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of the United States Patent and Trademark Office has received an application for a patent for a new and useful invention. The title and description of the invention are enclosed. The requirements of law have been complied with, and it has been determined shat a patent on the invention shall be granted under the law.

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Katherine Kelly Vidal C

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Patent Term Notice

If the application for this patent was filed on or after June 8, 1995, the term of this patent begins on the date on which this patent issues and ends twenty years from the filing date of the application or, if the application contains a specific reference to an earlier filed application or applications under 35 U.S.C. 120, 121, 365(c), or 386(c), twenty years from the filing date of the earliest such application ("the twenty-year term"), subject to the payment of maintenance fees as provided by 35 U.S.C. 41(b), and any extension as provided by 35 U.S.C. 154(b) or 156 or any disclaimer under 35 U.S.C. 253.

If this application was filed prior to June 8, 1995, the term of this patent begins on the date on which this patent issues and ends on the later of seventeen years from the date of the grant of this patent or the twenty-year term set forth above for patents resulting from applications filed on or after June 8, 1995, subject to the payment of maintenance fees as provided by 35 U.S.C. 41(b) and any extension as provided by 35 U.S.C. 156 or any disclaimer under 35 U.S.C. 253.

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(12) United States Patent

Clube et al.

(54) SELECTIVELY ALTERING MICROBIOTA FOR IMMUNE MODULATION

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- (72) Inventors: Jasper Clube, London (GB); Morten Sommer, London (GB); Christian Grøndahl, London (GB)
- (73) Assignee: **SNIPR Technologies Limited**, London (GB)
- (*) 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 disclaimer.

- (21) Appl. No.: 17/132,624
- (22) Filed: Dec. 23, 2020

(65) **Prior Publication Data**

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Related U.S. Application Data

(63) Continuation of application No. 16/453,598, filed on Jun. 26, 2019, which is a continuation of application No. 16/192,752, filed on Nov. 15, 2018, now Pat. No. 10,363,308, which is a continuation of application No. 15/820,296, filed on Nov. 21, 2017, now Pat. No. 10,195,273, which is a continuation of application No. PCT/EP2017/063593, filed on Jun. 4, 2017.

(30) Foreign Application Priority Data

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(56) **References Cited**

U.S. PATENT DOCUMENTS

4,626,504	Α	12/1986	Puhler
5,633,154	Α	5/1997	Schaefer

(10) Patent No.: US 11,471,531 B2

(45) **Date of Patent:** *Oct. 18, 2022

8,241,498	B2	8/2012	Summer	
8,252,576	B2	8/2012	Campbell	
8,906,682	B2	12/2014	June	
8,911,993	B2	12/2014	June	
8,916,381	B1	12/2014	June	
8,975,071	B1	3/2015	June	
9,101,584	B2	8/2015	June	
9,102,760	B2	8/2015	June	
9,102,761	B2	8/2015	June	
9,113,616	B2	8/2015	Stevens	
9,328,156	B2	5/2016	June	
9,464,140	B2	10/2016	June	
9,481,728	B2	11/2016	June	
9,499,629	B2	11/2016	June	
9,518,123	B2	12/2016	June	
9,540,445	B2	1/2017	June	
9,701,964	B2	7/2017	Clube	
9,758,583	B2	9/2017	Wang	
9,822,372	B2	11/2017	Zhang	
9,879,269	B2	1/2018	Barrangou	
10,066,233	B2	9/2018	Barrangou	
10,136,639	B2	11/2018	Wuest	
10,136,649	B2	11/2018	Barrangou	
10,195,273	B2 *	2/2019	Clube A61K 35/15	
(Continued)				

FOREIGN PATENT DOCUMENTS

CA	3010891 A1	7/2017
EP	2325332 A1	5/2011
	(Conti	nued)

OTHER PUBLICATIONS

How gut bacteria boost cancer immunotherapy; https://microbiomepost. com/how-gut-bacteria-boost-cancer-immunotherapy/ (Year: 2020).* Chen et al., Journal of Immunology Research 2020, vol. 2020:1-13 (Year: 2020).*

Kaiser, Science Mag, 2017; https://www.sciencemag.org/news/2017/ 11/your-gut-bacteria-could-determine-how-you-respond-cuttingedge-cancer-drugs (Year: 2017).*

Aklujkar et al. (2010) "Interference With Histidyl-tRNA Synthetase by a CRISPR Spacer Sequence as a Factor in the Evolution of Pelobacter Carbinolicus," BMC Evolutionary Biology 10:203, 15 pages.

American Lung Association (2019). "Preventing COPD," retrieved from https://www.lung.org/lung-health-and-diseases/lung-diseaselookup/copd/symptoms-causes-risk-factors/preventing-copd.html, last visited Aug. 5, 2019, 1 page.

(Continued)

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(57) ABSTRACT

The invention relates to methods of modulating immune cells in a patient by altering microbiota of the patient. The invention also relates to methods of modulating treatments or therapies in a subject organism by altering microbiota of the subject. The invention also relates to cell populations, systems, arrays, cells, RNA, kits and other means for effecting this. In an example, advantageously selective targeting of a particular species in a human gut microbiota using guided nucleic acid modification is carried out to effect the alteration.

16 Claims, 19 Drawing Sheets

Specification includes a Sequence Listing.

U.S. PATENT DOCUMENTS

10,300,138 B2	5/2019	Clube
	5/2019	Clube
10,363,308 B2	7/2019	Clube
10,463,049 B2	11/2019	Clube
10,506,812 B2	12/2019	Clube
10,524,477 B2	1/2020	Clube
10,596,255 B2	3/2020	Clube
10,603,379 B2	3/2020	Clube
10,760,065 B2	9/2020	Lu et al.
10,760,075 B2	9/2020	Sommer et al.
10,765,740 B2	9/2020	Clube et al.
10,920,222 B2	2/2021	Sommer et al.
10,953,090 B2	3/2021	Clube et al.
11,147,830 B2	10/2021	Clube
2003/0049841 A1	3/2003	Short
2004/0096974 A1	5/2004	Herron
2005/0118719 A1	6/2005	Schmidt
2009/0155768 A1	6/2009	Scholl
2010/0076057 A1	3/2010	Sontheimer
2010/0093617 A1	4/2010	Barrangou
2011/0002889 A1	1/2011	Barrangou
2011/0143997 A1	6/2011	Henry et al.
2012/0269859 A1	10/2012	Minato
2013/0011828 A1	1/2013	Barrangou
2013/0109053 A1	5/2013	Macdonald
		-
2013/0121968 A1	5/2013	Quay
2013/0287748 A1	10/2013	June
2013/0288368 A1	10/2013	June
2013/0309258 A1	11/2013	June
2014/0068797 A1	3/2014	Doudna
2014/0105912 A1	4/2014	Noelle
2014/0106449 A1		June
		Meyerson
2014/0179726 A1	6/2014	Bajaj
2014/0199767 A1	7/2014	Barrangou
2014/0234972 A1	8/2014	Zhang
2014/0341920 A1	11/2014	Noelle
2014/0349400 A1	11/2014	Jakimo et al.
2014/0370017 A1	12/2014	June
2015/0004705 A1		Lu et al.
2015/0031134 A1	1/2015	Zhang
2015/0032263 A1	1/2015	Keyl et al.
2015/0050699 A1		Siksnys
2015/0050729 A1	2/2015	June
2015/0064138 A1	3/2015	Lu
2015/0093822 A1	4/2015	June
2015/0099299 A1	4/2015	June
2015/0118202 A1	4/2015	June
2015/0125463 A1	5/2015	Cogswell
	5/2015	Liu et al.
2015/0132419 A1	5/2015	Arvik
2015/0139943 A1	5/2015	Campana
2015/0140001 A1	5/2015	Lee
2015/0184139 A1	7/2015	Zhang
2015/0225730 A1		Minshull et al.
2015/0232881 A1		Glucksmann
2015/0290244 A1		June
2015/0250211 A1		Weiss
2016/0009805 A1		Kowanetz
2016/0009813 A1		Themeli
2016/0024510 A1		Bikard
2016/0040215 A1		Henn et al.
2016/0081314 A1	3/2016	Thurston
2016/0115488 A1	4/2016	Zhang
2016/0115489 A1		Zhang
2016/0130355 A1		June
2016/0159905 A1		Abdiche
2016/0159907 A1		June
2016/0160186 A1		Parsley
2016/0194404 A1	7/2016	June
2016/0208012 A1	7/2016	June
2016/0244784 A1		Jacobson et al.
2016/0281053 A1		Sorek
2016/0324938 A1		Bikard
2016/0333348 A1	11/2/016	
2010/02/20/10/11	11/2010	Clube

2016/0339064 A1	11/2016	Kovarik et al.
2016/0345578 A1	12/2016	Barrangou
2016/0347836 A1	12/2016	Grosso
2016/0354416 A1*	12/2016	Gajewski A61P 43/00
2017/0022499 A1	1/2017	Lu
2017/0022499 A1 2017/0037416 A1	2/2017	Barrangou
2017/0106025 A1	4/2017	Kovarik
2017/0106026 A1	4/2017	Kovarik
2017/0114351 A1	4/2017	Mahfouz
2017/0143772 A1	5/2017	Mulder
2017/0173085 A1	6/2017	Kovarik
2017/0174713 A1	6/2017	Du
2017/0175142 A1	6/2017	Zhang
2017/0196225 A1	7/2017	Clube
2017/0246221 A1	8/2017	Clube
2017/0247690 A1	8/2017	Quake
2017/0304443 A1	10/2017	Lebwohl
2017/0327582 A1	11/2017	Bissonnette
2017/0340733 A1	11/2017	Cao
2018/0015131 A1*	1/2018	Gajewski A61K 45/06
2018/0055852 A1	3/2018	Kutok
2018/0064114 A1	3/2018	Clube
2018/0064115 A1	3/2018	Clube
2018/0070594 A1	3/2018	Clube
2018/0084785 A1	3/2018	Clube
2018/0084786 A1	3/2018	Clube
2018/0140698 A1	5/2018	Clube
2018/0146681 A1	5/2018	Clube
	5/2018	von Maltzahn A61P 35/04
2018/0155721 A1	6/2018	Lu
2018/0155729 A1	6/2018	Beisel
2018/0161368 A1	6/2018	Odegard
2018/0179547 A1	6/2018	Zhang
2018/0200342 A1	7/2018	Bikard
2018/0273937 A1	9/2018	Beisel et al.
2018/0273940 A1	9/2018	Sommer
2018/0303934 A1	10/2018	Clube
2018/0326057 A1	11/2018	Clube
2018/0325378 A1	12/2018	Krom et al.
2018/0371405 A1	12/2018	
		Barrangou
2019/0015441 A1	1/2019	Shachar
2019/0021343 A1	1/2019	Barrangou
2019/0070233 A1	3/2019	Yeung
2019/0117709 A1	- 4/2010	Kovarik
2019/011//09 /11	4/2019	KOVALIK
2019/0133135 A1	5/2019	Clube
2019/0133135 A1	5/2019	Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1	5/2019 5/2019	Clube Clube Sather
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1	5/2019 5/2019 5/2019 5/2019	Clube Clube Sather Turner et al.
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1	5/2019 5/2019 5/2019 5/2019 5/2019	Clube Clube Sather Turner et al. Haaber
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1	5/2019 5/2019 5/2019 5/2019 5/2019 5/2019 8/2019	Clube Clube Sather Turner et al. Haaber Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0142881 A1 2019/0230936 A1 2019/0240325 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019	Clube Clube Sather Turner et al. Haaber Clube Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1 2019/0255084 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/012020 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1 2019/0240326 A1 2019/0255084 A1 2019/0256900 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1 2019/0255084 A1 2019/0255080 A1 2019/0256900 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1 2019/0255084 A1 2019/0256900 A1 2019/0298779 A1 2019/0298779 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al.
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1 2019/025084 A1 2019/0255084 A1 2019/0258090 A1 2019/0228779 A1 2019/0321468 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al.
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1 2019/0240326 A1 2019/0255084 A1 2019/0255090 A1 2019/0321468 A1 2019/0321469 A1 2019/0321470 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1 2019/025084 A1 2019/0255084 A1 2019/0258090 A1 2019/0228779 A1 2019/0321468 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube Swee
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1 2019/0240326 A1 2019/0255084 A1 2019/0255090 A1 2019/0321468 A1 2019/0321469 A1 2019/0321470 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0256900 A1 2019/0256900 A1 2019/0321468 A1 2019/0321469 A1 2019/0321470 A1 2019/032193 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2019	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al.
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0240325 A1 2019/0240326 A1 2019/025084 A1 2019/025084 A1 2019/025084 A1 2019/0256900 A1 2019/0256900 A1 2019/0321468 A1 2019/0321468 A1 2019/0321469 A1 2019/0321470 A1 2019/0359933 A1 2020/0030444 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 11/2019 11/2019 2/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube Swee Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0240326 A1 2019/0240326 A1 2019/0255084 A1 2019/02550900 A1 2019/0298779 A1 2019/0321468 A1 2019/0321470 A1 2019/0321470 A1 2019/0321470 A1 2019/0321470 A1 2019/0321470 A1 2019/0321470 A1 2020/0030444 A1 2020/0046773 A1 2020/0085066 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 2/2020 3/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube Swee Clube Borody Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0142881 A1 2019/0142881 A1 2019/0142881 A1 2019/0230936 A1 2019/0240325 A1 2019/0240326 A1 2019/0240326 A1 2019/0240326 A1 2019/0255084 A1 2019/0255084 A1 2019/0228779 A1 2019/0321468 A1 2019/0321469 A1 2019/0321470 A1 2019/0321470 A1 2020/0030444 A1 2020/0046773 A1 2020/0085066 A1 2020/00870660 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 2/2020 3/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Sommer
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0256900 A1 2019/0321468 A1 2019/0321468 A1 2019/0321469 A1 2019/0321470 A1 2019/0321470 A1 2020/0030444 A1 2020/008766 A1 2020/0087660 A1 2020/012551 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 2/2020 3/2020 4/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Sommer Barrangou
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142831 A1 2019/0160120 A1 2019/0230936 A1 2019/0240325 A1 2019/0240325 A1 2019/0240326 A1 2019/025084 A1 2019/0255084 A1 2019/025468 A1 2019/0321468 A1 2019/0321469 A1 2020/0030444 A1 2020/0046773 A1 2020/0085066 A1 2020/0085066 A1 2020/012551 A1 2020/012551 A1 2020/01125716 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 2/2020 3/2020 3/2020 4/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Borody Clube Borody Clube Sommer Barrangou Martinez
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/014281 A1 2019/0160120 A1 2019/0160120 A1 2019/0240325 A1 2019/0240326 A1 2019/0255084 A1 2019/0321468 A1 2019/0321470 A1 2020/0030444 A1 2020/0085066 A1 2020/0085066 A1 2020/0012551 A1 2020/012551 A1 2020/012571 A1 2020/015716 A1 2020/015716 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2019 11/2019 11/2019 11/2020 2/2020 3/2020 3/2020 4/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Borody Clube Borody Clube Borody Clube Sommer Barrangou Martinez Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0160120 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0240326 A1 2019/0240326 A1 2019/0240326 A1 2019/0255084 A1 2019/0255080 A1 2019/0298779 A1 2019/0321468 A1 2019/0321470 A1 2019/0321470 A1 2019/0321470 A1 2020/0030444 A1 2020/0085066 A1 2020/0087660 A1 2020/012551 A1 2020/012551 A1 2020/012551 A1 2020/0121787 A1 2020/0121787 A1 2020/0121787 A1	5/2019 5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 2/2020 3/2020 3/2020 4/2020 4/2020 5/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Borody Clube Sommer Barrangou Martinez Clube Regev
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0142881 A1 2019/0142881 A1 2019/0142832 A1 2019/0240325 A1 2019/0240325 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0321468 A1 2019/0321470 A1 2020/0030444 A1 2020/0045773 A1 2020/0085066 A1 2020/012551 A1 2020/012551 A1 2020/012573 A1 2020/0157237 A1 2020/0157237 A1 2020/0157237 A1 2020/0154070 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 2/2020 3/2020 4/2020 4/2020 4/2020 5/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Sommer Barrangou Martinez Clube Regev Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0142881 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0321468 A1 2019/0321470 A1 2020/0030444 A1 2020/0035066 A1 2020/0187660 A1 2020/012551 A1 2020/012551 A1 2020/012757 A1 2020/012777 A1 2020/0157237 A1 2020/0157237 A1 2020/01579460 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 4/2020 4/2020 4/2020 5/2020 5/2020 6/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Sommer Barrangou Martinez Clube Regev Clube Kovarik
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0142881 A1 2019/0142881 A1 2019/0142832 A1 2019/0240325 A1 2019/0240326 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0321468 A1 2019/0321470 A1 2020/0030444 A1 2020/0045773 A1 2020/0085066 A1 2020/012551 A1 2020/012551 A1 2020/012571 A1 2020/0157237 A1 2020/0157237 A1 2020/0157237 A1 2020/0154070 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 2/2020 3/2020 4/2020 4/2020 4/2020 5/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Sommer Barrangou Martinez Clube Regev Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0142881 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0321468 A1 2019/0321470 A1 2020/0030444 A1 2020/0035066 A1 2020/0187660 A1 2020/012551 A1 2020/012551 A1 2020/012757 A1 2020/012777 A1 2020/0157237 A1 2020/0157237 A1 2020/01579460 A1	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 4/2020 4/2020 4/2020 5/2020 5/2020 6/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Sommer Barrangou Martinez Clube Regev Clube Kovarik
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142831 A1 2019/0160120 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/025084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/025084 A1 2019/025084 A1 2019/0321469 A1 2019/0321469 A1 2020/0030444 A1 2020/00457660 A1 2020/0087660 A1 2020/012551 A1 2020/012551 A1 2020/0125716 A1 2020/012787 A1 2020/0157237 A1 2020/0157237 A1 2020/0157460 A1 2020/0157460 <	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 4/2020 4/2020 5/2020 5/2020 5/2020 7/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube Borody Clube Borody Clube Sommer Barrangou Martinez Clube Regev Clube Kovarik Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0140230 A1 2019/0140281 A1 2019/0160120 A1 2019/0240325 A1 2019/0240326 A1 2019/0255084 A1 2019/0321468 A1 2019/0321470 A1 2020/0030444 A1 2020/0085066 A1 2020/012551 A1 2020/012571 A1 2020/012577 A1 2020/0157237 A1 2020/0157237 A1 2020/0157237 A1 2020/0157237 A1 2020/0157237 A1 2020/0157460	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2019 11/2019 11/2019 11/2019 11/2020 3/2020 3/2020 4/2020 5/2020 5/2020 6/2020 9/2020 10/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Sommer Barrangou Martinez Clube Regev Clube Regev Clube Barrangou Martinez Clube Bibard et al. Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0140281 A1 2019/0160120 A1 2019/0140281 A1 2019/0240325 A1 2019/0240325 A1 2019/0240326 A1 2019/0255084 A1 2019/0255084 A1 2019/0255080 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/025900 A1 2019/0259739 A1 2020/00321469 A1 2020/003444 A1 2020/0085066 A1 2020/0115716 A1 2020/0115716 A1 2020/0157237 A1 2020/0157237 A1 2020/0157237 A1 2020/0157237 A1 2020/0157237 A1 2020/0205416 A1 2020/0205406 A1 2020/0205406	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2020 3/2020 3/2020 3/2020 4/2020 4/2020 5/2020 5/2020 6/2020 7/2020 10/2020 11/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Sommer Barrangou Martinez Clube Regev Clube Regev Clube Bikard et al. Clube Bikard et al. Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0142881 A1 2019/0142881 A1 2019/0240325 A1 2019/0240325 A1 2019/0255084 A1 2019/0321469 A1 2019/0321470 A1 2020/0030444 A1 2020/0085066 A1 2020/0085066 A1 2020/012551 A1 2020/0157237 A1 2020/0157237 A1 2020/0157237 A1 2020/025416 A1 2020/025407 A1 2020/025407 A1 2020/0337313 A1 2020/0354690 A1 2020/0354690	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 4/2020 4/2020 5/2020 5/2020 6/2020 7/2020 10/2020 11/2020	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Borody Clube Sommer Barrangou Martinez Clube Regev Clube Borody Clube Borody Clube Sommer Barrangou Martinez Clube Barrangou Martinez Clube Regev Clube Bikard et al. Clube Bikard et al. Clube Bikard et al. Clube Clube Clube Regev Regev Regv Reg
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0142881 A1 2019/0142881 A1 2019/0142881 A1 2019/0240325 A1 2019/0240325 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0256900 A1 2019/0321469 A1 2020/003021470 A1 2020/00321470 A1 2020/0087660 A1 2020/012551 A1 2020/012777 A1 2020/015716 A1 2020/0157237 A1 2020/025416 A1 2020/025277 A1 2020/0354690 A1 2020/0354690 A1 2020/0354690	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 4/2020 4/2020 6/2020 5/2020 6/2020 7/2020 10/2020 11/2020 12/2020 1/2021	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Sommer Barrangou Martinez Clube Sommer Barrangou Martinez Clube Regev Clube Regev Clube Bikard et al. Clube Garofolo Clube Sigarofolo Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0140230 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0321469 A1 2019/0359933 A1 2020/0030444 A1 2020/008566 A1 2020/012551 A1 2020/012551 A1 2020/0125717 A1 2020/015716 A1 2020/0157237 A1 2020/0157237 A1 2020/025416 A1 2020/037313 A1 2020/0354690 A1 2020/0354690 A1 2020/0390886 <t< td=""><td>5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 4/2020 4/2020 5/2020 5/2020 5/2020 0/2020 11/2020 11/2020 12/2020 12/2021 5/2021</td><td>Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube al. Clube Borody Clube Sommer Barrangou Martinez Clube Sommer Barrangou Martinez Clube Regev Clube Regev Clube Baradou Barrangou Martinez Clube Regev Regev R</td></t<>	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 4/2020 4/2020 5/2020 5/2020 5/2020 0/2020 11/2020 11/2020 12/2020 12/2021 5/2021	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube al. Clube Borody Clube Sommer Barrangou Martinez Clube Sommer Barrangou Martinez Clube Regev Clube Regev Clube Baradou Barrangou Martinez Clube Regev Regev R
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0140230 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/025084 A1 2019/0255084 A1 2019/0321468 A1 2019/0359933 A1 2020/0030444 A1 2020/0087660 A1 2020/012551 A1 2020/015716 A1 2020/015716 A1 2020/0179460 A1 2020/025416 A1 2020/0337313 A1 2020/0354690 A1 2020/0354690 A1 2021/0145006 <	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 3/2020 4/2020 5/2020 5/2020 5/2020 10/2020 10/2020 10/2020 11/2020 12/2020 12/2021 5/2021 5/2021	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube al. Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Barrangou Martinez Clube Barrangou Martinez Clube Bikard et al. Clube Bikard et al. Clube Garofolo Clube Sommer Clube Garofolo Clube Clube Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0140230 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0255084 A1 2019/0321469 A1 2019/0359933 A1 2020/0030444 A1 2020/008566 A1 2020/012551 A1 2020/012551 A1 2020/0125717 A1 2020/015716 A1 2020/0157237 A1 2020/0157237 A1 2020/025416 A1 2020/037313 A1 2020/0354690 A1 2020/0354690 A1 2020/0390886 <t< td=""><td>5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 4/2020 4/2020 5/2020 5/2020 5/2020 0/2020 11/2020 11/2020 12/2020 12/2021 5/2021</td><td>Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Clube Borody Clube Clube Bikard et al. Clube Garofolo Clube Clube Clube Clube Clube</td></t<>	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 4/2020 4/2020 5/2020 5/2020 5/2020 0/2020 11/2020 11/2020 12/2020 12/2021 5/2021	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube Swee Clube Borody Clube Clube Borody Clube Clube Bikard et al. Clube Garofolo Clube Clube Clube Clube Clube
2019/0133135 A1 2019/0134194 A1 2019/0136230 A1 2019/0140230 A1 2019/0160120 A1 2019/0240325 A1 2019/0240325 A1 2019/0240325 A1 2019/025084 A1 2019/0255084 A1 2019/0321468 A1 2019/0359933 A1 2020/0030444 A1 2020/0087660 A1 2020/012551 A1 2020/015716 A1 2020/015716 A1 2020/0179460 A1 2020/025416 A1 2020/0337313 A1 2020/0354690 A1 2020/0354690 A1 2021/0145006 <	5/2019 5/2019 5/2019 5/2019 8/2019 8/2019 8/2019 8/2019 8/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 10/2019 11/2020 3/2020 3/2020 3/2020 4/2020 5/2020 5/2020 5/2020 10/2020 10/2020 10/2020 11/2020 12/2020 12/2021 5/2021 5/2021	Clube Clube Sather Turner et al. Haaber Clube Clube Clube Schentag Zhang Falb Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube et al. Clube al. Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Borody Clube Barrangou Martinez Clube Barrangou Martinez Clube Bikard et al. Clube Bikard et al. Clube Garofolo Clube Sommer Clube Garofolo Clube Clube Clube

U.S. PATENT DOCUMENTS

2021/0189406 A1	6/2021	Martinez et al.
2021/0198665 A1	7/2021	Sommer et al.
2021/0230559 A1	7/2021	Clube
2021/0283167 A1	9/2021	Clube
2021/0290654 A1	9/2021	Clube
2021/0386773 A1	12/2021	Clube

FOREIGN PATENT DOCUMENTS

ГD	2040140 41	2/2015
EP	2840140 A1	2/2015
EP	3461337 A1	4/2019
EP	3132035 B8	4/2020
EP	3132036 B8	4/2020
EP	3630975 A1	4/2020
EP	3633032 A2	4/2020
EP	3634442 A1	4/2020
ĒP	3634473 A1	4/2020
RU	2531343 C2	10/2014
WO	2005046579 A2	5/2005
WO	2005046579 A3	8/2005
WO	2007025097 A2	3/2007
WO	2007025097 A3	7/2007
WO	2008108989 A2	9/2008
WO	2008108989 A3	3/2009
WO	2010011961 A2	1/2010
WO	2010011961 A3	6/2010
WO	2010075424 A2	7/2010
WO	2010075424 A3	9/2010
WO	2012079000 A1	6/2012
WO	2012079000 A4	8/2012
WO	2012164565 A1	12/2012
WO	2013063361 A1	5/2013
WO	2013176772 A1	11/2013
WO	2014012001 A2	1/2014
WO	2014018423 A2	1/2014
WÕ	2014018423 A3	1/2014
WO	2014093595 A1	6/2014
WO	2014093661 A2	6/2014
WO	2014093661 A3	8/2014
WO		8/2014
WO	2014093661 A9	10/2014
WO	2014204725 A1	12/2014
wo	2015034872 A2	3/2015
WO	2014012001 A3	4/2015
WO	2015034872 A3	4/2015
WO	2015058018 A1	4/2015
WO	2015069682 A2	5/2015
WO	2015070083 A1	5/2015
WO	2015071474 A2	5/2015
WO	2015075688 A1	5/2015
WO	2015088643 A1	6/2015
WO	2015089351 A1	6/2015
wŏ		
	2015089419 A2	6/2015
WO	2015069682 A3	7/2015
WO	2015071474 A3	8/2015
WO	2015089419 A3	9/2015
WO	2015136541 A2	9/2015
WO	2015148680 A1	10/2015
WO	2015153940 A1	10/2015
WO	2015155686 A2	10/2015
WO	2015159068 A1	10/2015
WO	2015159086 A1	10/2015
		10/2015
WO	2015159087 A1	10/2015
WO	2015136541 A3	11/2015
WO	2016044745 A1	3/2016
wo	2016063263 A2	4/2016
WO	2016063263 A3	6/2016
WO	2016196361 A1	12/2016
WO	2016196605 A1	12/2016
WO	2016205276 A1	12/2016
WO	2017009399 A1	1/2017
WO	2017042347 A1	3/2017
wŏ	2017058751 A1	4/2017
WO	2017112620 A1	6/2017
WO	2017118598 A1	7/2017

WO	2017211753 A1	12/2017
WO	2018064165 A2	4/2018
WO	2018081502 A1	5/2018
WO	2018115519 A1	6/2018
WO	2018217351 A1	11/2018
WO	2018217981 A1	11/2018
WO	2018222969 A1	12/2018
WO	2018226853 A1	12/2018
WO	2018064165 A3	6/2019
WO	2020072248 A1	4/2020
WO	2020072250 A1	4/2020
WO	2020072253 A1	4/2020
WO	2020072254 A1	4/2020
WO	2020152369 A1	7/2020

OTHER PUBLICATIONS

Ang, Y.L.E. et al. (2015). "Best Practice in the Treatment of Advanced Squamous Cell Lung Cancer," Ther. Adv. Respir. Dis. 9(5):224-235.

Anonymous (Apr. 2016). "Checkpoint Inhibition: A Promising Immunotherapeutic Approach for Colorectal Cancer," Oncology, 5(3):1-5, retrieved from http://www.personalizedmedonc.com/ publications/prno/april-2016-vol-5-no-3/checkpoint-inhibition-aprornising-irmunotherapeutic-approach-for-colorectal-cancer-2/, last visited Aug. 27, 2019, 5 pages.

Arnold, I.C. et al. (Apr. 8, 2015, e-pub. Mar. 4, 2015). "Helicobacter Hepaticus Infection in BALB/c Mice Abolishes Subunit-Vaccine-Induced Protection Against *M. tuberculosis*," Vaccine 33(15):1808-1814.

Arslan, Z. et al. (May 7, 2013). "RcsB-BglJ-Mediated Activation of Cascade Operon Does Not Induce the Maturation of CRISPR RNAs in *E. coli* K12," RNA Biology 10(5):708-715.

Arumugam et al. (May 12, 2011). "Enterotypes of the human gut microbiome," Nature 473(7346):174-180, 16 pages.

Bae, T. et al. (2006). "Prophages of *Staphylococcus aureus* Newman and Their Contribution to Virulence," Molecular Microbiology pp. 1-13.

Barrangou, R. et al. (Mar. 2007). "CRISPR Provides Acquired Resistance Against Viruses in Prokaryotes," Science, 315:1709-1712.

Beisel, C.L. et al. (2014). "A CRISPR Design for Next-Generation Antimicrobials," Genome Biology 15:516, 4 pages.

Belizario, J.E. et al. (Oct. 6, 2015). "Human Microbiomes and Their Roles in Dysbiosis, Common Diseases, and Novel Therapeutic Approaches," Frontiers in Microbiology 6(1050):1-16.

Bikard, D. et al. (2013, e-pub. Jun. 12, 2013). "Programmable Repression and Activation of Bacterial Gene Expression Using an Engineered CRISPR-Cas System," Nucleic Acids Research 41(15):7429-7437.

Bikard, D. et al. (2017, e-pub. Sep. 6, 2017). "Using CRISPR-Cas Systems as Antimicrobials," Current Opinion in Microbiology 37:155-160.

Bikard, D. et al. (Aug. 16, 2012). "CRISPR Interference Can Prevent Natural Transformation and Virulence Acquisition during In Vivo Bacterial Infection," Cell Host & Microbe 12(2):177-186.

Bikard, D. et al. (Nov. 2014). "Development of Sequence-Specific Antimicrobials Based on Programmable CRISPR-Cas Nucleases," Nature Biotechnology 32(11):1146-1151, 16 pages.

Broaders, E. et al. (Jul./Aug. 2013). "Mobile Genetic Elements of the Human Gastrointestinal Tract," Gut Microbes 4(4):271-280.

Brouns, S.J.J. et al. (Aug. 15, 2008). Supplemental Material for "Small CRISPR RNAs Guide Antiviral Defense in Prokaryotes," Science 321:960-964.

Brouns, S.J.J. et al. (Aug. 15, 2008)."Small CRISPR RNAs Guide Antiviral Defense in Prokaryotes," Science 321:960-964.

Bryksin, A. V. et al. (Oct. 8, 2010). "Rational Design of a Plasmid Origin That Replicates Efficiently in Both Gram-Positive and Gram Negative Bacteria," PloS One 5(10):e13244, 9 pages.

Bugrysheva, J.V. et al. (Jul. 2011, E-Pub. Apr. 29, 2011). "The Histone-Like Protein Hip is Essential for Growth of *Streptococcus pyogenes*: Comparison of Genetic Approaches to Study Essential Genes," Appl. Environ. Microbiol. 77(13):4422-4428.

OTHER PUBLICATIONS

Bullman, S. et al. (Nov. 23, 2017). "Analysis of Fusobacterium Persistence and Antibiotic Response in Colorectal Cancer," Science pp. 1443-1448,10 pages.

Burns, M.B. et al. (2015). "Virulence Genes are a Signature of the Microbiome in the Colorectal Tumor Microenvironment," Genome Medicine 7:55, 12 pages.

Catalao, M.J. et al. (Jul. 2013, e-pub. Nov. 8, 2012). "Diversity in Bacterial Lysis Systems: Bacteriophages Show the Way," FEMS Microbiology Reviews 37(4):554-571.

Chan, B.K. et al. (2013). "Phage Cocktails and the Future of Phage Therapy," Future Microbiol. 8(6):769-783.

Chan, C.T.Y. et al. (Dec. 2015). "'Deadman' and 'Passcode' Microbial Kill Switches for Bacterial Containment," Nat. Chem. Biol. 12(2):82-86.

Cheadle, E.J. et al. (2012). "Chimeric Antigen Receptors for T-Cell Based Therapy," Methods Mol. Biol. 907:645-666, 36 pages.

Citorik, R.J. et al. (Nov. 2014, e-pub Sep. 21, 2014). "Sequence-Specific Antimicrobials Using Efficiently Delivered RNA-Guided Nucleases," Nat. Biotechnol. 32(11):1141-1145, 18 pages.

Cochrane, K. et al. (2016, e-pub. Nov. 3, 2015). "Complete Genome Sequences and Analysis of the *Fusobacterium nucleatum* Subspecies *animalis* 7-1 Bacteripophage offunul and offunu2," Anaerobe 38:125-129. Abstract Only.

Cong, L. et al. (Feb. 15, 2013, e-pub. Oct. 11, 2013). "Multiplex Genome Engineering Using CRISPR/Cas Systems," Science 339(6121):819-823, 9 pages.

Consumer Updates (2019). "Combating Antibiotic Resistance," retrieved from https://www.fda.gov/ForConsumers/ConsumerUpdates/ ucm092810.htm, last visited Jan. 28, 2019.

Coyne, M.J. et al. (2014). "Evidence of Extensive DNA Transfer between *Bacteroidales* Species Within the Human Gut," mBio 5(3):e01305-14, 12 pages.

Cui, L. et al. (2016, e-pub. Apr. 8, 2016). "Consequences of Cas9 Cleavage in the Chromosome of *Escherichia coli*," Nucleic Acids Research 44(9):4243-4251.

Daillere, R. et al. (Oct. 18, 2016). "Enterococcus hirae and Barnesiella intestinihominis Facilitate Cyclophosphamide-Induced Therapeutic Immunomodulatory Effects," Immunity 95:931-943.

Datsenko, K.A. et al. (Jul. 10, 2012). "Molecular Memory of Prior Infections Activates the CRISPR/Cas adaptive Bacterial Immunity System," Nature Communication 3:945, 7 pages.

De Filippo, C. et al. (Aug. 33 2010). "Impact of Diet in Shaping Gut Microbiota Revealed by a Comparative Study in Children From Europe and Rural Africa," Proc. Natl. Acad. Sci. USA 107(33):14691-14696, 6 pages.

De Paepe, M. et al. (Mar. 28, 2014). "Bacteriophages: An Underestimated Role in Human and Animal Health?" Frontiers in Cellular and Infection Microbiology 4(39):1-11.

Deeks, E.D. (2014, e-pub. Jul. 15, 2014). "Nivolumab: A Review of Its Use in Patients With Malignant Melanoma," Drugs 74:1233-1239.

Deghorain, M. et al. (Nov. 23, 2012). "The Staphylococci Phages Family: An Overview," Viruses 4:3316-3335.

Denham, J.D. et al. (2018). "Case Report: Treatment of Enteropathogenic *Escherichia coli* Diarrhea in Cancer Patients: A Series of Three Cases," Case Reports in Infectious Diseases Article ID 8438701:1-3.

Dhar, A.D. (Jul. 20, 2018). "Overview of Bacterial Skin Infections," Merck Manual retrieved from https://www.merckmanuals.com/home/ skin-disorders/bacterial-skin-infections/overview-of-bacterial-skininfections, last visited Jul. 20, 2018, 3 pages.

Dickson, R.P. et al. (Jan./Feb. 2017). "Bacterial Topography of the Healthy Human Lower Respiratory Tract," American Society for Microbiology 8(1):e02287-6, 12 pages.

Diez-Villasenor, C. et al. (May 2013). "CRISPR-Spacer Integration Reporter Plasmids Reveal Distinct Genuine Acquisition Specificities Among CROSPR-Cas 1-E Variants of *Escherichia coli*," RNA Biology 10(5):792-802. Dutilh, B.E. et al. (Jul. 24, 2014). "A Highly Abundant Bacteriophage Discovered in the Unknown Sequences of Human Faecal Metagenomes," Nature Communications 5(4498):1-10.

Edgar et al. (Dec. 2010). "The *Escherichia coli* CRISPR System Protects From λ Lysogenization, Lysogens, and Prophage Induction," Journal of Bacteriology 192(23):6291-6294, Supplemental Material, 2 pages.

Edgar et al. (Dec. 2010). "The *Escherichia coli* CRISPR System Protects From λ Lysogenization, Lysogens, and Prophage Induction," Journal of Bacteriology 192(23):6291-6294.

Esvelt, K.M. et al. (Nov. 2013). "Orthogonal Cas9 Proteins for RNA-Guided Gene Regulation and Editing," Nature Methods 10(11):1116-1123.

Ex Parte Re-Exam, mailed Apr. 21, 2021, for U.S. Appl. No. 90/014,705, filed Mar. 26, 2021, for U.S. Patent Reexamination 10,953,090, 15 pages. (2.34).

Ex Parte Re-Exam, mailed Apr. 30, 2021, for U.S. Appl. No. 90/014,681, filed Feb. 16, 2021, for U.S. Patent Reexamination 10,920,222, 25 pages.

Ex Parte Re-Exam, mailed Dec. 10, 2018, for U.S. Appl. No. 90/014,184, filed Aug. 10, 2018, for U.S. Patent Reexamination 9,701,964 102 pages.

Ex Parte Re-Exam, mailed Feb. 22, 2021, for U.S. Appl. No. 16/700,856, filed Dec. 2, 2019, for U.S. Patent Reexamination 10,920,222,459 pages.

Ex Parte Re-Exam, mailed Mar. 23, 2021, for U.S. Appl. No. 16/453,604, filed Jun. 26, 2019, for U.S. Patent Reexamination 10,953,090, 235 pages.

Ex Parte Re-Exam, mailed Mar. 24, 2021, for U.S. Appl. No. 90/014,681, filed Mar. 24, 2021, for U.S. Patent Reexamination 10,920,222, 18 pages.

Extended European Search Report, dated Jul. 27, 2020, for European Patent Application No. 20155001.9, 9 pages.

Extended European Search Report, dated Sep. 24, 2020, for European Patent Application No. 20154858.3, 12 pages.

