least 500% as compared to a control(such an amount of target RNA detected in a corresponding normal cell or sample). In one example, a control is a relative amount of

expression in a normal cell (e.g., a non-recombinant cell that does not include Cas13d or guide RNA).

Under conditions sufficient for: A phrase that is used to describe any environment that permits a desired activity. In one example the desired activity is expression of an Cas13d 5 protein disclosed herein, in combination with other necessary elements, for example to modulate a target RNA.

Vector: A nucleic acid molecule into which a foreign nucleic acid molecule can be introduced without disrupting the ability of the vector to replicate and/or integrate in a host 10 cell. Vectors include, but are not limited to, nucleic acid molecules that are single-stranded, double-stranded, or partially double-stranded; nucleic acid molecules that comprise one or more free ends, no free ends (e.g., circular); nucleic acid molecules that comprise DNA, RNA, or both; and other 15 varieties of polynucleotides known in the art.

A vector can include nucleic acid sequences that permit it to replicate in a host cell, such as an origin of replication. A vector can also include one or more selectable marker genes and other genetic elements known in the art. An integrating 20 vector is capable of integrating itself into a host nucleic acid. An expression vector is a vector that contains the necessary regulatory sequences to allow transcription and translation of inserted gene or genes.

One type of vector is a "plasmid," which refers to a 25 circular double stranded DNA loop into which additional DNA segments can be inserted, such as by standard molecular cloning techniques. Another type of vector is a viral vector, wherein virally-derived DNA or RNA sequences are present in the vector for packaging into a virus (e.g., 30 retroviruses, replication defective retroviruses, adenoviruses, replication defective adenoviruses, and adeno-associated viruses). Viral vectors also include polynucleotides carried by a virus for transfection into a host cell. In some embodiments, the vector is a lentivirus (such as an integration-deficient lentiviral vector) or adeno-associated viral (AAV) vector.

Certain vectors are capable of autonomous replication in a host cell into which they are introduced (e.g., bacterial vectors having a bacterial origin of replication and episomal 40 mammalian vectors). Other vectors (e.g., non-episomal mammalian vectors) are integrated into the genome of a host cell upon introduction into the host cell, and thereby are replicated along with the host genome.

Certain vectors are capable of directing the expression of 45 genes to which they are operatively-linked. Such vectors are referred to herein as "expression vectors." Common expression vectors are often in the form of plasmids. Recombinant expression vectors can comprise a nucleic acid provided herein (such as a guide RNA [which can be expressed from 50 an RNA sequence or a RNA sequence], nucleic acid encoding an Cas13d protein) in a form suitable for expression of the nucleic acid in a host cell, which means that the recombinant expression vectors include one or more regulatory elements, which may be selected on the basis of the 55 host cells to be used for expression, that is operativelylinked to the nucleic acid sequence to be expressed. Within a recombinant expression vector, "operably linked" is intended to mean that the nucleotide sequence of interest is linked to the regulatory element(s) in a manner that allows 60 for expression of the nucleotide sequence (e.g., in an in vitro transcription/translation system or in a host cell when the vector is introduced into the host cell). It will be appreciated by those skilled in the art that the design of the expression vector can depend on such factors as the choice of the host 65 cell to be transformed, the level of expression desired, etc. A vector can be introduced into host cells to thereby produce

transcripts, proteins, or peptides, including fusion proteins or peptides, encoded by nucleic acids as described herein (e.g., clustered regularly interspersed short palindromic repeats (CRISPR) transcripts, proteins, enzymes, mutant forms thereof, fusion proteins thereof, etc.).

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II. Overview of Several Embodiments

Class 2 CRISPR-Cas systems endow microbes with diverse mechanisms for adaptive immunity. Provided herein is an analysis of prokaryotic genome and metagenome sequences to identify an uncharacterized family of RNAguided, RNA-targeting CRISPR systems which is classified as Type VI-D. Biochemical characterization and protein engineering of seven distinct orthologs generated a ribonuclease effector derived from Ruminococcus flavefaciens XPD3002 (CasRx) with robust activity in human cells. CasRx-mediated knockdown exhibits high efficiency and specificity relative to RNA interference across diverse endogenous transcripts. As one of the most compact single effector Cas enzymes, CasRx can also be flexibly packaged into adeno-associated virus. Virally encoded, catalytically inactive CasRx can be targeted to cis-elements of premRNA to manipulate alternative splicing, alleviating dysregulated tau isoform ratios in a neuronal model of frontotemporal dementia. The results herein present CasRx as a programmable RNA-binding module for efficient targeting of cellular RNA, enabling a general platform for transcriptome engineering and therapeutic methods.

Class 2 CRISPR systems are found throughout diverse bacterial and archaeal life. Using a minimal definition of the CRISPR locus for bioinformatic mining of prokaryotic genome and metagenome sequences, which requires only a CRISPR repeat array and a nearby protein, provided herein is the identification of an uncharacterized, remarkably compact family of RNA-targeting class 2 CRISPR systems designated as Type VI CRISPR-Cas13d.

Because CRISPR systems generally exist as a functional operon within 20 kilobases of genome sequence, even fragmented metagenome reads may be sufficient to recover useful Cas enzymes for bioengineering purposes. CRISPR genome mining strategies described herein and by others (Shmakov et al., 2015), combined with ongoing efforts to profile microbial populations via next-generation sequencing, should contribute mechanistically diverse additions to the genome engineering toolbox.

Two distinct ribonuclease properties of the Cas13d effector, which processes a CRISPR repeat array into mature guides via a HEPN domain-independent mechanism followed by guide sequence-dependent recognition of a complementary activator RNA, were biochemically characterized. This triggers HEPN-mediated RNase activity, enabling Cas13d to cleave both activator and bystander RNAs, a property shared by other RNA-targeting CRISPR systems. Cas13d additionally exhibits no apparent flanking sequence requirements and was found to be active across crRNAs tiling a target RNA, suggesting the ability to target arbitrary single-stranded RNA sequences.

A comprehensive activity reporter screen in human cells of Cas13d orthologs sampled from distinct branches of the Cas13d family revealed that NLS fusions to Cas13d from *Ruminococcus flavefaciens* strain XPD3002 (CasRx) can be engineered for programmable RNA targeting in a eukaryotic context (FIG. 8D). CasRx fusions knocked down a diverse set of 14 endogenous mRNAs and lncRNAs, consistently achieving >90% knockdown with favorable efficiency relative to RNA interference, dCas9-mediated CRISPR interference, and other members of the Cas13 superfamily (FIGS. 10A-10C). Additionally, CasRx interference is mark-

edly more specific than spacer-matching shRNAs, with no detectable off-target changes compared with hundreds for RNA interference.

CasRx is a minimal two-component platform, including an engineered CRISPR-Cas13d effector and an associated 5 guide RNA, and can be fully genetically encoded. Because CasRx is an orthogonally delivered protein, HEPN-inactive dCasRx can be engineered as a flexible RNA-binding module to target specific RNA elements. Importantly, because CasRx uses a distinct ribonuclease activity to process guide 10 RNAs, dCasRx can still be paired with a repeat array for multiplexing applications. The utility of this concept is shown herein by creating a dCasRx splice effector fusion for tuning alternative splicing and resulting protein isoform ratios, applying it in a neuronal model of frontotemporal 15 dementia.

At an average size of 930 aa, Cas13d is the smallest class 2 CRISPR effector characterized in mammalian cells. This allows CasRx effector domain fusions to be paired with a CRISPR array encoding multiple guide RNAs while remain- 20 ing under the packaging size limit of the versatile adenoassociated virus (AAV) delivery vehicle (Naldini, 2015) for primary cell and in vivo delivery. Further, targeted AAV delivery of CasRx to specific postmitotic cell types such as neurons can mediate long-term expression of a corrective 25 payload that avoids permanent genetic modifications or frequent re-administration (Chiriboga et al., 2016), complementing other nucleic acid targeting technologies such as DNA nuclease editing or antisense oligonucleotides. RNA mis-splicing diseases have been estimated to account for up 30 to 15% of genetic diseases (Hammond and Wood, 2011), highlighting the potential for engineered splice effectors capable of multiplexed targeting. The materials provided herein can be used for RNA targeting for knockdown and splicing, such as live cell labeling and genetic screens to 35 detect a target RNA). transcript imaging, trafficking, or regulation. CRISPR-Cas13d and engineered variants such as CasRx collectively enable flexible nucleic acid engineering, transcriptome-related study, and therapeutics, expanding the genome editing toolbox beyond DNA to RNA.

Provided herein are methods of targeting (e.g., modifying, detecting) one or more target RNA molecules, such as a clustered regularly interspaced short palindromic repeats (CRISPR)-CRISPR associated (Cas) system-mediated RNA editing method. Such methods can include contacting one or 45 more target RNA molecules with a non-naturally occurring or engineered (e.g., does not naturally occur in the cell or system into which it is introduced) CRISPR-Cas system. Thus in some examples, the disclosed CRISPR-Cas system includes a naturally occurring Cas13d protein (or coding 50 sequence) and a naturally occurring gRNA, but is used in a system or cell in which the Cas1 protein (or coding sequence) and the gRNA are not naturally found. Furthermore, the spacer sequence within the gRNA molecule is not naturally occurring, and has been modified to be comple- 55 mentary to the target RNA molecule.

In some examples, a target RNA is a coding RNA. In some examples, the RNA is non-coding RNA.

The disclosed CRISPR-Cas system can include (1) at least one Cas13d protein or at least one Cas13d nucleic acid 60 coding sequence (such as a mRNA or a vector encoding the at least one Cas13d protein) and (2) at least one CRISPR-Cas system guide nucleic acid molecule (e.g., gRNA) (or at least one nucleic acid molecule encoding the gRNA) having sufficient complementary to a target RNA such that it can 65 hybridize to a target RNA molecule. The Cas13d protein forms a complex with the gRNA, and the gRNA directs the

complex to the one or more target RNA molecules. This targeting can allow the Cas13d-gRNA complex to modify or detect the one or more target RNA molecules. In some examples, the one or more target RNA molecules (or a cell containing the one or more target RNA molecules) are contacted with a complex comprising the at least one Cas13d protein and the at least one gRNA. In some examples, the system includes Mg²⁺. However, in some examples, the system does not require Mg²⁺, such as if

cleavage of the target RNA is not desired.

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In some examples, contacting the one or more target RNA molecules with the non-naturally occurring or engineered CRISPR-Cas system includes introducing into a cell (such as a eukaryotic or prokaryotic cell) containing the one or more target RNA molecules the non-naturally occurring or engineered CRISPR-Cas system, for example using endocytosis (e.g., receptor-mediated endocytosis, micropinocytosis), a liposome, a particle, an exosome, a microvesicle, a gene gun, electroporation, a virus, RNP-antibody fusion (e.g., by tethering an Cas13d RNP to an antibody, antibody fragment, or other targeting moiety [such as ScFv, aptamers, DARPins, nanobodies, affibodies, etc.], the RNP can be endocytosed into the cell, The RNP could conceivably be tethered to many things other than), or combinations thereof. Thus, cells can be transformed, transduced, transfected, or otherwise contacted with appropriate nucleic acid molecules of the disclosed CRISPR-Cas system. The resulting cells are recombinant cells. In some examples, contacting the one or more target RNA molecules with the non-naturally occurring or engineered CRISPR-Cas system includes contacting a cell-free system (such as a biological or environmental sample, or a cell lysate) containing the one or more target RNA molecules the non-naturally occurring or engineered CRISPR-Cas system (for example in a diagnostic method to

In some examples, at least 2, at least 3, at least 4, at least 5, at least 10, or at least 20 different gRNAs are used. For example, such a method could include targeting at least 2, at least 3, at least 4, at least 5, at least 10, or at least 20 different target RNA molecules, targeting at least 2, at least 3, at least 4, at least 5, at least 10, or at least 20 different regions of one or more RNA molecules, or combinations thereof.

Also provided are isolated nucleic acid molecules encoding such Cas13d proteins, such as a cDNA, genomic DNA, RNA, or mDNA. Such isolated nucleic acid molecules can be part of a vector (such as a plasmid or viral vector), and can be operably linked to a promoter. In some examples, an isolated nucleic acid molecule encoding a Cas13d protein has at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 124, 125, 126, 127, 128, 139, 140 or 141; or at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 114, 115, 116, 117, 118, 119, 120, 121, 122, 123, 142, 143, 144, or 145. In an additional example, an isolated nucleic acid encodes a Cas13d protein having at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104,

105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149,

153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253.

In some examples, an isolated nucleic acid molecule 5 encoding at least one Cas13d protein (which can be part of a vector) includes at least one Cas13d protein coding sequence that is codon optimized for expression in a eukaryotic cell, at least one Cas13d protein coding sequence codon optimized for expression in a human cell. In one example, 10 such an Cas13d coding sequence has at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 114, 115, 116, 117, 118, 119, 120, 121, 122, 123, 142, 143, 144, or 145, or has at least 80%, at 15 least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 124, 125, 126, 127, 128, 139, 140 or 141. In an additional example, a eukaryotic cell codon optimized nucleic acid sequence encodes a Cas13d 20 protein having at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 25 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 30 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253.

In some examples, the gRNA that hybridizes with the one or more target RNA molecules includes one or more direct repeat (DR) sequences, one or more spacer sequences, or one or more sequences comprising DR-spacer-DR-spacer. In 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 129, 130, 131, 132, 133, 134, 135, 136, 137, 148, 150, 151, 152, 154, 156, 157, 159, 161, 163, 165, 167, 169, 176, 178, 180, 182, 184, 186, 45 188, 190, 191, 192, 193, 199, 201, 203, 205, 207, 209, 211, 213, 215, 217, 219, 221, 223, 225, 227, 228, 230, 232, 234, 236, 238, 240, 242, 244, 246, 248, 250, 252, or 254. In one example, the gRNA includes additional sequences, such as an aptamer sequence.

In some examples, a plurality of gRNAs are processed from a single array transcript, wherein each gRNA can be different, for example to target different RNAs or target multiple regions of a single RNA.

In some examples, the DRs are truncated by 1-10 nucleo- 55 tides (such as 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides) at the 5'end, for example to be expressed as mature preprocessed guide RNAs.

Methods of targeting one or more target RNA molecules are provided. Targeting an RNA molecule can include one or 60 more of cutting or nicking one or more target RNA molecules, activating or upregulating one or more target RNA molecules, activating or suppressing translation the one or more target RNA molecules, deactivating the one or more target RNA molecules, visualizing, labeling, or detecting the one or more target RNA molecules, binding the one or more target RNA molecules, editing the one or more target RNA

molecules, trafficking the one or more target RNA molecules, and masking the one or more target RNA molecules. In some example, modifying one or more target RNA molecules includes one or more of an RNA base substitution, an RNA base deletion, an RNA base insertion, a break in the target RNA, methylating RNA, and demethylating

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In some examples, such methods are used to treat a disease, such as a disease in a human In such examples, the one or more target RNA molecules is associated with the disease

Also provided are isolated proteins, including non-naturally occurring proteins. in some examples, a protein has at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, $103,\,104,\,105,\,106,\,107,\,108,\,109,\,110,\,111,\,112,\,113,\,138,$ 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253. In some examples, an isolated protein is an Cas13d ortholog from a prokaryotic genome or metagenome, gut metagenome, an activated sludge metagenome, an anaerobic digester metagenome, a chicken gut metagenome, a human gut metagenome, a pig gut metagenome, a bovine gut metagenome, a sheep gut metagenome, a goat gut metagenome, a capybara gut metagenome, a primate gut metagenome, a termite gut metagenome, a fecal metagenome, a genome from the Order Clostridiales, or the Family Ruminococcaceae. In some examples, an Cas13d ortholog includes an Cas13d ortholog from Ruminococcus albus, Eubacterium siraeum, a Ruminococcus flavefaciens strain some examples, the one or more DR sequences have at least 40 XPD3002, Ruminococcus flavefaciens FD-1, uncultured Eubacterium sp TS28-c4095, uncultured Ruminococcus sp., Ruminococcus bicirculans, or Ruminococcus sp CAG57. Such proteins can include a subcellular localization signal. In some examples, such proteins include a mutation in at least one native HEPN domain

Also provided are isolated guide RNA (gRNA) molecules. In some examples, an isolated gRNA includes one or more direct repeat (DR) sequences, such as one having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 129, 130, 131, 132, 133, 134, 135, 136, 137, 148, 150, 151, 152, 154, 156, 157, 159, 161, 163, 165, 167, 169, 176, 178, 180, 182, 184, 186, 188, 190, 191, 192, 193, 199, 201, 203, 205, 207, 209, 211, 213, 215, 217, 219, 221, 223, 225, 227, 228, 230, 232, 234, 236, 238, 240, 242, 244, 246, 248, 250, 252, or 254. Such a gRNA can further include one or more spacer sequences specific for (e.g., is complementary to) the target RNA. Such guide RNAs can further be optionally truncated by 1-10 nucleotides (such as 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 nucleotides) at the 5'end of the DR, for example to generate pre-processed guide RNAs.

Also provided are ribonucleoprotein (RNP) complexes, which include one or more Cas13d proteins provided herein and one or more gRNAs provided herein.

Also provided are recombinant cells that include any Cas13d protein (or nucleic acid molecule encoding Cas13d),

any gRNA, any RNP complex, or any vector, provided herein. In one example, the cell is not a bacterial cell. In one example, the cell is a bacterial cell.

Also provided are compositions that include one or more of any Cas13d protein (or nucleic acid molecule encoding Cas13d), any gRNA or array, any RNP complex, any isolated nucleic acid molecule, any vector, or any cell, provided herein. Such compositions can include a pharmaceutically acceptable carrier.

Also provided are kits. Such kits can include one or more of any Cas13d protein (or nucleic acid molecule encoding Cas13d), any gRNA or array, any RNP complex, any isolated nucleic acid molecule, any vector, any cell, or any composition provided herein. Such reagents can be combined or in separate containers.

In some examples, a Cas13d protein is programmed toward its RNA target by combining the protein (or nucleic acid encoding the protein) with an engineered RNA guide (or nucleic acid encoding RNA guide) consisting of a full or 20 partial direct repeat sequence followed by a "spacer" sequence complementary to the RNA target(s) (or variations thereof, i.e. arrays (DR-spacer-DR-spacer-DR-spacer . . . etc.) or pre-guide RNAs (DR-spacer-DR). Cas13d Proteins can be catalytically inactivated and transformed into RNA 25 binding modules by mutating the conserved RNAse HEPN motif (RXXXXH). Exemplary Cas13d proteins and corresponding guides are provided herein (e.g., SEQ ID NOS: 147-170, 175-193 and SEQ ID NOS: 198-254).

A. Cas13d Proteins

Provided herein are novel Cas13d proteins, such as those as shown in in the sequence listing. SEQ ID NOS: 1, 3, 42, 62, 70, 82, 83, 92, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, and 253 provide distinct full length proteins, and SEQ ID NOS: 2, 4-41, 43-61, 63-69, 71-81, 84-91, and 93-113 and 194 provide Cas13d variants and fragments. 40 Such proteins can be used in the disclosed methods, compositions, and kits.