Fact Sheet (Oct. 2010). "Antimicrobial Resistance," National Institutes of Health, 1-2.

Foca, A. et al. (2015, e-pub. Apr. 7, 2015). Gut Inflammation and Immunity: What is the Role of the Human Gut Virome? Mediators of Inflammation 2015(326032):1-7.

Galperin, M.Y. (Dec. 2013). "Genome Diversity of Spore-Forming Firmicutes," Microbiology Spectrum 1(2):TBS-0015-2012, 27 pages. Garneau, J. E. et al. (Nov. 4, 2010). "The CRISPR/Cas Bacterial Immune System Cleaves Phage and Plasmid DNA," Nature 468(7320):67-71, 28 pages.

Garon, E.B. et al. (Oct. 2015). "Current Perspectives in Immunotherapy for Non-Small Cell Lung Cancer," Seminars in Oncology 42(5 Supp. 2):S11-S18.

Garrett W.S. et al. (Oct. 5, 2007). "Communicable Ulcerative Colitis Induced by T-Bet Deficiency in the Innate Immune System," Cell 131(1):33-45, 23 pages.

Gauer, R.L. et al. (Jul. 1, 2013). "Early Recognition and Management of Sepsis in Adults: The First Six Hours," American Family Physician 88(1):44-53.

Geller, L.T. et al. (Sep. 15, 2017). "Potential Role of Intratumor Bacteria in Mediating Tumor Resistance to the Chemotherapeutic Drug Gemcitabine," Cancer, 1156-1160, 6 pages.

Golubovskaya, V. et al. (Mar. 15, 2016). "Different Subsets of T Cells, Memory, Effector Functions, and CAR-T Immunotherapy," Cancers 8(36), 12 pages.

Gomaa et al. (Jan. 28, 2014). "Programmable Removal of Bacterial Strains by Use of Genome-Targeting CRISPR-Cas Systems," mBio, 5(1):e000928-13.

Gomaa, A.A. et al. (Jan./Feb. 2014). Supplemental Material to "Programmable Removal of Bacterial Strains by Use of GenomeTargeting CRISPR-Cas Systems," American Society for Microbiology 5(1):1-9.

Goodall, E.C.A. et al. (Feb. 20, 2018). "The Essential Genome of *Escherichia coli* K-12," Am. Society for Microbiology—mBio 9(1):e02096-17, 18 pages.

Gopalakrishnan, V. et al. (Jan. 5, 2018). "Gut Microbiome Modulates Response to Anti-PD-1 Immunotherapy in Melanoma Patients," Science 359:97-103, 20 pages.

OTHER PUBLICATIONS

Green, J. (Jul. 20, 2018). Colgate https://www.colgate.com/en-us/ oral-health/conditions/mouth-sores-and-infections/eight-commonoral-infections-0615, last visited Jul. 20, 2018, 4 pages.

Gudbergsdottir, S. et al. (2011, e-pub. Nov. 18, 2010). "Dynamic Properties of the Sulfolobus CRISPR/Cas and CRISPR/Cmr Systems When Challenged With Vector-Borne Viral and Plasmid Genes and Protospacers," Molecular Microbiology 79(1):35-49.

Guedan, S. et al. (Aug. 14, 2014). "ICOS-Based Chimeric Antigen Receptors Program Bipolar TH17/TH1 Cells," Blood 124(7):1070-1080.

Gupta, R. et al. (2011). "P-27/HP Endolysin as Antibacterial Agent for Antibiotic Resistant *Staphylococcus aureus* of Human Infections," Curr. Microbiol. 63:39-45.

Hansen, J.J. et al. (Mar. 2015). "Therapeutic Manipulation of the Microbiome in IBD: Current Results and Future Approaches," Curr. T. Options Gastroentrol. 13(1):1-18.

Hargreaves, K.R. et al. (Aug. 26, 2014). "Abundant and Diverse Clustered Regularly Interspaced Short Palindromic Repeat Spacers in Clostridium difficile Strains and Prophages Target Multiple Phage Types within This Pathogen," mBio 5(5):e01045-13. Harrington, L.E. (Nov. 2005, e-pub. Oct. 2, 2005). "Interleukin

Harrington, L.E. (Nov. 2005, e-pub. Oct. 2, 2005). "Interleukin 17-producing CD4+ Effector T Cells Develop Via a Lineage Distinct From the T Helper Type 1 and 2 Lineages," Nat. Immunol. 6(11):1123-1132.

Hartland, E.L. et al. (Apr. 30, 2013). "Enteropathogenic and Enterohemorrhagic *E. coli*: Ecology, Pathogenesis, and Evolution," Frontiers in Cellular and Infection Microbiology 3(15):1-3.

Healthline (2019). "Cystic Fibrosis," retrieved from https://www. healthline.com/health/cystic-fibrosis#prevention, last visited Aug. 5, 2019, 14 pages.

Hooper, L.V. et al. (Jun. 8, 2012). "Interactions Between the Microbiota and the Immune System," Science 336(6086):1268-1273, 16 pages.

Horvath, P. et al. (2008, e-pub. Dec. 7, 2007). "Diversity, Activity, and Evolution of CRISPR Loci in *Streptococcus thermophiles*," Journal of Bacteriology 190(4):1401-1412.

Hotta, K. et al. (2011, e-pub. Sep. 20, 2011). "Prognostic Significance of CD45RO+ Memory T Cells in Renal Cell Carcinoma," British Journal of Cancer 105:1191-1196.

Huddleston, J.R. (Jun. 20, 2014). "Horizontal Gene Transfer in the Human Gastrointestinal Tract: Potential Spread of Antibiotic Resistance Genes," Infection and Drug Resistance 7:167-176.

Huo, Y. et al. (Sep. 2014). "Structures of CRISPR Cas3 Offer Mechanistic Insights Into Cascade-Activated DNA Unwinding and Degradations," Nat. Struct. Mol. Biol. 21(9):771-777, 21 pages.

International Search Report and The Written Opinion of the International Searching Authority for PCT/EP2018/066954, dated Oct. 23, 2018, filed Jun. 25, 2018, 14 pages.

International Search Report and The Written Opinion of the International Searching Authority for PCT/EP2019/057453, dated Aug. 16, 2019, filed Mar. 25, 2019, 21 pages.

International Search Report for PCT/EP2016/059803, dated Jun. 30, 2016, filed May 3, 2016, 6 pages.

International Search Report for PCT/EP2018/082053, dated Mar. 14, 2019, filed Nov. 21, 2018, 9 pages.

Ivanov, I.I. et al. (May 2010). "Segmented Filamentous Bacteria Take the Stage," Muscosal Immunol. 3(3):209-212, 7 pages.

Jiang, W. et al. (Jan. 29, 2013). "RNA-Guided Editing of Bacterial Genomes Using CRISPR-Cas Systems," Nat. Biotechnology 31:233-241.

Jiang, W. et al. (Mar. 2013, e-pub. Sep. 1, 2013). "CRISPR-Assisted Editing of Bacterial Genomes," Nat. Biotechnol. 31(3):233-239.

Jiang, W. et al. (Nov. 2013). "Demonstration of CRISPR/Cas9/ sgRNA-Mediated Targeted Gene Modification in *Arabidopsis*, Tobacco, Sorghum and Rice," Nucleic Acids Research 41(20):e188, 12 pages. Jin, Y. et al. (2019, e-pub. Apr. 23, 2019). "The Diversity of Gut Microbiome is Associated With Favorable Responses to Anti-Programmed Death 1 Immunotherapy in Chinese Patients With NSCLC," Journal of Thoracic Oncology 14(8):1378-1389. Jinek et al. (Aug. 17, 2012). "A Programmable Dual-RNA-Guided DNA Endonuclease in Adaptive Bacterial Immunity," Science 337(6096):816-821.

Khoja, L. et al. (2015). "Pembrolizumab," Journal for ImmunoTherapy of Cancer 3(36):1-13.

Kochenderfer, J.N. et al. (Sep. 2009). "Construction and Pre-clinical Evaluation of an Anti-CD19 Chimeric Antigen Receptor," J. Immunother. 32(7):689-702, 26 pages.

Koonin, E.V. et al. (2017, e-pub. Jun. 9, 2017). "Diversity, Classification and Evolution of CRISPR-Cas Systems," Current Opinion in Microbiology 37:67-78.

Kosiewicz, M.M. et al. (2014, e-pub. Mar. 26, 2014). "Relationship Between Gut Microbiota and Development of T Cell Associated Disease," FEBS Lett. 588:4195-4206.

Kostic, A.D. et al. (Aug. 14, 2013). "Fusobacterium nucleatum Potentiates Intestinal Tumorigenesis and Modulates the Tumor-Immune Microenvironment," Cell Host Microbe. 14(2):207-215, 18 pages.

Krom, R.J. et al. (Jul. 5, 2015). "Engineered Phagemids for Nonlytic, Targeted Antibacterial Therapies," Nano Letters 15(7):4808-4813. La Scola, B. et al. (Sep. 4, 2008). "The Virophage as a Unique Parasite of the Giant Mimivirus," Nature Letters 455:100-104.

Lopez-Sanchez, M.-J. et al. (2012, e-pub. Jul. 27, 2012). "The Highly Dynamic CRISPR1 System of *Streptococcus agalactiae* Controls the Diversity of its Mobilome," Molecular Microbiology 85(6):1057-1071.

Lu, T.K. et al. (Jul. 3, 2007). "Dispersing Biofilms With Engineered Enzymatic Bacteriophage," PNAS 104(27):11197-11202. Ludwig, W. et al. (1985). "The Phylogenetic Position of *Strepto*-

Ludwig, W. et al. (1985). "The Phylogenetic Position of *Strepto-coccus* and Enterococcus," Journal of General Microbiology 131:543-551.

Luo, M.L. et al. (2015, e-pub. Oct. 17, 2014). "Repurposing Endogenous Type I CRISPR-Cas Systems for Programmable Gene Repression," Nucleic Acids Research 43(1):674-681.

López, P. et al. (Apr. 5, 2016). "Th17 Responses and Natural IgM Antibodies are Related to Gut Microbiota Composition in Systemic Lupus Erythematosus Patients," Sci. Rep. 6:24072, 12 pages. Macon, B.L. et al. (Jan. 2, 2018). "Acute Nephrities," retrieved from

Macon, B.L. et al. (Jan. 2, 2018). "Acute Nephrities," retrieved from healthline, https://www.healthline.com/health/acute-nephritic-syndrome#types, last visited Jul. 20, 2018, 13 pages.

Magee, M.S. et al. (Nov. 2014). "Challenges to Chimeric Antigen Receptor (CAR)-T Cell Therapy for Cancer," Discov. Med. 18(100):265-271, 6 pages.

Mahoney, K.M. et al. (2015). "The Next Immune-Checkpoint Inhibitors: PD-1/PD-L1 Blockade in Melanoma," Clinical Therapeutics 37(4):764-782.

Makarova, K.S. (Jun. 2011). "Evolution and Classification of the CRISPR-Cas Systems," Nat. Rev. Microbiol. 9(6):467-477, 23 pages.

Mali, P. et al. (Oct. 2013, e-pub. Sep. 27, 2013). "Cas9 as a Versatile Tool for Engineering Biology," Nature Methods 10(10):957-963. 16 pages.

Mancha-Agresti, P. et al. (Mar. 2017). "A New Broad Range Plasmid for DNA Delivery in Eukaryotic Cells Using Lactic Acid Bacteria: In Vitro and In Vivo Assays," Molecular Therapy: Methods & Clinical Development 4:83-91.

Manica, A. et al. (2011, e-pub. Mar. 8, 2011). "In vivo Activity of CRISPR-Mediated Virus Defence in a Hyperthermophilic Archaeon," Molecular Microbiology 80(2):481-491.

Marraffini, L.A. et al. (Dec. 19, 2008). "CRISPR Interference Limits Horizontal Gene Transfer in Staphylococci by Targeting DNA," Science 322(5909):1843-1845, 12 pages.

Martel, B. et al. (2014). "CRISPR-Cas: An Efficient Tool for Genome Engineering of Virulent Bacteriophages," Nucleic Acids Research 42(14):9504-9513.

Martinez, R.M. et al. (Aug. 12, 2016). "Bloodstream Infections," Microbial Spectrum 4(4):DMIH2-0031-2016, 34 pages.

Mayo Clinic (2019). "Pulmonary Embolism," retrieved from https:// www.nnayoclinic.org/diseases-conditions/pulmonary-ennbolism/ synnptonns-causes/syc-20354647, last visited Aug. 5, 2019, 8 pages. Mayo Clinic (2020). "Infectious Diseases," retrieved from https:// www.nnayoclinic.org/diseases-conditions/infectious-diseases/diagnosistreatnnent/drc-20351179, last visited Jan. 17, 2020, 5 pages.

OTHER PUBLICATIONS

Mayo Clinic (2020). "Malaria," retrieved from https://www. nnayoclinic.org/diseases-conditions/nnalaria/diagnosis-treatnnent/ drc-20351190, last visited Jan. 17, 2020, 3 pages.

Mayo Clinic (2020). "Sexually Transmitted Diseases (STDs)," retrieved from https://www.nnayoclinic.org/diseases-conditions/ sexually-transnnitted-diseases-stds/diagnosis-treatnnent/drc-20351246, last visited Jan. 17, 2020, 5 pages.

Mayo Clinic (Jul. 20, 2018). "Bacterial Vaginosis," retrieved from https://www.mayoclinic.org/diseases-conditions/bacterial-vaginosis/ symptoms-causes/syc-20352279, last visited Jul. 20, 2018, 3 pages. Mayo Clinic (Jul. 20, 2018). "Cystitis," retrieved from https://www. mayoclinic.org/diseases-conditions/cystitis/symptoms-causes/syc-20371306, last visited Jul. 20, 2018, 10 pages.

Mayo Clinic (Jul. 20, 2018). "Meningitis," retrieved from https:// www.mayoclinic.org/diseases-conditions/meningitis/symptomscauses/syc-20350508, last visited Jul. 20, 2018, 6 pages.

Mayo Clinic (Jul. 20, 2018). "Pneumonia," retrieved from https:// www.mayoclinic.org/diseases-conditions/pneumonia/symptomscauses/syc-20354204, last visited Jul. 20, 2018, 5 pages.

Mayo Clinic (Mar. 29, 2020). "Liver Disease," retrieved from https://www.mayoclinic.org/diseases-conditions/liver-problems/ diagnosis-treatment/drc-20374507, last visited Mar. 29, 2020, 8 pages.

Medina-Aparicio, L. et al. (May 2011, e-pub. Mar. 11, 2011). "The CRI SPR/Cas Immune System is an Operon Regulated by LeuO, H—NS, and Leucine-Responsive Regulatory Protein in *Salmonella enterica* Serovar Typhi," Journal of Bacteriology 193(10):2396-2407.

Mei, J.-M. et al. (1997). "Identification of *Staphylococcus aureus* Virulence Genes in a Murine Model of Bacteraemia Using Signature-Tagged Mutagenesis," Molecular Microbiology 26(2):399-407.

Mercenier, A. (1990). "Molecular Genetics of *Streptococcus* thermophiles," FEMS Microbiology Letters 87(1-2):61-77.

Mick, E. et al. (May 2013). "Holding a Grudge: Persisting Anti-Phage CRISPR Immunity in Multiple Human Gut Microbiomes," RNA Biology 10(5):900-906.

Mills, S. et al. (Jan./Feb. 2013). "Movers and Shakers: Influence of Bacteriophages in Shaping the Mammalian Gut Microbiota," Gut Microbes 4(1):4-16.

Mitsuhashi, K. et al. (Mar. 13, 2015). "Association of *fusobacterium* Species in Pancreatic Cancer Tissues With Molecular Features and Prognosis," Oncotarget 6(9):7209-7220.

Nakamura, S. et al. (Nov. 2008). "Metagenomic Diagnosis of Bacterial Infections," Emerging Infectious Diseases 14(11):1784-1786.

Nale, J.Y. et al. (2012). "Diverse Temperate Bacteriophage Carriage in Clostridium Difficile 027 Strains," PloS One 7(5):e37263, 9 pages.

Navarre, L. et al. (2007). "Silencing of Xenogeneic DNA by H—NS—Facilitation of Lateral Gene Transfer in Bacteria by a Defense System That Recognizes Foreign DNA," Genes & Development 21:1456-1471.

Nelson, M.H. et al. (2015). "Harnessing the Microbiome to Enhance Cancer Immunotherapy," Journal of Immunology Research 2015: Article 368736, 12 pages.

News (May 22, 2018). "UK Government and Bill & Melinda Gates Foundation Join Carb-X Partnership in Fight Against Superbugs: Millions Earmarked to Boost Research Into New Life-Saving Products to Address the Global Rise of Drug-Resistant Bacteria," Combating Antibiotic Resistant Bacteria, 7 pages.

Noonan, K.A. et al. (May 20, 2015). Adoptive Transfer of Activated Marrow-Infiltrating Lymphocytes Induces Measurable Antiumor Immunity in the Bone Marrow in Multiple Myeloma Science Translational Medicine 7(228):288ra78, 14 pages.

Norris, J.S. et al. (2000). "Prokaryotic Gene Therapy to Combat Multidrug Resistant Bacterial Infection," Gene Therapy 7:723-725. Notice of Intent to Issue Ex Parte Reexamination Certificate, mailed Aug. 12, 2019, for U.S. Appl. No. 90/014,184, filed Aug. 10, 2018, 26 pages.

Nowak, P. et al. (Nov. 28, 2015). "Gut Microbiota Diversity Predicts Immune Status in HIV-1 Infection," AIDS 29(18):2409-2418.

Park, A. (Oct. 18, 2011). "A Surprising Link Between Bacteria and Colon Cancer," Cancer retrieved from http://healthlande.time.com/2011/10/18/a-surprising-link-between-bacteria-and-colon-cancer/, last visited Aug. 27, 2019, 3 pages.

Park, H. et al. (2005). "A Distinct Lineage of CD4 T Cells Regulates Tissue Inflammation by Producing Interleukin 17," Nat. Immunol. 6(11):1133-1141, 24 pages.

Patterson, A.G. et al. (2017, e-pub. Mar. 27, 2017). "Regulation of CRISPR-Cas Adaptive Immune Systems," Current Opinion in Microbiology 37:1-7.

Patterson, A.G. et al. (Dec. 15, 2016). "Quorum Sensing Controls Adaptive Immunity Through the Regulation of Multiple CRISPR-Cas Systems," Mol. Cell 64(6):1102-1108.

Pawluk, A. et al. (Apr. 15, 2014). "A New Group of Phage Anti-CRISPR Genes Inhibits the Type I-E CRISPR-Cas System of Pseudomonas aeruginosa," mBio. 5(2):e00896.

Perez-Chanona, E. et al. (2016, e-pub. Jan. 26, 2016). "The Role of Microbiota in Cancer Therapy," Current Opinion in Immunology 39:75-81.

Pires, D.P. et al. (Sep. 2016, e-pub. Jun. 1, 2016). "Genetically Engineered Phages: A Review of Advances Over the Last Decade," Microbiology and Molecular Biology Reviews 80(3):523-543.

Pul, Ü. et al. (2010, e-pub. Feb. 17, 2010). "Identification and Characterization of *E. coli* CRISPR-cas Promoters and Their Silencing by H—NS," Molecular Microbiology 75(6):1495-1512.

Ramalingam, S.S. et al. (2014). "LB2-Metastatic Non-Small Cell Lung Cancer: Phase II Study of Nivolumab (Anti-PD-1, BMS-936558, ONO-4538) In Patients With Advanced, Refractory Squamous Non-Small Cell Lung Cancer," International Journal of Radiation Oncology Biology Physics Late Breaking Abstract (LB2).

Ran, F.A.et al. (Apr. 9, 2015). "In Vivo Genome Editing Using *Staphylococcus aureus* Cas9," Nature 570(7546):186-191, 28 pages. Rashid, T. et al. (2013). "The Role of Klebsiella in Crohn's Disease With a Potential for the Use of Antimicrobial Measures," International Journal of Rheumatology 2013(Article ID 610393):1-9.

Ray, K. (Jan. 2020). "Manipulating the Gut Microbiota to Combat Alcoholic Hepatitis," Nature Reviews Gastroenterology & Hepatology 17:3, 1 page.

Request for Ex Parte Reexamination mailed Aug. 10, 2018, for U.S. Appl. No. 15/160,405, now U.S. Pat. No. 9,701,964, 42 pages. Request for Ex Parte Reexamination mailed Nov. 1, 2018, for U.S.

Appl. No. 15/160,405, now U.S. Pat. No. 9,701,964, 35 pages.

Richter, C. et al. (2012, e-pub. Oct. 19, 2012). "Function and Regulation of Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR) / CRISPR Associated (Cas) Systems," Viruses 4(12):2291-2311.

Ridaura, V.K. et al. (Sep. 6, 2013). "Cultured Gut Microbiota From Twins Discordant for Obesity Modulate Adiposity and Metabolic Phenotypes in Mice," Science 341(6150):1241214, 22 pages.

Roberts, A.P. et al. (Jun. 2009, e-pub. May 20, 2009). "A Modular Master on the Move: The Tn916 Family of Mobile Genetic Elements," Trends Microbiol. 17(6):251-258. Abstract Only.

Rogers, L. et al. (2016). "*Escherichia coli* and Other Enterobacteriaceae: Occurrence and Detection," Encyclopedia of Food and Health pp. 545-551.

Rong, Z. et al. (Mar. 14, 2014). "Homologous Recombination in Human Embryonic Stem Cells Using CRISPR/Cas9 Nickase and a Long DNA Donor Template," Protein & Cell 5(4):258-260.

Routy, B. et al. (Jan. 5, 2018, e-pub. Nov. 2, 2017). "Gut Microbiome Influences Efficacy of PD-1-Based Immunotherapy Against Epithelial Tumors," Science 359(6371):91-97.

Samaržija, D. et al. (2001). "Taxonomy, Physiology and Growth of Lactococcus Lactis: A Review," Mljekarstvo 51(1):35-48.

Sapranauskas, R. et al. (Nov. 1, 2011, e-pub. Aug. 3, 2011). "The *Streptococcus thermophilus* CRISPR/Cas System Provides Immunity in *Escherichia coli*," Nucleic Acids Research 39(21):9275-9282.

Schnabi, B.G. (2020), "The Role of Enterococcus Faecalis in Alcoholic Liver Disease," retrieved from https://grantome.com/grant/NIH/O01-BX004594-01A2, last visited Oct. 20, 2020, 2 pages.

OTHER PUBLICATIONS

Seed, K.D. et al. (Feb. 27, 2013). "A Bacteriophage Encodes Its Own CRISPR/Cas Adaptive Response to Evade Host Innate Immunity," Nature 494(7438):489-491.

Selle, K. et al. (Apr. 1, 2015). "Harnessing CRISPR-Cas Systems for Bacterial Genome Editing," Trends in Microbiology 23(4):225-232.

Sepsis Alliance. (Dec. 14, 2017). "What Are Vaccines," Retrieved from https://www.sepsis.org/sepsisand/prevention-vaccinations/; last visited Jul. 8, 2019, 3 pages.

Sepsis Alliance. (Jul. 8, 2019). "Prevention," Retrieved from https:// www.sepsis.org/sepsisand/prevention/; accessed last visited Jul. 8, 2019, 5 pages.

Shoemaker, N.B. et al. (Feb. 2001). "Evidence for Extensive Resistance Gene Transfer Among *bacteroides* spp. and Among Bacteroides and Other Genera in the Human Colon," Appl. Environ. Microbiol. 67(2):561-68.

Sivan, A. et al. (Nov. 27, 2015, e-pub Nov. 5, 2015). "Commensal Bifidobacterium Promotes Antitumor Immunity and Facilitates Anti-PD-L1 Efficacy," Science 350(6264):1084-1089, 13 pages.

Sivan, A. et al. (Nov. 6, 2014). "Evidence Implicating the Commensal Microbiota in Shaping Anti-Tumor Immunity in Melanoma," Journal for ImmunoTherapy of Cancer 2(Suppl. 3):O11, 1 page.

Skennerton, C.T. et al. (May 2011). "Phage Encoded H-NS: A Potential Achilles Heel in the Bacterial Defence System," PLoS One 6(5):e20095.

Somkuti, G. A. et al. (Apr. 1988). "Genetic Transformation of *Streptococcus thermophilus* by Electroporation," Biochimie 70(4):579-585. Abstract Only.

Sorek, R. et al. (2013, e-pub. Mar. 11, 2013). "CRISPR-Mediated Adaptive Immune Systems in Bacteria and Archaea," Annual Review of Biochemistry 82:237-266.

Sorg, R. A. et al. (2014). "Gene Expression Platform for Synthetic Biology in the Human Pathogen *Streptococcus pneumoniae*," ACS Synthetic Biology 4(3):228-239. Abstract Only.

Soutourina, O.A. et al. (May 9, 2013). "Genome-Wide Identification of Regulatory RNAs in the Human Pathogen Clostridium difficile," PLos Genet. 9(5):e1003493, 20 pages.

Stern, A. et al. (2012). "CRISPR Targeting Reveals a Reservoir of Common Phages Associated With the Human Gut Microbiome," Genome Research 22(10):1985-1994.

Stern, A. et al. (Aug. 2010), Self-Targeting by CRISPR: Gene Regulation or Autoimmunity? Trends Genet. 26(8):335-340, 10 pages.

Stiefel, U. et al. (Aug. 2014, e-pub. May 27, 2014). "Gastrointestinal Colonization With a Cephalosporinase-Producing *Bacteroides* Species Preserves Colonization Resistance Against Vancomycin-Resistant Enterococcus and Clostridium Difficile in Cephalosporin-Treated Mice," Antimicrob. Agents Chemother. 58(8):4535-4542. Stoebel, D.M. et al. (2008). "Anti-Silencing: Overcoming H—NS-Mediated Repression of Transcription in Gramnegative Enteric Bacteria," Microbiology 154:2533-2545.

Suvorov, A. (1988). "Transformation of Group A Streptococci by Electroporation," FEMS Microbiology Letters 56(1):95-100.

Takaishi, H. et al. (2008). "Imbalance in Intestinal Microflora Constitution Could be Involved in the Pathogenesis of Inflammatory Bowel Disease," Int. J. Med. Microbiol.298:463-472.

Takeda, T. et al. (2011). "Distribution of Genes Encoding Nucleoid-Associated Protein Homologs in Plasmids," International Journal of Evolutionary Biology 2001:685015, 31 pages.

Tan, J. (Dec. 17, 2015). "Immunotherapy Meets Microbiota," Cell 163:1561.

Tarr, P.I. et al. (Mar. 19-25, 2005). "Shiga-Toxin-Producing *Escherichia coli* and Haemolytic Uraemic Syndrome," Lancet 365(9464):1073-1086.

Todar, K. (2012). "The Normal Bacterial Flora of Humans," Todar's Online Textbook of Bacteriology, 8 pages.

Topalian, S.L. et al. (Jun. 28, 2012). "Safety, Activity, and Immune Correlates of Anti-PD-1 Antibody in Cancer," N. Engl. J. Med. 366(26):2443-2454, 19 pages.

Turnbaugh, P.J. et al. (Dec. 2006). "An Obesity-Associated Gut Microbiome With Increased Capacity for Energy Harvest," Nature 444:1027-1131.

U.S. Appl. No. 62/168,355, filed May 29, 2015, Barrangou, R. et al.(Copy not submitted herewith pursuant to the waiver of 37 C.F.R. 1.98(a)(2)(iii) issued by the Office dated Sep. 21, 2004).

U.S. Appl. No. 62/296,853, filed Feb. 18, 2016, Barrangou, R. et al.(Copy not submitted herewith pursuant to the waiver of 37 C.F.R. 1.98(a)(2)(iii) issued by the Office dated Sep. 21, 2004).

Uchiyama, J. et al. (2013, e-pub. Mar. 8, 2013). "Characterization of Helicobacter pylori Bacteriophage KHP30," Applied and Environmental Microbiology 79(10):3176-3184.

USPTO Interference 106,123—Declaration to Declare Interference Jun. 11, 2020, 11 pages.

USPTO Interference 106,123—Junior Party Annotated Claims Jul. 9, 2020, 31 pages.

USPTO Interference 106,123—Junior Party List of Motions Jul. 16, 2020, 6 pages.

USPTO Interference 106,123—Redeclaration Jul. 21, 2020, 6 pages. USPTO Interference 106,123—Rockefeller Clean Claims Jun. 25, 2020, 7 pages.

USPTO Interference 106,123—Rockefeller Motion 2 (Indefiniteness), Oct. 16, 2020, 24 pages.

USPTO Interference 106,123—Rockefeller Notice of Lead and Backup Counsel Jun. 25, 2020, 3 pages.

USPTO Interference 106,123—Rockefeller Notice of Real Party in Interest Jun. 25, 2020, 3 pages.

USPTO Interference 106,123—Rockefeller Notice of Related Proceedings Jun. 25, 2020, 3 pages.

USPTO Interference 106,123—Rockefeller Power of Attorney Jun. 25, 2020, 3 pages.

USPTO Interference 106,123—Rockefeller Request for File Copies Jun. 25, 2020, 10 pages.

USPTO Interference 106,123—Rockefeller Revised List of Proposed Motions Aug. 13, 2020, 4 pages.

USPTO Interference 106,123—Senior Party List of Proposed Motions Jul. 16, 2020, 5 pages.

USPTO Interference 106,123—SNIPR Clean Claims Jun. 25, 2020, 27 pages.

USPTO Interference 106,123—SNIPR Motion 2 (Lack of Enablement and Written Description), Oct. 16, 2020, 32 pages.

USPTO Interference 106,123—SNIPR Motion 4 (Deny Benefit to Count 1), Oct. 16, 2020, 16 pages.

USPTO Interference 106,123—SNIPR Motion 5 (Substitute Count), Oct. 16, 2020, 41 pages.

USPTO Interference 106,123—SNIPR Motion 6 (Motion to Designate Claims as Not Corresponding to Count 1 or Proposed Count 2), Oct. 16, 2020, 24 pages.

USPTO Interference 106,123—SNIPR Notice of Lead and Backup Counsel Jun. 25, 2020, 4 pages.

USPTO Interference 106,123—SNIPR Notice of Related Proceedings Jun. 25, 2020, 4 pages.

USPTO Interference 106,123—SNIPR Real Party in Interest Jun. 25, 2020, 3 pages.

USPTO Interference 106,123—SNIPR Request for File Copies Jun. 25, 2020, 10 pages.

USPTO Interference 106,123—Standing Order Jun. 11, 2020, 81 pages.

USPTO Interference 106,123—Joint Stipulated Extension of Time, Sep. 4, 2020, 4 pages.

USPTO Interference 106,123—Junior Party Revised List of Motions Aug. 13, 2020, 6 pages.

USPTO Interference 106,123-Memorandum, Jan. 19, 2021, 6 pages.

USPTO Interference 106,123—Notice of Cross Examination—van der Oost, Dec. 1, 2020, 3 pages.

USPTO Interference 106,123—Order Additional Applications 37 C.F.R. § 41.104(a), Sep. 3, 2020, 6 pages.

USPTO Interference 106,123—Order Authorizing Motions and Setting Times 37 C.F.R. 11.104(c) and 121 Aug. 24, 2020, 10 pages.

OTHER PUBLICATIONS

USPTO Interference 106,123—Order—Additional Applications, Jan. 13, 2021, 6 pages.

USPTO Interference 106,123—Order—Bd.R. 109(b)—Authorizing Office Records Jul. 21, 2020, 3 pages.

USPTO Interference 106,123—Order—Video Dispositions 37 C.F. R. § 41.104(a), Sep. 25, 2020, 3 pages.

USPTO Interference 106,123—Rockefeller List of Exhibits, Oct. 16, 2020, 4 pages.

USPTO Interference 106,123—Rockefeller List of Exhibits, Nov. 13, 2020, 4 pages.

USPTO Interference 106,123—Rockefeller List of Exhibits, Feb. 19, 2021, 5 pages.

USPTO Interference 106,123—Rockefeller Motion 1 (Lack of Written Description), Oct. 16, 2020, 30 pages.

USPTO Interference 106,123—Rockefeller Motion 3 (To Add a Claim), Nov. 13, 2020, 36 pages.

USPTO Interference 106,123—Rockefeller Notice of Settlement Discussions, Oct. 21, 2020, 3 pages.

USPTO Interference 106,123—Rockefeller Order—Responsive Motion 37 C.F.R. § 41.121(a)(2), Nov. 2, 2020, 2 pages.

USPTO Interference 106,123—Rockefeller Reply 1, Feb. 19, 2021, 51 pages.

USPTO Interference 106,123—Rockefeller Reply 2, Feb. 19, 2021, 37 pages.

USPTO Interference 106,123—Rockefeller Reply 3, Feb. 19, 2021, 48 pages.

USPTO Interference 106,123—Rockefeller Updated Notice of Related Proceedings, Nov. 13, 2020, 3 pages.

USPTO Interference 106,123—SNIPR Exhibit List, Oct. 16, 2020, 7 pages.

USPTO Interference 106,123—SNIPR Exhibit List, Feb. 19, 2021, 8 pages.

USPTO Interference 106,123—SNIPR Motion 1 (Terminate Interference as Contrary to AIA), Oct. 16, 2020, 20 pages.

USPTO Interference 106,123—SNIPR Request for Oral Argument, Mar. 12, 2021, 4 pages.

USPTO Interference 106,123-Rockefeller Request for Oral Argument, Mar. 12, 2021, 3 pages.

USPTO Interference 106,123—SNIPR Reply 1, Feb. 19, 2021, 19 pages.

USPTO Interference 106,123—SNIPR Reply 2, Feb. 19, 2021, 42 pages.

USPTO Interference 106,123—SNIPR Reply 4, Feb. 19, 2021, 28 pages.

USPTO Interference 106,123—SNIPR Reply 5, Feb. 19, 2021, 44 pages.

USPTO Interference 106,123—SNIPR Reply 6, Feb. 19, 2021, 27 pages.

Veeranagouda, Y. et al. (Jun. 4, 2014). "Identification of Genes Required for the Survival of B. fragilis Using Massive Parallel Sequencing of a Saturated Transposon Mutant Library," BMC Genomics 15:429, 11 pages.

Vega, N.M. et al. (Oct. 2014). "Collective Antibiotic Resistence: Mechanisms and Implications," Curr. Opin. Microbiol. 21:28-34, 14 pages.

Vercoe, R.B. et al. (Apr. 18, 2013). "Cytotoxic Chromosomal Targeting by CRISPR/Cas Systems Can Reshape Bacterial Genomes and Expel or Remodel Pathogenicity Islands," PLOS Genetics 9(4):e1003454, 13 pages.

Villarino, N.F. et al. (Feb. 23, 2016, e-pub. Feb. 8, 2016). "Composition of the Gut Microbiota Modulates the Severity of Malaria," Proc. Natl. Acad. Sci. USA 113(8):2235-2240.

Vétizou, M. et al. (Nov. 27, 2015, e-pub Nov. 5, 2015). "Anticancer Immunotherapy by CTLA-4 Blockade Relies on the Gut Microbiota," Science 350(6264):1079-1084, 13 pages.

Walters, W.A. et al. (Nov. 17, 2014). "Meta-Analyses of Human Gut Microbes Associated With Obesity and IBD," FEBS Letters 588(22):4223-4233, 34 pages. Wang, I.-N. et al. (2000). "HOLINS: The Protein Clocks of Bacteriophage Infections," Annu. Rev. Microbiol. 54:799-825.

Wang, J. et al. (2019). "Core Gut Microbiota Analysis of Feces in Healthy Mouse Model," Supplementary Information, 12 pages.

Wang, J. et al. (Apr. 2019). "Core Gut Bacteria Analysis of Healthy Mice," Frontiers in Microbiology 10(887):1-14.

Wegmann, U. et al. (Apr. 2007). "Complete Genome Sequence of the Prototype Lactic Acid Bacterium *Lactococcus lactis* Subsp. *cremoris* MG 1363," Journal of Bacteriology 189(8):3256-3270.

Wei, Y. et al. (2015, e-pub. Jan. 14, 2015). "Sequences Spanning the Leader-Repeat Junction Mediate CRISPR Adaptation to Phage in *Streptococcus thermophiles*," Nucleic Acids Research 43(3):1749-1758.

Weir, T.L. et al. (Aug. 6, 2013). "Stool Microbiome and Metabolome Differences Between Colorectal Cancer Patients and Healthy Adults," PLOS One 8(8):e70803, 10 pages.

Westra, E.R. et al. (Jun. 8, 2012). "CRISPR Immunity Relies on the Consecutive Binding and Degradation of Negatively Supercoiled Invader DNA by Cascade and Cas3," Molecular Cell 46:595-605. Westra, E.R. et al. (Sep. 1, 2010, e-pub. Aug. 18, 2010). "H—NS-

Mediated Repression of CRISPR-Based mmunity in *Escherichia coli* K12 Can be Relieved by the Transcription Activator LeuO," Molecular Microbiology 77(6):1380-1393.

Westwater, C. et al. (2002). "Development of a P1 Phagemid System for the Delivery of DNA Into Gram-Negative Bacteria," Microbiology 148:943-950.

Westwater, C. et al. (Apr. 2003). "Use of Genetically Engineered Phage to Deliver Antimicrobial Agents to Bacteria: An Alternative Therapy for Treatment of Bacterial Infections," Antimicrobial Agents and Chemotherapy 47(4):1301-1307.

Wexler, H.M. (Oct. 2007). "Bacteroides: the Good, the Bad, and the Nitty-Gritty," Clinical Microbiology Reviews 20(4):593-621.

Written Opinion for PCT Application No. PCT/EP2016/059803, dated Jun. 30, 2016, filed May 3, 2016, 6 pages.

Written Opinion for PCT/EP2018/082053, dated Mar. 14, 2019, filed Nov. 21, 2018, 6 pages.

Xie, Z. et al. (2013, e-pub. Aug. 9, 2013). "Development of a Tunable Wide-Range Gene Induction System Useful for the Study of Streptococcal Toxin-Antitoxin Systems," Applied and Environmental Microbiology 79(20):6375-6384.

Xu, T. et al. (Jul. 2015). "Efficient Genome Editing in Clostridium cellulolyticum via CRISPR-Cas9 Nickase," Applied and Environmental Microbiology 81(13):4423-4431.

Yang, Y. et al. (Jun. 5, 2014, e-pub. Apr. 13, 2014). "Focused Specificity of Intestinal Th17 Cells Towards Commensal Bacterial Antigens," Nature 510(7503):152-156, 29 pages.

Yao, J. et al. (2016, e-pub. May 9, 2016). "A Pathogen-Selective Antibiotic Minimizes Disturbance to the Microbiome," Antimicrob. Agents Chemother., 24 pages.

Yosef, I. et al. (2011). "High-Temperature Protein G is Essential for Activity of the *Escherichia coli* Clustered Regularly Interspaced Palindromic Repeats (CRISPR)/Cas System," Proc. Natl. Acad. Sci. USA 108(50):20136-20141.

Yosef, I. et al. (Jun. 9, 2015). "Temperate and Lytic Bacteriophages Programmed to Sensitize and Kill Antibiotic-Resistant Bacteria," Proc. Natl. Acad. Sci. USA 112(23):7267-7272.