In some examples, an Cas13d protein includes one or more (such as 1 or 2) native HEPN domains. In some examples, an Cas13d protein includes one or more mutated 45 HEPN domains (such as mutant Cas13d protein can process the gRNA, but cannot modify the target RNA). In some examples, an Cas13d protein is no more than 150 kD, no more than 140 kD, no more than 130 kD, no more than 120 kD, such as about 90 to 120 kD, about 100 to 120 kD or 50 about 110 kD.

In addition to the Cas13d proteins provided in Table 1 and in Example 2, the disclosure encompasses Cas13d orthologs from a prokaryotic genome or metagenome, gut metagenome, an activated sludge metagenome, an anaerobic 55 digester metagenome, a chicken gut metagenome, a human gut metagenome, a pig gut metagenome, a bovine gut metagenome, a sheep gut metagenome, a goat gut metagenome, a capybara gut metagenome, a primate gut metagenome, a termite gut metagenome, a fecal metagenome, a 60 genome from the Order Clostridiales, or the Family Ruminococcaceae, such as an Cas13d ortholog from Ruminococcus albus, Eubacterium siraeum, a Ruminococcus flavefaciens strain XPD3002, Ruminococcus flavefaciens FD-1, uncultured Eubacterium sp TS28-c4095, uncultured Rumi- 65 nococcus sp., Ruminococcus bicirculans, 1or Ruminococcus sp CAG57.

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In some examples, an Cas13d protein is at least 800 aa, at least 900 aa, or at least 1000 aa, such as 800 to 1200 aa, 850 to 1050 aa, or 860-1040 aa.

1. Variant Cas13d Sequences

Cas13d proteins, including variants of the sequences provided herein (such as variants of SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) are encompassed within this disclosure. In some examples, Cas13d proteins provided herein can contain one or more mutations, such as a single insertion, a single deletion, a single substitution, or combinations thereof. In some examples, the Cas13d protein includes at least 1, at least 5, at least 10, at least 15, at least 20, at least 25, at least 30, at least 40, at least 50, at least 75, at least 100, at least 200 or at least 300 as insertions, such as 1-20 insertions (for example at the N- or C-terminus or within the protein, such as insertion of a whole small domain), at least 1, at least 5, at least 10, at least 15, at least 20, at least 25, at least 30, at least 40, at least 50, at least 75, at least 100, at least 200 or at least 300 aa deletions (such as deletion of a whole small domain), such as 1-20 deletions (for example at the N- or C-terminus or within the protein), at least 1, at least 5, at least 10, at least 15, at least 20, at least 25, at least 30 aa substitutions, such as 1-20 substitutions, or any combination thereof (e.g., single insertion together with 1-19 substitutions), but retain the ability to bind target RNA molecules complementary to the spacer sequence within the gRNA molecule and/or process an guide array RNA transcript into gRNA molecules and/or retain the ability to cleave target RNA. In some examples, the disclosure provides a variant of any disclosed Cas13d protein (such as SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) having 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29 or 30 amino acid changes, but retain the ability to bind target RNA molecules complementary to the spacer sequence within the gRNA molecule and/or process an guide array RNA transcript into gRNA molecules. In some examples, any disclosed Cas13d protein (such as SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175,

177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) further includes 1-8 amino acid insertions, 1-15 amino acid deletions, 1-10 amino acid substitutions, or any com- 5 bination thereof (e.g., 1-15, 1-4, or 1-5 amino acid deletions together with 1-10, 1-5 or 1-7 amino acid substitutions), with the retained ability to bind target RNA molecules complementary to the spacer sequence within the gRNA molecule and/or process an guide array RNA transcript into 10 gRNA molecules. In one example, such variant peptides are produced by manipulating the nucleotide sequence encoding a peptide using standard procedures such as site-directed mutagenesis or PCR. Such variants can also be chemically synthesized.

In some examples, an Cas13d protein includes a motif shown in SEQ ID NO: 195, 196 or 197. Thus, an Cas13d protein having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEO ID 20 NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 25 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 30 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253, in some examples includes at least one motif shown in SEQ ID NO: 195, 196 or 197.

One type of modification or mutation includes the substitution of amino acids for amino acid residues having a 35 similar biochemical property, that is, a conservative substitution (such as 1-4, 1-8, 1-10, or 1-20 conservative substitutions). Typically, conservative substitutions have little to no impact on the activity of a resulting peptide. For example, an Cas13d protein (such as SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 45 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 50 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) that does not substantially affect the ability of the Cas13d protein to bind target RNA molecules complementary to the spacer sequence within the gRNA molecule and/or process 55 an guide array RNA transcript into gRNA molecules. An alanine scan can be used to identify which amino acid residues in an Cas13d protein (such as SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, $22,\,23,\,24,\,25,\,26,\,27,\,28,\,29,\,30,\,31,\,32,\,33,\,34,\,35,\,36,\,37,\ \, 60$ 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 65 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202,

204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253), can tolerate an amino acid substitution. In one example, the ability of a variant Cas13d protein (such as 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) to modify gene expression in a CRISPR/ Cas system, is not altered by more than 25%, for example not more than 20%, for example not more than 10%, when an alanine, or other conservative amino acid, is substituted for 1-4, 1-8, 1-10, or 1-20 native amino acids. Examples of amino acids which may be substituted for an original amino acid in a protein and which are regarded as conservative substitutions include: Ser for Ala; Lys for Arg; Gln or His for Asn; Glu for Asp; Ser for Cys; Asn for Gln; Asp for Glu; Pro for Gly; Asn or Gln for His; Leu or Val for Ile; Ile or Val for Leu; Arg or Gln for Lys; Leu or Ile for Met; Met, Leu or Tyr for Phe; Thr for Ser; Ser for Thr; Tyr for Trp; Trp or Phe for Tyr; and Ile or Leu for Val.

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One method for identifying regions particularly amenable to insertions, substitutions or deletion is to target stretches of amino acids exhibiting low levels of conservation between orthologs. Such regions are indicated in the conservation graph of the alignment of Cas13d proteins provided in FIG. 1B. Conserved residues of Cas13d are further marked in the Cas13d protein alignment provided in FIG. 18A-18MMM (indicated by symbols "." ":" or "*" below aligned conserved residues). Examples of deletions and their functional testing are further provided in FIGS. 16A-16B.

Another type of substitution can be achieved by swapping a conservative substitution is an amino acid substitution in 40 out parts of one ortholog with the homologous region of another ortholog to obtain a combined "chimeric" protein. Such a chimeric protein may combine favorable properties of multiple Cas13d orthologs.

More substantial changes can be made by using substitutions that are less conservative, e.g., selecting residues that differ more significantly in their effect on maintaining: (a) the structure of the polypeptide backbone in the area of the substitution, for example, as a sheet or helical conformation; (b) the charge or hydrophobicity of the polypeptide at the target site; or (c) the bulk of the side chain. The substitutions that in general are expected to produce the greatest changes in polypeptide function are those in which: (a) a hydrophilic residue, e.g., serine or threonine, is substituted for (or by) a hydrophobic residue, e.g., leucine, isoleucine, phenylalanine, valine or alanine; (b) a cysteine or proline is substituted for (or by) any other residue; (c) a residue having an electropositive side chain, e.g., lysine, arginine, or histidine, is substituted for (or by) an electronegative residue, e.g., glutamic acid or aspartic acid; or (d) a residue having a bulky side chain, e.g., phenylalanine, is substituted for (or by) one not having a side chain, e.g., glycine.

Thus, the disclosure provides Cas13d proteins having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39,

40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 5 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253 or combinations (e.g., chimeras) thereof.

In one example, an Cas13d protein includes non-naturally occurring amino acids.

2. Cas13d proteins with Other Elements

An Cas13d protein (such as any of SEQ ID NOS: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 15 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 20 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, 25 or 253) can include other elements or domains, for example at the N- or C-terminus (or both). The resulting protein can be referred to as an Cas13d fusion protein. In one example, an Cas13d protein provided herein (such as a native Cas13d or an Cas13d with mutated HEPN domain(s)) includes a 30 subcellular localization signal. Exemplary subcellular localization signals include an organelle localization signal, such as a nuclear localization signal (NLS), nuclear export signal (NES), or mitochondrial localization signal. In one example, an Cas13d protein includes an NLS, such as 35 SPKKKRKVEAS (SEQ ID NO: 256; e.g., encoded by AGCCCCAAGAAgAAGAGaAAGGTGGAGGCCAGC, SEQ ID NO: 257) or GPKKKRKVAAA (SV40 large T antigen NLS, SEQ ID NO: 258; e.g., encoded by ggacctaagaaaaagaggaaggtggcggccgct, SEQ ID NO: 260). 40 Exemplary NES that can be part of an Cas13d protein include an adenovirus type 5 E1B nuclear export sequence, an HIV nuclear export sequence, a MAPK nuclear export sequence, or a PTK2 nuclear export sequence.

In some examples, the at least one Cas13d protein (such 45 as a native Cas13d or an Cas13d with mutated HEPN domain(s)) further includes one or more effector domains. Exemplary effector domains include proteins and/or enzymes, such that those can cleave RNA (e.g., a PIN endonuclease domain, an NYN domain, an SMR domain 50 from SOT1, or an RNase domain from Staphylococcal nuclease), those that can affect RNA stability (e.g., tristetraprolin (TTP) or domains from UPF1, EXOSC5, and STAU1), those that can edit a nucleotide or ribonucleotide (e.g., a cytidine deaminase, PPR protein, adenosine deami- 55 Cas13d or an Cas13d with mutated HEPN domain(s)) nase, ADAR family protein, or APOBEC family protein), those that can activate translation (e.g., eIF4E and other translation initiation factors, a domain of the yeast poly(A)binding protein or GLD2), those that can repress translation (e.g., Pumilio or FBF PUF proteins, deadenylases, CAF1, 60 Argonaute proteins), those that can methylate RNA (e.g., domains from m6A methyltransferase factors such as METTL14, METTL3, or WTAP), those that can demethylate RNA (e.g., human alkylation repair homolog 5 or Alkbh5), those that can affect splicing (e.g., the RS-rich 65 domain of SRSF1, the Gly-rich domain of hnRNP Al, the alanine-rich motif of RBM4, or the proline-rich motif of

DAZAP1), those that can enable affinity purification or immunoprecipitation (e.g., FLAG, HA, biotin, or HALO tags), and those that can enable proximity-based protein labeling and identification (e.g., a biotin ligase (such as BirA) or a peroxidase (such as APEX2) in order to biotinylate proteins that interact with the target RNA).

In some examples, the Cas13d protein and effector module combination can constitute a transcriptional sensor. For example, the transcriptional sensor can be comprised of at least one Cas13d protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4), at least one gRNA containing at least one spacer sequence specific for the target RNA, and an effector module such as an optionally split fluorescent protein or probe (e.g., a split Venus fluorescent protein, a split GFP, a split enhanced GFP, a split mCherry, a split super-folder mCherry, and other fluorescent protein variants such as ECFP, YFP, RFP, and derivatives or fragments thereof); an optionally split luminescent protein or probe (e.g. Gaussia, Firefly, NanoLuc, or Renilla variants); an optionally split enzyme (e.g., ubiquitin or TEV protease); a FRET-compatible protein pair; one or more transcription factor(s) fused to Cas13d via cleavable linkers (e.g., an artificial GAL4, zinc finger, transcriptional activator like effector (TALE), CRISPR-Cas9, CRISPR-Cpf1, or TetRbased transcription factor or an endogenous transcription factor); a split intein that trans-splices a protein to restore its function such as a transcription factor (e.g., an intein from Rhodothermus marinus or DnaE); a kinase-substrate pair that activates upon phosphorylation (e.g., TYK2-STAT3); one, two, or more monomers that activate upon dimerization or multimerization (e.g., caspase 9); or one or more proteins that induce conformational and functional change upon interaction. In one example, the spatial proximity of two or more Cas13d proteins and gRNAs due to binding a particular transcript would activate the effector module, resulting in a detectable signal or detectable activity in the cell.

In one example, the effector domain is fused to a protein that specifically recognizes and binds an RNA aptamer, such as one that can be appended to or inserted within a gRNA molecule (e.g., an MS2, PP7, Qβ, and other aptamers). This aptamer-effector domain fusion can be used to target the target RNA because the Cas13d and gRNA complex will guide the aptamer protein-effector domain in proximity to the target RNA.

In another example, the aptamer can be directly inserted into the gRNA molecule to permit detection of a target RNA, such as a fluorophore aptamer (e.g., Spinach, Mango, etc.).

In some examples, the Cas13d protein (such as a native Cas13d or an Cas13d with mutated HEPN domain(s)) includes a purification tag, such as an HA-tag, His-tag (such as 6-His), Myc-tag, E-tag, S-tag, calmodulin tag, FLAG-tag, GST-tag, MBP-tag, and the like. Such tags are in some examples at the N- or C-terminal end of the Cas13d protein.

In some examples, an Cas13d protein (such as a native includes one or more subcellular localization signals, effector domains, and purification tags.

In some examples, an Cas13d protein may be split into multiple fragments, which are then expressed individually. Such fragments of Cas13d may be optionally fused to other protein domains. In one example, an Cas13d can be split into two halves, which are then fused to two parts of an inducible heterodimer pair. Upon induction of heterodimer binding, the Cas13d halves are recruited to each other to form an active protein. Such a system would allow for the inducible control of Cas13d activity. Useful heterodimer pairs include two proteins that dimerize upon light illumination or through

administration of a small molecule compound, amongst others. Specific examples of heterodimer pair include but are not limited to: light inducible Magnets proteins, the light inducible iLID-SspB pair, the light inducible Cryptochrome2-CIB1 dimer and the small molecule inducible 5 FKBP protein. In another example of a split Cas13d design, two halved of the Cas13d protein may be fused to protein trans-splicing domains. Such a design would enable the separate expression of two halves which are reconstituted into a full length protein once expressed inside a cell. An 10 example of such transsplicing domains includes the Intein system.

One method for identifying regions particularly amenable to splitting of the protein, is to identify stretches of amino acids exhibiting low levels of conservation between orthologs. Such regions are indicated in the conservation graph of the alignment of Cas13d proteins provided in FIG.

18. Regions of conserved residues of Cas13d are further marked in the Cas13d protein alignment provided in FIG.

18. A-18MMM (indicated by symbols .: or * below aligned conserved residues).

at least 96%, at least 97%, at least 99% or a

3. Generation of Cas13d Proteins

In one example, the Cas13d protein is expressed in vitro, for example, in a prokaryotic cell (e.g., bacteria such as Lactobacillus, Lactococcus, Bacillus (such as B. subtilis), 25 Escherichia (such as E. coli), Salmonella typhimurium, and Clostridium), archea cell, plant or plant cell, fungal cell (e.g., Neurospora), yeast cell (e.g., Saccharomyces or Pichia (such as S. cerevisiae or P. pastoris), Kluvveromyces lactis), insect cell (e.g., SF9 cells), or mammalian cells (e.g., 293 30 cells, or immortalized mammalian myeloid and lymphoid cell lines). Once expressed, the Cas13d protein can be isolated and/or purified (e.g., using chromatography or immunological separation). In some examples, as tag on the Cas13d protein permits isolation of the protein from a 35 culture media. Exemplary procedures include ammonium sulfate precipitation, affinity columns, column chromatography, and the like (see, generally, R. Scopes, Protein Purification, Springer-Verlag, N.Y., 1982). Substantially pure compositions of at least about 90 to 95% homogeneity, 40 such as 98% to 99% homogeneity, can be used in the methods provided herein. For example, a purified preparation of an Cas13d protein can be used as an alternative to expressing the Cas13d protein from a nucleic acid molecule in the CRISPR/Cas system.

In addition to recombinant methods, Cas13d proteins disclosed herein can also be constructed in whole or in part using native chemical ligation and/or expressed protein ligation.

B. Nucleic Acid Molecules Encoding Cas13d Proteins Nucleic acid molecules encoding an Cas13d protein are encompassed by this disclosure. Nucleic acid molecules include DNA, genomic DNA, cDNA, mRNA, and RNA sequences which encode an Cas13d peptide. Such nucleic acid molecules can include naturally occurring or non- 55 naturally occurring nucleotides or ribonucleotides. Exemplary nucleic acid molecules that encode the novel Cas13d proteins shown in SEQ ID NOS: 1, 3, 42, 62, 70, 82, 83, 92 and 104, are shown in SEQ ID NOS: 124-128, 139, 140, and 141. Also provided are codon optimized nucleic acid mol- 60 ecules that encode the novel Cas13d proteins, for example those optimized for expression in a mammalian cells, such as a human cell (SEQ ID NOS: 114-123 and 142-145). For example, SEQ ID NOS: 114, 118, and 122 provide nucleic acid molecules optimized for expression in human cells. 65 SEQ ID NOS: 115, 119 and 123 provide nucleic acid molecules optimized for expression in human cells, and

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which encode for mutant HEPN sites. SEQ ID NOS: 116 and 120 provide nucleic acid molecules optimized for expression in human cells, and which includes an N-terminal nuclear localization (NLS) coding sequence (namely, SPKKKRKVEAS). SEQ ID NO: 117 and 121 provide nucleic acid molecules optimized for expression in human cells, and which include N-terminal and C-terminal NLS coding sequences (namely, SPKKKRKVEAS, SEQ ID NO: 256, and GPKKKRKVAAA SEQ ID NO: 258, respectively).

In one example, a nucleic acid sequence encodes an Cas13d protein having at least 60%, at least 70%, at least 75%, at least 80%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 99% or at least 99% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253. Such nucleic acid sequences can be generated based on the amino acid sequences provided herein, and the genetic code. In one example, an Cas13d nucleic acid sequence has at least 70%, at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 124, 125, 126, 127, 128, 139, 140 or 141. In one example, an Cas13d nucleic acid sequence is optimized for expression in mammalian cells, such as human cells, such as one having at least 70%, at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 114, 115, 116, 117, 118, 119, 120, 121, 122, 123, 142, 143, 144, or 145.

One of skill can readily construct a variety of clones containing functionally equivalent nucleic acids, such as nucleic acids which differ in sequence but which encode the same Cas13d protein sequence. Silent mutations in the coding sequence result from the degeneracy (i.e., redundancy) of the genetic code, whereby more than one codon can encode the same amino acid residue. Thus, for example, leucine can be encoded by CTT, CTC, CTA, CTG, TTA, or TTG; serine can be encoded by TCT, TCC, TCA, TCG, AGT, or AGC; asparagine can be encoded by AAT or AAC; aspartic acid can be encoded by GAT or GAC; cysteine can be encoded by TGT or TGC; alanine can be encoded by GCT, GCC, GCA, or GCG; glutamine can be encoded by CAA or CAG; tyrosine can be encoded by TAT or TAC; and isoleucine can be encoded by ATT, ATC, or ATA. Tables showing the standard genetic code can be found in various sources (see, for example, Stryer, 1988, Biochemistry, 3rd Edition, W.H. 5 Freeman and Co., NY).

Based on the genetic code, nucleic acid sequences coding for any Cas13d sequence can be generated. In some examples, such a sequence is optimized for expression in a host or target cell, such as a host cell used to express the Cas13d protein or a cell in which the disclosed methods are practice (such as in a mammalian cell, such as a human cell). Codon preferences and codon usage tables for a particular species can be used to engineer isolated nucleic acid molecules encoding an Cas13d (such as one encoding a protein having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4,

5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 5 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 10 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253 that takes advantage of the codon usage preferences of that particular species). For example, the Cas13d proteins disclosed herein can be designed to have codons that are preferentially used by a particular organism of interest. In 15 one example, an Cas13d nucleic acid sequence is optimized for expression in human cells, such as one having at least 70%, at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 98%, or at least 99% sequence identity to SEQ ID NO: 114, 115, 116, 117, 118, 119, 120, 121, 122, 20 123, 142, 143, 144, or 145.

A nucleic acid encoding an Cas13d protein (such as one encoding a protein having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to 25 SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 30 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 35 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) can be cloned or amplified by in vitro methods, such as the polymerase chain reaction (PCR), the ligase chain reaction (LCR), the transcriptionbased amplification system (TAS), the self-sustained 40 sequence replication system (3SR) and the Q β replicase amplification system (QB). In addition, nucleic acids encoding an Cas13d protein (such as one encoding a protein having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 45 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 50 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) can be prepared by cloning techniques. Examples of appropriate cloning and sequencing techniques, and instructions sufficient to direct persons of skill through cloning are 60 found in Sambrook et al. (ed.), Molecular Cloning: A Laboratory Manual 2nd ed., vol. 1-3, Cold Spring Harbor Laboratory Press, Cold Spring, Harbor, N.Y., 1989, and Ausubel et al., (1987) in "Current Protocols in Molecular Biology," John Wiley and Sons, New York, N.Y.

Nucleic acid sequences encoding an Cas13d protein (such as one encoding a protein having at least 80%, at least 85%,

at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) can be prepared by any suitable method including, for example, cloning of appropriate sequences or by direct chemical synthesis by methods such as the phosphotriester method of Narang et al., Meth. Enzymol. 68:90-99, 1979; the phosphodiester method of Brown et al., Meth. Enzymol. 68:109-151, 1979; the diethylphosphoramidite method of Beaucage et al., Tetra. Lett. 22:1859-1862, 1981; the solid phase phosphoramidite triester method described by Beaucage & Caruthers, Tetra. Letts. 22(20):1859-1862, 1981, for example, using an automated synthesizer as described in, for example, Needham-VanDevanter et al., Nucl. Acids Res. 12:6159-6168, 1984; and, the solid support method of U.S. Pat. No. 4,458,066. Chemical synthesis produces a single stranded oligonucleotide. This can be converted into double stranded DNA by hybridization with a complementary sequence, or by polymerization with a DNA polymerase using the single strand as a template. One of skill would recognize that while chemical synthesis of DNA is generally limited to sequences of about 100 bases, longer sequences may be obtained by the ligation of shorter sequences.

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In one example, an Cas13d protein (such as a protein having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) is prepared by inserting the cDNA which encodes the Cas13d protein into a plasmid or vector. The insertion can be made so that the Cas13d protein is read in frame so that the Cas13d protein is produced.

The Cas13d nucleic acid coding sequence (such as one 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 55 having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226,

229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) can be inserted into an expression vector including, but not limited to a plasmid, virus or other vehicle that can be manipulated to allow insertion or incorporation of sequences and can be expressed in either prokaryotes or 5 eukaryotes. Hosts can include microbial, yeast, insect, plant and mammalian organisms. The vector can encode a selectable marker, such as a thymidine kinase gene or antibiotic resistance gene.

Nucleic acid sequences encoding an Cas13d protein (such 10 as one encoding a protein having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 15 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 20 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) can be operatively 25 linked to expression control sequences. An expression control sequence operatively linked to an Cas13d coding sequence is ligated such that expression of the Cas13d protein coding sequence is achieved under conditions compatible with the expression control sequences. The expression control sequences include, but are not limited to appropriate promoters, enhancers, transcription terminators, a start codon (i.e., ATG) in front of an Cas13d proteinencoding gene, splicing signal for introns, maintenance of the correct reading frame of that gene to permit proper 35 translation of mRNA, and stop codons.

In one embodiment, vectors are used for expression in yeast such as S. cerevisiae, P. pastoris, or Kluyveromyces lactis. Exemplary promoters for use in yeast expression systems include but are not limited to: the constitutive 40 promoters plasma membrane H+-ATPase (PMA1), glyceraldehyde-3-phosphate dehydrogenase (GPD), phosphoglycerate kinase-1 (PGK1), alcohol dehydrogenase-1 (ADH1), and pleiotropic drug-resistant pump (PDR5). In addition, many inducible promoters are of use, such as GAL1-10 45 (induced by galactose), PHO5 (induced by low extracellular inorganic phosphate), and tandem heat shock HSE elements (induced by temperature elevation to 37° C.). Promoters that direct variable expression in response to a titratable inducer include the methionine-responsive MET3 and MET25 pro- 50 moters and copper-dependent CUP1 promoters. Any of these promoters may be cloned into multicopy (2μ) or single copy (CEN) plasmids to give an additional level of control in expression level. The plasmids can include nutritional markers (such as URA3, ADE3, HIS1, and others) for 55 selection in yeast and antibiotic resistance (AMP) for propagation in bacteria. Plasmids for expression on K. lactis are known, such as pKLAC1.

Viral vectors can also be prepared that encode an Cas13d (such as one encoding a protein having at least 80%, at least 60 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 65 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76,

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77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253). Exemplary viral vectors include polyoma, SV40, adenovirus, vaccinia virus, adeno-associated virus, herpes viruses including HSV and EBV, lentivirus, Sindbis viruses, alphaviruses and retroviruses of avian, murine, and human origin. Baculovirus (Autographa californica multinuclear polyhedrosis virus; AcMNPV) vectors can be used and obtained from commercial sources. Other suitable vectors include retrovirus vectors, orthopox vectors, avipox vectors, fowlpox vectors, capripox vectors, suipox vectors, adenoviral vectors, herpes virus vectors, alpha virus vectors, baculovirus vectors, Sindbis virus vectors, vaccinia virus vectors and poliovirus vectors. Specific exemplary vectors are poxvirus vectors such as vaccinia virus, fowlpox virus and a highly attenuated vaccinia virus (MVA), adenovirus, baculovirus and the like. Pox viruses of use include orthopox, suipox, avipox, and capripox virus. Orthopox include vaccinia, ectromelia, and raccoon pox. One example of an orthopox of use is vaccinia. Avipox includes fowlpox, canary pox and pigeon pox. Capripox include goatpox and sheeppox. In one example, the suipox is swinepox. Other viral vectors that can be used include other DNA viruses such as herpes simplex virus and adenoviruses, and RNA viruses such as retroviruses and polio.

Viral vectors that encode an Cas13d protein (such as one encoding a protein having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) can include at least one expression control element operationally linked to the nucleic acid sequence encoding the Cas13d protein. The expression control elements control and regulate the expression of the Cas13d nucleic acid sequence. Exemplary expression control elements that can be used include, but are not limited to, lac system, operator and promoter regions of phage lambda, yeast promoters and promoters derived from polyoma, adenovirus, retrovirus or SV40. In one example the promoter is CMV, U6, CBh, CMW, Cbh, EF1a. In one example, the promoter is a cell type specific promoter, such as synapsin or GFAP, or an inducible promoter, such as a tetracycline inducible promoter. Additional operational elements include, but are not limited to, leader sequence, termination codons, polyadenylation signals and any other sequences necessary for the appropriate transcription and subsequent translation of the nucleic acid sequence encoding the Cas13d protein in the host system. The expression vector can contain additional elements necessary for the transfer and subsequent replication of the expression vector containing the nucleic acid sequence in the host system. Examples

of such elements include, but are not limited to, origins of replication and selectable markers.

In one example, the vector includes a polyA signal after the Cas13d protein coding sequence, a WPRE signal for expression in viral vectors, or combinations thereof.

In one example, the method uses direct delivery of an mRNA that encodes for an Cas13d protein.

C. Guide Nucleic Acid Molecules

The disclosure provides guide nucleic acid molecules, such as guide RNA (gRNA or crRNA, CRISPR (guide) 10 RNA), which can be used in the methods, compositions, and kits provided herein. Such molecules can include naturally occurring or non-naturally occurring nucleotides or ribonucleotides (such as LNAs or other chemically modified nucleotides or ribonucleotides, for example to protect a guide RNA from degradation). In some examples, the guide sequence is RNA. The guide nucleic acid can include modified bases or chemical modifications (e.g., see Latorre et al., *Angewandte Chemie* 55:3548-50, 2016). A guide sequence directs an Cas13d protein to a target RNA, thereby 20 targeting the RNA (e.g., modifying or detecting the RNA).

Guide molecules include one or more regions referred to as spacers. A spacer has sufficient complementarity with a target RNA sequence to hybridize with the target RNA and direct sequence-specific binding of an Cas13d protein to the 25 target RNA. Thus, the spacer is the variable portion of the guide sequence. In some examples, a spacer has 100% complementarity to a target RNA (or region of the RNA to be target), but a spacer can have less than 100% complementarity to a target RNA, such as at least 80%, at least 30 85%, at least 90%, at least 95%, at least 98% or at least 99% complementarity to a target RNA.

A guide sequence can also include one or more direct repeats (DRs). The DR is the constant portion of the guide, which contains strong secondary structure (FIG. 3C), which 35 facilitate interaction between a Cas13d protein and the guide molecule. Each ortholog has a slightly different DR sequence (e.g., SEQ ID NOS: 129, 130, 131, 132, 133, 134, 135, 136, 137, 148, 150, 151, 152, 154, 156, 157, 159, 161, 163, 165, 167, and 169). In one example, the gRNA includes 40 at least one DR sequence having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 129, 130, 131, 132, 133, 134, 135, 136, 137, 148, 150, 151, 152, 154, 156, 157, 159, 161, 163, 165, 167, 45 169, 176, 178, 180, 182, 184, 186, 188, 190, 191, 192, 193, 199, 201, 203, 205, 207, 209, 211, 213, 215, 217, 219, 221, 223, 225, 227, 228, 230, 232, 234, 236, 238, 240, 242, 244, 246, 248, 250, 252, or 254 (such as 1, 2, 3, or 4 of such DR sequences).

In one example, a guide sequence includes a constant DR on its 5'-end and a variable spacer on its 3' end. In one example includes the sequence DR-spacer-DR-spacer. In some examples, the sequence DR-spacer is repeated two or more times, such as at least 3 times or at least 4 times. This 55 type of sequence is called a guide array.

Guide molecules generally exist in various states of processing. In one example, an unprocessed guide RNA is 36nt of DR followed by 30-32 nt of spacer. The guide RNA is processed (truncated/modified) by Cas13d itself or other 60 RNases into the shorter "mature" form. In some embodiments, an unprocessed guide sequence is about, or at least about 30, 35, 40, 45, 50, 55, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, or more nucleotides (nt) in length. In some embodiments, a processed guide sequence is about 44 to 60 nt (such as 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64,

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65, 66, 67, 68, 69, or 70 nt). In some embodiments, an unprocessed spacer is about 28-32 nt long (such as 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, or 35 nt) while the mature (processed) spacer can be about 10 to 30 nt, 10 to 25 nt, 14 to 25 nt, 20 to 22 nt, or 14-30 nt (such as 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, or 35 nt). In some embodiments, an unprocessed DR is about 36 nt (such as 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40 or 41 nt), while the processed DR is about 30 nt (such as 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, or 35 nt).

The ability of a guide sequence to direct sequence-specific binding of a CRISPR complex to a target RNA may be assessed by any suitable assay. For example, the components of a CRISPR system sufficient to form a CRISPR complex, including the guide sequence to be tested, may be provided to a host cell having the corresponding target RNA molecule, such as by transfection with vectors encoding the components of the CRISPR sequence, followed by an assessment of preferential cleavage within the target sequence. Similarly, cleavage of a target RNA sequence may be evaluated in a test tube by providing the target RNA, components of a CRISPR complex, including the guide sequence to be tested and a control guide sequence different from the test guide sequence, and comparing binding or rate of cleavage at the target RNA between the test and control guide sequence reactions. Other assays are possible, and will occur to those skilled in the art.

Also provided are vectors, such as a viral vector or plasmid (e.g., retrovirus, lentivirus, adenovirus, adeno-associated virus, or herpes simplex virus), that includes a guide nucleic acid molecule. Exemplary vectors are described herein. In some examples, the guide nucleic acid molecule is operably linked to a promoter or expression control element (examples of which are provided elsewhere in this application). As described elsewhere herein, such vectors can include other elements, such as a gene encoding a selectable marker, such as an antibiotic, such as puromycin, hygromycin, or a detectable marker such as GFP or other fluorophore.

In one example, a plurality of gRNAs are part of an array (which can be part of a vector, such as a viral vector or plasmid). For example, a guide array including the sequence DR-spacer-DR-spacer, can include three unique unprocessed gRNAs (one for each DR-spacer sequence).

45 Once introduced into a cell or cell-free system, the array is processed by the Cas13d protein into the three individual mature gRNAs. This allows for multiplexing, e.g. the delivery of multiple gRNAs to a cell or system to target multiple target RNAs or multiple positions within a single target RNA (or combinations thereof).

D. Vectors that encode Cas13d and Guide Nucleic Acid Molecules

The disclosure provides vectors, such as plasmids and viral vectors as described elsewhere herein, which include one or more guide molecule coding sequences (e.g., to permit targeting of one or more RNA molecules), and one or more Cas13d protein coding sequences. Such vectors can be used in the methods, compositions, and kits provided herein. Such vectors can include naturally occurring or non-naturally occurring nucleotides or ribonucleotides. Such vectors can include a single promoter operably linked to the guide molecule (which can be part of an array that includes at least two different guide molecules) and the Cas13d protein coding sequence. Alternatively, the guide molecule (which can be part of an array that includes at least two different guide molecules) and the Cas13d protein coding sequence can be operably linked to different promoters.

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E. Recombinant Cells and Cell-Free Systems

Cells that include a non-native Cas13d protein, a non-native Cas13d protein coding sequence, a guide molecule (or coding sequence), or combinations thereof, are provided. Such recombinant cells can be used in the methods, compositions, and kits provided herein. Nucleic acid molecules encoding an Cas13d protein disclosed herein and/or nucleic acid molecules encoding a guide molecule can be introduced into cells to generate transformed (e.g., recombinant) cells. In some examples, such cells are generated by introducing one or more non-native Cas13d proteins and one or more guide molecules (e.g., gRNAs) into the cell, for example as a ribonucleoprotein (RNP) complex.

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Similarly, cell free systems, such as those generated from lysed cells (or those that include an Cas13d RNP in a test 15 tube or other vessel, into which in vitro transcribed or chemically synthesized target RNAs are added), which include a, Cas13d protein, a Cas13d protein coding sequence, a guide molecule (or coding sequence), or combinations thereof, are provided. Such cell free systems can 20 be used in the methods, compositions, and kits provided herein. In some examples, one or more non-native Cas13d proteins and one or more guide molecules (e.g., gRNAs) are added to a cell free system, for example as a RNP complex.

Thus, cells and cell-free systems containing an Cas13d 25 protein (such as a protein having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 30 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 64, 65, 66, 67, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 88, 89, 90, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 103, 104, 105, 106, 107, 35 108, 109, 110, 111, 112, 113, 138, 147, 149, 153, 155, 158, 160, 162, 164, 166, 168, 170, 175, 177, 179, 181, 183, 185, 187, 189, 194, 198, 200, 202, 204, 206, 208, 210, 212, 214, 216, 218, 220, 222, 224, 226, 229, 231, 233, 235, 237, 239, 241, 243, 245, 247, 249, 251, or 253) are disclosed. Simi- 40 larly, cells and cell-free systems containing a guide molecule, such as one having at least one DR sequence having at least 80%, at least 85%, at least 90%, at least 92%, at least 95%, at least 96%, at least 97%, at least 98%, at least 99%, or 100% sequence identity to SEQ ID NO: 129, 130, 131, 45 132, 133, 134, 135, 136, or 137, and in some examples also at least one spacer sequence complementary to a target RNA, are provided.

Such recombinant cells (e.g., which can be used to generate a cell-free system) can be eukaryotic or prokaryotic. Examples of such cells include, but are not limited to bacteria, archaea, plant, fungal, yeast, insect, and mammalian cells, such as *Lactobacillus*, *Lactococcus*, *Bacillus* (such as *B. subtilis*), *Escherichia* (such as *E. coli*), *Clostridium*, *Saccharomyces* or *Pichia* (such as *S. cerevisiae* 55 or *P. pastoris*), *Kluyveromyces lactis*, *Salmonella typhimurium*, *Drosophila* cells, *C. elegans* cells, *Xenopus* cells, SF9
cells, C129 cells, 293 cells, *Neurospora*, and immortalized mammalian cell lines (e.g., Hela cells, myeloid cell lines, and lymphoid cell lines).

In one example, the cell is a prokaryotic cell, such as a bacterial cell, such as *E. coli*.

In one example, the cell is a eukaryotic cell, such as a mammalian cell, such as a human cell. In one example, the cell is primary eukaryotic cell, a stem cell, a tumor/cancer 65 cell, a circulating tumor cell (CTC), a blood cell (e.g., T cell, B cell, NK cell, Tregs, etc.), hematopoietic stem cell,

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specialized immune cell (e.g., tumor-infiltrating lymphocyte or tumor-suppressed lymphocytes), a stromal cell in the tumor microenvironment (e.g., cancer-associated fibroblasts, etc.) In one example, the cell is a brain cell (e.g., neurons, astrocytes, microglia, retinal ganglion cells, rods/cones, etc.) of the central or peripheral nervous system).

In one example, a cell is part of (or obtained from) a biological sample, such as a biological specimen containing genomic DNA, RNA (e.g., mRNA), protein, or combinations thereof, obtained from a subject. Examples include, but are not limited to, peripheral blood, serum, plasma, urine, saliva, sputum, tissue biopsy, fine needle aspirate, surgical specimen, and autopsy material. Such cells can also be used to generate a cell free system.

In one example the cell (or cell free system) is from a tumor, such as a hematological tumor (e.g., leukemias, including acute leukemias (such as acute lymphocytic leukemia, acute myelocytic leukemia, acute myelogenous leukemia and myeloblastic, promyelocytic, myelomonocytic, monocytic and erythroleukemia), chronic leukemias (such as chronic myelocytic (granulocytic) leukemia, chronic myelogenous leukemia, and chronic lymphocytic leukemia), polycythemia vera, lymphoma, Hodgkin's disease, non-Hodgkin's lymphoma (including low-, intermediate-, and high-grade), multiple myeloma, Waldenström's macroglobulinemia, heavy chain disease, myelodysplastic syndrome, mantle cell lymphoma and myelodysplasia) or solid tumor (e.g., sarcomas and carcinomas: fibrosarcoma, myxosarcoma, liposarcoma, chondrosarcoma, osteogenic sarcoma, and other sarcomas, synovioma, mesothelioma, Ewing's tumor, leiomyosarcoma, rhabdomyosarcoma, colon carcinoma, lymphoid malignancy, pancreatic cancer, breast cancer, lung cancers, ovarian cancer, prostate cancer, hepatocellular carcinoma, squamous cell carcinoma, basal cell carcinoma, adenocarcinoma, sweat gland carcinoma, sebaceous gland carcinoma, papillary carcinoma, papillary adenocarcinomas, medullary carcinoma, bronchogenic carcinoma, renal cell carcinoma, hepatoma, bile duct carcinoma, choriocarcinoma, Wilms' tumor, cervical cancer, testicular tumor, bladder carcinoma, and CNS tumors (such as a glioma, astrocytoma, medulloblastoma, craniopharyogioma, ependymoma, pinealoma, hemangioblastoma, acoustic neuroma, oligodendroglioma, menangioma, melanoma, neuroblastoma and retinoblastoma).