Young, R. et al. (1995). "Holins: Form and Function in Bacteriophage Lysis," FEMS Microbiology Reviews 17:191-205.

YourGenome: CRISPR/CAS9, retrieved from https://www.yourgenonne. org/facts/what-is-crispr-cas9, last visited Jan. 6, 2020, 8 pages.

Yu, Z. et al. (Mar. 21, 2014). "Various Applications of TALEN- and CRISPR/Cas9-Mediated Homologous Recombination to Modify the *Drosophila* Genome," Biology Open 3(4):271-280.

Zhang, R. et al. (2009, e-pub. Oct. 30, 2008). "DEG 5.0, A Database of Essential Genes in Both Prokaryotes and Eukaryotes," Nucleic Acids Research 37:D455-D458.

Zhang, T. et al. (Sep. 24, 2016). "The Efficacy and Safety of Anti-PD-1/PD-L1 Antibodies for Treatment of Advanced or Refractory Cancers: A Meta-Analysis," Oncotarget 7(45):73068-73079.

Zhang, X.Z. (2011). "Simple, Fast and High-Efficiency Transformation System for Directed Evolution of Cellulase in Bacillus Subtilis," Microbial Biotechnology 4(1):98-105.

OTHER PUBLICATIONS

Zitvogel, L. et al. (Jan. 2015), "Cancer and the Gut Microbiota: An Unexpected Link," Sci. Transl. Med. 7(271):271ps1, 10 pages.

Hase, K. (Nov. 2014). "Intestinal Microbiota and Immunity," Infectious Disease (in Japanese). 44(6):193-200 22 pages. English Translation.

Ichikawa, M. et al. (Dec. 2019). "The Relationship Between Gut Microbiome, Immune System, and Cancer," Jpn. J. Cancer Chemother 16(12):1807-1813. English Abstract.

Abedon, S.T. et al. (Dec. 2003). "Experimental Examination of Bacteriophage Latent-Period Evolution as a Response to Bacterial Availability," Applied and Environmental Microbiology 69(12):7499-7506.

Advisory Action, dated Dec. 9, 2021 for U.S. Appl. No. 90/014,705, filed Mar. 23, 2021, 9 pages.

Bellanger, X. et al. (Jul. 1, 2014, e-pub. Jan. 27, 2014). "Conjugative and Mobilizable Genomic Islands in Bacteria: Evolution and Diversity," FEMS Microbiology Reviews 38(20144):720-760.

Cronan, J.E. (Jan. 2013). "Improved Plasmid-Based System for Fully Regulated Off-To-On Gene Expression in *Escherichia coli*: Application to Production of Toxic Proteins," Plasmid 69(1):81-89, 17 pages.

European Office Action, dated Jun. 29, 2021, for European Patent Application No. 16719873.8, 24 pages.

European Search Report, dated Oct. 4, 2021, for European Patent Application No. 21170379.8, 6 pages.

European Search Report, dated Oct. 8, 2021, for European Patent Application No. 21170380.6, 7 pages.

Ex Parte Re-Exam Communication Transmittal Form, dated Jun. 30, 2021, for U.S. Appl. No. 90/014,705, for U.S. Patent Reexamination 10,953,090, 26 pages.

Ex Parte Re-Exam, mailed Dec. 16, 2021, for U.S. Appl. No. 90/014,877, filed Oct. 6, 2021, for U.S. Patent Reexamination 10,953,090, 12 pages.

Ex Parte Re-Exam, mailed Nov. 15, 2021, for U.S. Appl. No. 90/014,877, 12 pages.

Ex Parte Re-Exam, mailed Sep. 27, 2021, for U.S. Appl. No. 90/014,705, filed Mar. 23, 2021, for U.S. Patent Reexamination 10,953,090, 16 pages.

Gutierrez, B. et al. (Apr. 30, 2018). "Genome-Wide CRISPR-Cas9 Screen in *E. coli* Identifies Design Rules for Efficient Targeting," 22 pages.

Hamanishi, J. et al. (2016, e-pub. Feb. 22, 2016). "PD-1/PD-L1 Blockade in Cancer Treatment: Perspectives and Issues," International Journal of Clinical Oncology 21:462-473.

Johnson, C. M. et al. (Nov. 23, 2015). "Integrative and Conjugative Elements (ICEs): What They Do and How They Work," Annual Review of Genetics 49(1):577-601, 33 pages.

Kaulich, M. et al. (2015, e-pub. Jan. 13, 2015). "Efficient CRISPRrAAV Engineering of Endogenous Genes to Study Protein Function by Allele-Specific RNAi," Nucleic Acids Research 43(7):e45, 8 pages.

Keskin, H. et al. (Nov. 20, 2014). "Transcript-RNA-Templated DNA Recombination and Repair," Nature 515:436-439.

Kugelberg, E. et al. (Aug. 2005). "Establishment of a Superficial Skin Infection Model in Mice by Using *Staphylococcus aureus* and *Streptococcus pyogenes*," Antimicrob Agents Chemother 49(8):3435-3441.

Leshem, A. et al. (Sep. 29, 2020). "The Gut Microbiome and Individual-Specific Responses to Diet," mSystems 5(5):e00665-20, 12 pages.

Office Action, dated Nov. 4, 2021 for U.S. Appl. No. 90/014,705, filed Mar. 23, 2021, 7 pages.

Pastagia, N. et al. (Feb. 2011). "A Novel Chimeric Lysin Shows Superiority to Mupirocin for Skin Decolonization of Methicillin-Resistant and -Sensitive *Staphylococcus aureus* Strains," Antimicrobial Agents and Chemotherapy 55(2):738-744.

Rea, K. et al. (2020, e-pub. Nov. 14, 2019). "Gut Microbiota: A Perspective for Psychiatrists," Neuropsychobiology 79:50-62.

Request for Ex Parte Reexamination under 35 U.S. C. \S 302 and 37 C.F.R. \S 1.510, dated Feb. 16, 2021, 72 pages.

Roberts, A.P. et al. (Dec. 1, 2003). "Development of an Integrative Vector for the Expression of Antisense RNA in Clostridium difficile," Journal of Microbiological Methods 55(3):617-624.

Saito, H. et al. (Jun. 15, 2016, e-pub. Apr. 12, 2016). "Adoptive Transfer of CD8+ T Cells Generated From Inducted Pluripotent Stem Cells Triggers Regressions of Large Tumors Along With Immunological Memory," Cancer Research 76(12):3473-3483.

Svenningsen, S.L. et al. (Mar. 22, 2005). "On the Role of Cro in λ Prophage Induction," PNAS 102(12):4465-4469.

Tlaskalová-Hogenová, H. et al. (2011, e-pub. Jan. 31, 2011). "The Role of Gut Microbiota (Commensal Bacteria) and the Mucosal Barrier in the Pathogenesis of Inflammatory and Autoimmune Diseases and Cancer: Contribution of Germ-Free and Gnotobiotic Animal Models of Human Diseases," Cellular & Molecular Immunology 8:110-120.

USPTO Interference 106,123—Decision on Motions, Sep. 7, 2021, 18 pages.

USPTO Interference 106,123—Judgement, Nov. 19, 2021, 3 pages. USPTO Interference 106,123—SNIPR Notice of Appeal, Dec. 14, 2021, 28 pages.

USPTO Interference 106,123—Order—Show Cause, Aug. 19, 2021, 4 pages.

USPTO Interference 106,123—Rockefeller Notice, Aug. 13, 2021, 3 pages.

USPTO Interference 106,123—Rockefeller Response to Show Cause, Sep. 7, 2021, 7 pages.

USPTO Interference 106,123—Rockefeller Updated Notice of Related Proceedings, Jul. 15, 2021, 3 pages.

Waters, J. L. et al. (Nov./Dec. 2013). "Regulation of CTnDOT Conjugative Transfer is a Complex and Highly Coordinated Series of Events," MBIO 4(6):e00569-13, 8 pages.

Wu, J. et al. (Jun. 2019). "Fusobacterium nucleatum Contributes to the Carcinogenesis of Colorectal Cancer by Inducting Inflammation and Suppressing Host Immunity," Translational Oncology 12(6):846-851.

Ericsson, A.C. et al. (Sep. 10, 2015). "Differential Susceptibility to Corolectal Cancer Due to Naturally Occuring Gut Microbiota," Oncotarget 6(32):33689-33707.

Ex Parte Re-Exam Communication Transmittal Form, dated May 25, 2022, for Control No. 901014105, for U.S. Patent Reexamination of U.S. Pat. No. 10,953,090, 19 pages.

Gur, C. et al. (Feb. 17, 2015). "Binding of the Fap2 Protein of Fusobacterium nucleatum to Human Inhibitory Receptor TIGIT Protects Tumors from Immune Cell Attack," Immunity 42(2):344-355.

Mima, K. et al. (Aug. 1, 2015, e-pub. Jun. 4, 2014). "Fusobacterium nucleatum and T-Cells in Colorectal Carcinoma," JAMA Oncol. 1(5):653-661.

Novosiadly, R. et al. (2016, e-pub. Mar. 30, 2016). "High-Content Molecular Profiling of T-Cell Therapy in Oncology," Molecular Therapy—Oncolytics 3:16009, 6 pages.

Request for Ex Parte Re-Exam Under 35 U.S.C. § 302 and 37 C.F.R. § 1.510 dated Jun. 7, 2022, for U.S. Pat. No. 11,291,723, 328 pages. Request for Ex Parte Re-Exam Under 35 U.S.C. § 302 and 37 CPR.

 [§ 1.510 dated Jun. 7, 2022, for U.S. Pat. 11,351,252, 219 pages.
 West, N.R. et al. (Dec. 14, 2015). "Immunolherapy Not Working? Check Your Microbiola," Cancer cell 28:687-689.

* cited by examiner

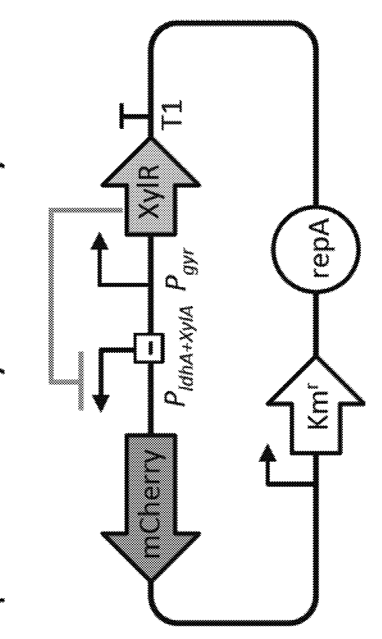


FIG. 1

pBAV1KT5-XylR-mCherry-PldhA

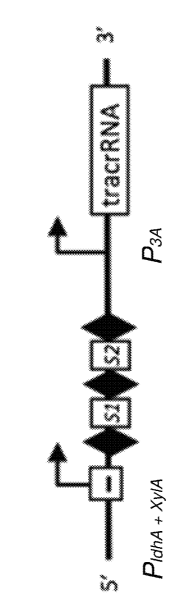


FIG. 2



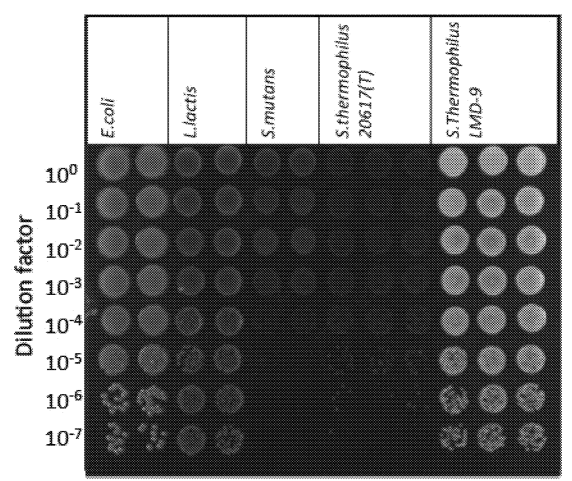


FIG. 4

Commensal gut bacteria

E.coli-MG1655 (40 hrs)

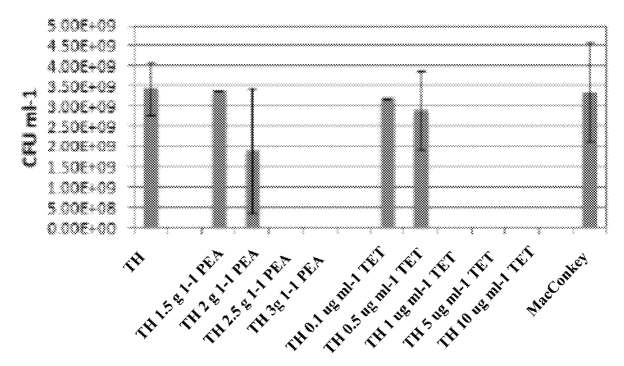


FIG. 4 (Continued)

'Relative' of target species



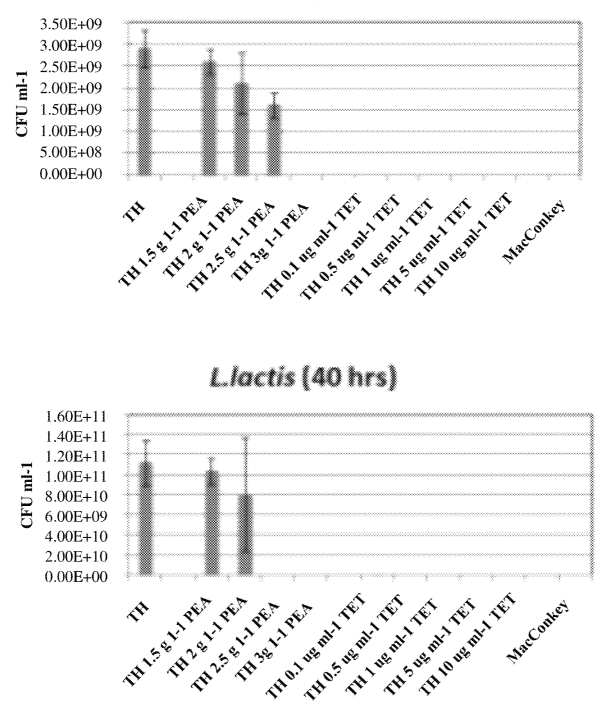
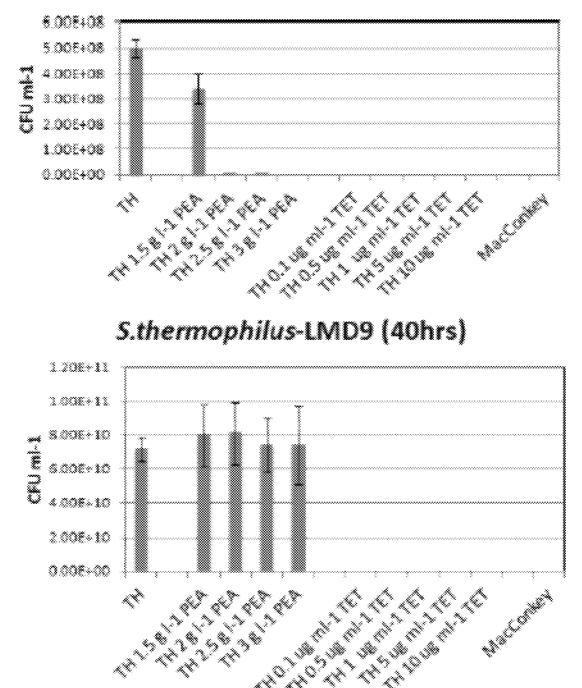
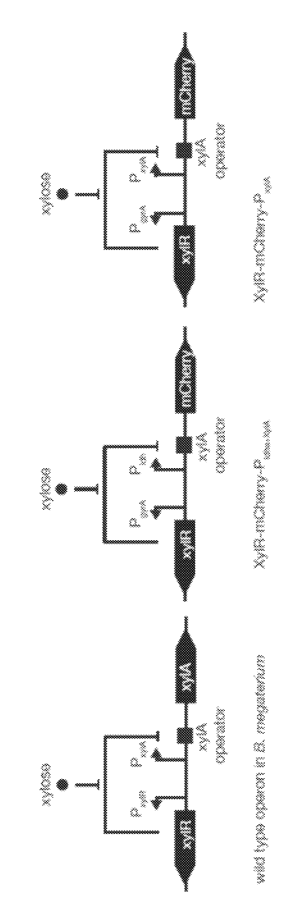


FIG. 4 (Continued)

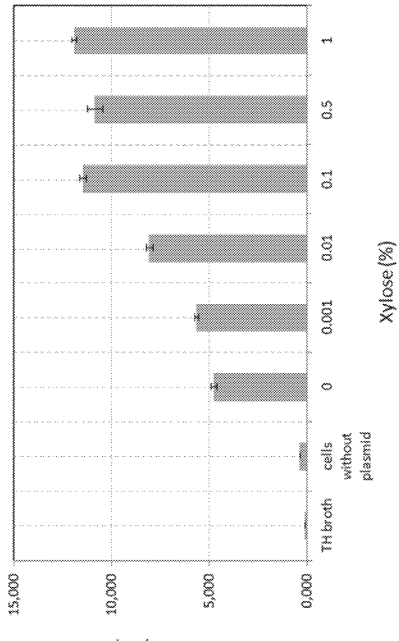
Target species

S.thermophilus-B (40 hrs)



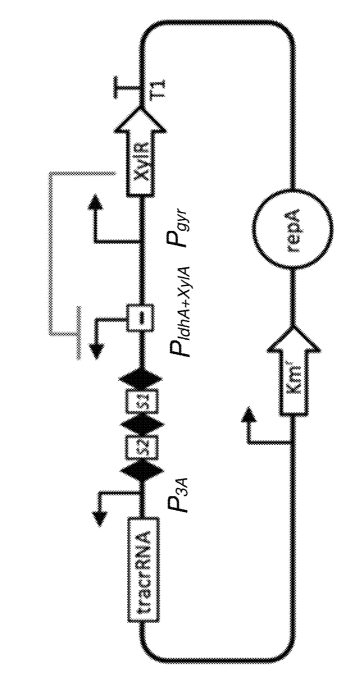








RFP (a.u)





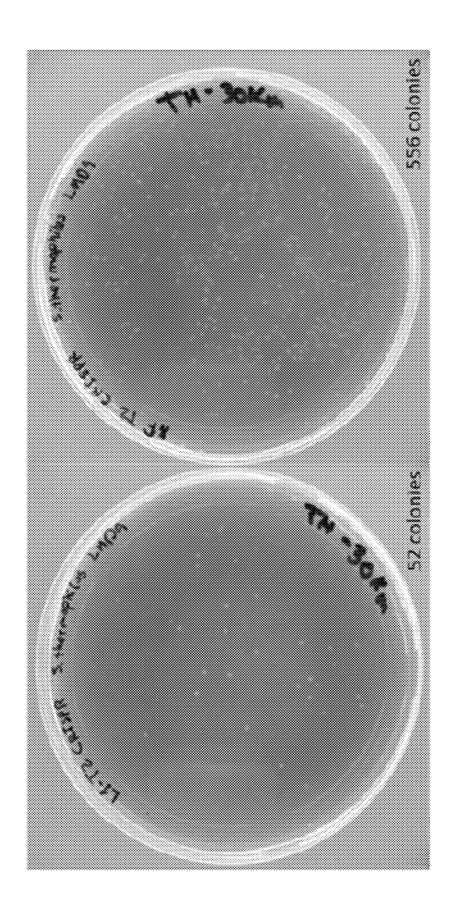
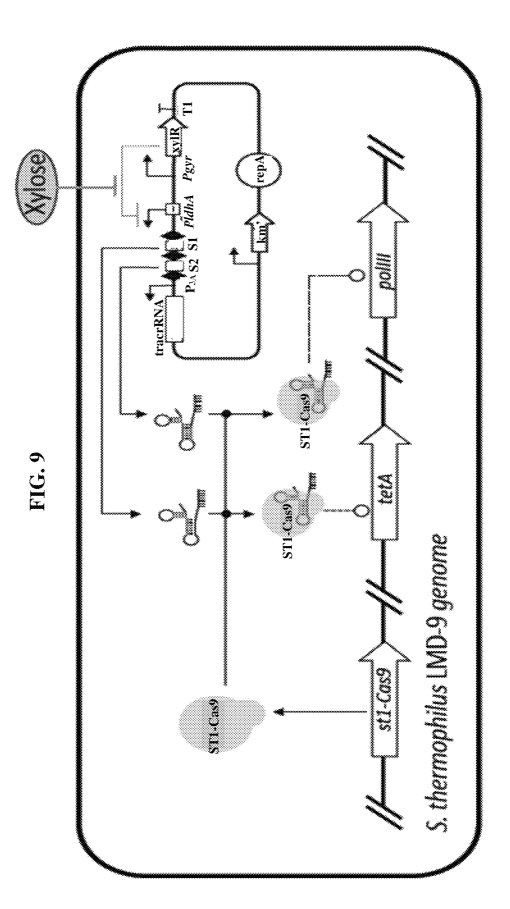
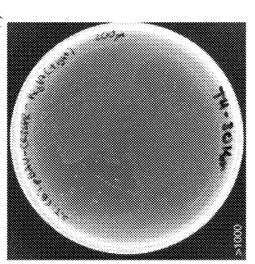


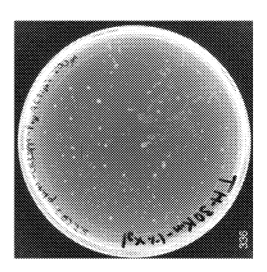
FIG. 8



Streptoccocus thermophilus pBAV1KT5-XyIR-CRISPR-P_{Xyik}



-xylose



ie Xai

a na star a s

+xylose

Streptoccocus thermophilus pBAV1KT5-XyIR-CRISPR-P_{Isma+Kylk}

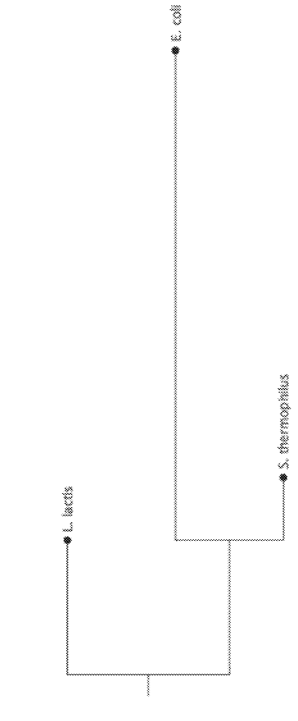
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Sheet 12 of 19

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FIG. 11



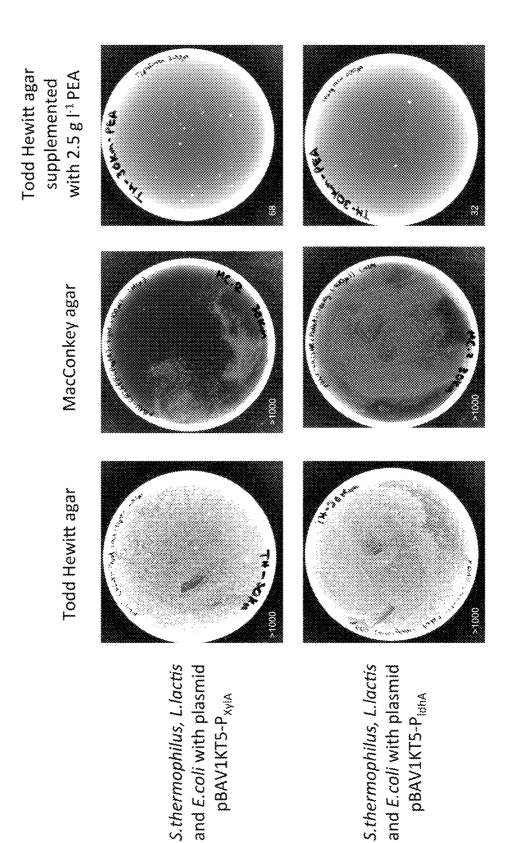
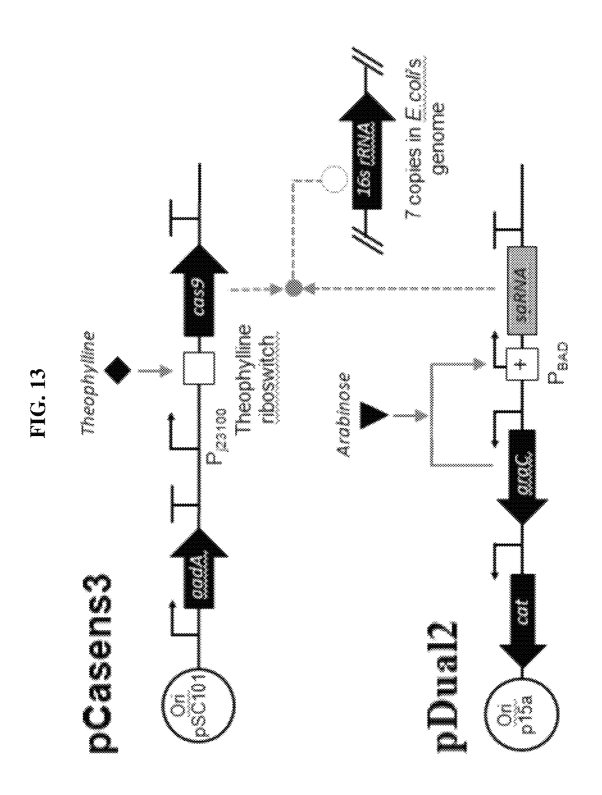
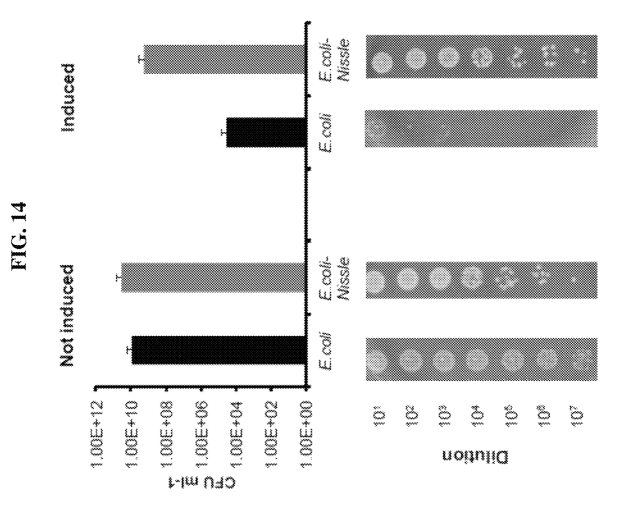
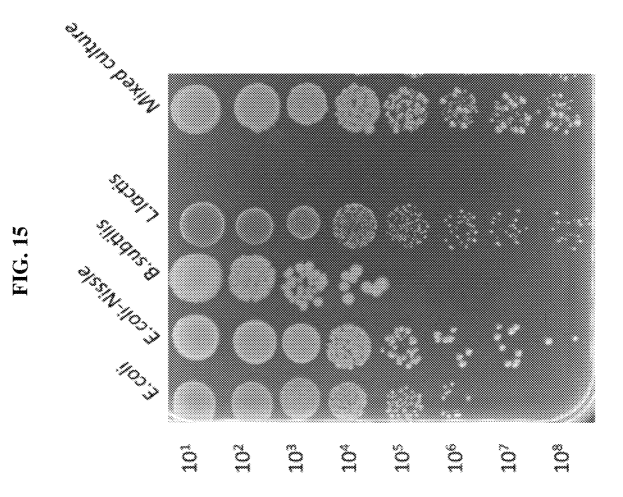


FIG. 12







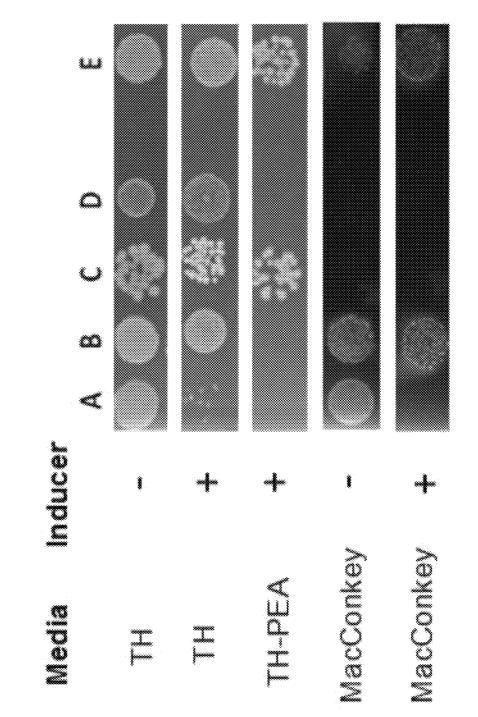
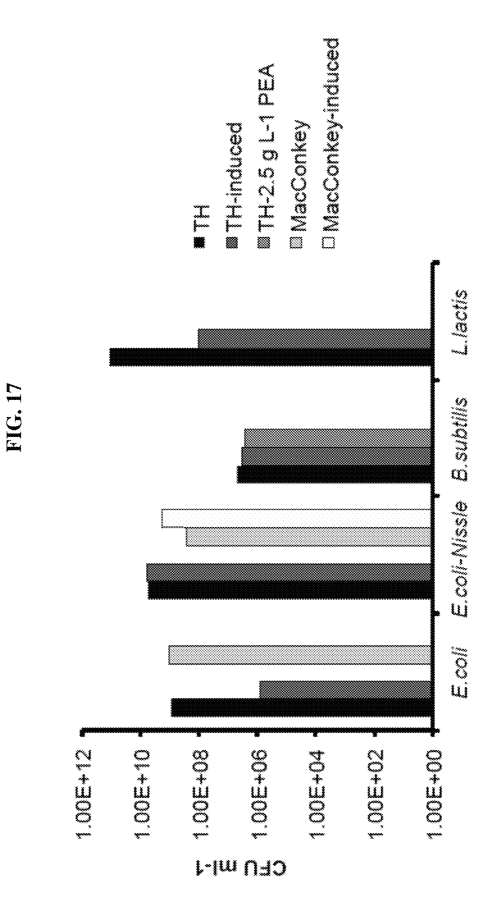


FIG. 16



SELECTIVELY ALTERING MICROBIOTA FOR IMMUNE MODULATION

CROSS REFERENCE TO RELATED APPLICATION

This application is a Continuation Application of U.S. patent application Ser. No. 16/453,598, which was filed on Jun. 26, 2019, which is a Continuation Application of U.S. patent application Ser. No. 16/192,752, which was filed on Nov. 15, 2018 (now U.S. Pat. No. 10,363,308), which is a Continuation Application of U.S. patent application Ser. No. 15/820,296, which was filed on Nov. 21, 2017 (now U.S. Pat. No. 10,195,273), which is a Continuation Application under 35 U.S.C. § 120 of International Patent Application No. PCT/EP2017/063593, which was filed on Jun. 4, 2017, which claims priority benefit to United Kingdom Patent Application No. GB1609811.3, which was filed on Jun. 5, 2016, the contents of each of which are incorporated herein by reference in their entirety.

SUBMISSION OF SEQUENCE LISTING ON ASCII TEXT FILE

The content of the following submission on ASCII text ²⁵ file is incorporated herein by reference in its entirety: a computer readable form (CRF) of the Sequence Listing (file name: 786212000233SEQLIST.TXT, date recorded: Dec. 22, 2020, size: 8 KB).

FIELD OF THE INVENTION

The invention relates to methods of modulating immune cells in a patient (endogenous cells of the patient and/or administered cells, such as via adoptive cell therapy) by 35 altering microbiota of the patient. The invention also relates to methods of modulating treatments or therapies in a subject organism by altering microbiota of the subject. The invention also relates cell populations, systems, kits and other means for effecting this. In an example, advanta- 40 geously selective targeting of a particular species in a human gut microbiota using guided nucleic acid modification is carried out to effect the alteration.

BACKGROUND OF THE INVENTION

One approach to immunotherapy involves engineering patients' own (or a donor's) immune cells to express cellsurface antigen receptors (CARs) that recognise and attack tumours. Although this approach, called adoptive cell trans- 50 fer (ACT), has been restricted to small clinical trials so far, treatments using these engineered immune cells have generated some remarkable responses in patients with advanced cancer.

The Chimeric Antigen Receptor (CAR) consists of an 55 antibody-derived targeting domain fused with T-cell signaling domains that, when expressed by a T-cell, endows the T-cell with antigen specificity determined by the targeting domain of the CAR. CARs can potentially redirect the effector functions of a T-cell towards any protein and non- 60 protein target expressed on the cell surface as long as an antibody-based targeting domain is available. This strategy thereby avoids the requirement of antigen processing and presentation by the target cell and is applicable to nonclassical T-cell targets like carbohydrates. This circumven- 65 tion of HLA-restriction means that the CAR T-cell approach can be used as a generic tool broadening the potential of

applicability of adoptive T-cell therapy. See, eg, Methods Mol Biol. 2012; 907:645-66. doi: 10.1007/978-1-61779-974-7_36, "Chimeric antigen receptors for T-cell based therapy", Cheadle E J et al.

The first CAR-T construct was described in a 1989 paper by immunotherapy pioneer Zelig Eshhar in PNAS. The structure of the CAR now comprises a transmembrane polypeptide chain which is a chimaera of different domains from different cellular proteins. For example, the CAR has an extracellular part joined (often by a linker and/or a hinge region) to an intracellular part, with a transmembrane portion of the CAR embedding the receptor in the membrane of an immune cell, normally a T-cell. The extracellular moiety includes an antibody binding site (usually in the form of an scFv, such as derived from a mouse mAb) that recognizes a target antigen, that commonly is a tumour associated antigen (TAA) on the surface of cancer cells. Antigen recognition in this way dispenses with the need to rely on TCRs that require MHC-restricted antigen presentation, and where binding affinities may be relatively low. The intracellular moiety of the CAR typically includes a CD3-zeta (CD3) domain for intracellular signaling when antigen is bound to the extracellular binding site. Later generation CARs also include a further domain that enhances T-cell mediated responses, which often is a 4-1BB (CD137) or CD28 intracellular domain. On encountering the cognate antigen ligand for the CAR binding site, the CAR can activate intracellular signaling and thus activation of the CAR T-cell to enhance tumour cell killing.

Most CAR-Ts expand in vivo so dose titration in a conventional sense is difficult, and in many cases the engineered T-cells appear to be active "forever"-i.e., the observation of on-going B-cell aplasia seen in most of the CD19 CAR-T clinical studies to date. This poses a serious problem for CAR T-cell approaches. Some observed risks are discussed in Discov Med. 2014 November; 18(100):265-71, "Challenges to chimeric antigen receptor (CAR)-T cell therapy for cancer", Magee M S & Snook A E, which explains that the first serious adverse event following CAR-T cell treatment occurred in a patient with colorectal cancer metastatic to the lung and liver (Morgan et al., 2010). This patient was treated with T cells expressing a thirdgeneration CAR targeting epidermal growth factor receptor 2 (ERBB2, HER2). The CAR contained an scFv derived 45 from the 4D5 antibody (trastuzumab) that is FDA approved for the treatment of HER2-positive breast cancers (Zhao et al., 2009). The patient developed respiratory distress within 15 minutes of receiving a single dose of 1010 CAR-T cells, followed by multiple cardiac arrests over the course of 5 days, eventually leading to death. Serum analysis four hours after treatment revealed marked increases in the cytokines IFNγ, GM-CSF, TNFα, IL-6, and IL-10. CAR-T cells were found in the lung and abdominal and mediastinal lymph nodes, but not in tumour metastases. The investigators attributed toxicity to recognition of HER2 in lung epithelium resulting in inflammatory cytokine release producing pulmonary toxicity and cytokine release syndrome (CRS) causing multi-organ failure (Morgan et al., 2010). Trials utilizing second-generation HER2-targeted CARs derived from a different antibody (FRP5) following conservative dose-escalation strategies are currently underway for a variety of HER2+ malignancies by other investigators (clinicaltrials. gov identifiers NCT01109095, NCT00889954, and NCT00902044).

A variation on the CAR T-cell theme are antibodycoupled T-cell receptor (ACTR) therapeutics, which use CD16 A (FCyRIIIA) to bind to Fc regions of tumour-specific IgG (see eg, WO2015/058018, US2015139943). The aim is to enable more control of CAR T-cell activity in vivo by titrating IgG administered to patients. The CD16 binding sites of the CAR-T-cells may be free, however, to also bind to endogenous IgG of the patients and this reduces the 5 attractiveness of the approach. The approach also needs to consider the inherently long half-life of IgG in the body (around 20 days for IgG in man), which may limit control of CAR-cell activity. Ongoing studies may assess the risk of this.

It would be desirable to provide an alternative way to modulate (downregulate or upregulate) immune cell-based therapies, like CAR-T-cell approaches and other cell-based approaches. It would also be desirable to provide a way to address diseases and conditions mediated by endogenous 15 immune cells, such as autoimmune, inflammatory and infectious diseases and conditions.

STATEMENTS OF INVENTION

The invention provides guided nucleases, host cell modifying (HM)-CRISPR/Cas systems, gRNAs, HM-arrays, HM-crRNA, HM-Cas, HM-TALENs, HM-meganucleases, HM-zinc fingers and methods as set out in the claims herein.

Medical practice often involves the administration of 25 antibiotics to patients. Such treatments can typically involve administration of broad-spectrum antibiotics, or antibiotics that target many gram-positive bacterial species or many gram-negative species without discrimination. Similarly, use of broad-spectrum antibiotics in farming and agriculture, for 30 example, raise environmental concerns, including entry of such antibiotics into the human and animal food chain which may be deleterious to health and may add to development of microbial resistance. Rather, the invention involves selective targeting of a first microbiota species or strain. As shown in 35 the worked examples herein, selective targeting of a particular bacterial species has been achieved using guided nuclease targeting of the genome of the selected species, whilst at the same time sparing phylogenetically related species and strains. Furthermore, the invention realises the 40 role that microbiota bacteria and archaea play in shaping immune function in humans and animals, as discussed further below.

Thus, the invention relates to methods of modulating immune cells in a patient (endogenous cells of the patient 45 and/or administered cells, such as via adoptive cell therapy) by altering microbiota of the patient. In an example, advantageously selective targeting of a species in a microbiota (eg, gut microbiota) is carried out to effect the alteration. Selective targeting may, for example, avoid targeting of related 50 species or strains, such as species of the same phylum or such as a different strain of the same species.

For example, the invention provides for modulating immune cell-based or other therapy of diseases and conditions in patients and subjects by altering microbiota, as well 55 as systems, kits and other means for effecting this.

For example, the invention provides for treating or reducing diseases and conditions in patients by altering microbiota, wherein the diseases and conditions are those mediated by immune cells (eg, T-cells) or addressed by altering 60 immune cell activities or populations in patients. Embodiments are cancers, autoimmune diseases or conditions, inflammatory diseases or conditions, viral infections (eg, HIV infection of human patients), or diseases or conditions mediated or caused by viral infections.

The invention also relates to methods of modulating treatments or therapies in a subject organism (eg, a plant,

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yeast, human or animal patient) by altering microbiota of the subject. Examples of therapies are adoptive cell therapy, antibody therapy (eg, immune checkpoint inhibition), radiation therapy, chemotherapy, eg, for treatment or prevention of a disease or condition in a patient.