In one example the cell (or cell free system) is obtained from an environmental sample, such as a water, soil, or air sample.

F. Compositions & Kits

Compositions and kits that include a Cas13d protein, a Cas13d protein coding sequence, a guide molecule (or coding sequence), or combinations thereof, are provided. In one example, the composition or kit includes an RNP complex composed of one or more Cas13d proteins and one or more guide molecules (e.g., gRNAs). In one example, the composition or kit includes a vector encoding an Cas13d protein, a guide molecule, or both. In one example, the composition or kit includes a cell, such as a bacterial cell or eukaryotic cell, that includes a non-native Cas13d protein, a non-native Cas13d protein coding sequence, a guide molecule (or coding sequence), or combinations thereof. In one example, the composition or kit includes a cell-free system that includes a non-native Cas13d protein, a non-native Cas13d protein coding sequence, a guide molecule (or coding sequence), or combinations thereof.

Such compositions can include a pharmaceutically acceptable carrier (e.g., saline, water, PBS). In some examples, the composition is a liquid, lyophilized powder, or cryopreserved.

In some examples, the kit includes a delivery system (e.g., 5 liposome, a particle, an exosome, a microvesicle, a viral vector, or a plasmid), and/or a label (e.g., a peptide or antibody that can be conjugated either directly to an Cas13d RNP or to a particle containing the Cas1 RNP to direct cell type specific uptake/enhance endosomal escape/enable 10 blood-brain barrier crossing etc.). In some examples, the kits further include cell culture or growth media, such as media appropriate for growing bacterial, plant, insect, or mammalian cells.

In some examples, such parts of a kit are in separate 15 containers.

G. Targeting RNA

The disclosed Cas13d proteins (and coding sequences), and guide molecules (e.g., gRNA and coding sequences) can be used in a CRISPR/Cas system to target one or more RNA 20 molecules, such as those present in a sample (such as a biological sample, environmental sample (e.g., soil, air or water sample), and the like. In one example, the target RNA is a coding RNA. In one example, the target RNA is a nuclear RNA. In other examples, the target RNA is non-coding RNA (such as functional RNA, siRNA, microRNA, snRNA, snoRNA, piRNA, scaRNA, tRNA, rRNA, lncRNA, or lincRNA). Such RNA targeting methods can be performed in vitro (such as in cell culture or in a cell-free system), or in vivo (such as in an organism, embryo, or 30 mammal)

The CRISPR/Cas system provided herein includes two general components: (1) an Cas13d protein or its coding sequence (whose expression can be driven by a promoter) and (2) a guide nucleic acid molecule, such as RNA (gRNA), 35 which is specific for the target RNA (whose expression can also be driven by promoter. When introduced into cells (or to a cell free system) for example (1) as Cas13d mRNA and Cas13d gRNA, (2) as part of a single vector or plasmid or divided into multiple vectors or plasmids, (3) as separate 40 Cas13d protein and guide molecules, or (4) as an RNP complex of the Cas13d protein and guide molecule, the guide molecule guides the Cas13d to the target RNA. If the Cas13d protein has a native HEPN domain(s) or is fused to an appropriate effector domain bearing RNase activity, the 45 RNA can be cut. If the Cas13d protein has a mutated HEPN domain(s), a guide array can be processed into mature gRNAs, but the target RNA is not cut. Using this system, RNA sequences are easily targeted, for example edited or detected, optionally with an effector domain.

1. Introduction of Cas13d Protein Directly into a Cell

In one example, the Cas13d protein is expressed in a recombinant cell, such as *E. coli*, and purified. The resulting purified Cas13d protein, along with an appropriate guide molecule specific for the target RNA, is then introduced into 55 a cell or organism where one or more RNAs can be targeted. In some examples, the Cas13d protein and guide nucleic acid molecule are introduced as separate components into the target cell/organism. In other examples, the purified Cas13d protein is complexed with the guide nucleic acid (e.g., gRNA), and this ribonucleoprotein (RNP) complex is introduced into target cells (e.g., using transfection or injection). In some examples, the Cas13d protein and guide molecule are injected into an embryo (such as a human, mouse, zebrafish, or *Xenopus* embryo).

Once the Cas13d protein and guide nucleic acid molecule are in the cell, one or more RNAs can be targeted.

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2. Expression of Cas13d from Nucleic Acids

In one example, the Cas13d protein is expressed from a nucleic acid molecule in a cell containing a target RNA, for example an RNA to be detected or modified. In some such examples, the Cas13d protein is expressed from a vector, such as a viral vector or plasmid introduced into a cell or into a cell-free system. This results in the production of the Cas13d protein in the cell, organism, or system. In addition, these nucleic acid molecules can be co-expressed in the cell/organism/system with the guide nucleic acid molecule (e.g., gRNA) specific for the target RNA.

In one example, multiple plasmids or vectors are used for RNA targeting. The nucleic acid molecule encoding the Cas13d can be provided for example on one vector or plasmid, and the guide nucleic acid molecule (e.g., gRNA) on another plasmid or vector. Multiple plasmids or viral vectors can be mixed and introduced into cells (or a cell free system) at the same time, or separately.

In some examples, multiple nucleic acid molecules are expressed from a single vector or plasmid. For example, a single vector can include the nucleic acid molecule encoding the Cas13d, and a separate vector can include the guide molecule.

In some examples a plurality of different guide molecules (e.g., gRNAs), one for each target (such as 1, 2, 3, 4, 5, or 10 different targets), are present on a single array and/or vector. In one example, the method includes delivering a plurality of gRNAs (such as at least 2, at least 3, at least 4, at least 5, at least 10, at least 20, or at least 50 different gRNAs), which are part of an array (which can be part of a vector, such as a viral vector or plasmid). Once introduced into a cell or cell-free system, the array is processed by the Cas13d protein into the individual mature gRNAs.

The nucleic acid molecules expressed from the vector can be under the control of a promoter and optionally contain selection markers (such as antibiotic resistance).

In some examples, the protein and guide molecules are expressed by an embryo (such as a zebrafish or *Xenopus* embryo). The Cas13d protein can be expressed from injected plasmid DNA, injected mRNA, or stably integrated copies into the animal genome. The gRNA can be directly injected or expressed from a vector or stably integrated copies into the animal genome.

3. Targets

One or more RNAs can be targeted by the disclosed methods, such as at least 1, at least 2, at least 3, at least 4 or at least 5 different RNAs in a cell, cell-free system, or organism, such as 1, 2, 3, 4, 5, 6, 7, 8, 9 or 10 different RNAs. In one example, the RNA is associated with a disease such as cystic fibrosis, Huntington's disease, Tay-Sachs, Fragile X syndrome, Fragile X-associated tremor/ataxia syndrome, Duchenne muscular dystrophy, myotonic dystrophy, spinal muscular atrophy, spinocerebellar ataxia, or familial ALS. In one example, the RNA is associated with cancer (e.g., a cancer of the lung, breast, colon, liver, pancreas, prostate, bone, brain, skin (e.g., melanoma), or kidney). Examples of target RNAs include, but are not limited to those associated with cancer (e.g., BCR-ABL, Ras, Raf, p53, BRCA1, BRCA2, CXCR4, beta-catenin, HER2, and CDK4).

In one example, the RNA is associated with viral infection, such as infection by a positive-strand RNA viruses, such as Picornaviruses (such as Aphthoviridae [for example foot-and-mouth-disease virus (FMDV)]), Cardioviridae; Enteroviridae (such as Coxsackie viruses, Echoviruses, Enteroviruses, and Polioviruses); Rhinoviridae (Rhinoviruses)); Hepataviridae (Hepatitis A viruses); Togaviruses

(examples of which include rubella; alphaviruses (such as Western equine encephalitis virus, Eastern equine encephalitis virus, and Venezuelan equine encephalitis virus)); Flaviviruses (examples of which include Dengue virus, West Nile virus, and Japanese encephalitis virus); Calciviridae 5 (which includes Norovirus and Sapovirus); or Coronaviruses (examples of which include SARS coronaviruses, such as the Urbani strain), or a negative-strand RNA virus, such as Orthomyxyoviruses (such as the influenza virus), Rhabdoviruses (such as Rabies virus), and Paramyxoviruses 10 (examples of which include measles virus, respiratory syncytial virus, or parainfluenza viruses), or a DNA viral infection (such as infection by Herpesviruses (such as Varicella-zoster virus, for example the Oka strain; cytomegalovirus; and Herpes simplex virus (HSV) types 1 and 2), 15 Adenoviruses (such as Adenovirus type 1 and Adenovirus type 41), Poxviruses (such as Vaccinia virus), or Parvoviruses (such as Parvovirus B19).

In one example, the RNA is associated with a bacterial infection or property of a bacterial infection, such as bac- 20 terial resistance, persistence, or antibiotic resistance. Detection of these RNAs can be used for diagnostic methods, while editing these RNAs in cell-based or cell-free systems can be used for therapeutic methods.

4. Methods of Detecting RNA

In one example, the method of targeting an RNA results in detecting, visualizing, or labeling a target RNA. For example, by using at least one Cas13d protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4), at least one gRNA containing at least one spacer sequence specific 30 for the target RNA, and an effector module, the target RNA will be recognized by Cas13d but will not be cut or nicked while the effector module becomes activated. In some examples, such a method is used to detect a target RNA. Such a method can be used in a cell or cell free system to 35 determine if a target RNA is present, such as in a tumor cell. In some examples, the cell or cell free system is obtained from a tissue sample, blood sample or saliva sample.

In one example, the method of detecting RNA comprises of an Cas13d protein fused to a fluorescent protein or other 40 detectable label along with a gRNA containing a spacer sequence specific for the target RNA. Binding of Cas13d to the target RNA can be visualized by microscopy or other methods of imaging. In another example, RNA aptamer sequences can be appended to or inserted within the gRNA 45 molecule, such as MS2, PP7, Qβ, and other aptamers. The introduction of proteins that specifically bind to these aptamers, e.g. the MS2 phage coat protein, fused to a fluorescent protein or other detectable label can be used to detect the target RNA because the Cas13d-gRNA-target RNA complex 50 will be labeled via the aptamer interaction.

In another example, the method of detecting RNA is a transcriptional sensor (e.g., as part of a synthetic circuit) for diagnostics or therapeutics. For example, the transcriptional sensor can be comprised of at least one Cas13d protein with 55 a fluorescent probe and quencher. The intact detectable RNA a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4), at least one gRNA containing at least one spacer sequence specific for the target RNA, and an effector module such as an optionally split fluorescent protein or probe; an optionally split luminescent protein or probe; an optionally split 60 enzyme that catalyzes a detectable reaction such as ubiquitin or TEV protease; a FRET-compatible protein pair; one or more transcription factor(s) fused to Cas13d via cleavable linkers; a split intein that trans-splices a protein to restore its function such as a transcription factor; a kinase-substrate 65 pair that activates upon phosphorylation; one, two, or more monomers that activate upon dimerization or multimeriza-

tion; or one or more proteins that induce conformational and functional change upon interaction. In one example, the spatial proximity of two or more Cas13d proteins and gRNAs due to binding a particular transcript would activate the effector module, resulting in a detectable signal or detectable activity in the cell.

For example, the transcriptional sensor could allow a cancer-specific transcript, inflammation-specific transcript, disease-specific transcript, or cell state-specific transcript to be detected. A synthetic circuit containing a Cas13d-based system that is able to sense particular transcripts could encode conditional logic, e.g. requiring target detection to up- or down-regulate a gene for therapeutic application.

In one example, the method results in a detectable agent being bound to the target RNA, which can be detected. For example, two separate Cas13d fusion proteins that each include part of a fluorophore (e.g., GFP), and two different gRNAs with different spacer sequences that target regions of an RNA in close proximity, can be used. When the two parts bind to the target RNA in proximity, the two parts of the fluorophore form a complete fluorophore, thereby generating a detectable signal.

In one example, the method results in RNA detection, for example by triggering a response such as expression of a second gene, modification of a protein, translocation of a protein or RNA to a different location, induction of cell death via suicide gene, induction of cell proliferation, induction of a transgene that enables a secondary function, induction of a permanent change in DNA sequence to enable storing a memory of past transcriptional events, or altering the RNA to enable pulldown.

In one embodiment, two halves of a transcription factor could be linked to two separate Cas13d via a split intein system. The Cas13d proteins are provided with two different gRNAs with different spacer sequences that target regions of an RNA in close proximity Upon binding to the target RNA in proximity, the split inteins trans-splice a reconstituted transcription factor (TF) so that it can translocate to the nucleus and turn on a target gene or cluster of target genes. In one example, the target gene could be an endogenous gene in the cell. In another example, the target gene could be a transgene expressed on a vector or introduced through genetic engineering, such as a fluorescent protein or toxin.

5. Methods of Detecting RNA in Cell-Free Systems

In one example, the method of detecting a target RNA in a cell-free system results in a detectable label or enzyme activity. For example, by using at least one Cas13d protein (e.g., SEQ ID NO: 3, 42, 62, 70, 82, 83, and 92), at least one gRNA containing at least one spacer sequence specific for the target RNA, and a detectable label, the target RNA will be recognized by Cas13d. The binding of the target RNA by Cas13d triggers its RNase activity, which can lead to the cleavage of the target RNA as well as the detectable label.

In one example, the detectable label is an RNA linked to links the fluorescent probe and quencher, suppressing fluorescence. Upon cleavage by Cas13d of the detectable RNA, the fluorescent probe is released from the quencher and displays fluorescent activity. Such a method can be used to determine if a target RNA is present in a lysed cell sample, lysed tissue sample, blood sample, saliva sample, environmental sample (such as a water, soil, or air sample), or other lysed cell or cell-free sample. Such a method can also be used to detect a pathogen, such as a virus or bacteria, or diagnose a disease state, such as a cancer.

In one example, the detection of the target RNA aids in the diagnosis of disease and/or pathological state, or the exis-

tence of a viral or bacterial infection. For example, Cas13d-mediated detection of non-coding RNAs such as PCA3 can be used to diagnose prostate cancer if detected in patient urine. In another example, Cas13d-mediated detection of the lncRNA-AA174084, which is a biomarker of gastric cancer, 5 can be used to diagnose gastric cancer.

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6. Methods of Editing Target RNA

In one example, the method of targeting an RNA results in editing the sequence of a target RNA. For example, by using an Cas13d protein with a non-mutated HEPN domain 10 (e.g., SEQ ID NOS: 1, 3, 42, 62, 70, 82, 83, and 92), and a gRNA containing at least one a spacer sequence specific for the target RNA, the target RNA can be cut or nicked at a precise location. In some examples, such a method is used to decrease expression of a target RNA, which will decrease 15 translation of the corresponding protein. Such a method can be used in a cell where increased expression of an RNA is not desired. In one example, the RNA is associated with a disease such as cystic fibrosis, Huntington's disease, Tay-Sachs, Fragile X syndrome, Fragile X-associated tremor/ 20 ataxia syndrome, muscular dystrophy, myotonic dystrophy, spinal muscular atrophy, spinocerebellar ataxia, or familial ALS. In another example, the RNA is associated with cancer (e.g., a cancer of the lung, breast, colon, liver, pancreas, prostate, bone, brain, skin (e.g., melanoma), or kidney). 25 Examples of target RNAs include, but are not limited to those associated with cancer (e.g., PD-L1, BCR-ABL, Ras, Raf, p53, BRCA1, BRCA2, CXCR4, beta-catenin, HER2, and CDK4). Editing such target RNAs can have a therapeu-

In another example, the RNA is expressed in an immune cell. The target RNA could, for example, code for a protein leading to the repression of a desirable immune response, such as infiltration of a tumor. Knock-down of such an RNA could enable progression of such a desirable immune 35 response (e.g., PD1, CTLA4, LAG3, TIM3). In another example, the target RNA encodes a protein resulting in the undesirable activation of an immune response, for example in the context of an autoimmune disease such as multiple sclerosis, Crohn's disease, lupus, or rheumatoid arthritis.

In one example, targeting the target RNA allows for decreasing expression of the target protein encoded by the RNA. For example, by using an Cas13d fusion protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4) and a translational repression domain (such as Pumilio or FBF 45 PUF proteins, deadenylases, CAF1, Argonaute proteins, and others), and a guide RNA containing at least one spacer sequence specific for the target RNA, expression of a target RNA can be decreased.

In some examples, Cas13d can be fused to a ribonuclease 50 (such as a PIN endonuclease domain, an NYN domain, an SMR domain from SOT1, or an RNase domain from Staphylococcal nuclease) or a domain that affects RNA stability (such as tristetraprolin or domains from UPF1, EXOSC5, and STAU1).

In another example, RNA aptamer sequences can be appended to or inserted within the gRNA molecule, such as MS2, PP7, Q β , and other aptamers. Proteins that specifically bind to these aptamers, e.g. the MS2 phage coat protein, can be fused to a translational repression domain, a ribonuclease, 60 or a domain that affects RNA stability. This aptamer-effector domain fusion can be used to target the target RNA because the Cas13d and gRNA complex will guide the aptamer protein-effector domain in proximity to the target RNA.

Such a method can be used in a cell where increased 65 expression of an RNA is not desired, such as when an expressed RNA is associated with a disease such as cystic

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fibrosis, Huntington's disease, Tay-Sachs, Fragile X syndrome, Fragile X-associated tremor/ataxia syndrome, muscular dystrophy, myotonic dystrophy, spinal muscular atrophy, spinocerebellar ataxia, or familial ALS. In another example, the target RNA is associated with cancer (e.g., a cancer of the lung, breast, colon, liver, pancreas, prostate, bone, brain, skin (e.g., melanoma), or kidney). Examples of target RNAs include, but are not limited to those associated with cancer (e.g., PD-L1, BCR-ABL, Ras, Raf, p53, BRCA1, BRCA2, CXCR4, beta-catenin, HER2, and CDK4). Editing such target RNAs would have a therapeutic effect.

In another example, the RNA is expressed in an immune cell. The target RNA could, for example, code for a protein leading to the repression of a desirable immune response, such as infiltration of a tumor. Knock-down of such an RNA could enable progression of such a desirable immune response (e.g., PD1, CTLA4, LAG3, TIM3). In another example, the target RNA could encode a protein resulting in the undesirable activation of an immune response, for example in the context of an autoimmune disease such as multiple sclerosis, Crohn's disease, lupus, or rheumatoid arthritis.

In one example, targeting the target RNA allows for activating or increasing expression of the target RNA. For example, by using an Cas13d fusion protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4) and a translational activation domain (such as eIF4E and other translation initiation factors, a domain of the yeast poly(A)-binding protein or GLD2), and a guide RNA containing at least one a spacer sequence specific for the target RNA, expression of a target RNA can be increased. Aptamer introduction into the gRNA with a cognate aptamer-binding protein fused to a translational activation domain can also be used. In one example, RNA aptamer sequences are appended to or inserted within the gRNA molecule, such as MS2, PP7, Qβ, and other aptamers. The introduction of proteins that specifically bind to these aptamers, e.g. the MS2 phage coat protein, fused to a translational activation domain can be used to target the target RNA because the Cas13d and gRNA complex will bring the aptamer protein-translational activation domain in proximity to the target RNA.