In a first configuration the invention provides

A method of modulating a therapy of a disease or condition in a patient, the method comprising

a. Carrying out the therapy in the patient; and

b. Causing gut bacterial microbiota dysbiosis in the patient, 10whereby said dysbiosis modulates the therapy in the patient by modulating immune cells in the patient.

In another aspect, the first configuration the invention provides

A method of modulating a therapy of a disease or condition in a human or animal patient, the method comprising

a. Carrying out the therapy in the patient; and

b. Causing bacterial (eg, gut bacterial) microbiota dysbiosis in the patient, whereby said dysbiosis modulates the therapy

20 in the patient by modulating immune cells in the patient; wherein the therapy comprises adoptive immune cell therapy (eg, adoptive T-cell therapy, eg, CAR-T cell administration to the patient).

In another aspect, the first configuration the invention provides

A method of modulating a therapy of a disease or condition in a human or animal patient, the method comprising

a. Carrying out the therapy in the patient; and

b. Causing bacterial (eg, gut bacterial) microbiota dysbiosis in the patient, whereby said dysbiosis modulates the therapy in the patient;

wherein the therapy comprises administering an immune checkpoint inhibitor (eg, an anti-PD-L1, anti-PD-1, anti-CTLA4 or anti-TIM3 inhibitor, eg, an antibody) to the patient.

In another aspect, the first configuration the invention provides

A method of modulating a therapy of a disease or condition in a human or animal patient, the method comprising

a. Carrying out the therapy in the patient; and

b. Causing bacterial (eg, gut bacterial) microbiota dysbiosis in the patient, whereby said dysbiosis modulates the therapy in the patient;

wherein the therapy comprises administering an antibody (eg, an anti-PD-L1, anti-PD-1, anti-CTLA4 or anti-TIM3 antibody; or an anti-TNFa superfamily member antibody, eg, an anti-TNFa, TNFR1 or BAFF antibody; or, an anti-IL6R or anti-IL-4Ra antibody; or an anti-PCSK9 antibody) to the patient.

In another aspect, the first configuration the invention provides

A method of modulating a treatment in a subject, the method comprising

a. Carrying out the treatment in the subject; and

b. Causing microbiota dysbiosis in the subject, whereby said dysbiosis modulates the treatment in the subject.

In an example, the subject or patient is a human. In an example, the subject or patient is a non-human animal. In an example, the subject is a plant, and optionally the treatment is a plant growth-promoting treatment, growth-inhibiting treatment, pesticide treatment, nitrogen fixing promotion treatment, herbicidal treatment or fertilizer treatment. In an example, the subject is a yeast, and optionally the treatment is a yeast growth-promoting treatment or growth-inhibiting treatment.

In an example, the modulating augments, upregulates, downregulates, inhibits, enhances or potentiates the treatment or therapy of the subject or patient. In an example, the treatment or therapy is effective in the subject or patient, wherein the treatment or therapy is not effective or has reduced or increased efficacy in the subject, patient or a control subject or patient that has not been subject to the 5 modulation. The control is of the same species as the subject or patient, and optionally the same age and/or sex. In an example, bacterial or archaeal host cells are killed or growth thereof is inhibited in the subject or patient using a method of an invention, wherein the control comprises cells of the 10 condition in a patient, the method comprising same bacterial or archaeal species and the cells are not killed or growth inhibited by a method of the invention.

In an example, steps (a) and (b) are carried out simultaneously. In an example, step (a) is carried out before step (b). In an example, step (b) is carried out before step (a), and 15 optionally step (b) is performed again after (a).

In an embodiment, the invention provides

A method of modulating a treatment in a plant or yeast, the method comprising

a. Carrying out the treatment in the plant or yeast; and b. Causing bacterial microbiota dysbiosis in the plant or yeast, whereby said dysbiosis modulates the treatment in the subject:

wherein the treatment is a growth-promoting treatment, growth-inhibiting treatment, pesticide treatment, nitrogen 25 fixing promotion treatment, herbicidal treatment or fertilizer treatment.

Causing microbial dysbiosis in the subject, patient, plant or yeast is, in an example comprises causing microbial dysbiosis on a surface of the subject, patient, plant or yeast, 30 eg, on a leaf surface (when the subject is a plant) or on skin, lung, ocular or mucosal surface (when the subject or patient is a human or animal).

Instead of or additionally to causing bacterial dysbiosis, the invention comprises in step (b) causing archaeal micro- 35 biota dysbiosis in said subject, patient, plant or yeast.

For example, the disease or condition is an autoimmune disease or condition (eg, SLE) and the therapy is a treatment therefor, eg, administration of a tumor necrosis factor ligand superfamily member antagonist, eg, an anti-B-cell activating 40 factor (BAFF) antibody, such as BENLYSTA® (belimumab) or a generic version thereof. For example, the disease or condition is an inflammatory disease or condition (eg, rheumatoid arthritis, IBD, Crohn's disease, colitis or psoriasis) and the therapy is a treatment therefor, eg, adminis- 45 tration of sarilumab, dupilumab, a tumor necrosis factor ligand superfamily member antagonist, eg. an anti-TNF alpha antibody or trap, such as HUMIRA® (adalimumab), REMICADE® (infliximab), SIMPONI® (golimumab) or ENBREL® (etanercept) or a generic version thereof. For 50 example, the disease or condition is a viral infection or mediated by a viral infection (eg, HIV infection) and the therapy is a treatment therefor, eg, administration of an anti-retroviral medicament or an anti-HIV vaccine. For example, the disease or condition is a cancer (eg, melanoma, 55 provides NSCLC, breast cancer or pancreatic cancer) and the therapy is a treatment therefor, eg, administration of a chemotherapeutic agent, eg, a checkpoint inhibitor or agonist antibody such as an anti-CTLA4, PD-1, PD-L1, PD-L2, LAG3, OX40, CD28, BTLA, CD137, CD27, HVEM, KIR, TIM-3, 60 VISTA, ICOS, GITR, TIGIT or SIRPa antibody. In an example, the antibody is a bispecific antibody that specifically binds first and second targets selected from CTLA4, PD-1, PD-L1, PD-L2, LAG3, OX40, CD28, BTLA, CD137, CD27, HVEM, KIR, TIM-3, VISTA, ICOS, GITR, TIGIT 65 and SIRPa, eg, wherein the first target is CTLA4 and the second target is LAG3 or PD-1. Optionally, the antibody is

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a human gamma-1 antibody and/or may be enhanced for ADCC or CDC. For example, the therapy is a vaccine therapy, eg, a cancer vaccine therapy or a vaccine therapy for treating or preventing an infection or infectious disease, such as malaria, HIV infection, tuberculosis infection, cholera, Salmonella typhimurium infection, C dificile infection, Bordetella pertussis infection or Chlamydia infection.

An embodiment of the first configuration provides

A method of modulating a cell therapy of a disease or

- a. Carrying out cell therapy in the patient, comprising administering a population of cells to the patient, wherein administration of said cells is capable of treating the disease or condition in the patient; and
- b. Causing gut bacterial microbiota dysbiosis in the patient, whereby said dysbiosis modulates the cell therapy in the patient.

In an example the cell therapy is an adoptive immune cell therapy, such as CAR-T or TILs therapy for the treatment of 20 a cancer.

In a second configuration the invention provides

A method of treating or reducing the risk of a disease or condition in a patient, wherein the disease or condition is mediated by immune cells (eg, T-cells) in the patient, the method comprising causing gut bacterial microbiota dysbiosis in the patient, whereby said dysbiosis modulates immune cells (eg, $T_H 17$ cells) in the patient, thereby treating or reducing the risk of said disease or condition in the patient.

For example, the disease or condition is an autoimmune disease or condition (eg, SLE), an inflammatory disease or condition (eg, rheumatoid arthritis, IBD, Crohn's disease, colitis or psoriasis), a viral infection or mediated by a viral infection (eg, HIV infection).

In an example, microbiota dysbiosis is effected by killing one or more target bacterial species in the microbiota or inhibiting growth of a population of said bacteria in the microbiota. In an example, microbiota dysbiosis is effected by killing one or more target archaeal species in the microbiota or inhibiting growth of a population of said archaea in the microbiota.

In a third configuration the invention provides

A method of modulating an adoptive immune cell therapy of a disease or condition in a patient, the method comprising

- a. Carrying out adoptive immune cell therapy in the patient, comprising administering a population of immune cells to the patient, wherein administration of said immune cells is capable of treating the disease or condition in the patient; and
- b. Altering the relative proportion of a sub-population of cells of a first bacterial species or strain, or archaeal species or strain, in a microbiota (eg, gut microbiota) of the patient, thereby producing an altered microbiota that modulates the immune cell therapy in the patient. In another aspect, the third configuration the invention

A method of modulating a therapy of a disease or condition in a human or animal patient, the method comprising

a. Carrying out the therapy in the patient; and

b. Altering the relative proportion of a sub-population of cells of a first bacterial species or strain, or archaeal species or strain, in a microbiota (eg, gut microbiota) of the patient, thereby producing an altered microbiota that modulates the therapy in the patient;

wherein the therapy comprises administering an immune checkpoint inhibitor (eg, an anti-PD-L1, anti-PD-1, anti-CTLA4 or anti-TIM3 inhibitor, eg, an antibody) to the patient.

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In another aspect, the third configuration the invention provides

A method of modulating a therapy of a disease or condition in a human or animal patient, the method comprising

a. Carrying out the therapy in the patient; and b. Altering the relative proportion of a sub-population of cells of a first bacterial species or strain, or archaeal species or strain, in a microbiota (eg, gut microbiota) of the patient, thereby producing an altered microbiota that modulates the therapy in the patient;

wherein the therapy comprises administering an antibody (eg, an anti-PD-L1, anti-PD-1, anti-CTLA4 or anti-TIM3 antibody; or an anti-TNFa superfamily member antibody, eg, an anti-TNFa, TNFR1 or BAFF antibody; or, an anti-IL6R or anti-IL-4Ra antibody; or an anti-PCSK9 antibody) 15 to the patient.

In another aspect, the third configuration the invention provides

A method of modulating a treatment in a subject, the method comprising

a. Carrying out the treatment in the subject; and

b. Altering the relative proportion of a sub-population of cells of a first bacterial species or strain, or archaeal species or strain, in a microbiota of the subject, whereby said dysbiosis modulates the treatment in the subject.

In an example, the subject or patient is a human. In an example, the subject or patient is a non-human animal. In an example, the subject is a plant, and optionally the treatment is a plant growth-promoting treatment, growth-inhibiting treatment, pesticide treatment, nitrogen fixing promotion 30 treatment, herbicidal treatment or fertilizer treatment. In an example, the subject is a yeast, and optionally the treatment is a yeast growth-promoting treatment or growth-inhibiting treatment.

In an example, the modulating augments, upregulates, 35 downregulates, inhibits, enhances or potentiates the treatment or therapy of the subject or patient. In an example, the treatment or therapy is effective in the subject or patient, wherein the treatment or therapy is not effective or has reduced or increased efficacy in the subject, patient or a 40 control subject or patient that has not been subject to the modulation. The control is of the same species as the subject or patient, and optionally the same age and/or sex. In an example, bacterial or archaeal host cells are killed or growth thereof is inhibited in the subject or patient using a method 45 of an invention, wherein the control comprises cells of the same bacterial or archaeal species and the cells are not killed or growth inhibited by a method of the invention.

In an example, steps (a) and (b) are carried out simultaneously. In an example, step (a) is carried out before step (b). 50 In an example, step (b) is carried out before step (a), and optionally step (b) is performed again after (a).

In an embodiment, the invention provides

A method of modulating a treatment in a plant or yeast, the method comprising

a. Carrying out the treatment in the plant or yeast; and b. Altering the relative proportion of a sub-population of cells of a first bacterial species or strain, or archaeal species or strain, in a microbiota of the plant or yeast, whereby said dysbiosis modulates the treatment in the plant or yeast; 60 wherein the treatment is a growth-promoting treatment, growth-inhibiting treatment, pesticide treatment, nitrogen fixing promotion treatment, herbicidal treatment or fertilizer treatment.

Said altering of the relative proportion of sub-population 65 of cells in the subject, patient, plant or yeast is, in an example comprises causing microbial dysbiosis on a surface

of the subject, patient, plant or yeast, eg, on a leaf surface (when the subject is a plant) or on skin, lung, ocular or mucosal surface (when the subject or patient is a human or animal).

The proportion of the first bacteria or archaea sub-population is increased or decreased. In an example, the relative ratio of first and second bacterial species or strains is altered (eg, increased or decreased); or the relative ratio of first and second archaeal species or strains is altered (eg, increased or 10 decreased).

In an example, the adoptive immune cell therapy is CAR-T therapy for the treatment of a cancer. In an example, the adoptive immune cell therapy is a TILs therapy for the treatment of a cancer.

In an example of the first or third configuration, the cells of step (a) are of a first type selected from the group consisting of CD4⁺ T-cells, CD8⁺ T-cells, T_H 1 cells or T_H 17 cells and step (b) upregulates cells of that type in the patient. This is useful for enhancing the cell based therapy. In another example the cells of step (a) are of a first type selected from the group consisting of CD4⁺ T-cells, CD8⁺ T-cells, $T_H 1$ cells or $T_H 17$ cells and step (b) downregulates cells of that type in the patient. This is useful for dampening down the cell based therapy or a side effect thereof (eg, CRS).

In an embodiment, the disbyosis or step (b) is carried out using selective targeting of a bacterial or archaeal microbiota sub-population using CRISPR/Cas targeting of microbiota (eg, gut microbiota) bacteria and/or archaea. In an example, the method comprises using guided nuclease (eg RNA-guided nuclease) cutting of a respective target sequence in host cells to modify the target sequences, whereby host cells are killed or the host cell population growth is reduced, thereby reducing the proportion of said sub-population in the microbiota. Suitable systems for carrying out the guided nuclease cutting are, for example, engineered CRISPR/Cas systems, TALENs, meganucleases and zinc finger systems.

To this end, the inventors believe that they have demonstrated for the first time inhibition of population growth of a specific bacterial strain in a mixed consortium of bacteria that naturally occur together in gut microbiota with one or more of the following features:

Population growth inhibition using an engineered CRISPR/Cas system by

targeting wild-type cells;

harnessing of wild-type endogenous Cas nuclease activity;

targeting essential and antibiotic resistance genes;

wherein the targets are wild-type sequences.

The inventors have demonstrated this in a mixed population of human gut microbiota bacteria with the following features:

- targeting bacterial growth inhibition in a mixed population of human gut microbiota species;
- wherein the population comprises three different species; comprising selective killing of one of those species and sparing cells of the other species;
- targeting cell growth inhibition in the presence of a phylogenetically-close other human gut microbiota species, which is spared such inhibition;
- targeting cell growth inhibition in a mixed population of human gut microbiota bacteria comprising target Fir*micutes* species and non-*Firmicutes* species;
- targeting cell growth inhibition of a specific Firmicutes species whilst sparing a different Firmicutes species in a mixed population of human gut microbiota bacteria;

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- targeting cell growth inhibition of a specific gram positive bacterial strain whilst sparing a different gram positive bacterial species in a mixed population of human gut microbiota bacteria;
- targeting a human gut microbiota bacterial species whilst 5 sparing a commensal human gut bacterial species;
- targeting a human gut microbiota bacterial species whilst sparing a priobiotic human gut bacterial species;
- targeting cell growth inhibition in a mixed population of human gut microbiota bacteria on a surface;
- achieving at least a 10-fold growth inhibition of a specific bacterial species alone or when mixed with a plurality of other bacterial species in a consortium of human gut microbiota bacteria; and
- achieving at least a 10-fold growth inhibition of two different strains of a specific human gut microbiota bacterial species.

The invention provides:

of adoptive cell therapy of a patient for treating or preventing a disease or condition in the patient, the method comprising

- a. Carrying out adoptive immune cell therapy in the tion to the patient, wherein administration of said immune cells is capable of treating the disease or condition in the patient; and
- b. Causing gut bacterial microbiota dysbiosis in the patient, whereby said dysbiosis modulates the immune 30 cell therapy in the patient and said disease or condition is treated or prevented.
- The invention provides

An ex vivo population of immune cells for use in a method of adoptive cell therapy of a patient for treating or prevent- 35 ing a disease or condition in the patient, the method comprising

- a. Carrying out adoptive immune cell therapy in the patient, comprising administering cells of said population to the patient, wherein administration of said 40 immune cells is capable of treating the disease or condition in the patient; and
- b. Altering the relative proportion of a sub-population of cells of a first bacterial species or strain, or archaeal species or strain, in the gut microbiota of the patient, 45 thereby producing an altered gut microbiota that modulates the immune cell therapy in the patient. The invention also provides CRISPR/Cas systems, arrays, cRNAs and kits for carrying out a method of the invention. 50

The invention also relates to systems, kits and other means for effecting the method.

Any features on one configuration herein are, in an example, combined with a different configuration of the invention for possible inclusion of such combination in one 55 or more claims herein.

BRIEF DESCRIPTION OF THE FIGURES

FIG. 1 shows s Xylose inducible system.

FIG. 2 shows a ST1-CRISPR array.

FIG. 3 shows a spot assay on TH-agar of the strains used in this work. All strains were grown on TH-agar at 37° C. for 20 hours. Serial dilutions of overnight cultures were done in duplicate for E. coli, L. lactis and S. mutans, and triplicate 65 for both strains of S. thermophilus in order to count individual colonies.

FIG. 4 shows selective growth of S. thermophilus, S. mutans, L. lactis and E. coli under different culture conditions. Tetracycline cannot be used to selectively grown S. thermophilus LMD-9. However, 3 g 1^{-1} of PEA proved to selectively grow S. thermophilus LMD-9 while limiting growth of E. coli.

FIG. 5 illustrates construction of two xylose induction cassettes.

FIG. 6 demonstrated characterization of the xylose inducible cassette in Streptoccocus thermophilus LMD-9 with the pBAV1KT5-XylR-mCherry-Pldha. plasmid Α clear response in fluorescence can be observed with increasing amount of xylose.

FIG. 7 illustrates the design of CRISPR array in 15 pBAV1KT5-XylR-mCherry-P_{ldha+XylA}. The array contains 2 spacer sequences that target S. thermophilus genes under an inducible xylose promoter and a tracrRNA under a strong constitutive promoter P_{3A} .

FIG. 8 shows transformation efficiency of Streptoccocus An ex vivo population of immune cells for use in a method 20 thermophilus LMD-9 with the plasmid pBAV1KT5-XyIR-CRISPR-P_{1dh+Xv1A} and with pBAV1KT5-XylR-CRISPR-P_{XylA}.

FIG. 9 shows a schematic of the xylose-inducible CRISPR device. Upon induction of xylose the CRISPR patient, comprising administering cells of said popula- 25 array targeting both polIII and tetA on the S. thermophiles LMD-9 genome are expressed. Together with the constitutively expressed tracrRNA a complex is formed with Cas9. This complex will introduce a double stranded break in the tetA and polIII genes in the S. thermophilus LMD-9 genome resulting in limited cell viability.

> FIG. 10 shows growth inhibition of Streptoccocus thermophilus DSM 20617(T) with the plasmid pBAV1KT5-XylR-CRISPR-PXylA or pBAV1KT5-XylR-CRISPR-Pldha+XylA, not induced and induced. Picture taken after 63H of incubation. Colony counts in bottom left corner (top row: >1000, >1000, bottom row: 336, 113).

> FIG. 11 shows a maximum-likelihood phylogenetic tree of 16S sequences from S. thermophilus, L. lactis and E. coli.

> FIG. 12 shows the selective S thermophilus growth inhibition in a co-culture of E. coli, L. lactis and S. thermophiles harboring either the pBAV1KT5-XylR-CRISPR-PxylA or the pBAV1KT5-XylR-CRISPR-PldhA+XylA plasmid. No growth difference is observed between E. coli harboring the pBAV1KT5-XylR-CRISPR-PxylA or the pBAV1KT5-XylR-CRISPR-PldhA+XylA plasmid. However, S. thermophiles (selectively grown on TH agar supplemented with 2.5 g1-1 PEA) shows a decrease in transformation efficiency between the pBAV1KT5-XylR-CRISPR-PxylA (strong) or the pBAV1KT5-XylR-CRISPR-PldhA+XylA (weak) plasmid as we expected. We thus demonstrated a selective growth inhibition of the target S. thermophilus sub-population in the mixed population of cells. Colony counts in bottom left corner (top row: >1000, >1000, 68, bottom row: >1000, >1000, 32).

> FIG. 13 shows regulators controlling the expression of spCas9 and the self-targeting sgRNA targeting the ribosomal RNA subunit 16s.

FIG. 14 shows specific targeting of E. coli strain by an exogenous CRISPR-Cas system. The sgRNA target the genome of K-12 derived E. coli strains, like E. coli TOP10, while the other strain tested was unaffected.

FIG. 15 shows spot assay with serial dilutions of individual bacterial species used in this study and mixed culture in TH agar without induction of CRISPR-Cas9 system.

FIG. 16 shows spot assay of the dilution 10^3 on different selective media. TH with 2.5 g l⁻¹ PEA is a selective media for B. subtilis alone. MacConkey supplemented with malt-

ose is a selective and differential culture medium for bacteria designed to selectively isolate Gram-negative and enteric bacilli and differentiate them based on maltose fermentation. Therefore TOP10 Δ malK mutant makes white colonies on the plates while Nissle makes pink colonies; A is *E coli* 5 Δ malK, B is *E coli* Nissile, C is *B subtilis*, D is *L lactis*, E is mixed culture; the images at MacConkey–/B and E appear pink; the images at MacConkey+/B and E appear pink.

FIG. **17** shows selective growth of the bacteria used in this study on different media and selective plates.

DETAILED DESCRIPTION

In the worked Example below, growth inhibition was addressed in a mixed population of human gut microbiota 15 bacterial species. A >10-fold population growth inhibition in a selectively targeted species (a gram positive Firmicutes population) was achieved, sparing non-targeted commensal bacteria in the consortium. The inventors have realised the useful application of this for altering microbiota, such as gut 20 microbiota, in situ in patients, thereby enabling immune cell modulation in the patient in response to the altered microbiota. The inventors also realised application to modulating treatments in subjects such as plants and yeast that comprise microbiota that can be altered. The inventors furthermore 25 realised the utility for modulating immune cell-based therapies in patients or for treating or preventing immune cellmediated diseases or conditions in patients, such as autoimmune diseases, inflammatory diseases and viral infections (eg, HIV infection of humans). The inventors realised the 30 utility of causing dysbiosis of gut, skin, vaginal, nasal, ocular, lung, GI tract, rectal, scrotal, ear, skin or hair microbiota for effecting such modulation in a human or animal subject or patient.

As used herein "dysbiosis" refers to a change of the 35 bacterial and/or archaeal balance of the microbiota, eg, gut microbiota. Change is relative to the balance prior to (eg, immediately prior or no more than a day before) carrying out the method. The change can be one or more of (i) an increase in the proportion of a first species (bacterial or archaeal 40 species) or strain in the microbiota (eg, gut microbiota) (eg, B fragalis or thetaiotamicron); (ii) an increase in the relative proportion of first and second species (eg, B fragalis versus C dificile; or S thermophilus v E coli or L lactis), first and second strains of the same species, or first and second phyla 45 which are different from each other (eg, Bacteriodetes versus Firmicutes); (iii) an addition of a species or strain that was not comprised by the microbiota prior to the treatment method; (iv) a decrease in the proportion of a first species (bacterial or archaeal species) or strain in the microbiota (eg, 50 C dificile or S thermophilus); (v) a decrease in the relative proportion of first and second species (eg, B fragalis versus C dificile; or S thermophilus v E coli or L lactis), first and second strains of the same species, or first and second phyla which are different from each other (eg, Bacteriodetes versus 55 Firmicutes); and (vi) a removal of a species or strain that was not comprised by the microbiota prior to the treatment method. Dysbiosis may be effected, for example, using one or more selective antibacterial agents (eg, CRISPR-based or other guided nucleases described herein) of by administering 60 one or more bacterial and/or archaeal transplants to the patient or subject to alter the balance of the microbiota, eg, gut microbiota.

The impact of the immune system on microbiota composition is suggested by several immune deficiencies that alter 65 microbial communities in ways that predispose to disease. For example, Garrett et al. studied mice that lack the

transcription factor T-bet (encoded by Tbx21), which governs inflammatory responses in cells of both the innate and the adaptive immune system (Cell. 2007 Oct. 5; 131(1):33-45, "Communicable ulcerative colitis induced by T-bet deficiency in the innate immune system", Garrett WS et al.). When Tbx21-/-mice were crossed onto Rag2-/-mice, which lack adaptive immunity, the Tbx21-/-/Rag2-/-progeny developed ulcerative colitis in a microbiota-dependent manner Remarkably, this colitis phenotype was transmissible to wild-type mice by adoptive transfer of the Tbx21-/-/Rag2-/-microbiota. This demonstrated that altered microbiota were sufficient to induce disease. Another example of immune-driven dysbiosis is seen in mice deficient for epithelial cell expression of the inflammasome component NLRP6. These mice develop an altered microbiota with increased abundance of members of the Bacteroidetes phylum associated with increased intestinal inflammatory cell recruitment and susceptibility to chemicallyinduced colitis.

It has become evident that individual commensal species influence the makeup of *Lamina propria* T lymphocyte subsets that have distinct effector functions. Homeostasis in the gut mucosa is maintained by a system of checks and balances between potentially pro-inflammatory cells, which include T_{H1} cells that produce interferon- γ , T_{H1} 7 cells that produce IL-17a, IL-17f, and IL-22, diverse innate lymphoid cells with cytokine effector features resembling T_{H2} and T_{H1} 7 cells, and anti-inflammatory Foxp3⁺ regulatory T cells (T_{regs}).

A particular application of the invention is found in the shaping of $T_H 17$ cell populations in patients. Such cells have been implicated in autoimmune and inflammatory disorders. These cells were described in: Harrington L E, Hatton R D, Mangan P R, et al., "Interleukin 17-producing CD41 effector T cells develop via a lineage distinct from the T helper type 1 and 2 lineages", Nat Immunol. 2005; 6(11): 1123-1132; and Park H, Li Z, Yang X O, et al., "A distinct lineage of CD4 T cells regulates tissue inflammation by producing interleukin 17", Nat Immunol. 2005; 6(11):1133-1141. In the case of autoimmune disorders, $T_H 17$ cell over activation can cause an inappropriate amount of inflammation, like in the case of multiple sclerosis, rheumatoid arthritis, and psoriasis. $T_H 17$ cells have also been shown to be necessary for maintenance of mucosal immunity. $T_H 17$ cells may contribute to the development of late phase asthmatic response due to increases in gene expression relative to T_{reg} cells.

In HIV, the loss of $T_{H}17$ cell populations can contribute to chronic infection. The depletion of $T_H 17$ cell populations in the intestine disrupts the intestinal barrier, increases levels of movement of bacteria out of the gut through microbial translocation, and contributes to chronic HIV infection and progression to AIDS. Microbial translocation results in bacteria moving from out of the gut lumen, into the Lamina propia, to the lymph nodes, and beyond into non-lymphatic tissues. It can cause the constant immune activation seen through the body in the late stages of HIV. Increasing $T_{H}17$ cell populations in the intestine has been shown to be both an effective treatment as well as possibly preventative. Although all CD4⁺ T cells gut are severely depleted by HIV, the loss of intestinal $T_H 17$ cells in particular has been linked to symptoms of chronic, pathogenic HIV and SIV infection. Microbial translocation is a major factor that contributes to chronic inflammation and immune activation in the context of HIV. In non-pathogenic cases of SIV, microbial translocation is not observed. $T_H 17$ cells prevent severe HIV infection by maintaining the intestinal epithelial barrier

during HIV infection in the gut. Because of their high levels of CCR5 expression, the coreceptor for HIV, they are preferentially infected and depleted. Thus, it is through $T_H 17$ cell depletion that microbial translocation occurs. Additionally, the loss of $T_H 17$ cells in the intestine leads to a loss of 5 balance between inflammatory $T_H 17$ cells and T_{reg} cells, their anti-inflammatory counterparts. Because of their immunosuppressive properties, they are thought to decrease the anti-viral response to HIV, contributing to pathogenesis. There is more T_{reg} activity compared to $T_H 17$ activity, and the immune response to the virus is less aggressive and effective. Revitalizing $T_H 17$ cells has been shown to decrease symptoms of chronic infection, including decreased inflammation, and results in improved responses to highly active anti-retroviral treatment (HAART). This is 15 an important finding-microbial translocation generally results in unresponsiveness to HAART. Patients continue to exhibit symptoms and do not show as reduced a viral load as expected. In an SIV-rhesus monkey model, It was found that administering IL-21, a cytokine shown to encourage 20 T_{H} 17 differentiation and proliferation, decreases microbial translocation by increasing $T_H 17$ cell populations.

In an example of the method, IL-21, IL-15 and/or IL-2 is administered to the patient sequentially or simultaneously with the cell population. This is useful for further modulat- 25 ing immune cell populations in the patient.

Yang et al. observed that the presence of $T_H 17$ cells in mice requires colonisation of mice with microbiota. Segmented filamentous bacteria (SFB) were sufficient to induce $T_H 17$ cells and promote $T_H 17$ -dependent autoimmune dis- 30 ease in animal models (Nature, 2014 Jun. 5; 510(7503):152-6. doi: 10.1038/nature13279. Epub 2014 Apr. 13, "Focused specificity of intestinal Th17 cells towards commensal bacterial antigens", Yang Y et al.). SFB appear able to penetrate the mucus layer overlying the intestinal epithelial cells in the 35 terminal ileum, and they interact closely with the epithelial cells, inducing host cell actin polymerization at the site of interaction and, presumably, signaling events that result in a $T_H 17$ polarizing environment within the Lamina propria.

In an example, the first bacteria are of a species or strain 40 comprising a 16s rDNA sequence that is at least 80, 85, 90, 95, 96, 97, 98 or 99% identical to a 16s rDNA sequence of a segmented filamentous bacterium. In an embodiment, the method increases the proportion of the first bacteria, wherein $T_H 17$ cells in the patient are upregulated, eg, wherein the 45 disease is a cancer or a viral infection (eg, HIV). In an embodiment, the method decreases the proportion of the first bacteria, wherein $T_H 17$ cells in the patient are downregulated, eg, wherein the disease or condition is an autoimmune or inflammatory disease or condition, or for reducing the risk 50 of CRS in a cancer patient receiving ACT.

In an example, the method treats or prevents an allergic disease or condition, eg, asthma. In an example, the method treats or prevents an IgE-mediated disease or condition, eg, asthma

In an example, the method reduces autotoxicity in the patient mediated by $T_{\mu}2$ cell cytokine release.

López et al. observed that intestinal dysbiosis, characterised by a reduced *Firmicutes*/Bacteroidetes ratio, has been reported in systemic lupus erythematosus (SLE) patients. In 60 their study, in vitro cultures revealed that microbiota isolated from SLE patient stool samples (SLE-M) promoted lymphocyte activation and $T_H 17$ differentiation from naïve CD4⁺ lymphocytes to a greater extent than healthy control microbiota. Enrichment of SLE-M with Tree-inducing bac-65 teria showed that a mixture of two Clostridia strains significantly reduced the $T_H 17/T_H 1$ balance, whereas Bifidobac-

supplementation prevented CD4+ terium bifidum lymphocyte over-activation. Ex vivo analyses of patient samples showed enlarged T_H17 and Foxp3* IL-17⁺ populations, suggesting a possible T_{reg} - T_H 17 trans-differentiation. Moreover, analyses of faecal microbiota revealed a negative correlation between IL-17⁺ populations and Firmicutes in healthy controls, whereas in SLE this phylum correlated directly with serum levels of IFN γ , a T_H1 cytokine slightly reduced in patients. (Sci Rep. 2016 Apr. 5; 6:24072. doi: 10.1038/srep24072, "Th17 responses and natural IgM antibodies are related to gut microbiota composition in systemic lupus erythematosus patients", López P et al.).

Other bacteria have been shown to enhance the antiinflammatory branches of the adaptive immune system by directing the differentiation of T_{regs} or by inducing IL-10 expression. For example, colonisation of gnotobiotic mice with a complex cocktail of 46 mouse Clostridial strains, originally isolated from mouse faeces and belonging mainly to cluster IV and XIVa of the Clostridium genus, results in the expansion of Lamina propria and systemic Trees.

Bacteroides fragilis polysaccharide-A (PSA) impacts the development of systemic T cell responses. Colonization of germ-free mice with PSA-producing B. fragilis results in higher numbers of circulating CD4⁺ T cells as compared to mice colonized with B. fragilis lacking PSA. PSA-producing *B. fragilis* also elicits higher T_H cell frequencies in the circulation. Together, these findings show that commensal bacteria have a general impact on immunity that reaches well beyond mucosal tissues.

The decrease in F. prausnitzii found in IBD patients is of interest because this bacteria is butyrate-producing, and its oral administration reduces the severity of TNBS-induced colitis in mice. In an example, the first species is a butyrateproducing bacterial species (eg, F. prausnitzii) and the proportion of the first species in the microbiota is reduced, wherein the method downregulates T-effector and/or T-helper cells in the patient, thereby treating or preventing said disease or condition (eg, an autoimmune or inflammatory disease or condition or CRS).

Archaea have traditionally been divided into five phyla, namely Crenarchaeota, Euryarchaeota, Korarchaeota, Nanoarchaeota and Thaumarchaeota. Based on the increasing wealth of whole genome data (mainly from environmental isolates), the archaeal phylogeny has been revisited recently: the four groups Korarchaeota, Crenarchaeota, Thaumarchaeota and the newly proposed Aigarchaeota have been comprised into one superphylum (the so-called TACKsuperphylum) to the exclusion of Euryarchaeota and Nanoarchaeota. The first species in the method of the invention can be any of the archaea mentioned in this paragraph.

T cells mature in the thymus, express TCR (T cell receptor), and can express either CD8 glycoprotein on their surface and are called CD⁸⁺ T cells (cytotoxic) or CD4 glycoprotein and are then called CD4 cells (helper T cells). CD^{4+} cells differentiate into different subsets: T_H (T helper) 1, $T_H 2$, $T_H 9$, $T_H 17$, $T_H 22$, T_{reg} (regulatory T cells) and T_{fh} (follicular helper T cells), which are characterized by different cytokine profiles. These different CD4⁺ subsets play a critical role in the immune and effector response functions of T cells. All CD4⁺ T_H subsets are differentiated from naive CD4⁺ T cells by specific cytokines: T_H 1 by IL-12 and IFN- γ (pro-inflammatory cytokine, with multiple roles such as increase of TLR (Toll-like receptor), induction of cytokine secretion or macrophage activation); $T_H 2$ by IL-4; T_{reg} by IL-2 and TGF-beta. Each T_H subset releases specific cytokines that can have either pro- or anti-inflammatory func-

tions, survival or protective functions. For example, T_{H1} releases IFN- γ and TNF; T_{H2} releases IL-4 (an important survival factor for B-type lymphocytes), IL-5 and IL-13; T_{H9} produces IL-9; T_{reg} secretes IL-10 (a cytokine with an immunosuppressive function, maintaining expression of 5 FOXP3 transcription factor needed for suppressive function of T_{reg} on other cells) and TGF- β ; T_{H17} produces IL-17 (a cytokine playing an important role in host defense against bacteria, and fungi).

An embodiment of the invention finds application for 10 modulating CAR-T and other adoptive immune-cell therapies (such as adoptive TILs therapy). Several reports have demonstrated differential roles of different types of cytokines released by CD4+ subsets, an important consideration when assessing CAR-T and other immune cell-based therapies. T_H 1 and T_H 2 CD4⁺ T cell subset cytokines were shown to drive different types of cytotoxicity generated by second generation CD28-containing CAR-T. Short-term toxicity was observed with high levels of T_H cytokines, while high doses of $T_H 2$ type cytokines generated chronic 20 autocytotoxicity in animals that received second generation CD19-specific CAR-T. CAR-T cells engineered to deliver inducible IL-12 modulated tumor stroma to destroy cancer. IL-12 release by engineered CAR-T cells increased anticancer activity by recruiting macrophages. IL-12 released by 25 CAR-T also induced reprogramming of suppressive cells, reversing their inhibitory functions suggesting its evaluation in clinical trials. The persistence of CAR-T therapy was shown to be dependent on the number of CD4⁺ cells and the number of central memory cells in the infused product. 30 CD8+ clones isolated from central memory T cells but not from CD8⁺ effector cells persisted long-term in vivo during adoptive T cell transfer in a nonhuman primate model, indicating the importance of specific T cell subset functions for effective adoptive immunotherapy. It has also been 35 shown that the combination of CD8⁺ subset with CD4⁺ subset significantly enhanced T cell adoptive transfer. CD4+ cells were shown to support development of CD8+ memory functions, demonstrating the importance of both subsets and combinations in immunotherapy trials. Several preclinical 40 models demonstrated the advantage of different T cell subsets for effective CAR-T therapy: CD8+ CD45RA+ CCR7+ CAR-T cells with closest to the T-memory stem cells phenotype cells produced greater anti-tumor activity of CAR-T cells; both CD8⁺ and CD4⁺ subsets expressed synergistic 45 anti-tumor CAR-T activities.

In an example, the administered cell population is a population of CAR-T cells comprising a combination of a CD8⁺ CAR-T subset with CD4⁺ CAR-T subset.

In an example of the invention, the cell therapy is an 50 adoptive T-cell therapy and optionally cells selected from the group consisting of CD4+ T-cells, CD8+ T-cells, TH1 cells and TH17 cells are administered to the patient. In an example, cell therapy is enhanced by the method of the invention, eg, immune cell cytotoxicity of cancer cells is 55 enhanced in the patient, or treatment of the disease or condition is enhanced. In an example, cell therapy is reduced by the method of the invention, eg, immune cell cytotoxicity of cancer cells is reduced in the patient, or the risk of CRS is reduced (eg, in a cancer patient). Thus, in an embodiment 60 the method reduces or prevents the risk of cytokine release syndrome (CRS) in the patient. In an embodiment the method reduces or prevents the risk of an unwanted sideeffect of the cell therapy (eg, a CAR-T therapy side effect in a human patient, such as CRS). 65

In an example, the immune cell population comprises CAR-T cells and/or T-cells expressing engineered T-cell

receptors (TCRs) and/or tumour infiltrating lymphocytes (TILs, eg, Engineered TILs). WO2013063361, U.S. Pat. No. 9.113.616. US20130109053. US20160081314 and WO2016044745 (whose disclosures are incorporated herein by reference) describe suitable transgenic in vivo platforms for generating CARs and TCRs for use in generating cells for use in the present invention. The immune cell population may comprise engineered autologous or allogeneic immune cells (transplant), eg, T-cells, NK cells and/or TILs, eg, wherein the cells and patient are human A benefit of autologous cells is that the modulation of the endogenous system is likely to be tuned similarly to modulation of the cell transplanted autologous cells. In an embodiment, the administered cells and patient are of the same species or strain, for example, human or rodent (eg, mouse), for example, HLA or MHC matched donor transplant and recipient patient.

In an example, the T-cells are CD4⁺ T-cells or $T_H 17$ cells. For example, the administered CAR-T cells comprise a chimaeric antigen receptor comprising an ICOS intracellular domain and optionally the cells are $T_H 17$ cells. In an embodiment, the administered T-cells are CD8⁺ ^{CD}45RA⁺ CCR7⁺ CAR-T cells.

Adoptive transfer experiments in mice indicate that $T_H 17$ cells have higher in vivo survival and self-renewal capacity than $T_H 1$ polarized cells. In an example, therefore, $T_H 17$ cells are modulated in the patient, eg, upregulated, eg, expanded in the patient, or downregulated. These may be endogenous T-cells of the patient and/or cells that have been administered to the patient or progeny thereof. In an embodiment, ROR γ t-expressing $T_H 17$ cells are upregulated, eg, expanded in the patient. In an embodiment expression of one or more $T_H 17$ -related genes is increased, eg, one or more of Rorc, Il22 and Il26. In an embodiment expression of one or more $T_H 1$ -related genes is increased, eg, one or more of IFng, Tnfa and Tbx21 (T-bet). In an embodiment, in this case the disease or condition is a cancer.

In an example, T_{reg} cells are modulated in the patient, eg, upregulated, eg, expanded in the patient, or downregulated. These may be endogenous T-cells of the patient and/or cells that have been administered to the patient or progeny thereof. In an embodiment, in this case the disease or condition is an autoimmune, inflammatory or infectious disease or condition when the T_{reg} cells are upregulated.

In an example, CD4⁺ cells are modulated in the patient, eg, upregulated, eg, expanded in the patient, or downregulated. These may be endogenous cells of the patient and/or cells that have been administered to the patient or progeny thereof.

In an example, CD8⁺ cells are modulated in the patient, eg, upregulated, eg, expanded in the patient, or downregulated. These may be endogenous cells of the patient and/or cells that have been administered to the patient or progeny thereof.

In an example, tumour infiltrating lymphocytes (TILs) are modulated in the patient, eg, upregulated, eg, expanded in the patient, or downregulated. These may be endogenous cells of the patient and/or cells that have been administered to the patient or progeny thereof.

In an example, memory cells, such as one or more of central memory T cells (T_{CM}), effector memory T cells (T_{EM}), stem cell memory cells (T_{SCM}) and effector cells (T_{eff}), are upregulated in the microbiota or patient, optionally wherein the cells are comprised by the immune cell population administered to the patient and/or are progeny thereof. In an embodiment, the memory cells are CD45RO⁺CD62L⁺ or CD25⁺ CD45RA⁻ CD45RO⁺ CD127⁺.

Upregulation of a cell population may, for example, be an increase in the population size or proportion of cells of that type (eg, species or strain) in the microbiota or patient or subject and/or an increase in the activity (eg, cytotoxicity, effector function or suppressor function) of cells of that type in the microbiota or patient or subject. Downregulation of a cell population may, for example, be an decrease in the population size or proportion of cells of that type (eg, species or strain) in the microbiota or patient or subject and/or a decrease in the activity (eg, cytotoxicity, effector function) of cells of that type (eg, species or strain) in the microbiota or patient or subject and/or a decrease in the activity (eg, cytotoxicity, effector function or suppressor function) of cells of that type in the microbiota or patient or subject.

In an example, the cell therapy population comprises CAR-T cells (ie, respectively T-cells engineered to surfaceexpress chimaeric antigen receptors (CARs). Alternatively, the cells are CAR-TIL or CAR-NK cells. A CAR comprises an extracellular receptor domain for binding to a target antigen (eg, a tumour cell antigen), a transmembrane moiety and an intracellular moiety comprising one or more (eg, first and second) signalling domains for signalling in the immune cell (eg, T-cell). Examples of suitable intracellular domains are well known, eg, a combination of a CD3ζ domain and one or more of an ICOS, CD28, OX40 or 4-1BB signalling domain, eg, a combination of an ICOS and CD28; or ICOS 25 and 41-BB; CD28 and 41-BB signalling domain.

Optionally, the cell population is comprised by a transplant that is administered to the patient to treat or prevent a disease (eg, a cancer, autoimmune disease, transplant rejection or GvHD) or the cell or transplant is for such use.

In an example, the patient is a human, eg, is a woman; or a man.

In an example, the patient or human has undergone lymphodepletion before administration of the immune cell (eg, CAR-T cell).

Techniques for producing CARs and CAR T-cells are known and routine in the art, and these can be generally applied to producing cells for use in the invention (eg, see WO2012079000A1; U.S. Pat. Nos. 8,906,682, 8,911,993, 8,916,381, 8,975,071, 9,101,584, 9,102,760, 9,102,761, 40 9,328,156, 9,464,140, 9,481,728, 9,499,629, 9,518,123, 9,540,445, US20130287748, US20130288368, US20130309258, US20140106449, US20140370017, US20150050729, US20150093822, US20150099299, US20150118202, US20160130355, US20160159907, 45 US20160194404, US20160208012; J Immunother. 2009 September: 32(7): 689-702. doi: 10.1097/ CJI.0b013e3181ac6138, "Construction and Pre-clinical Evaluation of an Anti-CD19 Chimeric Antigen Receptor", James N. Kochenderfer et al; also WO2014012001 and 50 US20150290244 for general methods applicable to the present invention). For example, use of electroporation, retroviral vectors or lentiviral vectors-as will be known by the skilled addressee-can be used to introduce nucleotide sequences encoding elements of the CAR into T-cells, NK 55 cells, TILs or other immune cells to produce the CAR-cells. Cells isolated from the patient (autologous cell sample) or from another donor of the same species (allogeneic sample) can be used to provide ancestor cells that are genetically engineered to include the CAR-encoding sequences. Expan- 60 sion of cells can be used in the process, as known in the art. For example, after engineering CAR-cells, the cell population can be massively expanded using routine techniques to produce a transplant that is administered (eg, transfused) into the patient. The patient can be a human on non-human 65 animal Nucleotide sequences for one or more of the CAR elements (eg, for one or more of three signalling domains)

can be cloned or sequenced using a cell obtained from the patient or from another donor.

For example, the CAR comprises a first intracellular signalling domain, which is a human CD3 ζ domain and the cells administered to the patient are human cells comprising an endogenous nucleotide sequence encoding said human CD3 ζ domain. In an example, the CD3 zeta signaling domain comprises SEQ ID NO: 1, i.e., the amino acid sequence of SEQ ID NO: 24 as disclosed in WO2012079000A1, which sequence is explicitly incorporated herein for use in the present invention and possible inclusion in one or more claims herein. In an example, the CD3 zeta signaling domain is encoded by SEQ ID NO: 2, i.e., the nucleic acid sequence of SEQ ID NO: 18 as disclosed in WO2012079000A1, which sequence is explicitly incorporated herein for use in the present invention and possible inclusion in one or more claims herein.

For example, the first signalling domain is a human CD28 domain and the cell population of the invention is a population of human cells each comprising an endogenous nucleotide sequence encoding said human CD28 domain.

For example, the first signalling domain is a human 4-1BB domain and the cell population of the invention is a population of human cells each comprising an endogenous nucleotide sequence encoding said human 4-1BB domain.

For example, the first signalling domain is a human OX40 domain and the cell population of the invention is a population of human cells each comprising an endogenous nucleotide sequence encoding said human OX40 domain.

In an example, the first signalling domain is a CD3' domain, and the first and second intracellular signalling domains do not naturally occur together in a single cell (eg, a human wild-type cell or a cell isolated from the patient), eg, the second domain is a CD28, CD27, OX40 or 4-1BB 35 domain.

In an example, the first intracellular domain is a CD3' domain, CD28 domain or 4-1BB domain.

In an example, the CAR is an engineered single polypeptide comprising (in N- to C-terminal direction) an antigen binding site (eg, an antibody scFv, which may be human); an optional hinge (eg, a human CD8 α hinge); a transmembrane domain (eg, a human CD8a or CD28 transmembrane domain); and a human CD3' domain. In an example, the CAR is a complex of two or more of said polypeptides. Optionally, the CAR comprises a further intracellular signalling domain (i) between the transmembrane and CD32 domains. Optionally, the CAR comprises a further intracellular signalling domain, wherein the CD35 domain is between the further signaling domain and the transmembrane domain. In an example, the further signalling domain is a human CD27 domain, CD28 domain, ICOS domain, OX40 domain, CD40 domain, 4-1BB domain, a FceRIy domain, CD64 domain or CD16 domain. In an alternative, instead of a single polypeptide, the CAR comprises an engineered complex of at least 2 polypeptides comprising said domains.

The immune cells may be administered either alone, or as a pharmaceutical composition in combination with diluents and/or with other components such as IL-2 or other cytokines or cell populations.

In an embodiment, the immune cells (eg, CAR cells or cells bearing TCRs) comprise cell surface binding sites (eg, provided by the CAR or TCR) that bind a TAA. Tumour antigens (TAA) are proteins that are produced by tumour cells that elicit an immune response, particularly T-cell mediated immune responses. The selection of the antigen binding specificity will depend on the particular type of cancer to be treated. Tumour antigens are well known in the art and include in the context of an embodiment of the invention, for example, a glioma-associated antigen, carcinoembryonic antigen (CEA), β -human chorionic gonadotropin, alphafetoprotein (AFP), lectin-reactive AFP, thyroglobulm, RAGE-1, MN-CA IX, human telomerase reverse transcriptase, RU1, RU2 (AS), intestinal carboxyi esterase, mut hsp70-2, M-CSF, prostase, prostate-specific antigen (PSA), PAP, NY-ESO-1, LAGE-la, p53, prostein, PSMA, Her2/neu, survivin and telomerase, prostate-carcinoma 10 tumour antigen-1 (PCTA-1), MAGE, ELF2M, neutrophil elastase, ephrinB2, CD22, insulin growth factor (IGF)-I, IGF-II, IGF-II receptor and mesothelin.

In one embodiment, the tumour antigen comprises one or more antigenic cancer epitopes associated with a malignant 15 tumour. Malignant tumours express a number of proteins that can serve as target antigens for an immune attack. These molecules include but are not limited to tissue-specific antigens such as MART-1, tyrosinase and GP 100 in melanoma and prostatic acid phosphatase (PAP) and prostate- 20 specific antigen (PSA) in prostate cancer. Other target molecules belong to the group of transformation-related molecules such as the oncogene HER-2/Neu ErbB-2. Yet another group of target antigens are onco-foetal antigens such as carcinoembryonic antigen (CEA). In B-cell lym- 25 phoma the tumour-specific idiotype immunoglobulin constitutes a truly tumour-specific immunoglobulin antigen that is unique to the individual tumour. B-cell differentiation antigens such as CD19, CD20 and CD37 are other candidates for target antigens in B-cell lymphoma. Some of these 30 antigens (CEA, HER-2, CD19, CD20, idiotype) have been used as targets for passive immunotherapy with monoclonal antibodies with limited success. The first antigen or fourth binding moiety can be any of these TAAs or can be an antigenic sequence of any of these TAAs.

Non-limiting examples of TAA antigens in an embodiment of the invention include the following: Differentiation antigens such as MART-1/MelanA (MART-1), g 1 OO (Pmel 17), tyrosinase, TRP-1, TRP-2 and tumour-specific multilineage antigens such as MAGE-1, MAGE-3, BAGE, 40 GAGE-1, GAGE-2, pi 5; overexpressed embryonic antigens such as CEA; overexpressed oncogenes and mutated tumour-suppressor genes such as p53, Ras, HER-2/neu; unique tumour antigens resulting from chromosomal translocations; such as BCR-ABL, E2A-PRL, H4-RET, 1GH- 45 IGK, MYL-RAR; and viral antigens, such as the Epstein Barr virus antigens EBVA and the human papillomavirus (HPV) antigens E6 and E7. Other large, protein-based antigens include TSP-180, MAGE-4, MAGE-5, MAGE-6, RAGE, NY-ESO, p1 85erbB2, p 1 80erbB-3, c-met, 50 nm-23H1, PSA, TAG-72, CA 19-9, CA 72-4, CAM 17.1, NuMa, K-ras, beta-Catenin, CDK4, Mum-1, p 15, p 16, 43-9F, 5T4(791Tgp72} alpha-fetoprotem, beta-HCG, BCA225, BTAA, CA 125, CA 15-3\CA 27.29\BCAA, CA 195, CA 242, CA-50, CAM43, CD68\ I, CO-029, FGF-5, 55 G250, Ga733VEpCAM, HTgp-175, M344, MA-50, MG7-Ag, MOV 18, NB/70K, NY-CO-1, RCAS 1, SDCCAG16, TA-90\Mac-2 binding proteiiAcyclophilin C-associated protein, TAAL6, TAG72, TLP, and TPS.

In one embodiment, the CAR or TCR comprises a binding 60 site for human CD 19, eg, for a CAR this can be provided by an anti-CD 19 scFV, optionally wherein the anti-CD19 scFV is encoded by SEQ ID NO: 3, i.e., SEQ ID: 14 disclosed in WO2012079000A1. In one embodiment, the anti-CD 19 scFV comprises SEQ ID NO: 4, i.e., the amino 65 acid sequence of SEQ ID NO: 20 disclosed in WO2012079000A1. The sequences in this paragraph appear

in WO2012079000A1 and are explicitly incorporated herein for use in the present invention and for possible inclusion in one or more claims herein.

In one embodiment, the transmembrane domain that naturally is associated with one of the domains in the CAR is used. In some instances, the transmembrane domain can be selected or modified by amino acid substitution to avoid binding of such domains to the transmembrane domains of the same or different surface membrane proteins to minimize interactions with other members of the receptor complex.

The transmembrane domain may be derived either from a natural or from a synthetic source. Where the source is natural, the domain may be derived from any membranebound or transmembrane protein. Transmembrane regions of particular use in this invention may be derived from (i.e. comprise at least the transmembrane region(s) of) the alpha, beta or zeta chain of the T-cell receptor, CD28, CD3 epsilon, CD45, CD4, CD5, CD5, CD9, CD16, CD22, CD33, CD37, CD64, CD80, CD86, CD 134, CD137 or CD 154. Alternatively the transmembrane domain may be synthetic, in which case it will comprise predominantly hydrophobic residues such as leucine and valine. Optionally, a triplet of phenylalanine, tryptophan and valine will be found at each end of a synthetic transmembrane domain.

Optionally, a short oligo- or polypeptide linker, preferably between 2 and 10 amino acids in length forms a linkage between the transmembrane domain and the intracellular part of the immune cell transmembrane protein, such as the CAR. A glycine-serine doublet provides a particularly suitable linker (eg, a (G_aS), linker as disclosed herein).

Optionally, the transmembrane domain is the CD8 transmembrane domain encoded by SEQ ID NO: 5, i.e., the nucleic acid sequence of SEQ ID NO: 16 disclosed in WO2012079000A1. In one embodiment, the CD8 transmembrane domain comprises SEQ ID NO: 6, i.e., the amino acid sequence of SEQ ID NO: 22 disclosed in WO2012079000A1. The sequences in this paragraph appear in WO2012079000A1 and are explicitly incorporated herein for use in the present invention and for possible inclusion in 40 one or more claims herein.

In some instances, the transmembrane domain comprises the CD8 hinge domain encoded by SEQ ID NO: 7, i.e., the nucleic acid sequence of SEQ ID NO: 15 disclosed in WO2012079000A1. In one embodiment, the CD8 hinge domain comprises SEQ ID NO: 8, i.e., the amino acid sequence of SEQ ID NO: 21 disclosed in WO2012079000A1. The sequences in this paragraph appear in WO2012079000A1 and are explicitly incorporated herein for use in the present invention and for possible inclusion in one or more claims herein.

The intracellular part or otherwise the intracellular signaling domain(s) of the transmembrane protein expressed by cells of the cell population administered to the patient is responsible for activation of at least one of the normal effector functions of the immune cell that expresses the transmembrane protein (eg, a T-cell function, such as leading to cytotoxicity (for T-effector cells for example) or suppression (for T-regulatory cells)). The term "effector function" refers to a specialized function of a cell. Effector function of a T cell, for example, may be cytolytic activity or helper activity including the secretion of cytokines. Thus the term "intracellular signaling domain" refers to the portion of a protein which transduces the effector function signal and directs the cell to perform a specialized function. While usually the entire intracellular signaling domain can be employed, in many cases it is not necessary to use the entire chain. To the extent that a truncated portion of the intracellular signaling domain is used, such truncated portion may be used in place of the intact chain as long as it transduces the effector function signal. The term "signaling domain" is thus meant to include any truncated portion of the intracellular signaling domain sufficient to transduce the 5 effector function signal. Examples of intracellular signaling domains for use in the transmembrane protein of the administered cells include the cytoplasmic sequences of the T cell receptor (TCR) and co-receptors that act in concert to initiate signal transduction following antigen receptor 10 engagement, as well as any derivative or variant of these sequences and any synthetic sequence that has the same functional capability.

It is known that signals generated through the TCR alone are insufficient for full activation of the T cell and that a 15 secondary or co-stimulatory signal is also required. Thus, T cell activation can be said to be mediated by two distinct classes of cytoplasmic signaling sequence: those that initiate antigen-dependent primary activation through the TCR (primary cytoplasmic signaling domain) and those that act in an 20 antigen-independent manner to provide a secondary or costimulatory signal (secondary cytoplasmic signaling domain). Primary cytoplasmic signaling sequences regulate primary activation of the TCR complex either in a stimulatory way, or in an inhibitory way. Primary cytoplasmic 25 signaling sequences that act in a stimulatory manner may contain signaling motifs which are known as immunoreceptor tyrosine-based activation motifs or ITAMs.

In an example, the first signalling domain is a primary cytoplasmic signaling domain (eg, CD3ζ domain) In an 30 example, the first signalling domain is a secondary cytoplasmic signaling domain (eg, CD28 or 4-1BB domain).

In an example, the first signalling domain comprises one or more ITAMs.

Examples of suitable ITAM containing primary cytoplas- 35 mic signaling domains that are of particular use in the invention include those derived from TCR zeta, FcR gamma, FcR beta, CD3 gamma, CD3 delta, CD3 epsilon, CDS, CD22, CD79a, CD79b, and CD66d. It is particularly preferred that cytoplasmic signaling molecule in the trans- 40 membrane protein of the invention comprises a cytoplasmic signaling sequence derived from CD3 zeta.

The intracellular part optionally comprises (eg, as the first signalling domain or a further intracellular domain) a domain of a costimulatory molecule. A costimulatory mol- 45 ecule is a cell surface molecule other than an antigen receptor or their ligands that is required for an efficient response of lymphocytes (eg, T- or NK cells) to an antigen. Examples of such molecules include CD27, CD28, 4-1BB (CD 137), OX40, CD30, CD40, PD-1, ICOS, lymphocyte 50 function-associated antigen-1 (LFA-1), CD2, CD7, LIGHT, NKG2C, B7-H3, and a ligand that specifically binds with CD83, and the like. Thus, these and other costimulatory elements are within the scope of the invention for use in the intracellular part of the transmembrane protein.

The intracellular moiety domains may be linked together by one or more linkers, eg, a $(G_4S)_n$ linker as disclosed herein.

In one embodiment, the intracellular moiety comprises the signaling domain of CD3-zeta and the signaling domain 60 of CD28. In another embodiment, the intracellular moiety comprises the signaling domain of CD3-zeta and the signaling domain of 4-1BB. In yet another embodiment, the intracellular moiety comprises the signaling domain of CD3zeta and the signaling domain of CD28 and 4-1BB.

In one embodiment, the intracellular moiety comprises the signaling domain of 4-1BB and the signaling domain of

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CD3-zeta, wherein the signaling domain of 4-1BB is encoded by SEQ ID NO: 9, i.e., the nucleic acid sequence set forth in SEQ ID NO: 17 disclosed in WO2012079000A1 and the signaling domain of CD3-zeta is encoded by SEQ ID NO: 2, i.e., the nucleic acid sequence set forth in SEO ID NO: 18 disclosed in WO2012079000A1. The sequences in this paragraph appear in WO2012079000A1 and are explicitly incorporated herein for use in the present invention and for possible inclusion in one or more claims herein.

In one embodiment, the intracellular moiety comprises the signaling domain of 4-1BB and the signaling domain of CD3-zeta, wherein the signaling domain of 4-1BB comprises SEQ ID NO: 10, i.e., the amino acid sequence of SEQ ID NO: 23 disclosed in WO2012079000A1 and the signaling domain of CD3-zeta comprises SEQ ID NO: 1, i.e., the amino acid sequence of SEQ ID NO: 24 disclosed in WO2012079000A1. The sequences in this paragraph appear in WO2012079000A1 and are explicitly incorporated herein for use in the present invention and for possible inclusion in one or more claims herein.

In one embodiment, the intracellular moiety comprises the signaling domain of 4-1BB and the signaling domain of CD3-zeta, wherein the signaling domain of 4-1BB comprises SEQ ID NO: 10, i.e., the amino acid sequence set forth in SEQ ID NO: 23 as disclosed in WO2012079000A1 and the signaling domain of CD3-zeta comprises SEQ ID NO: 1, i.e., the amino acid sequence set forth in SEQ ID NO: 24 disclosed in WO2012079000A1. The sequences in this paragraph appear in WO2012079000A1 and are explicitly incorporated herein for use in the present invention and for possible inclusion in one or more claims herein.

Sources of T-cells and other immune cells are disclosed in WO2012079000A1, U.S. Pat. Nos. 8,906,682, 8,911,993, 8,916,381, 8,975,071, 9,101,584, 9,102,760, 9,102,761, 9,328,156, 9,464,140, 9,481,728, 9,499,629, 9,518,123, 9.540.445. U.S. 20130287748, US20130288368, US20130309258, US20140106449, US20140370017, US20150050729, US20150093822, US20150099299, US20150118202, US20160130355, US20160159907, US20160194404, US20160208012, as well as methods of generating, activating and expanding these. These disclosures are referred to for possible use in working the present invention

Cancers for Treatment or Prevention by the Method

Cancers that may be treated include tumours that are not vascularized, or not substantially vascularized, as well as vascularized tumours. The cancers may comprise non-solid tumours (such as haematological tumours, for example, leukaemias and lymphomas) or may comprise solid tumours. Types of cancers to be treated with the invention include, but are not limited to, carcinoma, blastoma, and sarcoma, and certain leukaemia or lymphoid malignancies, benign and malignant tumours, and malignancies e.g., sar-55 comas, carcinomas, and melanomas. Adult tumours/cancers and paediatric tumours/cancers are also included

Haematologic cancers are cancers of the blood or bone marrow. Examples of haematological (or haematogenous) cancers include leukaemias, including acute leukaemias (such as acute lymphocytic leukaemia, acute myelocytic leukaemia, acute myelogenous leukaemia and myeloblasts, promyeiocytic, myelomonocytic, monocytic and erythroleukaemia), chronic leukaemias (such as chronic myelocytic (granulocytic) leukaemia, chronic myelogenous leukaemia, and chronic lymphocytic leukaemia), polycythemia vera, lymphoma, Hodgkin's disease, non-Hodgkin's lymphoma (indolent and high grade forms), multiple myeloma,

Waldenstrom's macroglobulinemia, heavy chain disease, myeiodysplastic syndrome, hairy cell leukaemia and myelodysplasia.

Solid tumours are abnormal masses of tissue that usually do not contain cysts or liquid areas. Solid tumours can be benign or malignant. Different types of solid tumours are named for the type of cells that form them (such as sarcomas, carcinomas, and lymphomas). Examples of solid tumours, such as sarcomas and carcinomas, include fibrosarcoma, myxosarcoma, liposarcoma, chondrosarcoma, osteosarcoma, and other sarcomas, synovioma, mesothelioma, Ewing's tumour, leiomyosarcoma, rhabdomyosarcoma, colon carcinoma, lymphoid malignancy, pancreatic cancer, breast cancer, lung cancers, ovarian cancer, prostate 15 cancer, hepatocellular carcinoma, squamous cell carcinoma, basal cell carcinoma, adenocarcinoma, sweat gland carcinoma, medullary thyroid carcinoma, papillary thyroid carcinoma, pheochromocytomas sebaceous gland carcinoma, papillary carcinoma, papillary adenocarcinomas, medullary 20 carcinoma, bronchogenic carcinoma, renal cell carcinoma, hepatoma, bile duct carcinoma, choriocarcinoma, Wilms' tumour, cervical cancer, testicular tumour, seminoma, bladder carcinoma, melanoma, and CNS tumours (such as a glioma (such as brainstem glioma and mixed gliomas), 25 glioblastoma (also known as glioblastoma multiforme) astrocytoma, CNS lymphoma, germinoma, medulloblastoma, Schwannoma craniopharyogioma, ependymoma, pineaioma, hemangioblastoma, acoustic neuroma, oligodendroglioma, menangioma, neuroblastoma, retinoblastoma and brain metastases).

In one embodiment, the administered cells express a first antigen binding site (eg, comprised by a CAR) that is designed to treat a particular cancer. For example, it spe-35 cifically binds to CD19 can be used to treat cancers and disorders, eg, pre-B ALL (paediatric indication), adult ALL, mantle cell lymphoma, diffuse large B-cell lymphoma or for salvage post allogenic bone marrow transplantation. In another embodiment, the first moiety or first binding site $_{40}$ specifically binds CD22 to treat diffuse large B-cell lymphoma.

In one embodiment, cancers and disorders include but are not limited to pre-B ALL (paediatric indication), adult ALL, mantle cell lymphoma, diffuse large B-cell lymphoma, sal- 45 vage post allogenic bone marrow transplantation, and the like can be treated using a combination of bridging agents (or binding moieties or sites comprised by a single agent) that target two or three of: CD19, CD20, CD22, and ROR1 (eg, CD19 and one of the other targets).

In an example, the cells comprises first and second transmembrane proteins (eg, CARs or a CAR and an engineered TCR expressed by a T-cell) that are different, eg that differ in their target antigens (and optionally otherwise are the same). Similarly, the invention may use a mixture of 55 immune cells (eg, a mixture of CAR-cells), eg comprised by the same transplant, wherein the mixture comprises cells comprising transmembrane proteins (eg, CARs or a CAR and an engineered TCR expressed by a T-cell) that are different, eg that differ in their target antigens (and option- 60 ally otherwise are the same). This may be useful for reducing resistance to treatment by cancers, for example, or more effectively targeting cell populations such as cancer cells that surface express a plurality of target antigens.

In one embodiment, the antigen binding site specifically 65 binds to mesothelin to treat or prevent mesothelioma, pancreatic cancer or ovarian cancer.

In one embodiment, the antigen binding site specifically binds to CD33/IL3Ra to treat or prevent acute myelogenous leukaemia.

In one embodiment, the antigen binding site specifically binds to c-Met to treat or prevent triple negative breast cancer or non-small cell lung cancer.

In one embodiment, the antigen binding site specifically binds to PSMA to treat or prevent prostate cancer.

In one embodiment, the antigen binding site specifically binds to Glycolipid F77 to treat or prevent prostate cancer.

In one embodiment, the antigen binding site specifically binds to EGFRvIII to treat or prevent gliobastoma.

In one embodiment, the antigen binding site specifically binds to GD-2 to treat or prevent neuroblastoma or melanoma.

In one embodiment, the antigen binding site specifically binds to NY-ESO-1 TCR to treat myeloma, sarcoma or melanoma.

In one embodiment, the antigen binding site specifically binds to MAGE A3 TCR to treat myeloma, sarcoma and melanoma.

Specific antigen binding is binding with a KD of 1 mM or lower (eg, 1 mM or lower, 100 nM or lower, 10 nM or lower, 1 nM or lower, 100 pM or lower, or 10 pM or lower) as determined by Surface Plasmon Resonance (SPR) in vitro at 25 degrees celcius or rtp.

In one example, said treatment using the method reduces progression of the disease or condition or a symptom thereof. In one example, said treatment using the method reduces incidence of the disease or condition or symptom thereof, eg, for at least 1, 2, 3, 4, or 5 years.

In an example, the method is in vivo in a mammal, eg, a human, man or woman, or male child or female child, or a human infant (eg, no more than 1, 2, 3 or 4 years of age). In an example, the patient is an adult human or a paediatric human patient.

The CAR or TCR is engineered, ie, comprises a nonnaturally-occurring combination of moieties and domains. In an example, the cell therapy targets a target cell, wherein the target cell is a cancer cell, eg, a leukaemic cell, lymphoma cell, adenocarcinoma cell or cancer stem cell. Optionally, the CAR or TCR of administered immune cells specifically binds to human CD19 (and optionally the target cell is a leukaemic or lymphoma cell), EpCAM (and optionally the target cell is a lung cancer cell, gastrointestinal cancer cell, an adenocarcinoma, cancer stem cell), CD20 (and optionally the target cell is a leukaemic cell), MCSP (and optionally the target cell is a melanoma cell), CEA, EGFR, EGFRvIII, sialyl Tn, CD133, CD33 (and optionally the target cell is a leukaemic cell, eg, AML cell), PMSA, WT1, CD22, L1CAM, ROR-1, MUC-16, CD30, CD47, CD52, gpA33, TAG-72, mucin, CIX, GD2, GD3, GM2, CD123, VEGFR, integrin, cMET, Her1, Her2, Her3, MAGE1, MAGE A3 TCR, NY-ESO-1, IGF1R, EPHA3, CD66e, EphA2, TRAILR1, TRAILR2, RANKL, FAP, Angiopoietin, mesothelin, Glycolipid F77 or tenascin.

Optionally, the CAR comprises the variable domains of an antibody selected from the group consisting of the CD19 binding site of blinatumomab or antibody HD37; EpCAM binding site of Catumaxomab; CD19 binding site of AFM11; CD20 binding site of Lymphomun; Her2 binding site of Ertumaxomab; CEA binding site of AMG211 (MEDI-565, MT111); PSMA binding site of Pasotuxizumab; EpCAM binding site of solitomab; VEGF or angiopoietin 2 binding site of RG7221 or RG7716; Her1 or Her3 binding site of RG7597; Her2 or Her3 binding site of MM111; IGF1R or Her3 binding site of MM141; CD123 binding site of MGD006; gpa33 binding site of MGD007; CEA binding site of TF2; CD30 binding site of AFM13; CD19 binding site of AFM11; and Her1 or cMet binding site of LY3164530.

Optionally, the CAR comprises the variable domains of an antigen binding site of an antibody selected from the group 5 consisting of REOPRO® (Abciximab); RITUXAN® (Rituximab); ZENAPAX® (Daclizumab); SIMULECT® (Basiliximab); SYNAGIS® (Palivizumab); REMICADE® (Infliximab); HERCEPTIN® (Trastuzumab); MYLOTARG® (Gemtuzumab ozogamicin); CAMPATH® (Alemtuzumab); 10 ZEVALIN® (Ibritumomab); HUMIRA® (Adalimumab); XOLAIR® (Omalizumab); BEXXAR® (Tositumomab); RAPTIVA® (Efalizumab); ERBITUX® (Cetuximab); AVASTIN® (Bevacizumab); TYSABRI® (Natalizumab); ACTEMRA® (Tocilizumab); VECTIBIX® (Panitu- 15 mumab); LUCENTIS® (Ranibizumab); SOLIRIS® (Eculizumab); CIMZIA® (Certolizumab); SIMPONI® (Golimumab): **ILARIS®** (Canakinumab); STELARA® (Ustekinumab); ARZERRA® (Ofatumumab); PROLIA® (Denosumab); NUMAX® (Motavizumab); ABTHRAX® 20 (Raxibacumab); BENLYSTA® (Belimumab); YERVOY® (Ipilimumab); ADCETRIS® (Brentuximab vedotin); PER-JETA® (Pertuzumab); KADCYLA® (Ado-trastuzumab); and GAZYVA® (Obinutuzumab).

In an example, the target cell is a blood cell, eg, a stem cell 25 or bone marrow cell of a human or animal. Optionally, the target cell is a B- or T-cell.

In an example, the CAR or TCR comprises an antigen binding site for an autoimmune disease target and the signaling down-regulates cytotoxic activity or proliferation 30 of the immune cells. The term "autoimmune disease" as used herein is defined as a disorder that results from an autoimmune response. An autoimmune disease is the result of an inappropriate and excessive response to a self-antigen. Examples of autoimmune diseases include but are not lim- 35 ited to, Addision's disease, alopecia greata, ankylosing spondylitis, autoimmune hepatitis, autoimmune parotitis, Crohn's disease, diabetes (Type I), dystrophic epidermolysis bullosa, epididymitis, glomerulonephritis, Graves' disease, Guillain-Barr syndrome, Hashimoto's disease, hemolytic 40 anemia, systemic lupus erythaematosus, multiple sclerosis, myasthenia gravis, pemphigus vulgaris, psoriasis, rheumatic fever, rheumatoid arthritis, sarcoidosis, scleroderma, Sjogren's syndrome, spondyloarthropathies, thyroiditis, vasculitis, vitiligo, myxedema, pernicious anemia, ulcerative 45 colitis, among others.

Within the overall memory T cell population, several distinct subpopulations have been described and can be recognised by the differential expression of chemokine receptor CCR7 and L-selectin (CD62L). Stem memory 50 T_{SCM} cells, like naive cells, are CD45RO-, CCR7+, CD45RA+, CD62L+ (L-selectin), CD27+, CD28+ and IL-7R α +, but they also express large amounts of CD95, IL-2Rβ, CXCR3, and LFA-1, and show numerous functional attributes distinctive of memory cells. Central 55 memory T_{CM} cells express L-selectin and the CCR7, they secrete IL-2, but not IFN γ or IL-4. Effector memory T_{EM} cells, however, do not express L-selectin or CCR7 but produce effector cytokines like IFNy and IL-4. Memory T-cells, such as T_{SCM} may be particularly useful for estab- 60 lishing a sustained population of engineered immune cells in the human.

Any immune cell, target cell or stem cell herein can, in an example, be a T_{SCM} , T_{CM} or T_{EM} cell, eg, a human T_{SCM} , T_{CM} or T_{EM} cell. In an example, the immune cells of the cell 65 therapy (eg, CAR-T cells) each is a progeny of a cell of a human suffering from an autoimmune disease, an inflam-

matory disease, a viral infection or a cancer, eg, wherein the human is suffering from lymphoblastic leukaemia, ALL (eg, T-ALL), CLL (eg, B-cell chronic lymphocytic leukaemia) or non-Hodgkin's lymphoma. The human may, for example, be the patient or a relative (eg, sibling or parent) thereof.

In an example, the administered immune cells have been engineered for enhanced signaling, wherein the signaling is selected from CD28, 4-1BB, OX40, ICOS and CD40 signaling.

Optionally, the target cells (eg, tumour cells) are killed. In an example, each target cell is a tumour cell and the method treats or reduces the risk of cancer, or treats or reduces the risk of cancer progression in the human.

Optionally, the human has cancer. In an example, the cancer is a haematological cancer. In an example, the human has a cancer of B-cell origin. In an example, the human has a cancer of T-cell origin. For example the cancer is lung cancer, melanoma, breast cancer, prostate cancer, colon cancer, renal cell carcinoma, ovarian cancer, neuroblastoma, rhabdomyosarcoma, leukaemia and lymphoma. Preferred cancer targets for use with the present invention are cancers of B cell origin, particularly including acute lymphoblastic leukaemia, B-cell chronic lymphocytic leukaemia or B-cell non-Hodgkin's lymphoma. In an example, the cancer is a cancer of T-cell or B-cell origin, eg, lymphoblastic leukaemia, ALL (eg, T-ALL), CLL (eg, B-cell chronic lymphocytic leukaemia) or non-Hodgkin's lymphoma.

Optionally, each administered immune cell (eg, CAR-cells) is a progeny of an immune cell of said human, eg, wherein the human is suffering from lymphoblastic leukaemia, Diffuse Large B-cell Lymphoma (DLBCL), ALL (eg, T-ALL or B-ALL), CLL (eg, B-cell chronic lymphocytic leukaemia) or non-Hodgkin's lymphoma. Optionally, each administered immune cell (eg, CAR-cells) is an autologous cell (eg, T-cell) of said human or is a progeny of such an autologous cell. As used herein, the term "autologous" is meant to refer to any material derived from the same individual. "Allogeneic" refers to a graft derived from a different animal of the same species.

Optionally, each administered immune cell (eg, CARcells) is derived from a blood or tumour sample of the human and activated and expanded in vitro before step (c). "Activation," as used herein, refers to the state of a T-cell or other immune cell that has been sufficiently stimulated to induce detectable cellular proliferation. Activation can also be associated with induced cytokine production, and detectable effector functions. The term "activated T cells" refers to, among other things, T cells that are undergoing cell division.

In an embodiment, the human has an autoimmune disease, wherein the immune cells that are administered (eg, CARcells) are anergic, or have reduced proliferation and/or cytotoxic activity when bound to target cells, whereby the cell transplant cells (and/or their progeny) compete with endogenous immune cells of said human that up-regulate said autoimmune disease.

The administration of immune cells in the method may be by cell infusion into the blood of the patient. The immune cells may be expanded to produce an expanded immune cell population that is administered to the patient. The immune cells may be activated produce an activated immune cell population that is administered to the patient. In methods herein, an effective amount of immune cells are administered. An "effective amount" as used herein, means an amount which provides a therapeutic or prophylactic benefit to treat or prevent the disease or condition.

In an embodiment of the method of the invention, the method treats or reduces the risk of cancer in a patient (eg, a human), wherein the patient has undergone lymphodepletion before administration of the immune cells to the patient.

In one embodiment, the human is resistant to at least one 5 chemotherapeutic agent.

In one embodiment, the chronic lymphocytic leukaemia is refractory CD 19+ leukaemia and lymphoma.

The invention also includes a method of generating a persisting population of genetically engineered T cells in a 10 human diagnosed with cancer, wherein the administered cells comprise T-cells and the persisting population comprises progeny thereof. In one embodiment, the method comprises administering to a human a T-cell population (eg, a CAR T-cell population), wherein the persisting population 15 of genetically engineered T-cells persists in the human for at least one month after administration. In one embodiment, the persisting population of genetically engineered T-cells comprises a memory T-cell. In one embodiment, the persisting population of genetically engineered T-cells persists 20 in the human for at least three months after administration. In another embodiment, the persisting population of genetically engineered T-cells persists in the human for at least four months, five months, six months, seven months, eight months, nine months, ten months, eleven months, twelve 25 months, two years, or three years after administration.

In one embodiment, the chronic lymphocytic leukaemia is treated. The invention also provides a method of expanding a population of the engineered T-cells or NK cells in a human diagnosed with cancer, wherein the administered 30 cells comprise T-cells and/or NK cells and the expanded population comprises progeny thereof.

Optionally, autologous lymphocyte infusion is used in the treatment. For example, autologous PBMCs are collected from a patient in need of treatment and CAR-T-cells are 35 engineered to express the CAR transmembrane protein, activated and expanded using the methods known in the art and then infused back into the patient in step (a).

In an example, the administered cells are pluripotent or multipotent.

The stem cell cannot develop into a human. In an embodiment, the stem cell cannot develop into a human embryo or zygote.

In an example, the administered cell population comprises bone marrow stem cells, eg, human autologous or allogeneic 45 cells.

In an example, the administered cell population comprises haematopoietic stem cells, eg, human autologous or allogeneic cells.