In some examples, such a method is used to increase the activity or expression of a target RNA, which will increase translation of the corresponding protein (if the RNA is a coding RNA). Such a method can be used in a cell where increased expression of an RNA is desired, such as a heterozygous genetic disease or disorders caused by copy number variation. Increasing translation of a desired protein product could be therapeutic in nature.

In another example, increasing the expression of a target RNA (such as Cyclin B1) can render the target cell (such as cancers) more sensitive to drugs (such as chemotherapeutic agents).

In one example, targeting the target RNA allows for one or more RNA base substitutions, RNA base edits, RNA base deletions, RNA base insertions, or combinations thereof, in the target RNA. In some examples, the Cas13d protein with a mutated HEPN domain is associated, either via direct fusion or a gRNA-aptamer modification, an effector domain that allows base edits (such as a cytidine deaminase, PPR protein, adenosine deaminase, ADAR family protein, or APOBEC family protein). In some examples, such a method is used to modify an RNA sequence, edit an RNA mutation, or modify an RNA transcript (e.g., gene therapy), for

example to treat diseases such as ALS and melanoma or genetic disorders caused by undesired splice sites, such as Leber congenital amaurosis.

In one example, targeting the target RNA allows for methylating the target RNA. Some examples may use a 5 chimeric Cas13d protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4) associated either via direct fusion or a gRNA-aptamer modification with a methylation domain (e.g., m6A), and a guide RNA containing at least one a spacer sequence specific for the target RNA. In some 10 examples, such a method is used to combat aberrant RNA demethylation. In one example, such a method is used modify the methylation levels of pluripotency transcripts such as NANOG or KLF4 for example to decrease their stability in breast cancer cells, which can suppress the 15 acquisition of breast cancer stem cell phenotypes that are associated with increased proliferation and cancer stem cell formation

In one example, targeting the target RNA allows for demethylating the target RNA. Some examples can use 20 Cas13d protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4), a guide RNA containing at least one a spacer sequence specific for the target RNA, and a demethylation domain (e.g., human alkylation repair homolog 5 or Alkbh5). The demethylation domain can be associated either 25 via direct fusion to the Cas13d protein or via a gRNA-aptamer modification. In some examples, such a method is used to reverse aberrant RNA methylation, for example to treat myeloid leukemia by decreasing m⁶A levels.

In one example, targeting the target RNA allows for 30 binding to the target RNA. For example, by using a Cas13d protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4) and a guide RNA containing at least one a spacer sequence specific for the target RNA, molecules can be bound or tethered to a target RNA. In some examples, such 35 a method is used to capture the target RNA (e.g., immunoprecipitation). This can be used as part of a kit to identify the proteins interacting with a specific RNA transcript. In one example, an epitope tagged Cas13d (e.g. FLAG, HA, biotin, HALO tag) can be targeted to specific target RNAs and 40 cross-linked via fixation (e.g. with paraformaldehyde or glutaraldehyde). Immunoprecipitation of Cas13d with an epitope-recognizing antibody allows for the identification of co-immunoprecipitated proteins via Western blot or mass spectrometry.

In another example, Cas13d can be fused to a biotin ligase (such as BirA) or a peroxidase (such as APEX2) in order to biotinylate proteins that interact with the target RNA. Labeled proteins can then be pulled down with streptavidin beads followed by mass spectrometry or Western blot.

In some examples, biotinylated Cas13d could be targeted to ribosomal RNA sequences with a gRNA. Streptavidin bead-mediated pulldown can be used to deplete rRNA for RNA sequencing library preparation.

In one example, targeting the target RNA allows for 55 masking the target RNA. For example, by using an Cas13d protein with a mutated or intact HEPN domain and a guide RNA containing at least one a spacer sequence specific for the target RNA, a target RNA can be masked from RNA-binding proteins or RNA-binding elements such as miR- 60 NAs.

In some examples, the Cas13d can be used to mask RNA binding sites from RNA-binding proteins (RBPs). In another example, Cas13d can mask miRNA binding sites. For example, the liver-specific miR-122 forms a complex with 65 Hepatitis C viral RNA which protects it from degradation. A HEPN-active Cas13d protein could be targeted to the

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miRNA-122 binding site on the viral RNA to synergistically combat HCV infections by simultaneously reversing miRNA-122-mediated protection and directly degrading HCV RNA. In some examples, such a method is used to preserve or protect the target RNA molecule, for example to protect the target RNA from degradation. For example, by targeting AU-rich elements in the 3' UTR of a target gene, a HEPN-mutated Cas13d can block binding of RNA-binding proteins such as tristetraprolin (TTR) or AUF1, which lead to degradation of the target transcript.

In one example, targeting the target RNA allows for changing splicing of the target RNA. Both the direct binding of splice acceptor and/or donor sites as well as splice effector domains can be used to manipulate splicing. For example, by using an Cas13d protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4), a guide RNA containing at least one a spacer sequence specific for the target RNA, and optionally an effector domain that affects splicing (such as the RS-rich domain of SRSF1, the Gly-rich domain of hnRNP Al, the alanine-rich motif of RBM4, or the proline-rich motif of DAZAP1), alternative splicing of the RNA can be achieved.

In some examples, such a method is used for exon inclusion, for example to include exon 2 of acid alphaglucosidase (GAA) to treat Pompe disease or to include exon 7 of SMN2 to treat spinal muscular atrophy (SMA). In some examples, such a method is used for exon exclusion, for example to restore the reading frame of dystrophin to treat Duchenne muscular dystrophy or to shift the splicing of the Bcl-x pre-mRNA from the antiapoptotic long isoform to the proapoptotic short isoform to treat cancer.

In some examples, the method uses the Cas13d protein with a mutated HEPN domain to mask splice acceptor or donor sites, for example to create neoantigens to make cold tumors hot. By affecting the splicing of certain target premRNAs, this method can generate novel exon-exon junctions that can lead to the creation of neo-epitopes in cancer cells. This can make a cancer cell vulnerable to the immune system due to the display of unnatural antigens. In other examples, this method can be used to dynamically manipulate isoform ratios or to restore reading frame of a protein (e.g., dystrophin for Duchenne's muscular dystrophy).

In one example, targeting the target RNA allows for controlling transcript trafficking of the target RNA. For example, by using an Cas13d fusion protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4) and a subcellular localization signal or export sequence, a guide RNA containing at least one a spacer sequence specific for the target RNA. In some examples, such a method is used to traffic the target RNA molecule to a particular organelle or cytosolic compartment, or even export the target RNA transcript, for example to endosomes for extracellular release

In another example, the method can affect RNA trafficking. For example, the zipcode binding protein ZBP1 specifically recognizes an RNA sequence 5'-CGGAC(C/A-CA-C/U) that leads to localization of certain transcripts to the leading edge of fibroblasts. By masking or manipulating particular RNA zipcodes or regulatory sequences from recognition by regulatory protein complexes, this method can affect RNA localization or trafficking within a cell.

In one example, the target RNA is a nuclearly localized RNA. For example, by using an Cas13d protein with a non-mutated HEPN domain (e.g., SEQ ID NOS: 1, 3, 42, 62, 70, 82, 83, and 92) fused to a nuclear localization signal and a guide RNA containing at least one a spacer sequence specific for the target nuclear RNA, the nuclearly localized RNA can be targeted and degraded. In some examples, such

a method is used to degrade the target nuclear RNA molecule, for example to knock-down a non-coding nuclear RNA such as HOTAIR, which is associated with metastatic progression in breast cancer.

In one example, the target RNA is viral RNA or transcript 5 of a DNA virus. For example, an Cas13d protein with a non-mutated HEPN domain (e.g., SEQ ID NOS: 1, 3, 42, 62, 70, 82, 83, and 92) and a guide RNA containing at least one spacer sequence specific for the target RNA can be used. In some examples, such a method is used to treat an RNA viral 10 infection (such as infection by a positive-strand RNA viruses, such as Picornaviruses (such as Aphthoviridae [for example foot-and-mouth-disease virus (FMDV)]), Cardioviridae; Enteroviridae (such as Coxsackie viruses, Echoviruses, Enteroviruses, and Polioviruses); Rhinoviridae (Rhi-15 Hepataviridae (Hepatitis A viruses): noviruses)); Togaviruses (examples of which include rubella; alphaviruses (such as Western equine encephalitis virus, Eastern equine encephalitis virus, and Venezuelan equine encephalitis virus)); Flaviviruses (examples of which include Den- 20 gue virus, West Nile virus, and Japanese encephalitis virus); Calciviridae (which includes Norovirus and Sapovirus); or Coronaviruses (examples of which include SARS coronaviruses, such as the Urbani strain), or a negative-strand RNA virus, such as Orthomyxyoviruses (such as the influenza 25 virus), Rhabdoviruses (such as Rabies virus), and Paramyxoviruses (examples of which include measles virus, respiratory syncytial virus, or parainfluenza viruses), or a DNA viral infection (such as infection by Herpesviruses (such as Varicella-zoster virus, for example the Oka strain; 30 cytomegalovirus; and Herpes simplex virus (HSV) types 1 and 2), Adenoviruses (such as Adenovirus type 1 and Adenovirus type 41), Poxviruses (such as Vaccinia virus), or Parvoviruses (such as Parvovirus B19), for example by cutting the viral RNA or transcript of a DNA virus. Thus, 35 such methods can be used as an RNA-based antiviral or antimicrobial.

EXAMPLE 1

Materials and Methods

This example describes the materials and methods used to obtain the results shown in Examples 2-7.

Cell Culture of Human Embryonic Kidney (HEK) Cell Line 45 293FT

Human embryonic kidney (HEK) cell line 293FT (Thermo Fisher) was maintained in DMEM (4.5 g/L glucose), supplemented with 10% FBS (GE Life Sciences) and 10 mM HEPES at 37° C. with 5% CO₂. Upon reaching 50 80-90% confluency, cells were dissociated using TrypLE Express (Life Technologies) and passaged at a ratio of 1:2. Cell Culture of Human Bone Osteosarcoma Epithelial Cell Line U2OS

Human bone osteosarcoma epithelial U2OS were maintained in DMEM (4.5 g/L glucose) supplemented with 10% FBS and 10 mM HEPES at 37° C. with 5% CO₂. Cells were passaged at a 1:3 ratio upon reaching 70% confluence. This cell line was not authenticated.

Maintenance of induced pluripotent stem cells and neuronal 60 differentiation

Stable human iPSC lines containing the FTDP-17 IVS10+16 mutation or age- and sex-matched control lines were obtained from the laboratory of Fen-Biao Gao (Biswas et al., 2016). Briefly, cells obtained from one male patient with the 65 MAPT IVS10+16 mutation and two separate lines from one male control patient were reprogrammed into hiPSCs

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(Almeida et al., 2012). iPSCs were transduced with lentivirus containing a doxycycline-inducible Ngn2 cassette. Lentiviral plasmids were a gift from S. Schafer and F. Gage. iPSCs were then passaged with Accutase and plated into a Matrigel-coated 6-well plate with mTESR media containing ROCK inhibitor Y-27632 (10 µM, Cayman) at 500,000 cells per well. On day 1, media was changed with mTESR. On day 2, media was changed to mTESR containing doxycycline (2 µg/ml, Sigma) to induce Ngn2 expression. On day 3, culture media was replaced with Neural Induction media (NIM, DMEM/F12 (Life Technologies) containing BSA (0.1 mg/ml, Sigma), apo-transferrin (0.1 mg/ml, Sigma), putrescine (16 µg/ml, Sigma), progesterone (0.0625 82 g/ml, Sigma), sodium selenite (0.0104 µg/ml, Sigma), insulin (5 μg/ml, Roche), BDNF (10 ng/ml, Peprotech), SB431542 (10 μM, Cayman), LDN-193189 (0.1 μM, Sigma), laminin (2 μg/ml, Life Technologies), doxycycline (2 μg/ml, Sigma) and puromycin (Life Technologies)). NIM media was changed daily. Following 3 days of puromycin selection, immature neuronal cells were passaged with Accumax (Innovative Cell Technologies) and plated onto 96-well plates coated with poly-D-lysine and Matrigel in Neural Maturation media (NMM; 1:1 Neurobasal/DMEM (Life Technologies) containing B27 (Life Technologies), BDNF (10 ng/ml, Peprotech), N-Acetyl cysteine (Sigma), laminin (2 µg/ml, Life Technologies), dbcAMP (49 µg/ml, Sigma) and doxycycline (2 μg/ml, Sigma). Media was replaced the next day (day 7) with NMM containing AraC (2 µg/ml, Sigma) to eliminate any remaining non-differentiated cells. On day 8, AraC was removed and astrocytes were plated on top of neurons to support neuron cultures in NMM containing hbEGF (5 ng/ml, Peprotech). Cells were transduced with AAV on day 10 and assayed on day 24.

Computational Pipeline for Cas13d Identification

We obtained whole genome, chromosome, and scaffoldlevel prokaryotic genome assemblies from NCBI Genome in June 2016 and compared CRISPRfinder, PILER-CR, and CRT for identifying CRISPR repeats. The 20 kilobase flanking regions around each putative CRISPR repeat was 40 extracted to identify nearby proteins and predicted proteins using Python. Candidate Cas proteins were required to be >750 aa in length and within 5 proteins of the repeat array, and extracted CRISPR loci were filtered out if they contained Cas genes associated with known CRISPR systems such as types I-III CRISPR. Putative effectors were clustered into families via all-by-all BLASTp analysis followed by single-linkage hierarchical clustering where a bit score of at least 60 was required for cluster assignment. Each cluster of at least 2 proteins was subjected to BLAST search against the NCBI non-redundant (nr) protein database, requiring a bit score >200 to assign similarity. The co-occurrence of homologous proteins in each expanded cluster to a CRISPR array was analyzed and required to be >70%. Protein families were sorted by average amino acid length and multiple sequence alignment for each cluster was performed using Clustal Omega and the Geneious aligner with a Blosum62 cost matrix. The RxxxxH HEPN motif was identified in the Cas13d family on the basis of this alignment. TBLASTN was performed on all predicted Cas13d effectors against public metagenome whole genome shotgun sequences without predicted open reading frames (ORFs). The Cas13d family was regularly updated via monthly BLAST search on genome and metagenome databases to identify any newly deposited sequences. New full-length homologs and homologous fragments were aligned using Clustal Omega and clustered using PhyML 3.2. CRISPRDetect was used to predict the direction of direct repeats in the Cas13d array and

DR fold predictions were performed using the Andronescu 2007 RNA energy model at 37° C. Sequence logos for Cas13d direct repeats were generated using Geneious 10. Protein Expression and Purification

Recombinant Cas13d proteins were PCR amplified from 5 genomic DNA extractions of cultured isolates or metagenomic samples and cloned into a pET-based vector with an N-terminal His-MBP fusion and TEV protease cleavage site. The resulting plasmids were transformed into Rosetta2 (DE3) cells (Novagen), induced with 200 µM IPTG at OD_{600} 0.5, and grown for 20 hours at 18° C. Cells were then pelleted, freeze-thawed, and resuspended in Lysis Buffer (50 mM HEPES, 500 mM NaCl, 2 mM MgCl2, 20 mM Imidazole, 1% v/v Triton X-100, 1 mM DTT) supplemented with 1× protease inhibitor tablets, 1 mg/mL lysozyme, 2.5 U/mL Turbo DNase (Life Technologies), and 2.5 U/mL salt active nuclease (Sigma Aldrich). Lysed samples were then sonicated and clarified via centrifugation (18,000×g for 1 hour at 4° C.), filtered with 0.45 μM PVDF filter and incubated with 20 50 mL of Ni-NTA Superflow resin (Qiagen) per 10 L of original bacterial culture for 1 hour. The bead-lysate mixture was applied to a chromatography column, washed with 5 column volumes of Lysis Buffer, and 3 column volumes of Elution Buffer (50 mM HEPES, 500 mM NaCl, 300 mM 25 Imidazole, 0.01% v/v Triton X-100, 10% glycerol, 1 mM DTT). The samples were then dialyzed overnight into TEV Cleavage Buffer (50 mM Tris-HCl, 250 mM KCl, 7.5% v/v glycerol, 0.2 mM TCEP, 0.8 mM DTT, TEV protease) before cation exchange (HiTrap SP, GE Life Sciences) and gel 30 filtration (Superdex 200 16/600, GE Life Sciences). Purified, eluted protein fractions were pooled and frozen at 4 mg/mL in Protein Storage Buffer (50 mM Tris-HCl, 1M NaCl, 10% glycerol, 2 mM DTT).

Preparation of Guide and Target RNAs

Oligonucleotides carrying the T7 promoter and appropriate downstream sequence were synthesized (IDT) and annealed with an antisense T7 oligo for crRNAs and PCRamplified for target and array templates. Homopolymer anneal and PCR templates were in vitro transcribed with the Hiscribe T7 High Yield RNA Synthesis kit (New England Biolabs) at 31° C. for 12 hours. For labeled targets, fluorescently labelled aminoallyl-UTP atto 680 (Jena Biosciences) was additionally added at 2 mM. Guide RNAs were 45 purified with RNA-grade Agencourt AMPure XP beads (Beckman Coulter) and arrays and targets were purified with MEGAclear Transcription Clean-Up Kit (Thermo Fisher) and frozen at -80° C. For ssDNA and dsDNA targets, corresponding oligonucleotide sequences were synthesized 50 (IDT) and either gel purified, or PCR amplified and then subsequently gel purified respectively.

Biochemical Cleavage Reactions

Purified EsCas13d protein and guide RNA were mixed (unless otherwise indicated) at 2:1 molar ratio in RNA 55 Cleavage Buffer (25 mM Tris pH 7.5, 15 mM Tris pH 7.0, 1 mM DTT, 6 mM MgCl₂). The reaction was prepared on ice and incubated at 37° C. for 15 minutes prior to the addition of target at 1:2 molar ratio relative to EsCas13d. The reaction was subsequently incubated at 37° C. for 45 min- 60 utes and quenched with 1 µL of enzyme stop solution (10 mg/mL Proteinase K, 4M Urea, 80 mM EDTA, 20 mM Tris pH 8.0) at 37° C. for 15 minutes. The reaction was then denatured with 2x RNA loading buffer (2x: 13 mM Ficoll, 8 M Urea, 25 mM EDTA), at 85° C. for 10 minutes, and 65 separated on a 10% TBE-Urea gel (Life Technologies). Gels containing labeled targets were visualized on the Odyssey

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Imager (Li-Cor); unlabeled array or target cleavage gels were stained with SYBR Gold prior to imaging via Gel Doc EZ system (Bio-Rad).

Transient transfection of Human Cell Lines

Engineered Cas13 coding sequences were cloned into a standardized plasmid expression backbone containing an EF1a promoter and prepared using the Nucleobond Xtra Midi EF Kit (Machery Nagel) according to the manufacturer's protocol. NLS-LwaCas13a-msfGFP and PspCas13b-NES-HIV were PCR amplified from Addgene #103854, and #103862, respectively, a gift from Feng Zhang. Cas13d pre-gRNAs and gRNAs were cloned into a minimal backbone containing a U6 promoter. shRNAs and guides for LwaCas13a were cloned into the same backbone and position matched to their corresponding guide RNA at the 3' of the target sequence. Matched gRNAs for PspCas13b were moved to the closest 5'-G nucleotide.