Modifying Microbiota

Medical practice often involves the administration of antibiotics to patients. Such treatments can typically involve administration of broad-spectrum antibiotics, or antibiotics that target many gram-positive bacterial species or many gram-negative species without discrimination. Similarly, use 55 of broad-spectrum antibiotics in farming and agriculture, for example, raise environmental concerns, including entry of such antibiotics into the human and animal food chain which may be deleterious to health and may add to development of microbial resistance. Rather, in an example, the invention 60 involves selective targeting of a first microbial (eg, bacterial or archaeal) species or strain of the microbiota. As shown in the worked examples herein, selective targeting of a particular bacterial species has been achieved using guided nuclease targeting of the genome of the selected species, 65 whilst at the same time sparing related species and strains, as well as species that co-reside (in the Examples species

that co-reside in human gut microbiota). Thus, in one example, the step of causing dysbiosis or step (b) comprises killing first cells of a microbiota sub-population or inhibiting growth of said sub-population by using guided nuclease (eg, RNA guided nuclease) targeting to the genome of first cells comprised by a microbiota sub-population. Suitable systems for carrying out the guided nuclease targeting are, for example, engineered CRISPR/Cas systems, TALENs, meganucleases and zinc finger systems. By way of example, CRISPR/Cas-mediated guided targeting of a selected human gut microbiota bacterial species in a consortium is demonstrated in the Examples herein. The targeting produces nuclease cutting of target species or strain DNA, for example, which reduces the relative proportion of the species in the microbiota or inhibits growth of the sub-population of said species in the microbiota. Selective targeting of species in the method is generally advantageous to enable finer control over change in the relative proportions of bacterial and/or archaeal species in the microbiota. In this way, the invention provides the ability to alter the microbiota with the aim of influencing the upregulation or downregulation of particular immune cell populations, such as $T_H 1$, $T_H 17$ and/or T_{reg} cells (be these cells endogenous to the patient and/or comprised by adoptive immune cell populations that are administered to the patient), or other outcomes of modulating the microbiota as described herein.

In an example first cell population growth is reduced by at least 5-fold compared to the growth before said dysbiosis or step (b). The method may comprise inhibiting first cell population growth on a gut surface. The method may comprise inhibiting first cell population growth on a plant (eg, leaf and/or stem) surface.

In an alternative, instead of being applied to a subject, the treatment is applied to an environment or soil (eg, the treatment is a fertiliser, plant growth promoting or inhibiting, herbicide or pesticide treatment), wherein the treatment is modulated by the invention.

It will be readily apparent to the skilled addressee how to determine changes in bacteria and archaea in a gut micro-40 biota or other microbiota. For example, this can be done by analyzing a facecal sample of the patient before and after the treatment. One may determine the types of different species or strains in each sample and the proportion of species or strains before and after treatment. Using conventional analysis of 16s ribosomal RNA-encoding DNA (16s rDNA) it is possible to identify species, for example. Additionally or alternatively, standard biochemical test can be used to identify strains or species, eg, also involving one or more of: staining, motility testing, serological testing, phage typing and identification disc testing (eg using a Kirby Baur disc diffusion method). Biochemical testing may involve one or more of: a (a) Catalase test (b) Coagulase test (c) Oxidase test (d) Sugar fermentation test (e) Indole test (f) Citrate test and (g) Urease test. Relative proportions may be determined by growing colonies on agar plates (as in the Examples herein) from each sample and counting colony numbers.

In an example, the dysbiosis or step (b) increases the proportion of Bacteroides (eg, B fragalis and/or B thetaiota*micron*) in the microbiota, eg, gut microbiota.

In an example, the dysbiosis or step (b) decreases the proportion of Bacteroides (eg, B fragalis and/or B thetaiota*micron*) in the microbiota, eg, gut microbiota. In an example, the dysbiosis or step (b) increases the proportion of Bacteroidetes to Firmicutes in the microbiota, eg, gut microbiota. In an example, the dysbiosis or step (b) decreases the proportion of Bacteroidetes to Firmicutes in the microbiota, eg, gut microbiota.

Accumulating evidence supports the role of commensal strains of Bifidobacterium and Clostridium spp. belonging to clusters IV and XIVa in the induction of Treg cells. See, eg, Lopez et al. In an example, the dysbiosis or step (b) reduces the proportion of one or more *Clostridium* species or strain 5 (eg, In an example, each species is a cluster IV or XIVa Clostridium species) in the gut microbiota. In an example, the dysbiosis or step (b) increases the proportion of one or more *Clostridium* species or strain (eg, In an example, each species is a cluster IV or XIVa Clostridium species) in the 10 gut microbiota.

In an example, the dysbiosis or step (b) reduces the proportion of Bifidobacterium (eg, B bifidum) in the gut microbiota. In an example, the dysbiosis or step (b) increases the proportion of Bifidobacterium (eg, B bifidum) 15 in the gut microbiota.

For example, by selectively altering the human gut microbiota the invention provides for upregulation of CAR-T or other ACT treatment (eg, wherein the altered microbiota downregulates T_{reg} cells in the patient that has received the 20 CAR-T or ACT administration and/or upregulates T_{H} and/ or $T_H 17$ cells in the patient—such cells being comprised by the CAR-T or ACT transplant for example). Downregulating T_{reg} cells may reduce suppression of T-effectors and/or T-helpers in the patient, thereby enhancing the CAR-T or 25 ACT cytotoxicity or other desirable activity against cancer or other disease-mediating cells. Upregulating $T_{H}1$ and/or T_H17 cells may increase T-effector activity, thereby enhancing the CAR-T or ACT cytotoxicity or other desirable activity against cancer or other disease-mediating cells.

In another example, alteration of the microbiota can be used as a switch to dampen down CAR-T or other ACT treatment (eg, wherein the altered microbiota upregulates Treg cells in the patient that has received the CAR-T or ACT administration and/or downregulates $T_H 1$ and/or $T_H 17$ cells 35 in the patient-such cells being comprised by the CAR-T or ACT transplant for example). Upregulating T_{reg} cells may increase suppression of T-effectors and/or T-helpers in the patient, thereby reducing the CAR-T or ACT ability to promote cytokine release or other undesirable activity. 40 Downregulating $T_H 1$ and/or $T_H 17$ cells may decrease T-effector activity, thereby reducing the CAR-T or ACT ability to promote cytokine release or other undesirable activity. This may be useful for limiting the risk of cytokine release syndrome (CRS) in the patient. Subsequent further modifi- 45 cation of the gut microbiota of the patient using the method of the invention can be performed to upregulate the CAR-T or ACT treatment when it is desired to use this once more to address the disease or condition at hand (eg, a cancer, such as a haematological cancer). In this instance, memory T-cell 50 CAR-T or ACT populations may be present in the patient from the earlier treatment, and the upregulation using microbiota alteration according to the invention may upregulate memory T-cells to differentiate into effector and/or helper cells to address the disease or condition. Thus, in one 55 example, the cell therapy of the invention comprises administering an immune cell population comprising immune memory cells (eg, memory T-cells, such as central memory T cells (T_{CM}) and/or stem cell memory cells (T_{SCM}) ; and/or the administered population comprises cells that spawn such 60 memory cells following the initial microbiota alteration.

Whilst one aspect of the invention recognizes utility for modulating cell-based therapy in a patient, another aspect recognizes utility for modulating (ie, treating or preventing) cell-mediated diseases and conditions in patients, such as autoimmune and inflammatory diseases and conditions which are mediated, for example by T-cells or other immune

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cells of the patient. In a further aspect, the invention recognizes utility as a means for modulating (eg, enhancing) another therapy of a disease or condition; for example, for enhancing or effecting therapy with an antibody or anti-viral medicine to treat or prevent the disease or condition. For example, the medicine can be an immune checkpoint antagonist or agonist (eg, for treating or preventing a cancer, such as melanoma or NSCLC). By "effecting therapy" it is contemplated that the patient does not respond or poorly responds to the medicine and the microbiota alteration according to the invention (eg, using selective guided nuclease targeting of a bacterial or archaeal species as described herein) brings about a response (or improved response) to the medicine by the patient. For example, the method of the invention upregulates $T_H 17$ cells in a patient suffering from HIV infection. In one aspect, this enhances anti-retroviral therapy or HIV vaccine therapy of the patient. The $T_H 17$ cells may be the patient's endogenous cells or cells provided by ACT of the patient. In another example, the method of the invention upregulates $T_H 17$ cells in a patient suffering from a cancer (eg, melanoma or lung cancer, such as NSCLC). In one aspect, this enhances immune checkpoint antagonism or agonism therapy of the patient. The $T_H 17$ cells may be the patient's endogenous cells or cells provided by ACT of the patient. For example, the therapy is antibody therapy using an antibody selected from ipilimumab (or YERVOY™), tremelimumab, nivolumab (or OPDIVOTM), pembrolizumab (or KEYTRUDA[™]), pidilizumab, BMS-936559, durvalumab and atezolizumab.

The invention relates to guided nuclease systems (eg, engineered CRISPR/Cas systems, TALENs, meganucleases and zinc finger systems), arrays (eg, CRISPR arrays), cRNAs, gRNAs and vectors (eg, phage comprising components of a said system) for use in a method of the invention for targeting the first cells or causing said dysbiosis by inhibiting bacterial or archaeal cell population growth or altering the relative proportion of one or more sub-populations of cells in plant, yeast, environmental, soil, human or animal microbiota, such as for the alteration of the proportion of Bacteroidetes (eg, Bacteroides), Firmicutes and/or gram positive or negative bacteria in gut microbiota of a human. The invention, for example, involves modifying (eg, cutting and/or mutating) one or more target genomic or episomal nucleotide sequences of a host bacterial cell, eg, a Bacteroidetes cell or Firmicutes cell, or a host archaeal cell. In an example, the first bacteria are pathogenic gut bacteria.

There have been a number of studies pointing out that the respective levels of the two main intestinal phyla, the Bacteroidetes and the Firmicutes, are linked to obesity, both in humans and in germ-free mice. The authors of the studies deduce that carbohydrate metabolism is the important factor. They observe that the microbiota of obese individuals are more heavily enriched with bacteria of the phylum Firmicutes and less with Bacteroidetes, and they surmise that this bacterial mix may be more efficient at extracting energy from a given diet than the microbiota of lean individuals (which have the opposite proportions). In some studies, they found that the relative abundance of Bacteroidetes increases as obese individuals lose weight and, further, that when the microbiota of obese mice are transferred to germfree mice, these mice gain more fat than a control group that received microbiota from lean mice. See, eg, Turnbaugh, P. J., R. E. Ley, M. A. Mahowald, V. Magrini, E. R. Mardis, and J. I. Gordon. 2006, "An obesity-associated gut microbiome with increased capacity for energy harvest", Nature 444:1027-1131. In a further aspect, the invention recognizes utility as

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a means for enhancing an anti-obesity therapy of a patient, eg, by increasing the ratio of Bacteroidetes versus *Firmicutes* in the microbiota.

Optionally the first cells are in the presence of cells of a different strain or species, wherein the different cells are 5 Enterobacteriaceae or bacteria that are probiotic, commensal or symbiotic with humans (eg, in the human gut). In an example, each first cell is a *Firmicutes*, eg, *Streptococcus*, cell.

In an example, the invention is able to selectively kill or 10 downregulate the target microbes in the microbiota whilst not targeting a second related strain of the same species or a different species that is nevertheless phylogenetically related (as indicated by 16s rDNA). For example, the microbiota comprises cells of a second bacterial species or 15 strain, or archaeal species or strain, wherein the second species or strain has a 16s ribosomal RNA-encoding DNA sequence that is at least 80, 85, 90, 95, 96, 97, 98 or 99% identical to an 16s ribosomal RNA-encoding DNA sequence of the first cell species or strain, wherein the growth of the 20 second cells in the microbiota is not inhibited by said method. In an embodiment, the growth of second strain or species is not inhibited; or the growth of said first cells is inhibited by at least $2\times$, $3\times$, $4\times$, $5\times$, $6\times$, $7\times$, $8\times$, $9\times$, $10\times$, $50\times$, $100\times$ or $1000\times$ the growth inhibition of the second cells. 25

In one aspect of the method, causing the dysbiosis or step (b) comprises altering the proportion of a sub-population of first cells (host cells) in the microbiota, eg, gut microbiota, of the patient, thereby producing an altered gut microbiota that modulates the immune cell therapy in the patient, 30 wherein the sub-population comprises host cells of said first species or strain, the method comprising using guided nuclease (eg RNA-guided nuclease) cutting of a respective target sequence in host cells to modify the target sequences, whereby host cells are killed or the host cell population 35 growth is reduced, thereby reducing the proportion of said sub-population in the microbiota. Suitable systems for carrying out the guided nuclease cutting are, for example, engineered CRISPR/Cas systems, TALENs, meganucleases and zinc finger systems. By way of example, CRISPR/Cas- 40 mediated guided cutting of a selected human gut microbiota bacterial species in a consortium is demonstrated in the Examples herein.

In an example, the target sequence modification is carried out by

- a. combining the microbiota with multiple copies of engineered nucleic acid sequences encoding host modifying (HM) crRNAs, and
- b. expressing HM-crRNAs in host cells,
- wherein each engineered nucleic acid sequence is oper- 50 able with a Cas nuclease in a respective host cell to form a HM-CRISPR/Cas system and the engineered sequence comprises
- (i) spacer and repeat sequences encoding a HM-crRNA;
- (ii) the HM-crRNA comprising a sequence that is capable 55 of hybridizing to a host cell target sequence to guide Cas nuclease to the target sequence in the host cell; and
- optionally the HM-system comprises a tracrRNA sequence or a DNA sequence expressing a tracrRNA sequence;
- whereby HM-crRNAs guide Cas modification of host target sequences in host cells, whereby host cells are killed or the host cell population growth is reduced, thereby reducing the proportion of said sub-population in the microbiota.

In an alternative, HM-crRNA and tracrRNA are comprised by a single guide RNA (gRNA). In an example, each engineered nucleic acid sequence is comprised by a respective vector, wherein each vector is optionally a plasmid (eg, a conjugative plasmid capable of transfer into a host cell), phage, phagemid or prophage. The phage is capable of infecting a said host cell.

In an example, endogenous Cas nuclease of host cells is used for modification of target nucleotide sequences. In an embodiment, therefore, each vector lacks a Cas (eg, a Cas9) nuclease-encoding sequence. By harnessing endogenous Cas nuclease, embodiments of the invention use endogenous Cas nuclease activity (ie, without the need for prior genetic modification of the host cell to activate or enhance the nuclease activity). Thus, in an example, the Cas nuclease is encoded by a wild-type gene of the host cell. In an example, the nuclease is active to achieve the cell killing or growth reduction without inhibition of an endogenous Cas nuclease (or Cas nuclease gene) repressor in the host cell. Thus, the invention can address wild-type bacterial populations without the need for prior manipulation to make bring about effective Cas-mediated cell killing or growth reduction. Thus, the population can be exposed to the cRNA when the population is in its wild-type environment (such as comprised by a plant, yeast, environmental, soil, human or animal microbiome).

In an example, the cRNA or gRNA is for administration to (or administered to) a human or non-human animal patient by mucosal, gut, oral, intranasal, intrarectal or buccal administration.

Optionally said Cas nuclease is provided by an endogenous Type II CRISPR/Cas system of each first cell. Optionally, the tracrRNA sequence or DNA sequence expressing a tracrRNA sequence is endogenous to each host cell. Optionally, each target sequence is comprised by an antibiotic resistance gene, virulence gene or essential gene of the respective host cell, for example the target sequences are identical between the host cells. Optionally, the engineered nucleic acid sequences are comprised by an antibiotic composition, wherein the sequences are in combination with an antibiotic agent (first antibiotic), and in an example the target sequences are comprised by an antibiotic resistance gene wherein the antibiotic is said first antibiotic. The antibiotic composition is administered to the patient or subject to effect said dysbiosis or step (b).

Optionally, each host cell comprises a deoxyribonucleic acid strand with a free end (HM-DNA) encoding a HM-sequence of interest and/or wherein the method comprising into the host cells such a sequence encoding the HM-DNA, wherein the HM-DNA comprises a sequence or sequences that are homologous respectively to a sequence or sequences in or flanking the target sequence for inserting the HM-DNA into the host genome (eg, into a chromosomal or episomal site).

The invention also provides vectors for introducing into first cells (host cells) for carrying out the treatment or 55 prevention method of the invention, wherein each vector is: An engineered nucleic acid vector for modifying a bacterial or archaeal host cell comprising an endogenous CRISPR/ Cas system, the vector comprising nucleic acid sequences for expressing a plurality of different crRNAs (eg, gRNAs) 60 for use in causing the dysbiosis or for use in step (b) of the method; and optionally lacking a nucleic acid sequence encoding a Cas nuclease,

wherein a first of said crRNAs is capable of hybridising to a first nucleic acid sequence in said host cell; and a second of said crRNAs is capable of hybridising to a second nucleic acid sequence in said host cell, wherein said second sequence is different from said first sequence; and

- a. the first sequence is comprised by an antibiotic resistance gene (or RNA thereof) and the second sequence is comprised by an antibiotic resistance gene (or RNA thereof); optionally wherein the genes are different;
- b. the first sequence is comprised by an antibiotic resis- 5 tance gene (or RNA thereof) and the second sequence is comprised by an essential or virulence gene (or RNA thereof);
- c. the first sequence is comprised by an essential gene (or RNA thereof) and the second sequence is comprised by 10 an essential or virulence gene (or RNA thereof); or
- d. the first sequence is comprised by a virulence gene (or RNA thereof) and the second sequence is comprised by an essential or virulence gene (or RNA thereof).

Each vector may be as described above, eg, a phage 15 capable of infecting a host cell or conjugative plasmid capable of introduction into a host cell. In an example, the vectors are in combination with an antibiotic agent (eg, a beta-lactam antibiotic).

Each first cell (host cell) may be a Staphylococcus, 20 Streptococcus, Pseudomonas, Salmonella, Listeria, E coli, Desulfovibrio or Clostridium host cell. In an example, each first cell (host cell) is a Firmicutes cell, eg, a Staphylococcus, Streptococcus, Listeria or Clostridium cell.

In an example, each engineered nucleic acid sequence 25 comprises a sequence R1-S1-R1' for expression and production of the respective crRNA (eg, comprised by a single guide RNA) in the host cell, (i) wherein R1 is a first CRISPR repeat, R1' is a second CRISPR repeat, and R1 or R1' is optional; and (ii) S1 is a first CRISPR spacer that comprises 30 or consists of a nucleotide sequence that is 95% or more identical to said target sequence.

In an example, R1 and R1' are at least 95% identical respectively to the first and second repeat sequences of a CRISPR array of the host cell species. In an example, R1 35 and R1' are at least 95% (eg, 96, 97, 98, 99 or 100%) identical respectively to the first (5'-most) and second (the repeat immediately 3' of the first repeat) repeat sequences of a CRISPR array of said species, eg, of a said host cell of said species. In an example, R1 and R1' are functional with a 40 Type II Cas9 nuclease (eg, a S thermophilus, S pyogenes or S aureus Cas9) or Type I Cas3 to modify the target in a said host cell.

In one aspect, the method involves the following use, as demonstrated by the worked experimental Example:

The use of wild-type endogenous Cas nuclease activity of the first cell (host cell) population to inhibit growth of the population, wherein each host cell has an endogenous CRISPR/Cas system having wild-type Cas nuclease activity, the use comprising transforming host cells of the population, 50 wherein each transformed host cell is transformed with an engineered nucleotide sequence for providing host modifying (HM) cRNA or guide RNA (gRNA) in the host cell, the HM-cRNA or gRNA comprising a sequence that is capable of hybridising to a host cell target protospacer sequence for 55 guiding endogenous Cas to the target, wherein the cRNA or gRNA is cognate to an endogenous Cas nuclease of the host cell that has said wild-type nuclease activity and following transformation of the host cells growth of the population is inhibited. 60

By "cognate to" it is intended that the endogenous Cas is operable with crRNA or gRNA sequence to be guided to the target in the host cell. The skilled addressee will understand that such Cas guiding is generally a feature of CRISPR/Cas activity in bacterial and archaeal cells, eg, wild-type 65 CRISPR/Cas activity in bacterial or archaeal cells having endogenous active wild-type CRISPR/Cas systems.

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By "wild-type" Cas activity it is intended, as will be clear to the skilled addressee, that the endogenous Cas is not an engineered Cas or the cell has not been engineered to de-repress the endogenous Cas activity. This is in contrast to certain bacteria where Cas nuclease activity is naturally repressed (ie, there is no wild-type Cas nuclease activity or none that is useful for the present invention, which on the contrary is applicable to addressing wild-type host cells in situ for example where the endogenous Cas activity can be harnessed to effect cell population growth inhibition).

In the worked Examples below, inhibition was addressed in a bacterial population (a gram positive Firmicutes) on a solid surface. A >10-fold inhibition of host cell population growth was achieved. Targeting was directed to an antibiotic resistance gene and an essential gene. The demonstration of the invention's ability to inhibit host cell growth on a surface is important and desirable in embodiments where the invention is for treating or preventing diseases or conditions mediated or caused by microbiota as disclosed herein in a human or animal subject. Such microbiota are typically in contact with tissue of the subject (eg, gut, tissue) and thus we believe that the demonstration of activity to inhibit growth of a microbiota bacterial species (exemplified by Streptococcus) on a surface supports this utility. Targeting microbiota on plant surfaces is also a desired application.

In an example, inhibition of first cell (host cell) population growth is at least 2, 3, 4, 5, 6, 7, 8, 9 or 10-fold compared to the growth of cells of the same species or strain not exposed to said engineered nucleotide sequence. For example, growth inhibition is indicated by a lower bacterial colony number of a first sample of host cells (alone or in a mixed bacterial population, eg, a microbiota or faecal sample of the patient after treatment) by at least 2, 3, 4, 5, 6, 7, 8, 9 or 10-fold compared to the colony number of a second sample of the host cells (alone or in a mixed bacterial population, eg, a microbiota or faecal sample of the patient before treatment), wherein the first cells have been transformed by said engineered nucleotide sequence but the second sample has not been exposed to said engineered nucleotide sequence. In an embodiment, the colony count is determined 12, 24, 36 or 48 hours after the first sample has been exposed to the engineered sequence. In an embodiment, the colonies are grown on solid agar in vitro (eg, in a petri dish). It will be understood, therefore, that growth inhibition can be indicated by a reduction (<100% growth compared to no treatment, ie, control sample growth) in growth of first (host) cells or populations, or can be a complete elimination of such growth. In an example, growth of the host cell population is reduced by at least 10, 20, 30, 40, 50, 60, 70, 80, 90 or 95%, ie, over a predetermined time period (eg, 24 hours or 48 hours following combination with the cRNA or gRNA, TALEN, meganuclease, zinc finger etc in the host cells), ie, growth of the host cell population is at least such percent lower than growth of a control host cell population that has not been exposed to said cRNA or gRNA etc but otherwise has been kept in the same conditions for the duration of said predetermined period. In an example, percent reduction of growth is determined by comparing colony number in a sample of each population at the end of said period (eg, at a time of mid-exponential growth phase of the control sample). For example, after exposing the test population to the crRNA or gRNA etc at time zero, a sample of the test and control populations is taken and each sample is plated on an agar plate and incubated under identical conditions for said predetermined period. At the end of the period, the colony number of each sample is counted and the percentage difference (ie, test colony number divided by

control colony number and then times by 100, and then the result is subtracted from 100 to give percentage growth reduction). The fold difference is calculated by dividing the control colony number by the test colony number.

Inhibition of population growth can be indicated, there- 5 fore, by a reduction in proliferation of first (host) cell number in the population. This may be due to cell killing by the nuclease and/or by downregulation of host cell proliferation (division and/or cell growth) by the action of the nuclease on the target protospacer sequence. In an embodi- 10 ment of a treatment or prevention as disclosed herein, host cell burden of the human or animal subject is reduced, whereby the disease or condition is treated (eg, reduced or eliminated) or prevented (ie, the risk of the subject developing the disease or condition) is reduced or eliminated.

In an example, wild-type host cell endogenous Cas9 or cfp1 activity is used. In an example, wild-type host cell endogenous Cas3 and/or CASCADE activity is used. The engineered nucleotide sequence may not be in combination with an exogenous Cas nuclease-encoding sequence. 20 Optionally, said Cas nuclease is a nickase.

In an example, the formation of bacterial colonies of said host cells is inhibited following said dysbiosis or step (b). In an example, proliferation of host cells is inhibited following said dysbiosis or step (b). In an example, host cells are killed 25 following said dysbiosis or step (b).

In another aspect, the method comprises producing ex vivo a medicament for administration to the patient for causing said dysbiosis or step (b) for treating or preventing the disease or condition, wherein the medicament comprises 30 a modified mixed bacterial population (eg, obtained from faeces or gut microbiota of one or more human donors or said patient), wherein the modified population is administered to the patient to cause said dysbiosis or in step (b) to alter the balance of species or strains in the patient's gut 35 microbiota, thereby altering the proportion of the first cells in the gut microbiota. The modified mixed population can be produced ex vivo using guided nuclease modification techniques as described herein. Thus, for example, the method can be used to reduce the proportion of a specific Firmicutes 40 sub-population and spare Bacteroidetes in the mixed population, eg, for producing a medicament for treating or preventing a metabolic or GI condition or disease disclosed herein. In this way, the invention can use a modified bacterial transplant (eg, a modified faecal transplant) medica- 45 ment for such use or for said treatment or prevention in a human or animal. For example, the method can be used to modify one or more microbiota in vitro to produce a modified collection of bacteria for administration to a human or animal for medical use (eg, treatment or prevention of a 50 metabolic condition (such as obesity or diabetes) or a GI tract condition (eg, any such condition mentioned herein) or a cancer (eg, a GI tract cancer). In an example, the transformation of bacterial cells with phage or plasmid vectors comprising engineered nucleic acid sequences as described 55 herein is carried out in vitro, or the engineered nucleotide sequence is comprised by nucleic acid that is electroporated into host cells. In an example, the nucleic acid are RNA (eg, copies of the gRNA). In another example, the nucleic acid are DNA encoding the crRNA or gRNA for expression 60 thereof in host cells.

Thus, in an example, the invention provides an engineered nucleotide sequence for providing host cell modifying (HM) cRNA or guide RNA (gRNA) in a population of wild-type bacterial host cells comprised by a microbiota of a plant, 65 yeast, environmental, soil, human or animal subject for use in the method of the invention, the cRNA or gRNA com-

prising a sequence that is capable of hybridising to a host cell target protospacer sequence for guiding Cas to the target, wherein the cRNA or gRNA is cognate to an endogenous host cell Cas nuclease that has wild-type nuclease activity, wherein following transformation of host cells growth of the population is inhibited and the disease or condition is treated or prevented, or the therapy or treatment is modulated.

In an example, the engineered nucleotide sequence comprises a HM-CRISPR array. In an example, the engineered nucleotide sequence encodes a single guide RNA. In an example, the engineered nucleotide sequence is a guide RNA (eg, a single guide RNA) or crRNA. In an example, the engineered sequence is comprised by a bacteriophage that is capable of infecting the host cells, wherein the transformation comprises transduction of the host cells by the bacteriophage. The bacteriophage can be a phage as described herein. In an example, the engineered nucleotide sequence is comprised by a plasmid (eg, a conjugative plasmid) that is capable of transforming host cells. The plasmid can be a plasmid as described herein. In an example, the engineered nucleotide sequence is comprised by a transposon that is capable of transfer into and/or between host cells. The transposon can be a transposon as described herein.

Any method of the invention can comprise transforming host cells with nucleic acid vectors for producing cRNA or gRNA in the cells. For example, the vectors or nucleic acid comprising the engineered nucleotide sequence are administered orally, intravenously, topically, ocularly, intranasally, by inhalation, by rectal administration, or by any other route of administration disclosed herein or otherwise to the patient, wherein the administration transforms the first (host) cells with the vectors or nucleic acid.

In an example, the first are mixed with second bacteria in the microbiota of the patient or subject. Optionally, the second bacteria species is E coli, L lactis or S thermophilus, as shown in the worked Example below, such are strains that co-exist symbiotically in human and animal gut microbiota. The Example also addresses targeting in a mixed gram positive and gram negative bacterial population. Additionally, the Example addresses a population of *Firmicutes* (S thermophilus) and a population of Enterobacteriaceae (E coli), both of which are found in human microbiota. Other examples of Enterobacteriaceae are Salmonella, Yersinia pestis, Klebsiella, Shigella, Proteus, Enterobacter, Serratia, and Citrobacter.

In an example, the condition or disease is a metabolic or gastrointestinal disease or condition, eg, obesity, IBD, IBS, Crohn's disease or ulcerative colitis. In an example, the condition or disease is a cancer, eg, a solid tumour or a GI cancer (eg, stomach cancer), liver cancer or pancreatic cancer. In an example, the condition is resistance or reduced responsiveness to an antibiotic (eg, any antibiotic disclosed herein).

In an example, each first (host) cell comprises an endogenous RNase III that is operable with the engineered sequence in the production of said HM-crRNA in the cell. In an alternative, one or more of the vectors comprises a nucleotide sequence encoding such a RNase III for expression of the RNase III in the host cell.

In an example, the essential gene (comprising the target) encodes a DNA polymerase of the cell. This is exemplified below.

In an example, the cRNA or gRNA comprises a sequence that is capable of hybridising to a host cell target protospacer sequence that is a adjacent a NGG, NAG, NGA, NGC, NGGNG, NNGRRT or NNAGAAW protospacer adjacent motif (PAM), eg, a AAAGAAA or TAAGAAA PAM (these sequences are written 5' to 3'). In an embodiment, the PAM is immediately adjacent the 3' end of the protospacer sequence. In an example, the Cas is a S aureus, S theromophilus or S pyogenes Cas. In an example, the Cas is Cpf1 5 and/or the PAM is TTN or CTA.

In an example, each engineered nucleotide sequence or vector comprises a said CRISPR array or a sequence encoding a said crRNA or gRNA and further comprises an antibiotic resistance gene (eg, kanamycin resistance), 10 wherein the HM-crRNA or gRNA does not target the antibiotic resistance gene. In an example, the target sequence is comprised by an antibiotic resistance gene of the host cell, wherein the antibiotic is different from the first antibiotic (eg, kanamycin). In this way, the engineered sequence or 15 vector is able to target the host without targeting itself. By exposing the host cells to the first antibiotic (eg, by orally or intravenously administering it to the patient), one can promote retention of the engineered sequence or vector therein by positive selection pressure since cells containing the first 20 antibiotic resistance gene will have a survival advantage in the presence of the first antibiotic (when host cells that are not transformed by the engineered sequence or vectors are not resistant to the first antibiotic). Thus, an example provides: The method of the invention comprising exposing the 25 first (host) cell population to said antibiotic (eg, kanamycin) and said engineered sequence or vector(s), for promoting maintenance of cRNA or gRNA-encoding sequences in host cells; or the engineered sequence, array or vector of the invention is in combination with said antibiotic.

In an example the sequence encoding the cRNA or gRNA is under a constitutive promoter (eg, a strong promoter) operable in the host cell species, or an inducible promoter.

In an example, the or each first (host) cell is a gram positive cell. In another example, the or each first (host) cell 35 pathogens of humans, an animal (eg, non-human animal) or is a gram positive cell.

The invention also provides: An ex vivo mixed population of microbiota bacteria obtainable by the method by isolation of a gut microbiota sample from the patient after carrying out the method, or by isolation of a faecal sample of the 40 patient after carrying out the method. In an example, the mixed population is in a container for medical or nutritional use. For example, the container is a sterilised container, eg, an inhaler, intranasal delivery device or connected to a syringe or IV needle. In an example, the mixed population 45 is useful for administration to a human or animal to populate a microbiome thereof for treating a disease or condition (eg. a disease or condition disclosed herein).

Herein, in an example the Bacteroides is a species selected from caccae, capillosus, cellulosilyticus, copro- 50 cola, coprophilus, coprosuis, distasonis, dorei, eggerthii, faecis, finegoldii, fluxus, fragalis, intestinalis, melaninogenicus, nordii, oleiciplenus, oralis, ovatus, pectinophilus, plebeius, stercoris, thetaiotaomicron, uniformis, vulgatus and xylanisolvens. For example, the Bacteroides is thetaiotao- 55 micron. In an example, the first (host cell) population or second bacteria comprise a plurality of different Bacteroidetes species, or a plurality of Bacteroides species (eg, comprising B thetaiotaomicron and B fragalis), or Bacteroides and Prevotella species. Herein, in an example, the 60 Prevotella is a species selected from bergensis, bivia, buccae, buccalis, copri, melaninogenica, oris, ruminicola, tannerae, timonensis and veroralis. In an alternative, the first (host) cells or second bacteria are Firmicutes cells, for example comprising or consisting of one or more Firmicutes 65 selected from Anaerotruncus, Acetanaerobacterium, Acetitomaculum, Acetivibrio, Anaerococcus, Anaerofilum,

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Anaerosinus, Anaerostipes, Anaerovorax, Butyrivibrio, Clostridium, Capracoccus, Dehalobacter, Dialister, Dorea, Enterococcus, Ethanoligenens, Faecalibacterium, Fusobacterium, Gracilibacter, Guggenheimella, Hespellia, Lachnobacterium, Lachnospira, Lactobacillus, Leuconostoc, Megamonas, Mitsuokella, Oribacterium, Oxobacter, Papillibacter, Proprionispira, Pseudobutyrivibrio, Pseudoramibacter, Roseburia, Ruminococcus, Sarcina, Seinonella, Shuttleworthia, Sporobacter, Sporobacterium, Streptococcus, Subdoligranulum, Syntrophococcus, Thermobacillus, Turibacter and Weisella. In an example, the first (host) cells or second bacteria comprise or consist of Clostridium (eg, dificile) cells (and optionally the other sub-population consists of Bacteroides (eg, thetaiotaomicron) cells). In an example, the first (host) cells or second bacteria comprise or consist of Enterococcus cells (and optionally the other cells consist of Bacteroides (eg, thetaiotaomicron) cells). In an example, the first (host) cells or second bacteria comprise or consist of Ruminococcus cells (and optionally the other cells consist of Bacteroides (eg. thetaiotaomicron) cells). In an example, the first (host) cells or second bacteria comprise or consist of Streptococcus cells (and optionally the other cells consist of Bacteroides (eg, thetaiotaomicron) cells). In an example, the first (host) cells or second bacteria comprise or consist of *Faecalibacterium* cells (and optionally the other cells consist of Bacteroides (eg, thetaiotaomicron) cells). For example, the Faecalibacterium is a Faecalibacterium prausnitzii (eg, A2-165, L2-6, M21/2 or SL3/3).

In an example, the first (host) cells or second bacteria consist of Streptococcus cells (optionally S thermophilus and/or *pyogenes* cells) and the second bacteria consists of Bacteroides (eg, thetaiotaomicron) and/or Enterobacteriaceae (eg, E coli) cells.

In an example, the first (host) cells are infectious disease a plant.

In an example, the first (host) cells are selected from a species of Escherichia (eg, E coli O157:H7 or 0104: H4), Shigella (eg, dysenteriae), Salmonella (eg, typhi or enterica, eg, serotype typhimurium, eg, DT 104), Erwinia, Yersinia (eg, pestis), Bacillus, Vibrio, Legionella (eg, pneumophilia), Pseudomonas (eg, aeruginosa), Neisseria (eg, gonnorrhoea or meningitidis), Bordetella (eg, pertussus), Helicobacter (eg, pylori), Listeria (eg, monocytogenes), Agrobacterium, Staphylococcus (eg, aureus, eg, MRSA), Streptococcus (eg, pyogenes or thermophilus), Enterococcus, Clostridium (eg, dificile or botulinum), Corynebacterium (eg, amycolatum), Mycobacterium (eg, tuberculosis), Treponema, Borrelia (eg, burgdorferi), Francisella, Brucella, Campylobacter (eg, jejuni), Klebsiella (eg, pneumoniae), Frankia, Bartonella, Rickettsia, Shewanella, Serratia, Enterobacter, Proteus, Providencia, Brochothrix, Bifidobacterium, Brevibacterium, Propionibacterium, Lactococcus, Lactobacillus, Pediococcus, Leuconostoc, Vibrio (eg, cholera, eg, O139, or vulnificus), Haemophilus (eg, influenzae), Brucella (eg, abortus), Franciscella, Xanthomonas, Erlichia (eg, chaffeensis), Chlamydia (eg, pneumoniae), Parachlamydia, Enterococcus (eg, faecalis or faceim, eg, linezolid-resistant), Oenococcus and Acinetoebacter (eg, baumannii, eg, multiple drug resistant). In an example, the first (host) cells are Staphylococcus

aureus cells, eg, resistant to an antibiotic selected from methicillin, vancomycin-resistant and teicoplanin.

In an example, the first (host) cells are Pseudomonas aeuroginosa cells, eg, resistant to an antibiotic selected from cephalosporins (eg, ceftazidime), carbapenems (eg, imipenem or meropenem), fluoroquinolones, aminoglycosides (eg, gentamicin or tobramycin) and colistin.

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In an example, the first (host) cells are *Klebsiella* (eg, *pneumoniae*) cells, eg, resistant to carbapenem.

In an example, the first (host) cells are *Streptoccocus* (eg, *pneumoniae* or *pyogenes*) cells, eg, resistant to an antibiotic selected from erythromycin, clindamycin, beta-lactam, mac-⁵ rolide, amoxicillin, azithromycin and penicillin.

In an example, the first (host) cells are *Salmonella* (eg, serotype *Typhi*) cells, eg, resistant to an antibiotic selected from ceftriaxone, azithromycin and ciprofloxacin.

In an example, the first (host) cells are *Shigella* cells, eg, ¹⁰ resistant to an antibiotic selected from ciprofloxacin and azithromycin.

In an example, the first (host) cells are *Mycobacterium tuberculosis* cells, eg, resistant to an antibiotic selected from Resistance to isoniazid (INH), rifampicin (RMP), fluoroquinolone, amikacin, kanamycin and capreomycin.

In an example, the first (host) cells are *Enterococcus* cells, eg, resistant to vancomycin.

The method of claim 13, wherein all of the host cells are $_{20}$ Enterobacteriaceae cells, eg, resistant to an antibiotic selected from a cephalosporin and carbapenem.

In an example, the first (host) cells are *E. coli* cells, eg, resistant to an antibiotic selected from trimethoprim, itro-furantoin, cefalexin and amoxicillin.

In an example, the first (host) cells are *Clostridium* (eg, *dificile*) cells, eg, resistant to an antibiotic selected from fluoroquinolone antibiotic and carbapenem.

In an example, the first (host) cells are *Neisseria gonnorrhoea* cells, eg, resistant to an antibiotic selected from 30 cefixime (eg, an oral cephalosporin), ceftriaxone (an injectable cephalosporin), azithromycin and tetracycline.

In an example, the first (host) cells are *Acinetoebacter baumannii* cells, eg, resistant to an antibiotic selected from beta-lactam, meropenem and a carbapenem.

In an example, the first (host) cells are *Campylobacter* cells, eg, resistant to an antibiotic selected from ciprofloxacin and azithromycin.