For transient transfection, HEK 293FT cells were plated at a density of 20,000 cells per well in a 96-well plate and transfected at >90% confluence with 200 ng of Cas13 expression plasmid and 200 ng of gRNA expression plasmid using Lipofectamine 2000 (Life Technologies) according to the manufacturer's protocol. Transfected cells were harvested 48-72 hours post-transfection for flow cytometry, gene expression analysis, or other downstream processing.

For reporter assays, HEK 293FT cells were transfected in 96-well format with 192 ng of Cas13d expression plasmid, 192 ng of guide expression plasmid, and 12 ng of mCherry expression plasmid with Lipofectamine 2000 (Life Technologies). Cells were harvested after 48 hours and analyzed by flow cytometry.

U2OS cells were plated at a density of 20,000 cells per well in a 96-well plate and transfected at >90% confluence with 100 ng of Cas13d expression plasmid using Lipo-35 fectamine 3000 (Life Technologies) according to the manufacturer's protocol and processed for immunocytochemistry after 48 h.

Flow Cytometry

Cells were dissociated 48 hours post-transfection with target RNAs were synthesized by Synthego. The oligo 40 TrypLE Express and resuspended in FACS Buffer (1x DPBS^{-/-}, 0.2% BSA, 2 mM EDTA). Flow cytometry was performed in 96-well plate format using a MACSQuant VYB (Miltenyi Biotec) and analyzed using FlowJo 10. RG6 was a gift from Thomas Cooper (Addgene plasmid # 80167) and modified to replace EGFP with mTagBFP2. All represented samples were assayed with three biological replicates. In the mCherry reporter assay, data is representative of at least 20,000 gated events per condition. In the splicing reporter assay, data is representative of at least 2,500 gated events per condition.

Gene Expression Analysis

Cells were lysed 48 hours post-transfection with DTTsupplemented RLT buffer and total RNA was extracted using RNeasy Mini Plus columns (Qiagen). 200 ng of total RNA was then reverse transcribed using random hexamer primers and Revertaid Reverse Transcriptase (Thermo Fisher) at 25° C. for 10 min, 37° C. for 60 min, and 95° C. for 5 min followed by qPCR using 2× Taqman Fast Advanced Master Mix (Life Technologies) and Taqman probes for GAPDH and the target gene as appropriate (Life Technologies and IDT). Taqman probe and primer sets were generally selected to amplify cDNA across the Cas13 or shRNA target site position to prevent detection of cleaved transcript fragments (see Table S4 of Konermann et al., Cell 173:1-12, 2018, herein incorporated by reference in its entirety). qPCR was carried out in 5 µL multiplexed reactions and 384-well format using the LightCycler 480 Instrument II (Roche).

Fold-change was calculated relative to GFP-transfected vehicle controls using the ddCt method. One-way or two-way ANOVA with multiple comparison correction was used to assess statistical significance of transcript changes using Prism 7.

Immunohistochemistry

For immunohistochemical analysis, U2OS cells were cultured on 96-well optically clear plates (Greiner Bio-One), transfected as previously described, then fixed in 4% PFA (Electron Microscopy Sciences) diluted in PBS (Gibco) and washed with 0.3M glycine (Sigma) in PBS to quench PFA. Samples were blocked and permeabilized in a PBS solution containing 8% donkey serum (Jackson ImmunoResearch), 8% goat serum (Cell Signaling Technologies), and 0.3% $_{15}$ Triton-X 100 (Sigma) for one hour, followed by primary antibody incubation in 1% BSA (Fisher Bioreagents), 1% goat serum, and 0.25% Triton-X overnight at 4° C. Samples were washed 3 times with PBS containing 0.1% BSA and 0.1% Triton-X 100 before incubating with fluorophore- 20 conjugated secondary antibodies in PBS with 0.05% Triton-X 100 and 1% BSA at room temperature for one hour. Cells were washed with PBS with 0.1% Triton-X, stained with DAPI, and then covered with Mounting Media (Ibidi) before imaging. Primary antibody, HA-Tag 6E2 (Cell Sig- 25 naling, 2367), was used at a 1:100 dilution as per manufacturer's instructions. Secondary antibodies used were goat anti-mouse IgG1-Alexa-Fluor 647 (Thermo Fisher, A21240) and Anti-Mouse IgG1 CF 633 (Sigma, SAB4600335). Confocal images were taken using a Zeiss Airyscan LSM 880 30 followed by image processing in Zen 2.3 (Zeiss). Bacterial Small RNA Sequencing and Analysis

E. coli DH5α cells were transformed with pACYC184 carrying the CRISPR-Cas13d locus derived from an uncultured Ruminococcus sp. strain. Cells were harvested in 35 stationary phase, rinsed in PBS, resuspended in TRIzol (Life Technologies), transferred to Lysing Matrix B tubes containing 0.1 mm silica beads (MP Biomedicals), and homogenized on a Bead Mill 24 (Fisher Scientific) for three 30-second cycles. Total RNA was isolated by phenol-chlo- 40 roform extraction, then purified using the DirectZol Miniprep Kit (Zymo Research). RNA quality was assessed on an Agilent 2200 Tapestation followed by Turbo DNase treatment (Ambion). Total RNA was treated with T4 Polynucleotide Kinase (NEB) and rRNA-depleted using the Ribo-Zero 45 rRNA Removal Kit for bacteria (Illumina). RNA was treated with RNA 5' polyphosphatase, poly(A)-tailed with E. coli poly(A) polymerase, and ligated with 5' RNA sequencing adapters using T4 RNA ligase 1 (NEB). cDNA was generated via reverse transcription using an oligo-dT primer and 50 M-MLV RT/RNase Block (AffinityScript, Agilent) followed by PCR amplification and barcoding. Resulting libraries were sequenced on Illumina MiSeq, demultiplexed using custom Python scripts, and aligned to the Cas13d CRISPR locus using Bowtie 2. Alignments were visualized with 55 Geneious.

Ngn2 Lentivirus Preparation

Low passage HEK 293FT cells were transfected with Polyethylenimine Max (PEI, Polysciences) and Ngn2 target plasmid plus pMDG.2 and psPAX2 packaging plasmids (a 60 gift from Didier Trono, Addgene #12259 and #12260) in DMEM +10% FBS media during plating. The following day, media was changed to serum-free chemically defined minimal medium (Ultraculture supplemented with Glutamax, Lonza). Viral supernatant was harvested 48 h later, 65 clarified through a 0.45 micron PVDF filter (Millipore) and concentrated using ultracentrifugation.

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AAV preparation

Low passage HEK 293FT cells were transfected with Polyethylenimine Max (PEI, Polysciences) and AAV target plasmid plus AAV1 serotype and pAdDeltaF6 helper packaging plasmids (UPenn Vector Core) in DMEM+10% FBS media during plating. The following day, 60% of the media was changed to chemically defined minimal medium (Ultraculture supplemented with Glutamax, Lonza). 48 h later, AAV-containing supernatant was harvested and clarified through a 0.45 μm PVDF filter (Millipore) and concentrated using precipitation by polyethylene glycol (PEG virus precipitation kit #K904, Biovision) following the manufacturer's protocol.

RNA-Seq Library Preparation and Sequencing

48 h after transfection, total RNA was extracted from 293FT cells using the RNeasy Plus Mini kit from Qiagen. Stranded mRNA libraries were prepared using the NEBNext II Ultra Directional RNA Library Prep Kit from New England Biolabs (Cat# E7760S) and sequenced on an Illumina NextSeq500 with 42 nt paired end reads. ~15M total reads were demultiplexed per condition.

RNA-Seq Analysis

Sequenced reads were quality-tested using FASTQC and aligned to the hg19 human genome using the 2.5.1b STAR aligner (Dobin et al., 2013). Mapping was carried out using default parameters (up to 10 mismatches per read, and up to 9 multi-mapping locations per read). The genome index was constructed using the gene annotation supplied with the hg19 Illumina iGenomes collection (Illumina) and sidbOverhang value of 100. Uniquely mapped reads were quantified across all gene exons using the top-expressed isoform as proxy for gene expression with the HOMER analysis suite (Heinz et al., 2010), and differential gene expression was carried out with DESeq2 v 1.14.1 (Love et al., 2014) using triplicates to compute within-group dispersion and contrasts to compare between targeting and nontargeting conditions. Significant differentially expressed genes were defined as having a false discovery rate (FDR) <0.01 and a log2 fold change >0.75. Volcano plots were generated in R 3.3.2 using included plotting libraries and the alpha() color function from the scales 0.5.0 package. Statistics

All values are reported as mean±SD or mean±SEM as indicated in the appropriate figure legends. For comparing two groups, a one-tailed student's t-test was used and statistical significance was determined using the Holm-Sidak method with alpha=0.05. A one-way ANOVA with Tukey multiple hypothesis correction was used to assess significance between more than two groups. Two-way ANOVA was used when comparing across two factors (i.e., RNA targeting modality and guide position) and adjusted for multiple hypothesis correction by Sidak's multiple comparisons test. For comparing groups that were found to not meet the assumption of a normal distribution by a D'Agostino and Pearson normality test, the non-parametric Friedman test with Dunn's multiple comparison adjustment was performed. PRISM 7.0 was used for all statistical analysis. Sample sizes were not determined a priori. At least three biological replicates were used for each experiment, as indicated specifically in each figure.

Sequencing data reported herein can be found in the NCBI Gene Expression Omnibus under GEO Series accession number GSE108519.

Additional details on the materials and methods used, such as sequences (e.g., Tables S1 to S5), can be found in Konermann et al., Cell 173:1-12, 2018, herein incorporated by reference in its entirety.

EXAMPLE 2

Computational Identification of a Type VI-Like Cas Ribonuclease Family

This example describes methods used to identify previously undetected or uncharacterized RNA-targeting CRISPR-Cas systems by developing a computational pipeline for class 2 CRISPR-Cas loci, which require only a single nuclease for CRISPR interference such as Cas9, 10 Cas12a (formerly Cpf1), or Cas13a (formerly C2c2) (Makarova et al., 2015; Shmakov et al., 2015). To improve upon previous strategies for bioinformatic mining of CRISPR systems, which focus on discovering sets of conserved Cas genes involved in spacer acquisition (Shmakov et al., 2015), 15 the minimal requirements for a CRISPR locus to be the presence of a CRISPR repeat array and a nearby effector nuclease were defined. Using the CRISPR array as a search anchor, all prokaryotic genome assemblies and scaffolds were obtained from the NCBI WGS database and adapted 20 algorithms for de novo CRISPR array detection (Bland et al., 2007; Edgar, 2007; Grissa et al., 2007) to identify 21,175 putative CRISPR repeat arrays (FIG. 1A).

Up to 20 kilobases (kb) of genomic DNA sequence flanking each CRISPR array was extracted to identify predicted protein-coding genes in the immediate vicinity. Candidate loci containing signature genes of known class 1 and class 2 CRISPR-Cas systems such as Cas3 or Cas9 were excluded from further analysis, except for Cas12a and Cas13a to judge the ability of the pipeline to detect and 30 cluster these known class 2 effector families To identify new class 2 Cas effectors, it was required that candidate proteins be >750 residues in length and within 5 protein-coding genes of the repeat array, as large proteins closely associated with CRISPR repeats are key characteristics of known single 35 effectors. The resulting proteins were classified into 408 putative protein families using single-linkage hierarchical clustering based on homology.

To discard protein clusters that reside in close proximity to CRISPR arrays due to chance or overall abundance in the 40 genome, additional homologous proteins to each cluster were identified from the NCBI non-redundant protein database and their proximity to a CRISPR array determined. Reasoning that true Cas genes would have a high cooccurrence rate with CRISPR repeats, >70% of the proteins 45 for each expanded cluster were required to exist within 20 kb of a CRISPR repeat. These remaining protein families were analyzed for nuclease domains and motifs.

Among the candidates, which include the recently described Cas13b system (Smargon et al., 2017), a family of 50 uncharacterized putative class 2 CRISPR-Cas systems encoding a candidate CRISPR-associated ribonuclease containing 2 predicted HEPN ribonuclease motifs (Anantharaman et al., 2013) were identified (FIG. 2A). Importantly, they are among the smallest class 2 CRISPR effectors 55 described to date (~930 aa). The Type VI CRISPR-Cas13 superfamily is exemplified by sequence-divergent, single-effector signature nucleases and the presence of two HEPN domains. Other than these two RxxxxH HEPN motifs (FIG. 3A), the candidate effectors have no significant sequence 60 similarity to previously described Cas13 enzymes, so his family of putative CRISPR ribonucleases is designated as Type VI Cas13d, or Type VI-D (FIG. 3B).

CRISPR-Cas13d systems are derived from gut-resident microbes, so we sought to expand the Cas13d family via 65 alignment to metagenomic contigs from recent large-scale microbiome sequencing efforts. Comparison of Cas13d pro-

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teins against public metagenome sequences without predicted open reading frames (ORFs) identified additional full-length systems as well as multiple effector and array fragments that cluster in several distinct branches (FIG. 1B). To generate full-length Cas13d ortholog proteins and loci from the different branches of the Cas13d protein family, genomic DNA samples we43 obtained from associated assemblies and performed targeted Sanger sequencing to fill in gaps due to incomplete sequencing coverage, such as for the metagenomic ortholog 'Anaerobic digester metagenome' (Adm) (Treu et al., 2016).

Cas13d CRISPR loci are largely clustered within benign, Gram-positive gut bacteria of the genus Ruminococcus, and exhibit a surprising diversity of CRISPR locus architectures (FIG. 2A). With the exception of the metagenomic AdmCas13d system, Cas13d systems lack the key spacer acquisition protein Cas1 (Yosef et al., 2012) within their CRISPR locus, highlighting the utility of a class 2 CRISPR discovery pipeline without Cas1 or Cas2 gene requirements. Cas13d direct repeats (DRs) are highly conserved in length and predicted secondary structure (FIG. 3C), with a 36 nt length, an 8-10 nt stem with A/U-rich loop, and a 5'-AAAAC motif at the 3' end of the direct repeat (FIG. 3D). This conserved 5'-AAAAC motif has been shown to be specifically recognized by a type II Cast/2 spacer acquisition complex (Wright and Doudna, 2016). In fact, Cas1 can be found in relative proximity to some Cas13d systems (within 10-30 kb for P1E0 and Rfx) while the remaining Cas13dcontaining bacteria contain Cas1 elsewhere in their genomes, likely as part of another CRISPR locus.

EXAMPLE 3

CRISPR-Cas13d Possesses Dual RNase Activities

To demonstrate that the Cas13d repeat array is transcribed and processed into CRISPR guide RNAs (gRNA), the Cas13d CRISPR locus was cloned from an uncultured *Ruminococcus* sp. sample (Ur) into a bacterial expression plasmid. CRISPR systems tend to form self-contained operons with the necessary regulatory sequences for independent expression, facilitating heterologous expression in *E. coli* (Gasiunas et al., 2012). RNA sequencing (Heidrich et al., 2015) revealed processing of the array into ~52 nt mature gRNAs, with a 30 nt 5' direct repeat followed by a variable 3' spacer that ranged from 14-26 nt in length (FIG. 2B).

To characterize Cas13d properties in vitro, Eubacterium siraeum Cas13d protein (EsCas13d) was purified based on its robust recombinant expression in E. coli (FIGS. 4A-4C) and found that EsCas13d was solely sufficient to process its matching CRISPR array into constituent guides without additional helper ribonucleases (FIG. 2C, Table S1 of Konermann et al., "Transcriptome Engineering with RNA-Targeting Type VI-D CRISPR Effectors," Cell 173:1-12, 2018, herein incorporated by reference in its entirety), a property shared by some class 2 CRISPR-Cas systems (East-Seletsky et al., 2016; Fonfara et al., 2016; Smargon et al., 2017). Furthermore, inactivating the positively charged catalytic residues of the HEPN motifs (Anantharaman et al., 2013) (dCas13d: R295A, H300A, R849A, H854A) did not affect array processing, indicating a distinct RNase activity dictating gRNA biogenesis analogous to Cas13a (East-Seletsky et al., 2016; Liu et al., 2017).

Cas effector proteins typically form a binary complex with mature gRNA to generate an RNA-guided surveillance ribonucleoprotein capable of cleaving foreign nucleic acids for immune defense (van der Oost et al., 2014). To assess if

Cas13d has programmable RNA targeting activity as indicated by the presence of two HEPN motifs, EsCas13d protein was paired with an array or a mature gRNA along with a cognate in vitro-transcribed target. Based on the RNA sequencing results, a mature gRNA containing a 30 nt direct repeat and an intermediate spacer length of 22 nt was selected (nucleotides 6-36 of SEQ ID NO: 129, followed by 22 bases complementary to the RNA target).

Cas13d was able to efficiently cleave the complementary target ssRNA with both the unprocessed array and mature gRNA in a guide-sequence dependent manner, while non-matching spacer sequences abolished Cas13d activity (FIG. 5A). Substitution with dCas13d or the addition of EDTA to the cleavage reaction also abolished guide-dependent RNA targeting, indicating that Cas13d targeting is HEPN- and Mg²⁺-dependent (FIG. 5B). To determine the minimal spacer length for efficient Cas13d targeting, a series of spacer truncations ranging from the unprocessed 30 nt length down to 10 nt were generated (FIG. 6A). Cleavage 20 activity dropped significantly below a 21 nt spacer length, confirming the choice of a 22 nt spacer (FIG. 6B).

RNA-targeting class 2 CRISPR systems have been proposed to act as sensors of foreign RNAs (Abudayyeh et al., 2016; East-Seletsky et al., 2016), where general RNase 25 activity of the effector nuclease is triggered by a guidematching target. To assay for a similar property in Cas13d, RNase activity of the binary EsCas13d:gRNA complex was monitored in the presence of a matching RNA target. It was observed that EsCas13d can be activated by target RNA to cleave bystander RNA targets (FIG. 3C), albeit inefficiently relative to its activity on the complementary ssRNA target. Bystander cleavage is guide sequence- and HEPN-dependent, as the presence of non-matching bystander target alone was insufficient to induce cleavage while substitution of dCas13d or addition of EDTA abolished activity. These results indicate that bystander RNase activity may be a general property of RNA-targeting class 2 systems in CRISPR adaptive bacterial immunity (FIG. 3D).

To assess the generalizability of Cas13d reprogramming, twelve guides tiling a complementary RNA target were generated and efficient cleavage in all cases was observed (FIG. 7A). Cas13d was unable to cleave a ssDNA (FIG. 6C) or dsDNA (FIG. 6D) version of the ssRNA target, indicating 45 that Cas13d is an RNA-specific nuclease. Further, RNA target cleavage did not depend on the protospacer flanking sequence (PFS) (FIG. 7A) in contrast to other RNA-targeting class 2 systems, which require a 3'-H (Abudayyeh et al., 2016) or a double-sided, DR-proximal 5'-D and 3'-NAN or 50 NNA (Smargon et al., 2017). Although a slight bias against an adenine PFS was initially observed (FIG. 6E), varying the target PFS base with a constant guide sequence resulted in no significant differences (P=0.768) in targeting efficiency (FIG. 6F).