In an example, the second species or strain is a human gut commensal species or strain and/or a human gut probiotic 40 species or strain.

In an example, the second species or strain is a Bacteroidetes (eg, *Bacteroides*) and optionally the host cells are gram positive bacterial cells.

In an example, the first cells are *Firmicutes* cells.

In an example, causing said dysbiosis or step (b) is carried out by targeting the sub-population of first cells by administering thereto an anti-bacterial or anti-archaeal agent simultaneously or sequentially with said immune cell population, whereby first cells are killed or the sub-population 50 growth is reduced, thereby reducing the proportion of said sub-population in the gut microbiota of the patient.

In an example, the method reduces first (host) cell population growth by at least 5, 10, 20, 50 or 100-fold compared to the growth of a control population of host cells that have 55 not received said guided nuclease (eg, Cas) modification.

In an example, the method inhibits host cell population growth on a gut surface.

In an example, for each host cell the system comprises components according to (i) to (iv):—

- (i) at least one nucleic acid sequence encoding a Cas nuclease;
- (ii) an engineered HM-CRISPR array comprising a spacer sequence and repeats encoding a HM-crRNA, the HMcrRNA comprising a sequence that hybridises to a host 65 cell target sequence to guide said Cas to the target in the host cell to modify the target sequence;

- (iii) an optional tracrRNA sequence or a DNA sequence expressing a tracrRNA sequence;
- (iv) wherein said components of the system are split between the host cell and at least one nucleic acid vector that transforms the host cell, whereby the HMcrRNA guides Cas to the target to modify the host target sequence in the host cell; and
- wherein the target sequence is modified by the Cas whereby the host cell is killed or host cell growth is reduced;
- the method comprising introducing the vectors of (iv) into host cells and expressing said HM-crRNA in the host cells, allowing HM-cRNA to hybridise to host cell target sequences to guide Cas to the targets in the host cells to modify target sequences, whereby host cells are killed or host cell growth is reduced, thereby altering the proportion of said sub-population in the microbiota. In an example, component (i) is endogenous to each host

cell.

In an example, each vector is a virus or phage.

In an example, each target sequence is adjacent a NNA-GAAW or NGGNG protospacer adjacent motif (PAM).

In an example, alternatively HM-crRNA and tracrRNA are comprised by a single guide RNA (gRNA), the method comprising introducing said gRNA into host cells or expressing the gRNA in host cells.

In an example, the microbiota comprises a second bacterial or archaeal species, wherein each of the first and second species is a respective species of the same phylum (eg, both *Firmicutes* species) and the growth of the second bacteria is not inhibited by the HM-system; or wherein the microbiota comprises a second bacterial or archaeal strain, wherein each of the first and second bacteria or archaea is a respective strain of the same species and the growth of the second bacteria or archaea is not inhibited by the HM-system.

In an example, the microbiota comprises a second bacterial species, wherein each of the first and second species is a respective gram-positive species and the growth of the second bacteria is not inhibited by the HM-system.

In an example, each target sequence is comprised by an antibiotic resistance gene, virulence gene or essential gene of the host cell.

In an example, each first cell is a *Staphylococcus, Streptococcus, Pseudomonas, Salmonella, Listeria, E coli, Des-*45 *ulfovibrio, Vibrio* or *Clostridium* cell.

In an example, the dysbiosis or step (b) comprises stimulating Paneth cells of the patient by gut *Bacteroides* (eg, *B thetaiotamicron*), wherein the altered microbiota produced by the dysbiosis or step (b) comprises an increased proportion of *Bacteroides* first bacteria compared with the microbiota before the dysbiosis or step (b), whereby Paneth cells are stimulated and the cell therapy is modulated.

Bacteroides can affect expression of Paneth cell proteins. Small intestinal crypts house stem cells that serve to constantly replenish epithelial cells that die and are lost from the villi. Paneth cells (immune systems cells similar to neutrophils), located adjacent to these stem cells, protect them against microbes by secreting a number of antimicrobial molecules (defensins) into the lumen of the crypt, and it is possible that their protective effect even extends to the mature cells that have migrated onto the villi. In animal models, *B. thetaiotaomicron* can stimulate production of an antibiotic Paneth cell protein (Ang4) that can kill certain pathogenic organisms (e.g., *Listeria monocytogenes*). *Listeria monocytogenes* is a strong T_{H1} cell inducer, and thus by stimulating Paneth cells in certain aspects of the invention, this may be useful to skew immunity from TH1 to other cell

types, such as $T_H 17$. This may be useful for increasing or enhancing the immune cell therapy of the invention. For example, this may be useful when the invention comprises administering $T_H 17$ -based cell therapy (eg, CAR- $T_H 17$) cells) to the patient.

In an example, the dysbiosis or step (b) comprises developing an immune response in the patient to gut Bacteroides (eg, B thetaiotamicron), wherein the altered microbiota produced by the dysbiosis or step (b) comprises an increased proportion of Bacteroides first bacteria compared with the 10 microbiota before the dysbiosis or step (b), whereby the cell therapy is modulated.

In an example, the dysbiosis or step (b) comprises altering the relative proportion of a or said sub-population of first cells in the gut microbiota of the patient, thereby producing 15 an altered gut microbiota that modulates the immune cell therapy in the patient, wherein the dysbiosis or step (b) comprises killing first cells of said sub-population or inhibiting growth of said sub-population by using guided nuclease targeting to the genome of first cells comprised by the 20 sub-population.

The invention also provides: A bacterial or archaeal transplant for administration to a patient for therapy of a disease or condition in the patient using the method of the invention. Optionally the transplant comprises cells of said 25 first species. Optionally the transplant comprises cells of said second species (and eg, does not comprise cells of the first species).

The invention also provides: A bacterial or archaeal transplant for administration to a subject (eg, a plant or 30 yeast) or to soil or an environment for modulating a treatment thereof using the method of the invention. Optionally the transplant comprises cells of said first species. Optionally the transplant comprises cells of said second species (and eg, does not comprise cells of the first species).

In an example, the yeast comprises a population of yeast cells of the same species or strain. For example, the yeast is comprised by the microbiota, eg, the yeast comprises a sub-population of cells of the microbiota, such as where the targeted cells are bacteria or archaea cells also comprised by 40 the microbiota. In an embodiment, the method kills the targeted cells (eg, bacteria) or reduces growth of the target cell sub-population comprised by the microbiota, wherein release of (or the concentration of) one or more chemicals or messengers by target cells in the microbiota is reduced. For 45 example, the one or more chemicals or messengers that mediate quorum sensing in microbes (eg, bacteria) of the microbiota is reduced. This may be useful to shape the growth of target cell and/or other microbial cell populations in the microbiota. In an example, the one or more chemicals 50 or messengers inhibit growth of the yeast or kill the yeast, and thus reduction of the release or concentration of the chemical or messenger(s) in the microbiota is advantageous to promote growth and/or maintenance of the yeast in the microbiota. In an embodiment of these examples, the treat- 55 ment is nutrition of the yeast or treatment of the yeast with a growth promoter (eg, wherein the yeast are for foodstuff or beverage production, or for production of a biotechnology or medical product, such as an antibody eg, the yeast is Saccharaomyces). In another example, the chemical or 60 messenger(s) promote yeast growth, wherein the yeast treatment is treatment with a yeast killing agent (eg, an fungicide). This is useful to promote efficacy of the killing treatment when the yeast are undesirable (eg, an undesirable mould). In an example when the subject is a plant, the one 65 or more chemicals or messengers inhibit growth of the plant or kill the plant, and thus reduction of the release or

concentration of the chemical or messenger(s) in the microbiota is advantageous to promote growth of the plant and/or inhibit killing of the plant. In an embodiment of these examples, the treatment is nutrition of the plant or treatment of the yeast with a fertilizer or nitrogen fixing agent. In an embodiment, the treatment is a pesticide treatment of the plant, such as a treatment that is inhibited or reduced in the presence of the targeted microbiota cells.

In an example, the plant is a crop plant (eg, wheat, barley, cereal or livestock fodder plant), fruit plant (eg, apple, orange, citrus, lime, lemon, raspberry, strawberry, berry or banana plant), legume plant, sugar cane plant, spice plant, garden plant, vegetable plant, grass, tree or flowering plant. For example, the plant is a tuber plant, eg, a potato or a sweet potato (eg, and the first, host or targeted bacteria are Pectobacterium atrosepticum cells, optionally wherein the treatment is storage (eg, cold storage), washing, a pesticide or herbicide treatment, fertilising or hydrating of the plant of a crop thereof (eg, a potato crop)). For example, the plant is a tobacco plant (eg, and the first, host or targeted bacteria are Ralstonia solanacearum cells, optionally wherein the treatment is storage (eg, cold storage), washing, a pesticide or herbicide treatment, fertilising or hydrating of the plant of a crop thereof (eg, a tobacco leaf crop)).

In an example, the subject is a protozoa. In an example, the subject or patient is a fish. In an example, the subject or patient is a bird. In an example, the subject or patient is a reptile. In an example, the subject or patient is an arachnid. In an example, the subject is a yeast cell (eg, a Saccharomyces cell). In an example, the subject or patient is an animal (eg, a rodent, mouse or rat). In an example, the subject or patient is a human (eg, an embryo or not an embryo). In an example, the subject or patient is a companion pet animal (eg, a bird, horse, dog, cat or rabbit). In an example, the subject or patient is an insect (an insect at any stage of its lifecycle, eg, egg, larva or pupa, eg, a fly or crawling insect or a beetle). In an example, the subject or patient is a cephalopod or nematode. In an example, the subject or patient is a plant or animal pest. In an example, the treatment of an animal or human may be a nutritional treatment, therapy of a disease or condition, prophylais of a disease or condition, ocular treatment, pesticide treatment, dental treatment, topical treatment or digestion treatment. In an example, the method enhances immunity of the subject or patient against a pathogen (eg a pathogenic parasite, protozoan, virus or bacteria). In an example, the treatment or therapy is a combination treatment or therapy practised on the human or animal, wherein the human or animal is administered a first medicament in combination with a second medicament or radiation. The first and/or second medicament may be an antibody therapy or immune cell transplant (eg, CAR-T or TILs transplant) therapy (and the immune modulation aspect of the invention may be advantageous for modulating such therapies). In an example, each medicament or treatment is selected from Table 2.

In an example, the disease or condition is diabetes (eg. Type I or II diabetes, insulin-resistant diabetes (eg, insulinresistant Type II diabetes), onset of insulin resistance in a diabetic or pre-diabetic patient or reduction in insulin responsiveness in a diabetic or pre-diabetic patient. Optionally additionally in this example, the first or host cells that are targeted are Prevotella copri or Bacteroides vulgatus cells. In an embodiment, both Prevotella copri and Bacteroides vulgatus cells are targeted (ie, killed and/or population growth reduced, or reduced in the microbiota following

administration of a transplant as described herein) in the patient, wherein said disease or condition is treated or prevented.

The invention also provides: the HM-CRISPR/Cas system, HM-array or HM-crRNA for administration to a patient 5 for therapy of a disease or condition in the patient using the method of the invention.

The invention also provides: A kit comprising an ex vivo population of immune cells for adoptive cell therapy of a patient, wherein the kit further comprises the transplant, 10 system, array or crRNA, optionally wherein the immune cells are selected from CAR-T cells, T-cells expressing engineered T-cell receptors (TCRs), tumour infiltrating lymphocytes (TILs) and NK cells.

Mobile Genetic Elements (MGEs)

In an example, each vector is a nucleic acid vector comprising or consisting of a mobile genetic element (MGE), wherein the MGE comprises an origin of transfer (oriT) and a CRISPR array for modifying a target sequence of the genome of a host cell or the genome of a virus (eg, 20 prophage) in a host cell. Examples of MGEs are ICEs, transposons, plasmids and bacteriophage. An origin of transfer (oriT) is a short sequence (eg, up to 500 bp) that is necessary for transfer of the DNA that contains it from a bacterial host to recipient during conjugation. The oriT is 25 cis-acting—it is found on the same DNA that is being transferred, and it is transferred along with the DNA. A typical origin of transfer comprises three functionally defined domains: a nicking domain, a transfer domain, and a termination domain. 30

Reference is made to the ICEberg database (http://db-mml sjtu.edu.cn/ICEberg/), which provides examples of suitable ICEs for the invention and sources for suitable oriT. In an example, the ICE is a member of an ICE family comprising an ICE selected from the group 1 to 28, or the oriT is an oriT 35 of a member of such a family: 1=SXT/R391; 2=Tn916; 3=Tn4371; 4=CTnDOT/ERL; 5=ICEclc; 6=ICEBs1; 7=ICEHin1056; 8=PAPI-1; 9=ICEM1Sym(R7A); 11=SPI-7; 12=ICE6013; 10=ICESt1: 13=ICEKp1; 14=TnGBS1; 15=Tn5253; 16=ICESa2603; 17=ICEYe1; 40 18=10270-RD.2: 19=Tn1207.3; 20=Tn1806; 21=ICEA5632; 22=ICEF-I/II; 23=ICEAPG2; 24=ICEM; 25=10270-RD.1; 26=Tn5801; 27=PPI-1; 28=ICEF-III. Family descriptions are found in the ICEberg database. For example, the Tn916 family was defined by Roberts et al 45 (2009) (Trends Microbiol. 2009 June; 17(6):251-8. doi: 10.1016/j.tim.2009.03.002. Epub 2009 May 20; "A modular master on the move: the Tn916 family of mobile genetic elements", Roberts A, Mullany P). Elements belonging to the Tn916 family are defined by the following criteria: they 50 must have the general organization shown in Roberts et al, and they must have a core region (conjugation and regulation module) that is similar in sequence and structure to the original Tn916 at the DNA level. Exceptions are some conjugative transposons, such as Tn1549 which have been 55 previously classified in this family and those with a high degree of protein similarity as described in corresponding references. Optionally, the ICE is a transposon, eg, a conjugative transposon. In an example, the MGE is a mobilisable transposon that is mobilisable in the presence of a 60 functional helper element, optionally wherein the transposon is in combination with a said helper element.

Optionally the vector is a plasmid, optionally wherein the MGE is a transposon comprised by the plasmid. For example, the transposon is a conjugative transposon. In an 65 example the transposon is a mobilisable transposon (eg, mobilisable using one or more factors encoded by the

plasmid, eg, by genes outside the transposon sequence of the plasmid). Optionally, the transposon is a Type I transposon. Optionally, the transposon is a Type II transposon. Optionally, the vector oriT is an oriT of a Bacteroidetes (eg, Bacteroidales or *Bacteroides*) or *Prevotella* transposon. This useful when the first (host) cells are Bacteroidetes (eg, Bacteroidales or *Bacteroides*) or *Prevotella* respectively. Optionally, the vector oriT is a CTnDot, CTnERL SXT/R391, Tn916 or Tn4371 family transposon oriT.

Optionally, the method comprises exposing the patient's microbiota to a vector or MGE (eg, one described above) that comprises a toxin-antioxin module that comprises an anti-toxin gene that is operable in the second bacteria, but is not operable or has reduced operation in first (host) cells. Thus, first cells are killed and second bacteria are spared, thereby altering the proportion of first cells in the patient's microbiota.

Split CRISPR/Cas System

In one aspect, endogenous Cas of the first (host) cells is harnessed and operates with the engineered sequences comprised by vectors (eg, phage) that are introduced into host cells. This aspect is advantageous to free up space in vectors, for example viruses or phage that have restricted capacity for carrying exogenous sequence. By freeing up space, one is able to include more targeting spacers or arrays, which is useful for evading host resistance. It is advantageous, for example to harness the endogenous Cas endonuclease rather than encode it in the vector-especially for bulky Cas sequences such as sp or saCas9. Additionally, there is not chance of inferior compatibility as may be seen with some exogenous Cas from non-host sources. The ability to reduce virus, eg, phage genome size, may also be beneficial for promoting host cell uptake (infection and/or maintenance of the virus in host cells). In some examples, an advantage is that invasion of the host by the vector (eg, phage) may upregulate host CRISPR/Cas activity, including increased expression of host Cas nucleases-in an attempt of the host to combat invading nucleic acid. This, however, is also useful to provide endogenous Cas for use with the invention when these use cRNA or gRNA that are recognised by the host Cas. In the case where the invention involves cRNA or gRNA targeting a host CRISPR array, this then promotes inactivation of the host CRISPR array itself, akin to a "suicidal" host cell which then uses its own Cas nuclease to inactivate its own CRISPR systems.

Thus, the vectors may lack a Cas nuclease (eg, aCas9)encoding sequence.

Optionally, the endogenous first (host) cell system is a CRISPR/Cas9 system. Optionally, the nuclease is a Type I Cas nuclease. Optionally, the nuclease is a Type II Cas nuclease (eg, a Cas9). Optionally, the nuclease is a Type III Cas nuclease.

To save even more space, optionally a tracrRNA sequence is not provided by the vectors, but is a tracrRNA sequence of an endogenous host cell CRISPR/Cas system, wherein the tracrRNA is capable of hybridising with the HM-crRNA in the cell for subsequent processing into mature crRNA for guiding Cas to the target in the host cell.

Generally Applicable Features

The following features apply to any configuration (eg, in any of its aspects, embodiments, concepts, paragraphs or examples) of the invention:—

In an example, the target sequence is a chromosomal sequence, an endogenous host cell sequence, a wild-type host cell sequence, a non-viral chromosomal host cell sequence and/or a non-phage sequence (ie, one more or all of these), eg, the sequence is a wild-type host chromosomal cell sequence such as antibiotic resistance gene or essential gene sequence comprised by a host cell chromosome. In an example, the sequence is a host cell plasmid sequence, eg, an antibiotic resistance gene sequence.

In an example, at least two target sequences are modified 5 by Cas, for example an antibiotic resistance gene and an essential gene. Multiple targeting in this way may be useful to reduce evolution of escape mutant host cells.

In an example, the Cas is a wild-type endogenous host cell Cas nuclease. In an example, each host cell has constitutive 10 Cas nuclease activity, eg, constitutive wild-type Cas nuclease activity. In an example, the host cell is a bacterial cell; in another example the host cell is an archael cell. Use of an endogenous Cas is advantageous as this enables space to be freed in vectors cRNA or gRNA. For example, Type II Cas9 15 nucleotide sequence is large and the use of endogenous Cas of the host cell instead is advantageous in that instance when a Type II CRISPR/Cas system is used for host cell modification in the present invention. The most commonly employed Cas9, measuring in at 4.2 kilobases (kb), comes 20 from S pyogenes. While it is an efficient nuclease, the molecule's length pushes the limit of how much genetic material a vector can accommodate, creating a barrier to using CRISPR in the tissues of living animals and other settings described herein (see F. A. Ran et al., "In vivo 25 genome editing using Staphylococcus aureus Cas9," Nature, doi:10.1038/nature14299, 2015). Thus, in an embodiment, the vector of the invention is a AAV vector or has an exogenous DNA insertion capacity no more than an AAV vector, and the Cas is an endogenous Cas of the host cell, 30 wherein the cell is a bacterial or archaeal cell. S thermophilus Cas9 (UniProtKB-G3ECR1 (CAS9_STRTR)) nucleotide sequence has a size of 1.4 kb.

In an example, the vector is a viral vector. Viral vectors have a particularly limited capacity for exogenous DNA 35 insertion, thus virus packaging capacity needs to be considered. Room needs to be left for sequences encoding vital viral functions, such as for expressing coat proteins and polymerase. In an example, the vector is a phage vector or an AAV or lentiviral vector. Phage vectors are useful where 40 the host is a bacterial cell.

By use of the term "engineered" it will be readily apparent to the skilled addressee that the array, sequence, vector, cRNA, gRNA, MGE or any other configuration, concept, aspect, embodiment, paragraph or example etc of the invention is non-naturally occurring. For example, the MGE, vector, sequence or array comprises one or more sequences or components not naturally found together with other sequences or components of the MGE, vector, sequence or array. For example, the array or sequence is recombinant, 50 artificial, synthetic or exogenous (ie, non-endogenous or not wild-type) to the or each host cell.

In an example, the array or sequence of the invention is an engineered version of an array or sequence isolated, for example isolated from a host cell. In an example, the 55 engineered array or sequence is not in combination with a Cas endonuclease-encoding sequence that is naturally found in a cell.

Studies suggest that *Bacteroides* have a role in preventing infection with *Clostridium difficile*. The development of the 60 immune response that limits entry and proliferation of potential pathogens is profoundly dependent upon *B fragilis*. Also, Paneth cell proteins may produce antibacterial peptides in response to stimulation by *B thetaiotomicron*, and these molecules may prevent pathogens from colonizing the 65 gut. In addition, *B thetaiotomicron* can induce Paneth cells to produce a bactericidal lectin, RegIII, which exerts its

antimicrobial effect by binding to the peptidoglycan of gram-positive organisms. Thus, the use of the invention in any of its configurations for increasing the proportion of *Bacteroides* (eg, *thetaiotomicron* and/or *fragalis*) in the patient's microbiota is useful for limiting pathogenic bacterial colonisation of the population or a gut of a human or non-human animal.

Hooper et al demonstrated that *B thetaiotomicron* can modify intestinal fucosylation in a complex interaction mediated by a fucose repressor gene and a signaling system. Using transcriptional analysis it was demonstrated that *B thetaiotaomicron* can modulate expression of a variety of host genes, including those involved in nutrient absorption, mucosal barrier fortification, and production of angiogenic factors.

Optionally, the host (or first and/or second bacteria) is a gram negative bacterium (eg, a spirilla or vibrio). Optionally, the host (or first and/or second bacteria) is a gram positive bacterium. Optionally, the host (or first and/or second bacteria) is a mycoplasma, chlamydiae, spirochete or Mycobacterium. Optionally, the host (or first and/or second bacteria) is a Streptococcus (eg, pyogenes or thermophilus) host. Optionally, the host (or first and/or second bacteria) is a Staphylococcus (eg, aureus, eg, MRSA) host. Optionally, the host (or first and/or second bacteria) is an E. coli (eg, O157: H7) host, eg, wherein the Cas is encoded by the vecor or an endogenous host Cas nuclease activity is de-repressed. Optionally, the host (or first and/or second bacteria) is a Pseudomonas (eg, aeruginosa) host. Optionally, the host (or first and/or second bacteria) is a Vibro (eg, cholerae (eg, O139) or vulnificus) host. Optionally, the host (or first and/or second bacteria) is a Neisseria (eg, gonnorrhoeae or meningitidis) host. Optionally, the host (or first and/or second bacteria) is a Bordetella (eg, pertussis) host. Optionally, the host (or first and/or second bacteria) is a Haemophilus (eg, influenzae) host. Optionally, the host (or first and/or second bacteria) is a Shigella (eg, dysenteriae) host. Optionally, the host (or first and/or second bacteria) is a Brucella (eg, abortus) host. Optionally, the host (or first and/or second bacteria) is a Francisella host. Optionally, the host (or first and/or second bacteria) is a Xanthomonas host. Optionally, the host (or first and/or second bacteria) is a Agrobacterium host. Optionally, the host (or first and/or second bacteria) is a Erwinia host. Optionally, the host (or first and/or second bacteria) is a Legionella (eg, pneumophila) host. Optionally, the host (or first and/or second bacteria) is a Listeria (eg. monocytogenes) host. Optionally, the host (or first and/or second bacteria) is a Campylobacter (eg, jejuni) host. Optionally, the host (or first and/or second bacteria) is a Yersinia (eg, pestis) host. Optionally, the host (or first and/or second bacteria) is a Borelia (eg, burgdorferi) host. Optionally, the host (or first and/or second bacteria) is a Helicobacter (eg, pylori) host. Optionally, the host (or first and/or second bacteria) is a Clostridium (eg, dificile or botulinum) host. Optionally, the host (or first and/or second bacteria) is a Erlichia (eg, chaffeensis) host. Optionally, the host (or first and/or second bacteria) is a Salmonella (eg, typhi or enterica, eg, serotype typhimurium, eg, DT 104) host. Optionally, the host (or first and/or second bacteria) is a Chlamydia (eg, pneumoniae) host. Optionally, the host (or first and/or second bacteria) is a Parachlamydia host. Optionally, the host (or first and/or second bacteria) is a Corynebacterium (eg, amycolatum) host. Optionally, the host (or first and/or second bacteria) is a Klebsiella (eg, pneumoniae) host. Optionally, the host (or first and/or second bacteria) is a Enterococcus (eg, faecalis or faecim, eg,

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linezolid-resistant) host. Optionally, the host (or first and/or second bacteria) is a Acinetobacter (eg, baumannii, eg, multiple drug resistant) host.

A tracrRNA sequence may be omitted from a array or vector of the invention, for example for Cas systems of a 5 Type that does not use tracrRNA.

In an example, the Cas guided to the target is an exonuclease. Optionally a nickase as mentioned herein is a double nickase. An example of a nickase is a Cas9 nickase, ie, a Cas9 that has one of the two nuclease domains inactivated— 10 either the RuvC and/or HNH domain.

Mention herein of using vector DNA can also in an alternative embodiment apply mutatis mutandis to vector RNA where the context allows. For example, where the vector is an RNA vector. All features of the invention are 15 therefore in the alternative disclosed and to be read as "RNA" instead of "DNA" when referring to vector DNA herein when the context allows. In an example, the or each vector also encodes a reverse transcriptase.

In an example, the or each array or engineered nucleotide 20 sequence is provided by a nanoparticle vector or in liposomes.

In an example, the Cas is a Cas nuclease for cutting, dead Cas (dCas) for interrupting or a dCas conjugated to a transcription activator for activating the target.

In an example, the or each array or engineered sequence comprises an exogenous promoter functional for transcription of the crRNA or gRNA in the host.

In an embodiment the array or engineered sequence is contained in a virophage vector and the host is alternatively a virus which can infect a cell. For example, the host is a large virus that may have infected an amoeba cell. For example, the host is a Sputnik virus, Pithovirus, mimivirus, mamavirus, Megavirus or Pandoravirus, eg, wherein the host virus is in water. In an example of this embodiment, the 35 6. The method of concept 4 or 5, wherein $T_{H}17$ cells are invention is for water or sewage treatment (eg, purification, eg, waterway, river, lake, pond or sea treatment).

In an embodiment the or each vector or engineered sequence is or is comprised by a Φ NM1 phage, eg, wherein the host cell(s) is a S. aureus (eg, MRSA) cell.

For example the method is practised on a mammalian subject, eg, a human, rodent, mouse or rat. For example the method is practised on a vertebrate, reptile, bird or fish.

The cell population can be administered to the patient in one or more doses. For example, the method comprises 45 administering an antibacterial agent to cause said dysbiosis, or administering a bacterial transplant to the patient to cause said dysbiosis.

Wherein the method reduces the cell therapy, the therapy can be downregulated, dampened or switched off, eg, to 50 reduce or prevent an unwanted side-effect of the cell therapy (eg, a CAR-T therapy side effect in a human patient, such as CRS). Wherein the method increases the cell therapy, the therapy can be upregulated, enhance or switched on, eg, to enhance cytotoxicity against one target cells. 55

The method treats or prevents (ie, reduces the risk of) the disease or condition. This may be complete or partial treatment or prevention, ie, a reduction but not complete reduction of the disease/condition or symptoms thereof; or a reducing of the risk but not total prevention of the disease/ 60 condition or a symptom thereof. Similarly, the method treats or prevents (ie, reduces the risk of) an undesirable symptom of the disease or condition or the therapy (eg, CRS).

- Concepts:
- 1. A method of modulating an adoptive immune cell therapy 65 of a disease or condition in a patient, the method comprising

- a. Carrying out adoptive immune cell therapy in the patient, comprising administering a population of immune cells to the patient, wherein administration of said immune cells is capable of treating the disease or condition in the patient; and
- b. Causing bacterial microbiota (eg, gut microbiota) dysbiosis in the patient, whereby said dysbiosis modulates the immune cell therapy in the patient.
- 2. A method of modulating an adoptive immune cell therapy of a disease or condition in a patient, the method comprising
 - a. Carrying out adoptive immune cell therapy in the patient, comprising administering a population of immune cells to the patient, wherein administration of said immune cells is capable of treating the disease or condition in the patient; and
 - b. Altering the relative proportion of a sub-population of cells of a first bacterial species or strain, or archaeal species or strain, in a microbiota (eg, gut microbiota) of the patient, thereby producing an altered microbiota that modulates the immune cell therapy in the patient.
- 3. The method of concept 2, wherein step (b) is carried out by targeting the sub-population of first cells by administering thereto an anti-bacterial or anti-archaeal agent simultaneously or sequentially with said immune cell population, whereby first cells are killed or the subpopulation growth is reduced, thereby reducing the proportion of said sub-population in the microbiota of the patient.
- 30 4. The method of any preceding concept, wherein the cell therapy is an adoptive T-cell therapy.
 - 5. The method of concept 4, wherein cells selected from the group consisting of CD4⁺ T-cells, CD8⁺ T-cells, T_H1 cells and $T_H 17$ cells are administered to the patient in step (a).
 - modulated in the patient.
 - 7. The method of concept 4, 5 or 6, wherein T_{reg} cells are modulated in the patient.
 - 8. The method of any preceding concept, wherein the cell therapy is enhanced.
 - 9. The method of any preceding concept wherein $T_H 17$ cells are upregulated in the patient and/or T_{reg} cells are downregulated in the patient.
 - 10. The method of any preceding concept, wherein CD4⁺ T-cells are upregulated in the patient.
 - 11. The method of any preceding concept, wherein CD8⁺ T-cells are upregulated in the patient.
 - 12. The method of any preceding concept, wherein one or more of central memory T cells (T_{CM}) , effector memory T cells (T_{EM}), stem cell memory cells (T_{SCM}) and effector cells (T_{eff}) are upregulated in the patient, wherein the cells are comprised by the immune cell population administered in step (a) and/or are progeny thereof.
 - 13. The method of any one of concepts 1 to 7, wherein the cell therapy is reduced.
 - 14. The method of concept 13, wherein the method reduces or prevents the risk of cytokine release syndrome (CRS) in the patient.
 - 15. The method of any one of concepts 1 to 7, 13 and 14, wherein, $T_H 17$ cells are downregulated in the patient and/or T_{reg} cells are upregulated in the patient.
 - 16. The method of any one of concepts 1 to 7 and 13 to 15, wherein CD4⁺ T-cells are downregulated in the patient.
 - 17. The method of any one of concepts 1 to 7 and 13 to 16, wherein CD8⁺ T-cells are downregulated in the patient.
 - 18. The method of any one of concepts 1 to 7 and 13 to 17, wherein one or more of central memory T cells (T_{CM}) ,

effector memory T cells (T_{EM}) , stem cell memory cells (T_{SCM}) and effector cells (T_{eff}) are downregulated in the patient, wherein the cells are comprised by the immune cell population administered in step (a) and/or are progenv thereof.

- 19. The method of any preceding concept, wherein the immune cell population comprises CAR-T cells and/or T-cells expressing engineered T-cell receptors (TCRs) and/or tumour infiltrating lymphocytes (TILs).
- 20. The method of any preceding concept, wherein the immune cell population comprises engineered autologous or allogeneic immune cells, eg, T-cells, NK cells and/or TILs.
- 21. The method of concept 19 or 20, wherein the T-cells are 15 $CD4^+$ T-cells or T_H 17 cells.
- 22. The method of any preceding concept, wherein step (b) increases the proportion of Bacteroides (eg, B fragalis and/or *B* thetaiotamicron) in the microbiota.
- 23. The method of any one of concepts 1 to 21, wherein step 20 (b) decreases the proportion of *Bacteroides* (eg, *B fragalis* and/or B thetaiotamicron) in the microbiota.
- 24. The method of any preceding concept, wherein step (b) increases the proportion of Bacteroidetes to Firmicutes in the microbiota.
- 25. The method of any one of concepts 1 to 23, wherein step (b) decreases the proportion of Bacteroidetes to *Firmic*utes in the microbiota.
- 26. The method of any preceding concept, wherein step (b) reduces the proportion of one or more Clostridium species 30 or strain (eg, wherein each species is a cluster IV or XIVa Clostridium species) in the microbiota.
- 27. The method of any one of concepts 1 to 25, wherein step (b) increases the proportion of one or more Clostridium species or strain (eg, wherein each species is a cluster IV 35 or XIVa *Clostridium* species) in the microbiota.
- 28. The method of any preceding concept, wherein step (b) reduces the proportion of *Bifidobacterium* (eg, *B bifidum*) in the microbiota.
- 29. The method of any one of concepts 1 to 27, wherein step 40 (b) increases the proportion of *Bifidobacterium* (eg, Bbifidum) in the microbiota.
- 30. The method of any preceding concept, wherein step (b) comprises altering the relative proportion of a or said sub-population of first cells in the microbiota of the 45 patient, thereby producing an altered microbiota that modulates the immune cell therapy in the patient, wherein the sub-population comprises host cells of said first species or strain, the method comprising
 - a. combining the microbiota with multiple copies of 50 engineered nucleic acid sequences encoding host modifying (HM) crRNAs, and
 - b. expressing HM-crRNAs in host cells,

wherein each engineered nucleic acid sequence is operable with a Cas nuclease in a respective host cell to form a 55 40. The method of concept 38 or 39, wherein each vector is HM-CRISPR/Cas system and the engineered sequence comprises

- (iii) spacer and repeat sequences encoding a HM-crRNA; (iv) the HM-crRNA comprising a sequence that is capable
- Cas nuclease to the target sequence in the host cell; and optionally the HM-system comprises a tracrRNA sequence or a DNA sequence expressing a tracrRNA sequence; whereby HM-crRNAs guide Cas modification of host target sequences in host cells, whereby host cells are killed or the host cell population growth is reduced, thereby reducing the

proportion of said sub-population in the microbiota.

- 31. The method of concept 30, comprising using endogenous Cas nuclease of host cells for modification of target nucleotide sequences.
- 32. The method of concept 30 or 31, comprising reducing host cell population growth by at least 5-fold compared to the growth of a control population of host cells that have not received said Cas modification.
- 33. The method of any one of concepts 30 to 32, comprising inhibiting host cell population growth on a gut surface.
- 10 34. The method of any one of concepts 30 to 33, wherein the microbiota comprises cells of a second bacterial species or strain, or archaeal species or strain, wherein the second species or strain has a 16s ribosomal RNA-encoding DNA sequence that is at least 80% identical to an 16s ribosomal RNA-encoding DNA sequence of the host cell species or strain, wherein the growth of the second cells in the microbiota is not inhibited by said HM-system.
 - 35. The method of concept 34, wherein the second species or strain is a human gut commensal species or strain and/or a human gut probiotic species or strain.
 - 36. The method of concept 34 or 35, wherein the second species or strain is a Bacteroidetes (eg, Bacteroides) and optionally the host cells are gram positive bacterial cells.
 - 37. The method of any one of concepts 30 to 36, wherein the first cells are *Firmicutes* cells.
 - 38. The method of any one of concepts 30 to 37, wherein for each host cell the system comprises components according to (i) to (iv):-
 - (i) at least one nucleic acid sequence encoding a Cas nuclease:
 - (ii) an engineered HM-CRISPR array comprising a spacer sequence and repeats encoding a HM-crRNA, the HMcrRNA comprising a sequence that hybridises to a host cell target sequence to guide said Cas to the target in the host cell to modify the target sequence;
 - (iii) an optional tracrRNA sequence or a DNA sequence expressing a tracrRNA sequence;
 - (iv) wherein said components of the system are split between the host cell and at least one nucleic acid vector that transforms the host cell, whereby the HMcrRNA guides Cas to the target to modify the host target sequence in the host cell; and
 - wherein the target sequence is modified by the Cas whereby the host cell is killed or host cell growth is reduced:
 - the method comprising introducing the vectors of (iv) into host cells and expressing said HM-crRNA in the host cells, allowing HM-cRNA to hybridise to host cell target sequences to guide Cas to the targets in the host cells to modify target sequences, whereby host cells are killed or host cell growth is reduced, thereby altering the proportion of said sub-population in the microbiota.
 - 39. The method of concept 38, wherein component (i) is endogenous to each host cell.
 - a virus or phage.
 - 41. The method of any one of concepts 30 to 40, wherein each target sequence is adjacent a NNAGAAW or NGGNG protospacer adjacent motif (PAM).
- of hybridizing to a host cell target sequence to guide 60 42. The method of any one of concepts 30 to 41, wherein alternatively HM-crRNA and tracrRNA are comprised by a single guide RNA (gRNA), the method comprising introducing said gRNA into host cells or expressing the gRNA in host cells.
 - 65 43. The method of any one of concepts 30 to 35 and 37 to 42, wherein the microbiota comprises a second bacterial or archaeal species, wherein each of the first and second

species is a respective species of the same phylum (eg, both Firmicutes species) and the growth of the second bacteria is not inhibited by the HM-system; or wherein the microbiota comprises a second bacterial or archaeal strain, wherein each of the first and second bacteria or 5 archaea is a respective strain of the same species and the growth of the second bacteria or archaea is not inhibited by the HM-system.

44. The method of any one of concepts 30 to 43, wherein the microbiota comprises a second bacterial species, wherein each of the first and second species is a respective gram-positive species and the growth of the second bacteria is not inhibited by the HM-system.