While DNA-targeting class 2 CRISPR systems (Gasiunas et al., 2012; Jinek et al., 2012; Zetsche et al., 2015) and some RNA-targeting class 1 systems tend to cleave at defined positions relative to the target-guide duplex (Samai et al., 2015; Zhang et al., 2016), the Cas13d cleavage pattern ovaries for different targets (FIGS. 5A, 5C, 6H) and remains remarkably similar despite the guide sequence position (FIG. 7A). This indicates that Cas13d may preferentially cleave specific sequences or structurally accessible regions in the target RNA. Cas13d was tested activity on targets of containing variable homopolymer repeats in the loop region of a hairpin or as a linear single-stranded repeat. EsCas13d

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exhibited significant preference for uracil bases in both target structures, with lower but detectable activity at all other bases (FIG. 7B).

Cas enzymes are found in nearly all archaea and about half of bacteria (Hsu et al., 2014; van der Oost et al., 2014), spanning a wide range of environmental temperatures. To determine the optimal temperature range for Cas13d activity, a spectrum of cleavage temperature conditions from 16-62° C. was tested and observed maximal activity in the 24-41° C. range (FIG. 6G, 6H). This temperature range is compatible with a wide range of prokaryotic and eukaryotic hosts, indicating Cas13d can be adapted for RNA targeting in different cells and organisms.

EXAMPLE 4

Cell-Based Activity Screen of Engineered Orthologs

The Cas13d nuclease was used as a flexible tool for programmable RNA targeting in mammalian cells. CRISPR orthologs from distinct bacterial species commonly exhibit variable activity (Abudayyeh et al., 2017; East-Seletsky et al., 2017), especially upon heterologous expression in human cells (Ran et al., 2015; Zetsche et al., 2017). Highly active Cas13d orthologs were identified in a eukaryotic cell-based mCherry reporter screen.

By synthesizing human codon-optimized versions of 7 orthologs from distinct branches within the Cas13d family (FIG. 1B), mammalian expression plasmids carrying the catalytically active and HEPN-inactive proteins were generated. Each protein was then optionally fused to N- and C-terminal nuclear localization signals (NLS). These Cas13d effector designs were HA-tagged and paired with two distinct guide RNA architectures, either with a 30 nt spacer flanked by two direct repeat sequences to mimic an unprocessed guide RNA (pre-gRNA) or a 30 nt direct repeat with 22 nt spacer (gRNA) predicted to mimic mature guide RNAs (FIG. 8A). For each guide design, four distinct spacer sequences complementary to the mCherry transcript were then pooled to minimize potential spacer-dependent variability in targeting efficiency. The ability of Cas13d to knockdown mCherry protein levels was determined in a human embryonic kidney (HEK) 293FT cell-based reporter assay.

48 hours post-transfection, flow cytometry indicated that RfxCas13d and AdmCas13d efficiently knocked down mCherry protein levels by up to 92% and 87% (P<0.0003), respectively, relative to a non-targeting control guide (FIG. 8B). In contrast, EsCas13d along with RaCas13d and RffCas13d exhibited limited activity in human cells. Furthermore, none of the HEPN-inactive Rfx-dCas13d constructs significantly affected mCherry fluorescence, indicating HEPN-dependent knockdown (P>0.43 for all cases). Robust nuclear translocation of the Rfx and AdmCas13d NLS fusion constructs was observed via immunocytochemistry, while the wild-type effectors remain primarily extranuclear (FIG. 8C).

Proceeding with RfxCas13d and AdmCas13d as lead candidates, we next compared their ability to knockdown endogenous transcripts. To determine the optimal ortholog and guide architecture, the capability of Rfx and AdmCas13d construct variants to target β -1,4-N-acetylgalactosaminyl transferase 1 (B4GALNT1) transcripts was systematically assayed. In each condition, four guides containing distinct spacer sequences tiling the B4GALNT1 transcript were pooled. The RfxCas13d-NLS fusion targeted

B4GALNT1 more efficiently than wild-type RfxCas13d and both variants of AdmCas13d, with both the gRNA and pre-gRNA mediating potent knockdown (~82%, P<0.0001) (FIG. 8D). Cas13d-NLS from *Ruminococcus flavefaciens* strain XPD3002 was therefore selected for the remaining 5 experiments (CasRx).

EXAMPLE 5

Programmable RNA Knockdown in Human Cells with CasRx

Because Cas13d is capable of processing its own CRISPR array, this property was leveraged for the simultaneous delivery of multiple targeting guides in a simple single-vector system (FIG. 9A). Arrays encoding four spacers that each tile the transcripts of mRNAs (B4GALNT1 and ANXA4) or nuclear localized lncRNAs (HOTTIP and MALAT1) consistently facilitated robust (>90%) RNA knockdown by CasRx (P<0.0001) (FIG. 9B).

CasRx was compared to more established technologies for transcript knockdown or repression, by comparing CasRx-mediated RNA interference to dCas9-mediated CRISPR interference (Gilbert et al., 2014; Gilbert et al., 2013) and spacer sequence-matched shRNAs via transient 25 transfection (FIG. 9C). For CRISPRi-based repression, the most potent dCas9 guide for B4GALNT1 from previous reports was analyzed (Gilbert et al., 2014; Zalatan et al., 2015). Across 3 endogenous transcripts, CasRx outperformed shRNAs (11/11) and CRISPRi (4/4) in each case 30 (FIG. 9D), exhibiting a median knockdown of 96% compared to 65% for shRNA and 53% for CRISPRi after 48 hours. In addition, knockdown by CasRx was compared to two recently described Cas13a and Cas13b effectors (Abudayyeh et al., 2017; Cox et al., 2017) (FIG. 10A). Across 35 three genes and eight guide RNAs, CasRx mediated significantly greater transcript knockdown than both LwaCas13amsfGFP-NLS and PspCas13b-NES (median: 97% compared to 80% and 66% respectively, P<0.0001) (FIG. 10B).

RNAi has been widely used to disrupt any gene of interest 40 due to a combination of simple re-targeting principles, scalable synthesis, knockdown potency, and ease of reagent delivery. However, widespread off-target transcript silencing has been a consistent concern (Jackson et al., 2003; Sigoillot et al., 2012), possibly due to the entry of RNAi reagents into 45 the endogenous miRNA pathway (Doench et al., 2003; Smith et al., 2017). Consistent with these reports, upon RNA sequencing of human cells transfected with a B4GALNT1targeting shRNA, widespread off-target transcriptional changes relative to a non-targeting shRNA were observed 50 (>500 significant off-target changes, P<0.01, FIGS. 9E, 9G). In contrast, transcriptome profiling of spacer-matched CasRx guide RNAs revealed no significant off-target changes other than the targeted transcript (FIG. 9F). This indicates that the moderate bystander cleavage observed in 55 vitro (FIG. 5C) may not result in observable off-target transcriptome perturbation in mammalian cells. A similar pattern was observed when targeting ANXA4 (FIGS. 11A-11B), with over 900 significant off-target changes resulting from shRNA targeting compared to zero with CasRx (FIG. 60

To confirm that CasRx interference is broadly applicable, a panel of 11 additional genes with diverse roles in cancer, cell signaling, and epigenetic regulation were selected and 3 guides per gene were screened. CasRx consistently mediated 65 high levels of transcript knockdown across genes with a median reduction of 96% (FIG. 9H). Each tested guide

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mediated at least 80% knockdown, underscoring the consistency of the CasRx system for RNA interference.

EXAMPLE 6

Splice Isoform Engineering with dCasRx

The experiments on RNA targeting with CasRx revealed that target RNA and protein knockdown is dependent on the catalytic activity of the HEPN domains (FIGS. 8B, 5B). The same guide sequences mediating efficient knockdown with CasRx failed to significantly reduce mCherry levels when paired with catalytically inactive dCasRx (FIG. 8B), indicating that targeting of dCasRx to the coding portion of mRNA does not necessarily perturb protein translation. This observation indicated the possibility of utilizing dCasRx for targeting of specific coding and non-coding elements within a transcript to study and manipulate RNA. To validate this concept, the utility of the dCasRx system was expanded by creating a splice effector.

Alternative splicing is generally regulated by the interaction of cis-acting elements in the pre-mRNA with positive or negative trans-acting splicing factors, which can mediate exon inclusion or exclusion (Matera and Wang, 2014; Wang et al., 2015). It was reasoned that dCasRx binding to such motifs may be sufficient for targeted isoform perturbation. For proof-of-concept, distinct splice elements were identified in a bichromatic splicing reporter containing DsRed upstream of mTagBFP2 in two different reading frames following an alternatively spliced exon (Orengo et al., 2006) (FIG. 12A). Inclusion or exclusion of this second exon toggles the reading frame and resulting fluorescence, facilitating quantitative readout of splicing patterns by flow cytometry. To mediate exon skipping, four guide RNAs were designed to target the intronic branchpoint nucleotide, splice acceptor site, putative exonic splice enhancer, and splice donor of exon 2.

One widespread family of negative splice factors are the highly conserved heterogeneous nuclear ribonucleoproteins (hnRNPs), which typically inhibit exon inclusion via a C-terminal, glycine-rich domain (Wang et al., 2015). The splicing reporter was targeted with dCasRx and engineered fusions to the Gly-rich C-terminal domain of hnRNPa1, one of the most abundant hnRNP family members (FIG. 12B).

Guide position appears to be a major determinant of the efficiency of engineered exon skipping. While each guide position mediated a significant increase in exon exclusion (P<0.0001 in all cases) relative to the non-targeting guide, targeting the splice acceptor resulted in the most potent exon exclusion (increase from 8% basal skipping to 65% for dCasRx alone and 75% with hnRNPa1 fusion). By comparison, dLwaCas13a-msfGFP-NLS mediated significantly lower levels of exon skipping across all four positions (19% skipping for splice acceptor guide) (FIGS. 10C and 10D, P<0.0001).

Targeting all 4 positions simultaneously with a CRISPR array achieved higher levels of exon skipping than individual guides alone (81% for dCasRx and 85% for hnRNPa1 fusion, P<0.006 compared to SA guide) (FIG. 12B). These results indicate that dCasRx allows for tuning of isoform ratios through varying guide placement and suggest that it can be leveraged as an efficient RNA binding module in human cells for targeting and manipulation of specific RNA elements.

EXAMPLE 7

Viral Delivery of dCasRx to a Neuronal Model of Frontotemporal Dementia

The Cas13d family averages 930 amino acids in length, in contrast to Cas9 (~1100 aa to ~1400 aa depending on subtype, with compact outliers such as CjCas9 or SaCas9), Cas13a (1250 aa), Cas13b (1150 aa), and Cas13c (1120 aa) (FIG. 3B) (Chylinski et al., 2013; Cox et al., 2017; Hsu et 10 al., 2014; Kim et al., 2017; Shmakov et al., 2015; Smargon et al., 2017). Although adeno-associated virus (AAV) is a versatile vehicle for transgene delivery and gene therapy due to its broad range of capsid serotypes, low levels of insertional mutagenesis, and lack of apparent pathogenicity, its 15 limited packaging capacity (~4.7 kb) makes it challenging to effectively deliver many single effector CRISPR enzymes (Abudayyeh et al., 2017; Ran et al., 2015; Swiech et al., 2015). The remarkably small size of Cas13d effectors render them uniquely suited for all-in-one AAV delivery with a 20 CRISPR array, an optional effector domain, and requisite expression or regulatory elements (FIG. 12C).

Frontotemporal Dementia with Parkinsonism linked to Chromosome 17 (FTDP-17) is an autosomal dominant major neurodegenerative disease caused by diverse point 25 to uncultured Ruminococcus sp. Cas13d. mutations in MAPT, the gene encoding for tau. Tau exists as two major isoforms in human neurons, 4R and 3R, which are distinguished by the presence or absence of tau exon 10 and thus contain 4 or 3 microtubule binding domains. The balance of these two isoforms is generally perturbed in 30 FTDP-17 as well as other tauopathies, driving the progression of neurodegeneration (Boeve and Hutton, 2008). Some forms of FTD are caused by mutations in the intron following MAPT exon 10 which disrupt an intronic splice silencer and elevate the expression of 4R tau (Kar et al., 2005), 35 thereby inducing pathological changes (Schoch et al., 2016).

It was reasoned that dCasRx targeted to MAPT exon 10 could induce exon exclusion to alleviate dysregulated 4R/3R tau ratios. Patient-derived human induced pluripotent stem cells (hiPSCs) were differentiated into cortical neurons via 40 Neurogenin-2 directed differentiation for 2 weeks (Zhang et al., 2013). Postmitotic neurons were then transduced with AAV1 carrying dCasRx (FIG. 12D) paired with a repeat array containing 3 spacers that target the exon 10 splice acceptor and two putative exonic splice enhancers (FIG. 45 to mediate target knock-down. 12E). dCasRx-mediated exon exclusion was able to reduce the relative 4R/3R tau ratio by nearly 50% relative to a BFP vehicle control (FIG. 12F) and to a level similar to unaffected control neurons, demonstrating that CasRx can be exploited for transcriptional modulation in primary cell 50 types via AAV delivery.

EXAMPLE 8

RNA Targeting in Human Cells Using Cas13d

RNA can be targeted in human cells using the active Cas13d nuclease. As a proof of concept, human U-2 OS bone osteosarcoma cells were stably integrated with an mCherry reporter and transfected with plasmids encoding 60 human codon optimized Cas13d and guide RNAs targeting the mCherry transcript (FIG. 13).

Cas13d proteins were also fused with N- and C-terminal NLS sequences (SPKKKRKVEAS, SEQ ID NO: 256, for the N-terminal NLS and GPKKKRKVAAA, SEQ ID NO: 65 258, C-terminal NLS) to understand if nuclear localization can affect mCherry knockdown (these are denoted the 2x

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NLS constructs). Guide RNAs were either provided in a vector with a U6 promoter operably linked to a 36 nt DR-30 nt spacer-36 nt DR sequence, which mimics the unprocessed CRISPR guide array (denoted DR36), or to a 30nt DR-22 nt spacer to mimic the processed, mature gRNA (denoted gRNA). The DR36 construct is presumed to be processed by Cas13d into mature gRNAs within the cell. The spacer sequences within the DR36 or gRNA molecules were either complementary to the mCherry target RNA (on-target mCherry) or computationally optimized to avoid complementarity to mCherry or any endogenous human transcript (non-targeting mCherry).

mCherry knockdown was quantified via flow cytometry and normalized to a transfection control. The non-targeting mCherry guides do not affect mCherry protein levels via flow cytometry, presumably because the mCherry transcripts are not targeted. However, the on-target mCherry guides paired with 4 different Cas13d orthologues exhibited significant mCherry knockdown (FIG. 13). "XPD" refers to Ruminococcus flavefaciens XPD3002 Cas13d (SEQ ID NO: 92); "P1E0" refers to Gut metagenome P1E0 Cas13d (SEQ ID NO: 83); "AnDig" refers to Anaerobic digester gut metagenome Cas13d (SEQ ID NO: 42); "Uncultured" refers

EXAMPLE 9

In Vivo RNA Targeting Using Cas13d

RNA can be targeted in mouse models of cancer. To observe which cells in the mouse are expressing EGFR, a guide RNA is designed that includes one or more spacer regions complementary to mouse EGFR and is combined with Cas13d having a mutated HEPN domain (such as SEQ ID NO: 2 or 4), and a biotin label. The gRNA and Cas13d coding sequence are cloned into a viral vector (such as a lentivirus) which is used to infect the mice by tail vein injection at a titer to insure 100% infection rates. A fluorescent streptavidin label is administered to the mice. Cells expressing EGFR are visualized and detected with the appropriate excitation frequency for the fluorescent label. Alternatively, Cas13d is delivered in its active form in vivo

EXAMPLE 10

Treatment of Cancer

Human subjects with histologically confirmed stage 1, EGFR+ breast cancer can be treated with the disclosed methods. Each subject is administered a complex comprising an active Cas13d or a Cas13d protein mutated in the 55 HEPN domain (such as SEQ ID NO: 2 or 4), a guide RNA targeting EGFR, and a toxin, after receiving lumpectomy surgery. Treated individuals are monitored for breast cancer recurrence.

EXAMPLE 11

Treatment of HIV Infection

Human subjects with HIV infection can be treated with the disclosed methods. Each subject is administered a construct comprising an active Cas13d or a Cas13d protein mutated in the HEPN domain (such as SEQ ID NO: 2 or 4),

a guide RNA targeting HIV Nef protein, and a toxin. Treated individuals are monitored for HIV progression.

EXAMPLE 12

Treatment of Huntington's Disease

Human subjects with Huntington's Disease can be treated with the disclosed methods. Each individual is administered a construct comprising Cas13d, a guide RNA targeting the ¹⁰ Huntington mutation. Treated individuals are monitored for disease progression.

EXAMPLE 13

Alternative Splicing Using Cas13d

Cas13d splice effectors can be used for therapeutic protein restoration (for example that results from a mutation or deletion), gene knockdown via frameshift induction, tuning 20 or restoring a desired isoform ratio, or inducing a desired dominant splice isoform (FIG. 14). Alternative splicing is generally regulated by the interaction of cis-acting elements in the pre-mRNA with positive or negative trans-acting splicing factors, which can mediate exon inclusion or exclu- 25 sion. dCas13d and Cas13d, with optional fusion to positive or negative splicing factors, can be used as splicing effectors that target to said cis-acting elements in the pre-mRNA to manipulate splicing. Such elements can include exonic splicing enhancers or suppressors, intronic splicing enhanc- 30 ers or suppressors, splice acceptor and splice donor sites, and more generally protein- or RNA-interacting motifs or elements on that particular pre-mRNA, mRNA, or other RNA species, such as a non-coding RNA, tRNA, miRNA, and the like.

Additionally, that the effects of Cas13d-based splice effectors can be guide position-dependent. This can be exploited to perturb or discover particular motifs or sites in an RNA transcript such as protein-binding sites, via steric hindrance, blocking, recruitment, or effector-mediated interaction. For example, the interaction between non-coding RNAs and particular chromatin remodeling complexes can be perturbed. Access of the ribosomal binding site and other elements can be blocked in a 5' or 3' UTR (or to recruit appropriate effector domains) to decrease, increase, or otherwise manipulate translation.

Targeting or tiling Cas13d guides along a pre-mRNA can be used to discover or map new cis-acting elements such as intronic or exonic splice enhancers. This has been exploited in a therapeutic context in the case of the dystrophin gene for optimal antisense oligonucleotide positioning and can also be used for optimal Cas13d positioning. This can be also used to map, mask, or otherwise perturb RNA zipcodes or other cis-acting elements to affect trafficking and localization, chromatin remodeling, polyadenylation, RNA stability 55 and half-life, or levels of nonsense-mediated decay.

In one example, targeting an RNA allows for changing splicing of the target RNA. Both the direct binding of splice acceptor and/or donor sites as well as splice effector domains can be used to manipulate splicing. For example, by using a 60 dCas13d protein with a mutated HEPN domain (e.g., SEQ ID NO: 2 or 4), a guide RNA containing at least one a spacer sequence specific for the target RNA, and optionally an effector domain that affects splicing (such as the RS-rich domain of SRSF1, the Gly-rich domain of hnRNPA1, the 65 alanine-rich motif of RBM4, or the proline-rich motif of DAZAP1), alternative splicing of the RNA can be achieved.

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In some examples, such a method is used for exon inclusion, for example to include exon 2 of acid alphaglucosidase (GAA) to treat Pompe disease or to include exon 7 of SMN2 to treat spinal muscular atrophy (SMA). In some examples, such a method is used for exon exclusion, for example to restore the reading frame of dystrophin to treat Duchenne muscular dystrophy, to shift the splicing of the Bcl-x pre-mRNA from the antiapoptotic long isoform to the proapoptotic short isoform to treat cancer, to shift the splicing of the MAPT transcript to affect ratios of 3R and 4R tau, or to manipulate the splicing of the lamin A transcript in the case of Hutchinson-Gilford progeria syndrome or other genetic diseases of aging.

In some examples, the method uses an Cas13d protein,
optionally with a mutated HEPN domain, to mask splice
acceptor or donor sites, for example to create neoantigens to
make cold tumors hot. By affecting the splicing of certain
target pre-mRNAs, this method can generate novel exonexon junctions that can lead to the creation of neo-epitopes
in cancer cells. This can make a cancer cell vulnerable to the
immune system due to the display of unnatural antigens. In
other examples, this method can be used to dynamically
manipulate isoform ratios or to restore reading frame of a
protein (e.g., dystrophin for Duchenne's muscular dystrophy).

EXAMPLE 14

AAV Delivery of Cas13d

As described in the examples above, Cas13d can be effectively packaged into AAV to mediate expression in cell types that are not amenable to plasmid delivery or for in vivo delivery of Cas13d. AAV delivery of nuclease active Cas13d can be used to mediate RNA target knock-down in the cell type of interest. Due to its small size compared to other single-effector CRISPR nucleases, Cas13d can be packaged together with a guide RNA or an array containing multiple guide RNAs in a single AAV vector.

EXAMPLE 15

Nucleic Acid-Based Diagnostics with Cas13d

Cas13d enzymes can be exploited for nucleic acid-based diagnostics within the context of a cell, using cell-free lysate derived from a cell, or a cell-free system containing an engineered Cas13d enzyme and guide RNA to facilitate formation of a ribonucleoprotein complex. Said guide RNA can be provided in the form of a pre-guide RNA, a mature guide RNA, or an array containing one or more spacer sequences. The components can also be provided in the form of a DNA or RNA precursor encoding for the Cas13d enzyme and appropriate guide RNA design via an in vitro transcription/translation system to facilitate the generation of the necessary components. These components of the diagnostic kit comprise the "sensor" module.

Such a method can be used to determine if a target RNA is present in a test sample. Such a method can also be used to detect a pathogen, such as a virus or bacteria, or diagnose a disease state, such as a cancer (e.g., wherein the target RNA is specific for a particular microbe or disease). Such a method can also be used to test the purity or identity of an environmental sample or agricultural sample, such as seed or soil

The "sensor" module will then be challenged with a test sample in the form of RNA. Said test sample can be, but is

not limited to, a genomic DNA sample that is converted into RNA—for example via in vitro transcription—or a direct RNA sample. These samples can be extracted from biological material such as patient samples (e.g., cells, tissue, blood, plasma, serum, saliva, urine, tumor biopsy, cell free 5 DNA or RNA, exosomes, carrier vesicles or particles) and environmental samples (e.g., soil, water, air, seed, or plant samples). In one embodiment to improve diagnostic sensitivity, nucleic acid molecules in the sample are amplified using amplification techniques, such as polymerase chain 10 reaction, recombinase polymerase amplification, loop mediated isothermal amplification, nucleic acid sequence based amplification, strand displacement amplification, rolling circle amplification, ligase chain reaction, and others (e.g., those that use isothermal amplification). Said amplification 15 techniques can optionally employ nucleic acid conversion techniques such as transcription or reverse transcription with randomized primers or targeted primers.

If the sensor module recognizes a cognate target in the test sample, it will activate an RNase activity. This RNase 20 activity can be detected by using a detectable label. In one example, the detectable label includes an RNA linked to a fluorophore and quencher. The intact detectable RNA links the fluorophore and quencher, suppressing fluorescence. Upon cleavage by Cas13d of the detectable RNA, the 25 fluorophore is released from the quencher and displays detectable fluorescent activity.

In another example, cleavage of the reporter RNA releases a non-fluorescent molecule, which can be converted into a visible signal (e.g., visible by eye). In one example, cleavage 30 of the reporter RNA releases a molecule that can be detected via lateral flow. A molecule that can be detected by lateral flow is any molecule that can be bound specifically by antibodies. In one example, the Cas13d protein along with the guide RNA detecting the target and the reporter RNA 35 conjugated to the reporter molecule can be delivered as a single system in the form of a dry test strip. Upon incubation with the test sample, Cas1Cas13d protein a3d, guide RNA and the reporter RNA are rehydrated and in the presence of the RNA target. The Cas13d protein will cleave the reporter 40 RNA, resulting in the migration of the reporter molecule in the test strip via lateral flow and a resulting positive test line signal by binding to antibodies localized there. Such a shelf stable dry detection system not requiring special (frozen) storage could for example prove advantageous in situations 45 where detection of a target RNA or DNA is performed outside a centralized laboratory facility such as a doctor's office, a hospital, a pharmacy, during field work, in an agricultural setting and so forth.

Cas13d is active over a broad range of temperatures 50 making such an application outside of a controlled laboratory environment feasible.

EXAMPLE 16

Cas13d as a Diagnostic for RNA or DNA Transcribed into RNA In Vitro

Cas13d is capable of converting the presence of a matched target RNA into a visible signal in a minimal 60 diagnostic in vitro system. FIGS. **15**A-**15**D (A) Cas13d is converted into an active RNase complex upon binding a target matching the spacer sequence of the guide RNA. It is capable of cleaving gRNA-complementary target RNA or non-complementary bystander RNAs. (B) Cas13d target-65 dependent RNase activity can be converted into a detectable signal, for example through cleavage of a labeled detector

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RNA that is cleaved only in the presence of a target matching the spacer of the Cas13d guide RNA. In this example, the detector RNA contains a fluorophore, 'F', and a quencher 'Q', that abolishes fluorescence. Only upon bystander RNA cleavage is the fluorophore liberated from the quencher and fluorescence is generated. (C) Cas13d from *E. siraeum* produces a visible signal only in the presence of a perfectly matched target and not in the presence of different mismatched targets. (D) Cas13d from *R. flavefaciens* strain XPD3002 produces a visible signal only in the presence of a perfectly matched target and not in the presence of different mismatched target and not in the presence of different mismatched targets.

Thus, the system disclosed herein can be part of a lateral flow device (or other solid support), that can be used in diagnostics. The presence of an RNA or DNA sequence can be converted into a signal than can then be detected by conventional lateral flow.

EXAMPLE 17

Cas13d Modifications

FIGS. **16**A-**16**B, Cas13d is amenable to modifications including truncations of regions with low conservation among orthologs. The alignment of Cas13d orthologs in FIG. **16**A shows regions with high (green bars) and low (red bars) conservation.

EXAMPLE 18

Targeting Transcripts In Vivo

The ccdB gene in bacterial cells was targeted in vivo using different nCas1 orthologues, *Eubacterium siraeum* nCas1 (Es_nCas1; SEQ ID NO: 1); *Eubacterium siraeum* nCas1 with mutated HEPN domains (Es_nCas1 HEPN -/-; SEQ ID NO: 2); uncultured *Ruminococcus* sp. nCas1 (uncul_nCas1; SEQ ID NO: 3) and uncultured *Ruminococcus* sp. nCas1 with mutated HEPN domains (uncul_nCas1 HEPN -/-; SEQ ID NO: 4) (FIG. 17A). Chemically competent *E. coli* (strain BW25141-DE3) cells were transformed with (1) an arabinose-inducible ccdB plasmid, and (2) a second plasmid (targeting vector) carrying a compatible origin of replication, nCas1 protein coding sequence, and nCas1 guide array containing 4 spacer sequences targeting the ccdB transcript (FIG. 17A).

1. Aartificial $Eubacterium\ siraeum\ nCas1$ Array Targeting ccdB

a. (SEQ ID NO: 261)
GAACUACACCCGUGCAAAAAUGCAGGGGUCUAAAACUAACGGC
UCUCUUUUUAUAGGUGUAAACCGAACUACACCCGUGCAAAAAU
GCAGGGGUCUAAAACCUUUAUCUGACAGCAGACGUGCACUGGCC
AGAACUACACCCGUGCAAAAAUGCAGGGGUCUAAAACCAUCAU
GCGCCAGCUUUCAUCCCCGAUAUGGAACUACACCCGUGCAAAAA
UGCAGGGGUCUAAAACUAAUGGCGUUUUUGAUGUCAUUUUCGCG
GUCCGCUGA

i. Full 36 nt direct repeat:

55

(SEQ ID NO: 262)

GAACUACACCCGUGCAAAAAUGCAGGGGUCUAAAAC

76-continued

15	
-continued ii. Spacer 1:	
(SEQ ID NO: 263)	
iii. Spacer 2:	5
(SEQ ID NO: 264) CUUUAUCUGACAGCAGACGUGCACUGGCCA	
iv. Spacer 3:	
(SEQ ID NO: 265) CAUCAUGCGCCAGCUUUCAUCCCCGAUAUG	10
v. Spacer 4:	
(SEQ ID NO: 266) UAAUGGCGUUUUUGAUGUCAUUUUCGCGGUCCGCUGA	
2. Artificial Uncultured $\it Ruminoccus$ sp. nCas1 Array Targeting ccdB	15
a.	
(SEQ ID NO: 267) CUACUACACUGGUGCAAAUUUGCACUAGUCUAAAACUAACGGCU	20
CUCUCUUUUAAAGGUGUAAACCCUACUACACUGGUGCAAAUUUG	
CACUAGUCUAAAACCUUUAUCUGACAGCAGACGUGCACUGGCCAC	
UACUACACUGGUGCAAAUUUGCACUAGUCUAAAACCAUCAUGCG	25
CCAGCUUUCAUCCCCGAUAUGCUACUACACUGGUGCAAAUUUGC	
ACUAGUCUAAAACUAAUGGCGUUUUUGAUGUCAUUUUCGCGGUC	
CGC	30
i. Full 36 nt direct repeat:	
(SEQ ID NO: 268)	
ii. Spacer 1:	35
UAACGGCUCUCUUUUAUAGGUGUAAACC (SEQ ID NO: 269)	
iii.	
Spacer 2: (SEQ ID NO: 270)	40
CUUUAUCUGACAGCAGACGUGCACUGGCCA	70
iv. Spacer 3: (SEQ ID NO: 271)	
CAUCAUGCGCCAGCUUUCAUCCCCGAUAUG	
v. Spacer 4: (SEO ID NO: 272)	45
UAAUGGCGUUUUUGAUGUCAUUUUCGCGGUCCGCUGA	
3. Target RNA (ccdB Sequence)	
	50
a. (SEQ ID NO: 273)	
AUGCAGUUUAA GGUUUACACCUAUAAAAGAGAGACCCG<u>UUA</u>UC G	
<u>UCUGUUUGUGGAUG</u> UACAGAGUGAUAUUAUUGACACGCCCGGGCG	55
ACGGAUGGUGAUCCCCCUGGCCAGUGCACGUCUGCUGUCAGAUA	
AAGUCUCCCGUGAACUUUACCCGGUGGUGCAUAUCGGGGAUGAA	
AGCUGGCGCAUGAUGACCACCGAUAUGGCCAGUGUGCCGGUCUC	60
CGUUAUCGGGGAAGAAGUGGCUGAUCUCAGCCACCGCGAAAAUG	
ACAUCAAAAACGCCAUUAACCUGAUGUUUUGGGGAAUA	
i spacer 1 target:	

(SEQ ID NO: 274) 65

i. spacer 1 target:

GGUUUACACCUAUAAAAGAGAGAGCCGUUA

ii. spacer 2 target:

UGGCCAGUGCAGUCUGCUGUCAGAUAAAG

iii. spacer 3 target:

(SEQ ID NO: 276)

CAUAUCGGGGAUGAAAGCUGGCGCAUGAUG

ACCGCGAAAAUGACAUCAAAAACGCCAUUA

v. dCas9 target (underlined): (SEQ ID NO: 259)

The transformed bacteria were plated on 2 mM arabinose plates to induce ccdB expression and harvested after 24 hours. Total RNA was extracted with Trizol followed by random hexamer-mediated reverse transcription and Taqman probe-based qPCR.

Mutation of the two HEPN domains in each nCas1 protein demonstrate no targeting of ccdB (FIG. 17B, HEPN -/-), while active wild-type nCas1 proteins knock downed the expression of ccdB (FIG. 17B, Es_nCas1; uncul_nCas1). Because dCas9-(catalytically inactive, or "dead" Cas9)-mediated transcriptional repression through binding of the DNA sequence downstream of the transcriptional start site is a current standard in the field, we validate the assay by targeting dCas9 to the ccdB promoter to repress transcription of the ccdB gene. Taken together, this data demonstrates guide RNA-specific degradation of target transcripts in vivo within prokaryotic cells in a HEPN domain-dependent manner.

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In view of the many possible embodiments to which the principles of the disclosure may be applied, it should be recognized that the illustrated embodiments are only examples of the invention and should not be taken as limiting the scope of the invention. Rather, the scope of the invention is defined by the following claims. We therefore claim as our invention all that comes within the scope and spirit of these claims.

SEQUENCE LISTING

The patent contains a lengthy "Sequence Listing" section. A copy of the "Sequence Listing" is available in electronic form from the USPTO web site (https://seqdata.uspto.gov/?pageRequest=docDetail&DocID=US11316812B2). An electronic copy of the "Sequence Listing" will also be available from the USPTO upon request and payment of the fee set forth in 37 CFR 1.19(b)(3).

We claim:

- 1. A non-naturally occurring or engineered clustered regularly interspaced short palindromic repeats (CRISPR)-associated (Cas) system for targeting one or more target RNA molecules, comprising:
 - (a) at least one Cas13d protein comprising at least 95% sequence identity to SEQ ID NO: 42, or a nucleic acid molecule encoding the at least one Cas13d protein; and
 - (b) at least one CRISPR-Cas system guide RNA (gRNA) comprising one or more direct repeat (DR) sequences 10 and one or more spacer sequences, which hybridizes with the one or more target RNA molecules, or at least one nucleic acid molecule encoding the gRNA,
 - whereby the Cas13d protein forms a complex with the gRNA, wherein the gRNA directs the complex to the 15 one or more target RNA molecules and targets the one or more target RNA molecules.
 - 2. An isolated cell comprising the system of claim 1.
- 3. A ribonucleoprotein complex comprising the system of claim 1, wherein the ribonucleoprotein complex comprises 20 the (a) at least one Cas13d protein comprising at least 95% sequence identity to SEQ ID NO: 42 and the (b) at least one CRISPR-Cas system gRNA comprising one or more DR sequences and one or more spacer sequences, which hybridizes with the one or more target RNA molecules, 25
 - wherein the gRNA directs the ribonucleoprotein complex to the one or more target RNA molecules and targets the one or more target RNA molecules.
- 4. The system of claim 1, wherein targeting one or more target RNA molecules comprises one or more of cutting the 30 one or more target RNA molecules, nicking the one or more target RNA molecules, increasing expression of the one or more target RNA molecules as compared to an amount of target RNA in a corresponding cell or sample not contacted with the at least one Cas13d protein, decreasing expression 35 of the one or more target RNA molecules as compared to an amount of target RNA in a corresponding cell or sample not contacted with the at least one Cas13d protein, visualizing or detecting the one or more target RNA molecules, labeling the one or more target RNA molecules, capturing molecules 40 bound or tethered to the one or more target RNA molecules, enriching the one or more target RNA molecules, depleting the one or more target RNA molecules, editing the one or more target RNA molecules, trafficking the one or more target RNA molecules, splicing or perturbing the splicing of 45 the one or more target RNA molecules, and masking the one or more target RNA molecules.
- 5. The system of claim 1, wherein the at least one Cas13d protein comprises at least 96% sequence identity to SEQ ID NO: 42.
- **6**. The system of claim **1**, wherein the at least one Cas13d protein comprises at least 97% sequence identity to SEQ ID NO: 42.
- 7. The system of claim 1, wherein the at least one Cas13d protein comprises at least 98% sequence identity to SEQ ID 55 NO: 42.
- **8**. The system of claim **1**, wherein the at least one Cas13d protein comprises at least 99% sequence identity to SEQ ID NO: 42.
- **9**. The system of claim **1**, wherein the at least one Cas13d 60 protein comprises SEQ ID NO: 42.

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- 10. The system of claim 1, wherein the at least one Cas13d protein further comprises one or more subcellular localization signals, effector domains, purification tags, affinity tags, or combinations thereof.
- 11. The system of claim 1, wherein the at least one gRNA comprises one or more DR sequences comprising at least 95% sequence identity to SEO ID NO: 135.
- 12. The system of claim 1, wherein the at least one gRNA comprises one or more DR sequences comprising at least 96% sequence identity to SEQ ID NO: 135.
- 13. The system of claim 1, wherein the at least one gRNA comprises one or more DR sequences comprising at least 97% sequence identity to SEQ ID NO: 135.
- 14. The system of claim 1, wherein the at least one gRNA comprises one or more DR sequences comprising at least 98% sequence identity to SEQ ID NO: 135.
- **15**. The system of claim **1**, wherein the at least one gRNA comprises one or more DR sequences comprising at least 99% sequence identity to SEQ ID NO: 135.
- **16**. The system of claim **1**, wherein the at least one gRNA comprises one or more DR sequences comprising SEQ ID NO: 135.
- 17. The system of claim 1, wherein the at least one gRNA comprises a nucleic acid aptamer.
- **18**. The system of claim **17** wherein the nucleic acid aptamer inserted into the gRNA comprises MS2, PP7, or Oβ.
- 19. The system of claim 1, wherein the at least one gRNA comprises modified nucleotides.
- 20. The isolated cell of claim 2, wherein the cell is a eukaryotic cell or a bacterial cell.
- 21. The isolated cell of claim 20, wherein the eukaryotic cell is a mammalian cell.
- 22. The isolated cell of claim 20, wherein the eukaryotic cell is a plant cell.
- 23. The system of claim 1, wherein the at least one Cas13d protein further comprises a nuclear localization signal (NLS).
- **24**. The system of claim **23**, wherein the NLS comprises SEQ ID NO: 256 or SEQ ID NO: 258.
- 25. The system of claim 1, wherein the at least one Cas13d protein further comprises a nuclear export signal (NES).
- **26**. The system of claim **25**, wherein the NES comprises an adenovirus type 5 E1B nuclear export sequence, an HIV nuclear export sequence, a MAPK nuclear export sequence, or a PTK2 nuclear export sequence.
- 27. The system of claim 1, wherein the at least one Cas13d protein further comprises an effector domain that can cleave RNA, edit a nucleotide, edit a ribonucleotide, methylate RNA, or demethylate RNA.
- **28**. The system of claim **1**, wherein the at least one Cas13d protein is catalytically inactive for ribonuclease activity.
- 29. The system of claim 28, wherein the catalytically inactive Cas13d protein comprises one or more mutated HEPN domains.
- **30**. The system of claim **28**, wherein the catalytically inactive Cas13d protein comprises a mutated HEPN1 domain or a mutated HEPN2 domain.

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