- 45. The method of any one of concepts 30 to 44, wherein each target sequence is comprised by an antibiotic resis- 15 tance gene, virulence gene or essential gene of the host cell.
- 46. The method of any preceding concept, wherein each first cell is a Staphylococcus, Streptococcus, Pseudomonas, Salmonella, Listeria, E coli, Desulfovibrio, Vibrio or 20 Clostridium cell.
- 47. The method of any preceding concept, wherein step (b) comprises stimulating Paneth cells of the patient by gut Bacteroides (eg, B thetaiotamicron), wherein the altered microbiota produced by step (b) comprises an increased 25 proportion of Bacteroides first bacteria compared with the microbiota before step (b), whereby Paneth cells are stimulated and the cell therapy is modulated.
- 48. The method of any preceding concept, wherein step (b) comprises developing an immune response in the patient 30 to gut Bacteroides (eg, B thetaiotamicron), wherein the altered microbiota produced by step (b) comprises an increased proportion of Bacteroides first bacteria compared with the microbiota before step (b), whereby the cell therapy is modulated. 35
- 49. The method of any preceding concept, wherein step (b) comprises altering the relative proportion of a or said sub-population of first cells in the microbiota of the patient, thereby producing an altered microbiota that modulates the immune cell therapy in the patient, wherein 40 step (b) comprises killing first cells of said sub-population or inhibiting growth of said sub-population by using guided nuclease targeting to the genome of first cells comprised by the sub-population.
- 50. A bacterial or archaeal transplant for administration to a 45 patient for therapy of a disease or condition in the patient using the method of any preceding concept, optionally wherein the transplant comprises cells of said first species.
- recited in any one of concepts 30 to 45 for administration to a patient for therapy of a disease or condition in the patient using the method of any one of concepts 1 to 49.
- 52. A kit comprising an ex vivo population of immune cells for adoptive cell therapy of a patient, wherein the kit 55 further comprises a transplant, system, array or crRNA of concept 50 or 51, optionally wherein the immune cells are selected from CAR-T cells, T-cells expressing engineered T-cell receptors (TCRs), tumour infiltrating lymphocytes (TILs) and NK cells. 60 Aspects:
- 1. An ex vivo population of immune cells for use in a method of adoptive cell therapy of a patient for treating or preventing a disease or condition in the patient, the method comprising
 - a. Carrying out adoptive immune cell therapy in the patient, comprising administering cells of said popula-

tion to the patient, wherein administration of said immune cells is capable of treating the disease or condition in the patient; and

- b. Causing gut bacterial microbiota dysbiosis in the patient, whereby said dysbiosis modulates the immune cell therapy in the patient and said disease or condition is treated or prevented.
- 2. An ex vivo population of immune cells for use in a method of adoptive cell therapy of a patient for treating or preventing a disease or condition in the patient, the method comprising
 - a. Carrying out adoptive immune cell therapy in the patient, comprising administering cells of said population to the patient, wherein administration of said immune cells is capable of treating, the disease or condition in the patient; and
 - b. Altering the relative proportion of a sub-population of cells of a first bacterial species or strain, or archaeal species or strain, in the gut microbiota of the patient, thereby producing an altered gut microbiota that modulates the immune cell therapy in the patient.
- 3. The immune cell population of Aspect 2, wherein step (b) is carried out by targeting the sub-population of first cells by administering thereto an anti-bacterial or anti-archaeal agent (eg, a guided nuclease) simultaneously or sequentially with said immune cell population, whereby first cells are killed or the sub-population growth is reduced, thereby reducing the proportion of said sub-population in the gut microbiota of the patient.
- 4. The immune cell population of any preceding Aspect, wherein the cell therapy is an adoptive T-cell therapy.
- 5. The immune cell population of Aspect 4, wherein cells selected from the group consisting of CD4+ T-cells, CD8+ T-cells, $T_H 1$ cells and $T_H 17$ cells are administered to the patient in step (a).
- 6. The immune cell population of Aspect 4 or 5, wherein $T_H 17$ cells are modulated in the patient.
- 7. The immune cell population of Aspect 4, 5 or 6, wherein T_{reg} cells are modulated in the patient.
- 8. The immune cell population of any preceding Aspect, wherein the cell therapy is enhanced.
- 9. The immune cell population of any preceding Aspect wherein $T_H 17$ cells are upregulated in the patient and/or T_{reg} cells are downregulated in the patient.
- 10. The immune cell population of any preceding Aspect, wherein CD4⁺ T-cell are upregulated in the patient.
- 11. The immune cell population of any preceding Aspect, wherein CD8⁺ T-cells are upregulated in the patient.
- 51. A HM-CRISPR/Cas system, HM-array or HM-crRNA as 50 12. The immune cell population of any preceding Aspect, wherein one or more of central memory T cells (T_{CM}) , effector memory T cells (T_{EM}) , stem cell memory cells (T_{SCM}) and effector cells (T_{eff}) are upregulated in the patient, wherein the cells are comprised by the immune cell population administered in step (a) and/or are progeny thereof.
 - 13. The immune cell population of any one of Aspects 1 to 7, wherein the cell therapy is reduced.
 - 14. The immune cell population of Aspect 13, wherein the method reduces or prevents the risk of cytokine release syndrome (CRS) in the patient.
 - 15. The immune cell population of any one of Aspects 1 to 7, 13 and 14, wherein, T₁₃17 cells are downregulated in the patient and/or T_{reg} cells are upregulated in the patient.
 - 65 16. The immune cell population of any one of Aspects 1 to 7 and 13 to 15, wherein CD4+ T-cells are downregulated in the patient.

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- 17. The immune cell population of any one of Aspects 1 to 7 and 13 to 16, wherein CD8+ T-cells are downregulated in the patient.
- 18. The immune cell population of any one of Aspects 1 to 7 and 13 to 17, wherein one or more of central memory T cells (T_{CM}), effector memory T cells (T_{EM}), stem cell memory cells (T_{SCM}) and effector cells (T_{eff}) are downregulated in the patient, wherein the cells are comprised by the immune cell population administered in step (a) and/or are progeny thereof.
- 19. The immune cell population of any preceding Aspect, wherein the immune cell population comprises or consists of CAR-T cells and/or T-cells expressing engineered T-cell receptors (TCRs) and/or tumour infiltrating lym-15 32. The immune cell population of Aspect 30 or 31, com-
- 20. The immune cell population of any preceding Aspect, wherein the immune cell population comprises or consists of engineered autologous or allogeneic immune cells, eg, T-cells, NK cells and/or TILs.
- 21. The immune cell population of Aspect 19 or 20, wherein the T-cells are $CD4^{+T-cells or T}_{H}17$ cells.
- 22. The immune cell population of any preceding Aspect, wherein step (b) increases the proportion of Bacteroides (eg, B fragalis and/or B thetaiotamicron) in the gut 25 microbiota.
- 23. The immune cell population of any one of Aspects 1 to 21, wherein step (b) decreases the proportion of Bacteroides (eg, B fragalis and/or B thetaiotamicron) in the gut microbiota.
- 24. The immune cell population of any preceding Aspect, wherein step (b) increases the proportion of Bacteroidetes to Firmicutes in the gut microbiota.
- 25. The immune cell population of any one of Aspects 1 to 23, wherein step (b) decreases the proportion of Bacte- 35 36. The immune cell population of Aspect 34 or 35, wherein roidetes to *Firmicutes* in the gut microbiota.
- 26. The immune cell population of any preceding Aspect, wherein step (b) reduces the proportion of one or more Clostridium species or strain (eg, wherein each species is a cluster IV or XIVa Clostridium species) in the gut 40 microbiota.
- 27. The immune cell population of any one of Aspects 1 to 25, wherein step (b) increases the proportion of one or more Clostridium species or strain (eg, wherein each species is a cluster IV or XIVa Clostridium species) in the 45 gut microbiota.
- 28. The immune cell population of any preceding Aspect, wherein step (b) reduces the proportion of Bifidobacterium (eg, B bifidum) in the gut microbiota.
- 29. The immune cell population of any one of Aspects 1 to 50 27, wherein step (b) increases the proportion of Bifidobacterium (eg, B bifidum) in the gut microbiota.
- 30. The immune cell population of any preceding Aspect, wherein step (b) comprises altering the relative proportion of a or said sub-population of first cells in the gut 55 microbiota of the patient, thereby producing an altered gut microbiota that modulates the immune cell therapy in the patient, wherein the sub-population comprises host cells of said first species or strain, the method comprising
 - a. combining the microbiota with multiple copies of 60 engineered nucleic acid sequences encoding host modifying (HM) crRNAs, and
 - b. expressing HM-crRNAs in host cells,

wherein each engineered nucleic acid sequence is operable with a Cas nuclease in a respective host cell to form 65 a HM-CRISPR/Cas system and the engineered sequence comprises

- (i) spacer and repeat sequences encoding a HM-crRNA;
- (ii) the HM-crRNA comprising a sequence that is capable of hybridizing to a host cell target sequence to guide Cas nuclease to the target sequence in the host cell; and
- optionally the HM-system comprises a tracrRNA sequence or a DNA sequence expressing a tracrRNA sequence:
- whereby HM-crRNAs guide Cas modification of host target sequences in host cells, whereby host cells are killed or the host cell population growth is reduced, thereby reducing the proportion of said sub-population in the microbiota.
- 31. The immune cell population of Aspect 30, comprising using endogenous Cas nuclease of host cells for modifi-
- prising reducing host cell population growth by at least 5-fold compared to the growth of a control population of host cells that have not received said Cas modification.
- 33. The immune cell population of any one of Aspects 30 to 32, comprising inhibiting host cell population growth on a gut surface.
- 34. The immune cell population of any one of Aspects 30 to 33, wherein the microbiota comprises cells of a second bacterial species or strain, or archaeal species or strain, wherein the second species or strain has a 16s ribosomal RNA-encoding DNA sequence that is at least 80% identical to an 16s ribosomal RNA-encoding DNA sequence of the host cell species or strain, wherein the growth of the second cells in the microbiota is not inhibited by said HM-system.
- 35. The immune cell population of Aspect 34, wherein the second species or strain is a human gut commensal species or strain and/or a human gut probiotic species or strain.
- the second species or strain is a Bacteroidetes (eg, Bacteroides) and optionally the host cells are gram positive bacterial cells.
- 37. The immune cell population of any one of Aspects 30 to 36, wherein the first cells are Firmicutes cells.
- 38. The immune cell population of any one of Aspects 30 to 37, wherein for each host cell the system comprises components according to (i) to (iv):
 - (i) at least one nucleic acid sequence encoding a Cas nuclease;
 - (ii) an engineered HM-CRISPR array comprising a spacer sequence and repeats encoding a HM-crRNA, the HMcrRNA comprising a sequence that hybridises to a host cell target sequence to guide said Cas to the target in the host cell to modify the target sequence;
 - (iii) an optional tracrRNA sequence or a DNA sequence expressing a tracrRNA sequence;
 - (iv) wherein said components of the system are split between the host cell and at least one nucleic acid vector that transforms the host cell, whereby the HMcrRNA guides Cas to the target to modify the host target sequence in the host cell; and
 - wherein the target sequence is modified by the Cas whereby the host cell is killed or host cell growth is reduced:
 - the method comprising introducing the vectors of (iv) into host cells and expressing said HM-crRNA in the host cells, allowing HM-cRNA to hybridise to host cell target sequences to guide Cas to the targets in the host cells to modify target sequences, whereby host cells are killed or host cell growth is reduced, thereby altering the proportion of said sub-population in the microbiota.

39. The immune cell population of Aspect 38, wherein component (i) is endogenous to each host cell.

40. The immune cell population of Aspect 38 or 39, wherein each vector is a virus or phage.

- 41. The immune cell population of any one of Aspects 30 to 5 40, wherein each target sequence is adjacent a NNA-GAAW or NGGNG protospacer adjacent motif (PAM).
- 42. The immune cell population of any one of Aspects 30 to 41, wherein alternatively HM-crRNA and tracrRNA are comprised by a single guide RNA (gRNA), the method 10 comprising introducing said gRNA into host cells or expressing the gRNA in host cells.
- 43. The immune cell population of any one of Aspects 30 to 35 and 37 to 42, wherein the microbiota comprises a second bacterial or archaeal species, wherein each of the 15 first and second species is a respective species of the same phylum (eg, both *Firmicutes* species) and the growth of the second bacteria is not inhibited by the HM-system; or wherein the microbiota comprises a second bacterial or archaeal strain, wherein each of the first and second 20 bacteria or archaea is a respective strain of the same species and the growth of the second bacteria or archaea is not inhibited by the HM-system.
- 44. The immune cell population of any one of Aspects 30 to 43, wherein the microbiota comprises a second bacterial 25 species, wherein each of the first and second species is a respective gram-positive species and the growth of the second bacteria is not inhibited by the HM-system.
- 45. The immune cell population of any one of Aspects 30 to 44, wherein each target sequence is comprised by an 30 antibiotic resistance gene, virulence gene or essential gene of the host cell.
- 46. The method of any preceding Aspect, wherein each first cell is a *Staphylococcus, Streptococcus, Pseudomonas, Salmonella, Listeria, E coli, Desulfovibrio, Vibrio* or 35 *Clostridium* cell.
- 47. The immune cell population of any preceding Aspect, wherein step (b) comprises stimulating Paneth cells of the patient by gut *Bacteroides* (eg, *B thetaiotamicron*), wherein the altered microbiota produced by step (b) 40 comprises an increased proportion of *Bacteroides* first bacteria compared with the microbiota before step (b), whereby Paneth cells are stimulated and the cell therapy is modulated.
- 48. The immune cell population of any preceding Aspect, 45 wherein step (b) comprises developing an immune response in the patient to gut *Bacteroides* (eg, *B thetaiota-micron*), wherein the altered microbiota produced by step (b) comprises an increased proportion of *Bacteroides* first bacteria compared with the microbiota before step (b), 50 whereby the cell therapy is modulated.
- 49. The immune cell population of any preceding Aspect, wherein step (b) comprises altering the relative proportion of a or said sub-population of first cells in the gut microbiota of the patient, thereby producing an altered gut 55 microbiota that modulates the immune cell therapy in the patient, wherein step (b) comprises killing first cells of said sub-population or inhibiting growth of said subpopulation by using guided nuclease targeting to the genome of first cells comprised by the sub-population. 60
- 50. A bacterial or archaeal transplant for administration to a patient for therapy of a disease or condition in the patient using the method recited in any preceding Aspect, optionally wherein the transplant comprises cells of said first species. 65
- 51. A HM-CRISPR/Cas system, HM-array or HM-crRNA as recited in any one of Aspects 30 to 45 for administration

to a patient for therapy of a disease or condition in the patient using the method recited in any one of Aspects 1 to 49.

- 52. A kit comprising an ex vivo population of immune cells according to any one of Aspects 1 to 49 for adoptive cell therapy of a patient to treat said disease or condition, wherein the kit further comprises a transplant, system, array or crRNA of Aspect 50 or 51, optionally wherein the immune cells are selected from CAR-T cells, T-cells expressing engineered T-cell receptors (TCRs), tumour infiltrating lymphocytes (TILs) and NK cells.
- Optionally, the host cell(s), first cell(s), second cell(s) or mixed bacterial population is comprised by a human or a non-human animal subject, eg, the population is comprised by a gut microbiota, skin microbiota, oral cavity microbiota, throat microbiota, hair microbiota, armpit microbiota, vaginal microbiota, rectal microbiota, anal microbiota, ocular microbiota, nasal microbiota, tongue microbiota, lung microbiota, liver microbiota, kidney microbiota, genital microbiota, penile microbiota, scrotal microbiota, mammary gland microbiota, ear microbiota, urethra microbiota, labial microbiota, organ microbiota or dental microbiota. Optionally, the mixed bacterial population is comprised by a plant (eg, a tobacco, crop plant, fruit plant, vegetable plant or tobacco, eg on the surface of a plant or contained in a plant) or by an environment (eg, soil or water or a waterway or acqueous liquid).

Optionally, the disease or condition of a human or animal subject or patient is selected from

- (a) A neurodegenerative disease or condition;
- (b) A brain disease or condition;
- (c) A CNS disease or condition;
- (d) Memory loss or impairment;
- (e) A heart or cardiovascular disease or condition, eg, heart attack, stroke or atrial fibrillation;
- (f) A liver disease or condition;
- (g) A kidney disease or condition, eg, chronic kidney disease (CKD);
- (h) A pancreas disease or condition;
- (i) A lung disease or condition, eg, cystic fibrosis or COPD;
- (j) A gastrointestinal disease or condition;
- (k) A throat or oral cavity disease or condition;
- (l) An ocular disease or condition;
- (m) A genital disease or condition, eg, a vaginal, labial, penile or scrotal disease or condition;
- (n) A sexually-transmissible disease or condition, eg, gonorrhea, HIV infection, syphilis or *Chlamydia* infection;
- (o) An ear disease or condition;
- (p) A skin disease or condition;
- (q) A heart disease or condition;
- (r) A nasal disease or condition
- (s) A haematological disease or condition, eg, anaemia, eg, anaemia of chronic disease or cancer;
- (t) A viral infection;
- (u) A pathogenic bacterial infection;
- (v) A cancer;
- (w) An autoimmune disease or condition, eg, SLE;
- (x) An inflammatory disease or condition, eg, rheumatoid arthritis, psoriasis, eczema, asthma, ulcerative colitis, colitis, Crohn's disease or IBD;
- (y) Autism;
- (z) ADHD;
- (aa) Bipolar disorder;
- (bb) ALS [Amyotrophic Lateral Sclerosis];
- (cc) Osteoarthritis;

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- (dd) A congenital or development defect or condition; (ee) Miscarriage;
- (ff) A blood clotting condition;
- (gg) Bronchitis;
- (hh) Dry or wet AMD;
- (ii) Neovascularisation (eg, of a tumour or in the eye);
- (jj) Common cold;
- (kk) Epilepsy;
- (11) Fibrosis, eg, liver or lung fibrosis;
- (mm) A fungal disease or condition, eg, thrush;
- (nn) A metabolic disease or condition, eg, obesity, anorexia, diabetes, Type I or Type II diabetes.
- (oo) Ulcer(s), eg, gastric ulceration or skin ulceration; (pp) Dry skin;
- (qq) Sjogren's syndrome;
- (rr) Cytokine storm;
- (ss) Deafness, hearing loss or impairment;
- (tt) Slow or fast metabolism (ie, slower or faster than average for the weight, sex and age of the subject);
- (uu) Conception disorder, eg, infertility or low fertility; 20 (vv) Jaundice;
- (ww) Skin rash;
- (xx) Kawasaki Disease;
- (yy) Lyme Disease;
- (zz) An allergy, eg, a nut, grass, pollen, dust mite, cat or 25 dog fur or dander allergy;
- (aaa) Malaria, typhoid fever, tuberculosis or cholera;
- (bbb) Depression;
- (ccc) Mental retardation;
- (ddd) Microcephaly;
- (eee) Malnutrition;
- (fff) Conjunctivitis;
- (ggg) Pneumonia;
- (hhh) Pulmonary embolism;
- (iii) Pulmonary hypertension;
- (jjj) A bone disorder;
- (kkk) Sepsis or septic shock;
- (111) Sinusitus;
- (mmm) Stress (eg, occupational stress);
- (nnn) Thalassaemia, anaemia, von Willebrand Disease, or 40 haemophilia;
- (000) Shingles or cold sore;
- (ppp) Menstruation;
- (qqq) Low sperm count.
- Neurodegenerative or CNS Diseases or Conditions for 45 Treatment or Prevention by the Method
- In an example, the neurodegenerative or CNS disease or condition is selected from the group consisting of Alzheimer disease, geriopsychosis, Down syndrome, Parkinson's disease, Creutzfeldt-Jakob disease, diabetic neuropathy, Par- 50 kinson syndrome, Huntington's disease, Machado-Joseph disease, amyotrophic lateral sclerosis, diabetic neuropathy, and Creutzfeldt Creutzfeldt-Jakob disease. For example, the disease is Alzheimer disease. For example, the disease is Parkinson syndrome.

In an example, wherein the method of the invention is practised on a human or animal subject for treating a CNS or neurodegenerative disease or condition, the method causes downregulation of Treg cells in the subject, thereby promoting entry of systemic monocyte-derived macro- 60 phages and/or Treg cells across the choroid plexus into the brain of the subject, whereby the disease or condition (eg, Alzheimer's disease) is treated, prevented or progression thereof is reduced. In an embodiment the method causes an increase of IFN-gamma in the CNS system (eg, in the brain 65 and/or CSF) of the subject. In an example, the method restores nerve fibre and/or reduces the progression of nerve

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fibre damage. In an example, the method restores nerve myelin and/or reduces the progression of nerve myelin damage. In an example, the method of the invention treats or prevents a disease or condition disclosed in WO2015136541 and/or the method can be used with any method disclosed in WO2015136541 (the disclosure of this document is incorporated by reference herein in its entirety, eg, for providing disclosure of such methods, diseases, conditions and potential therapeutic agents that can be administered to the subject for effecting treatment and/or prevention of CNS and neurodegenerative diseases and conditions, eg, agents such as immune checkpoint inhibitors, eg, anti-PD-1, anti-PD-L1, anti-TIM3 or other antibodies disclosed therein).

Cancers for Treatment or Prevention by the Method 15

Cancers that may be treated include tumours that are not vascularized, or not substantially vascularized, as well as vascularized tumours. The cancers may comprise non-solid tumours (such as haematological tumours, for example, leukaemias and lymphomas) or may comprise solid tumours. Types of cancers to be treated with the invention include, but are not limited to, carcinoma, blastoma, and sarcoma, and certain leukaemia or lymphoid malignancies, benign and malignant tumours, and malignancies e.g., sarcomas, carcinomas, and melanomas. Adult tumours/cancers and paediatric tumours/cancers are also included.

Haematologic cancers are cancers of the blood or bone marrow. Examples of haematological (or haematogenous) cancers include leukaemias, including acute leukaemias 30 (such as acute lymphocytic leukaemia, acute myelocytic leukaemia, acute myelogenous leukaemia and myeloblasts, promyeiocytic, myelomonocytic, monocytic and erythroleukaemia), chronic leukaemias (such as chronic myelocytic (granulocytic) leukaemia, chronic myelogenous leukaemia, 35 and chronic lymphocytic leukaemia), polycythemia vera, lymphoma, Hodgkin's disease, non-Hodgkin's lymphoma (indolent and high grade forms), multiple myeloma, Waldenstrom's macroglobulinemia, heavy chain disease, myeiodysplastic syndrome, hairy cell leukaemia and myelodysplasia.

Solid tumours are abnormal masses of tissue that usually do not contain cysts or liquid areas. Solid tumours can be benign or malignant. Different types of solid tumours are named for the type of cells that form them (such as sarcomas, carcinomas, and lymphomas). Examples of solid tumours, such as sarcomas and carcinomas, include fibrosarcoma, myxosarcoma, liposarcoma, chondrosarcoma, osteosarcoma, and other sarcomas, synovioma, mesothelioma, Ewing's tumour, 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, medullary thyroid carcinoma, papillary thyroid carcinoma, pheochromocytomas sebaceous gland carcinoma, papillary carcinoma, papillary adenocarcinomas, medullary carcinoma, bronchogenic carcinoma, renal cell carcinoma, hepatoma, bile duct carcinoma, choriocarcinoma, Wilms' tumour, cervical cancer, testicular tumour, seminoma, bladder carcinoma, melanoma, and CNS tumours (such as a glioma (such as brainstem glioma and mixed gliomas), glioblastoma (also known as glioblastoma multiforme) astrocytoma, CNS lymphoma, germinoma, medulloblastoma, Schwannoma craniopharyogioma, ependymoma, pineaioma, hemangioblastoma, acoustic neuroma, oligodendroglioma, menangioma, neuroblastoma, retinoblastoma and brain metastases).

Autoimmune Diseases for Treatment or Prevention by the Method Acute Disseminated Encephalomyelitis (ADEM) Acute necrotizing hemorrhagic leukoencephalitis Addison's disease 5 Agammaglobulinemia Alopecia areata Amyloidosis Ankylosing spondylitis Anti-GBM/Anti-TBM nephritis 10 Antiphospholipid syndrome (APS) Autoimmune angioedema Autoimmune aplastic anemia Autoimmune dysautonomia Autoimmune hepatitis 15 Autoimmune hyperlipidemia Autoimmune immunodeficiency Autoimmune inner ear disease (AIED) Autoimmune myocarditis Autoimmune oophoritis 20 Autoimmune pancreatitis Autoimmune retinopathy Autoimmune thrombocytopenic purpura (ATP) Autoimmune thyroid disease Autoimmune urticaria 25 Axonal & neuronal neuropathies Balo disease Behcet's disease Bullous pemphigoid Cardiomyopathy 30 Castleman disease Celiac disease Chagas disease Chronic fatigue syndrome Chronic inflammatory demyelinating polyneuropathy 35 (CIDP) Chronic recurrent multifocal ostomyelitis (CRMO) Churg-Strauss syndrome Cicatricial pemphigoid/benign mucosal pemphigoid Crohn's disease 40 Cogans syndrome Cold agglutinin disease Congenital heart block Coxsackie myocarditis CREST disease 45 Essential mixed cryoglobulinemia Demvelinating neuropathies Dermatitis herpetiformis Dermatomyositis Devic's disease (neuromyelitis optica) 50 Discoid lupus Dressler's syndrome Endometriosis Eosinophilic esophagitis Eosinophilic fasciitis 55 Erythema nodosum Experimental allergic encephalomyelitis Evans syndrome Fibromyalgia Fibrosing alveolitis 60 Giant cell arteritis (temporal arteritis) Giant cell myocarditis Glomerulonephritis Goodpasture's syndrome Granulomatosis with Polyangiitis (GPA) (formerly called 65 Wegener's Granulomatosis) Graves' disease

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Guillain-Barre syndrome Hashimoto's encephalitis Hashimoto's thyroiditis Hemolytic anemia Henoch-Schonlein purpura Herpes gestationis Hypogammaglobulinemia Idiopathic thrombocytopenic purpura (ITP) IgA nephropathy IgG4-related sclerosing disease Immunoregulatory lipoproteins Inclusion body myositis Interstitial cystitis Juvenile arthritis Juvenile diabetes (Type 1 diabetes) Juvenile myositis Kawasaki syndrome Lambert-Eaton syndrome Leukocytoclastic vasculitis Lichen planus Lichen sclerosus Ligneous conjunctivitis Linear IgA disease (LAD) Lupus (SLE) Lyme disease, chronic Meniere's disease Microscopic polyangiitis Mixed connective tissue disease (MCTD) Mooren's ulcer Mucha-Habermann disease Multiple sclerosis Myasthenia gravis Myositis Narcolepsy Neuromyelitis optica (Devic's) Neutropenia Ocular cicatricial pemphigoid Optic neuritis Palindromic rheumatism PANDAS (Pediatric Autoimmune Neuropsychiatric Disorders Associated with Streptococcus) Paraneoplastic cerebellar degeneration Paroxysmal nocturnal hemoglobinuria (PNH) Parry Romberg syndrome Parsonnage-Turner syndrome Pars planitis (peripheral uveitis) Pemphigus Peripheral neuropathy Perivenous encephalomyelitis Pernicious anemia POEMS syndrome Polyarteritis nodosa Type I, II, & III autoimmune polyglandular syndromes Polymyalgia rheumatica Polymyositis Postmyocardial infarction syndrome Postpericardiotomy syndrome Progesterone dermatitis Primary biliary cirrhosis Primary sclerosing cholangitis Psoriasis Psoriatic arthritis Idiopathic pulmonary fibrosis Pyoderma gangrenosum Pure red cell aplasia Raynauds phenomenon **Reactive Arthritis**

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Reflex sympathetic dystrophy Reiter's syndrome Relapsing polychondritis Restless legs syndrome Retroperitoneal fibrosis Rheumatic fever Rheumatoid arthritis Sarcoidosis Schmidt syndrome Scleritis Scleroderma Sjogren's syndrome Sperm & testicular autoimmunity Stiff person syndrome Subacute bacterial endocarditis (SBE) Susac's syndrome Sympathetic ophthalmia Takayasu's arteritis Temporal arteritis/Giant cell arteritis Thrombocytopenic purpura (TTP) Tolosa-Hunt syndrome Transverse myelitis Type 1 diabetes Ulcerative colitis Undifferentiated connective tissue disease (UCTD) Uveitis Vasculitis Vesiculobullous dermatosis Vitiligo Wegener's granulomatosis (now termed Granulomatosis 30 with Polyangiitis (GPA). Inflammatory Diseases for Treatment or Prevention by the Method Alzheimer's ankylosing spondylitis arthritis (osteoarthritis, rheumatoid arthritis (RA), psoriatic arthritis) asthma atherosclerosis Crohn's disease colitis dermatitis diverticulitis fibromyalgia hepatitis irritable bowel syndrome (IBS) systemic lupus erythematous (SLE) nephritis Parkinson's disease ulcerative colitis. In an example (eg, in the method of the invention involv-

ing a mixed bacterial population), the host cell (or first cell or second cell) genus or species is selected from a genus or species listed in Table 1. In examples of the present invention, the Cas (eg, Cas nuclease such as a Type I, II or III Cas, 55 eg, a Cas3 or 9) is a Cas comprised by bacteria of a genus or species that is selected from a genus or species listed in Table 1, and optionally the host cell (or first cell or second cell) is of the same genus or species. In an example of this, the Cas is endogenous to said host cell (or first or second 60 cell), which is useful for embodiments herein wherein endogenous Cas is used to modify a target sequence. In this case, the HM-array may comprise one or more repeat nucleotide (eg, DNA or RNA) sequences that is at least 90, 95, 96, 97, 98 or 99% identical (or is 100% identical) to a 65 repeat sequence of said cell, genus or species, whereby the Cas is operable with cRNA encoded by the HM-array for

modifying one or more target sequences in the cell. In an example, the Cas is a Type I Cas3 and is used with a Type I CASCADE, wherein one or both of the Cas3 and CAS-CADE are endogenous to the host or first cells, or are vector-borne (ie, exogenous to the host or first cells).

In an example, the method of the invention selectively kills first cells in the microbiota whilst not targeting second cells, eg, wherein the second cells are (a) of a related strain to the strain of the first species or (b) of a species that is 10 different to the first species and is phylogenetically related to the first species, wherein the second species or strain has a 16s ribosomal RNA-encoding DNA sequence that is at least 80% identical to an 16s ribosomal RNA-encoding DNA sequence of the first cell species or strain. In an embodiment, 15 the first cells are of a first species selected from Table 1 and

the second cells are of a different species selected from Table 1. In an example, the species are of the same genus or are of different genera.

It will be understood that particular embodiments 20 described herein are shown by way of illustration and not as limitations of the invention. The principal features of this invention can be employed in various embodiments without departing from the scope of the invention. Those skilled in the art will recognize, or be able to ascertain using no more 25 than routine study, numerous equivalents to the specific procedures described herein. Such equivalents are considered to be within the scope of this invention and are covered by the claims. All publications and patent applications mentioned in the specification are indicative of the level of skill of those skilled in the art to which this invention pertains. All publications and patent applications and all US equivalent patent applications and patents are herein incorporated by reference to the same extent as if each individual publication or patent application was specifically and indi-35 vidually indicated to be incorporated by reference. The use of the word "a" or "an" when used in conjunction with the term "comprising" in the claims and/or the specification may mean "one," but it is also consistent with the meaning of "one or more," "at least one," and "one or more than one." 40 The use of the term "or" in the claims is used to mean "and/or" unless explicitly indicated to refer to alternatives only or the alternatives are mutually exclusive, although the disclosure supports a definition that refers to only alternatives and "and/or." Throughout this application, the term "about" is used to indicate that a value includes the inherent 45 variation of error for the device, the method being employed to determine the value, or the variation that exists among the study subjects.

As used in this specification and claim(s), the words "comprising" (and any form of comprising, such as "comprise" and "comprises"), "having" (and any form of having, such as "have" and "has"), "including" (and any form of including, such as "includes" and "include") or "containing" (and any form of containing, such as "contains" and "contain") are inclusive or open-ended and do not exclude additional, unrecited elements or method steps

The term "or combinations thereof" or similar as used herein refers to all permutations and combinations of the listed items preceding the term. For example, "A, B, C, or combinations thereof is intended to include at least one of: A, B, C, AB, AC, BC, or ABC, and if order is important in a particular context, also BA, CA, CB, CBA, BCA, ACB, BAC, or CAB. Continuing with this example, expressly included are combinations that contain repeats of one or more item or term, such as BB, AAA, MB, BBC, AAABCCCC, CBBAAA, CABABB, and so forth. The skilled artisan will understand that typically there is no limit

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on the number of items or terms in any combination, unless otherwise apparent from the context.

Any part of this disclosure may be read in combination with any other part of the disclosure, unless otherwise apparent from the context.

All of the compositions and/or methods disclosed and claimed herein can be made and executed without undue experimentation in light of the present disclosure. While the compositions and methods of this invention have been described in terms of preferred embodiments, it will be ¹⁰ apparent to those of skill in the art that variations may be applied to the compositions and/or methods and in the steps or in the sequence of steps of the method described herein without departing from the concept, spirit and scope of the invention. All such similar substitutes and modifications ¹⁵ apparent to those skilled in the art are deemed to be within the spirit, scope and concept of the invention as defined by the appended claims.

The present invention is described in more detail in the following non limiting Examples.

EXAMPLES

Example 1: Specific Microbiota Bacterial Population Growth Inhibition by Harnessing Wild-Type Endogenous Cas

1. Material and Methods

1.1. Strains

The following strains were used in the course of this 30 Example and Examples 2 and 3: *E. coli* MG1655, *E. coli* TOP10, *Streptococcus thermophilus* LMD-9 (ATCC BAA-491, Manassas, Va.), *Streptococcus thermophilus* DSM 20617(T) (DSMZ, Braunschweig, Germany), *Lactococcus lactis* MG1363 and *Streptococcus mutans* Clarke 1924 DSM 35 20523 (DSMZ, Braunschweig, Germany).

During the course of media selection and testing of the genetic constructs different Streptocccci strains were used. *Streptococcus thermophilus* LMD-9 (ATCC BAA-491) and *Escherichia coli* TOP10 were considered because of their 40 compatible growth requirements. All strains were cultivated in Todd-Hewitt broth (TH) (T1438 Sigma-Aldrich), in aerobic conditions and at 37° C., unless elsewhere indicated. The strains were stored in 25% glycerol at -80° C.

1.2. Differential Growth Media

All strains were grown on TH media at 37° C. for 20 hours. Selective media for *S. thermophilus* was TH media supplemented with 3 g l⁻¹ of 2-phenylethanol (PEA). PEA was added to the media and autoclaved at 121° C. for 15 minutes at 15 psi. Agar plates were prepared by adding 1.5% 50 (wt/vol) agar to the corresponding media. When necessary for selection or plasmid maintenance 30 µg ml⁻¹ kanamycin was used for both *S. thermophilus* strains and *E. coli*, and 500 µg ml⁻¹ for *S. mutans*.

In some cases, depending on the strain and plasmid, a 55 longer incubation, up to 48 hours, may be needed to see growth on media supplemented with PEA. In order to control for the viability of the organisms used, a control TH agar must be done in parallel.

1.3. Cloning

E. coli (One Shot® ThermoFischer TOP10 Chemically Competent cells) was used in all subcloning procedures. PCR was carried out using Phusion polymerase. All PCR products were purified with Nucleospin Gel and PCR Cleanup by Macherey-Nagel following the manufacturer's protocol. The purified fragments were digested with restriction enzyme DpnI in 1 λ FD buffer with 1 μ l enzyme in a total

volume of 34 μ l. The digested reaction was again purified with Nucleospin Gel and PCR Clean-up by Macherey-Nagel following the manufacturer's protocol. Gibson assembly was performed in 10 μ l reactions following the manufacturer's protocol (NewEngland Biolab).

Plasmid DNA was prepared using Qiagen kits according to the manufacturer's instructions. Modifications for Grampositive strains included growing bacteria in a medium supplemented with 0.5% glycine and lysozyme to facilitate cell lysis.

1.4. Transformation

1.4.1 Electro-Competent *E. coli* Cells and Transformation Commercially electrocompetent cells were used for cloning and the experiments (One Shot® ThermoFischer TOP10
15 Chemically Competent *E. coli*). Electroporation was done using standard settings: 1800 V, 25 μF and 200Ω using an Electro Cell Manipulator (BTX Harvard Apparatus ECM630). Following the pulse, 1 ml LB-SOC media was added and the cells were incubated at 37° C. for 1 hour. The
20 transformed cells were plated in LB-agar containing 50 μg ml⁻¹ of kanamycin.

1.4.2 Preparation of Electro-Competent S. thermophilus Cells

The electroporation protocol was modified from Somkuti
and Steinberg, 1988. An overnight culture of *Streptococcus* thermophilus in TH Broth supplemented with 40 mM DL-threonine (T8375 Sigma-Aldrich) was diluted 100-fold in 5 ml of the same media and grown to an OD₆₀₀ between 0.3-0.5 (approximately 2.5 hours after inoculation). The
cells were collected by centrifugation at 10,000×g for 10 min at 4° C. and washed three times with 5 ml of ice cold wash buffer (0.5 M sucrose+10% glycerol). After the cells were washed, they were suspended to an OD₆₀₀ of 15-30 in electroporation buffer (0.5 M sucrose, 10% glycerol and 1
mM MgCl₂). The cells in the electroporation buffer may be kept at 4° C. until use (within one hour) or aliquot 50 µl in eppendorf tubes, freezing them in liquid nitrogen and stored at -80° C. for later use.

1.4.3 Electroporation S. thermophilus Cells

1 μl of purified plasmid DNA was added to 50 μl of the cell suspension and electroporation was carried out in 2 mm-gap electroporation cuvettes pre-cooled. The electroporation setting were 2500 V, 25 μF and 200Ω using an Electro Cell Manipulator (BTX Harvard Apparatus ECM630) Immediately after the electric pulse, 1 ml of TH broth was added to the cells and the suspension was kept on ice for 10 minutes, subsequently the cells were incubated for 3 h at 37° C. After allowing time for expression of the resistance gene the cells were plated onto TH-agar plates containing 30 μg ml⁻¹ of kanamycin. Depending on the construct, colonies were visible between 12 and 48 h of incubation at 37° C.

1.5. Construction of XylS Plasmid

All the plasmids used in this work were based on pBAV1K-T5, which is a broad-host range expression vector derived from the a cryptic plasmid pWV01 from *Streptococcus cremoris* (Bryksin & Matsumura, 2010), the backbone was amplified using that contain overhangs for assembly with the other fragments using Gibson's method.

The xylose inducible system was constructed by cloning the promoter gyrA in front of the XylR repressor (FIG. 1). The XylR repressor was amplified from *Bacillus Subtilis* strain SCK6 (Zhang et al. 2011) with the a reverse primer that includes an overhang for Gibson assembly and a forward primer, that is an ultramer used to introduce the gyrA promoter (Xie et al. 2013) and the corresponding overhang for assembly into pBAV1KT5 backbone. The resulting fragment was flanked by an mCherry amplified from pCL002 (unpublished work) with an ultramer that include Pldha+ PxylA hybrid promoter (Xie et al. 2013). The three resulting PCR products were assembled in a Gibson Master Mix® (NewEngland Biolab) according to manufacturer's instructions. The product was finally transformed in *E. coli* TOP10 5 electrocompetent cells. See FIG. **1**.

1.6. Design and Construction of CRISPR Array Plasmid *Streptococcus thermophilus* has 4 distinct CRISPR systems (Sapranauskas, et al. 2011), for this work the type II CRISPR1 (ST1-CRISPR) system was chosen. The design of 10 the target sequence was based on the available genome sequence of LMD-9 (GenBank: CP000419.1). The ST1-CRISPR array was designed to contain only the CRISPR array repeats and spacers under a xylose inducible promoter (Xie et al. 2013), followed by the corresponding tracrRNA 15 under a strong constitutive promoter for Streptococci species (Sorg et al. 2014) (FIG. 2).

The tracrRNA plays a role in the maturation of crRNA and it is processed by *S. thermophilus* endogenous RNase III, forming a complex with crRNA. This complex acts as a 20 guide for the endonuclease ST1-Cas9 (Horvath & Barrangou, 2010). After transcription of the synthetic array from the xylose inducible promoter, the endogenous Cas9 and RNAses will process it into a functional gRNA. The gRNA/ Cas9 complex will cause a double stranded break at the 25 target location.

The design of the array used 2 specific target sequences high on GC content and a reduced portion of the tracrRNA (ie, a less than complete tracrRNA sequence), which has been suggested not to be necessary for proper maturation of 30 crRNA (Horvath & Barrangou, 2010).

The 2 targets were an essential gene (DNA polymerase III subunit alpha) and an antibiotic resistance gene (tetA-like gene).

Primers were used to amplify pBAV1KT5-XylR-PldhA 35 backbone. The CRISPR array gBlock and the backbone with overhangs were assembled in a Gibson Master Mix® according to manufacturer's instructions (NewEngland Biolabs). The product was finally transformed in *E. coli* TOP10 electrocompetent cells. 40

1.7. Characterization of Xylose Inducible System in *Streptoccocus thermophilus* LMD-9

Overnight stationary-phase cultures were diluted 1:100 into TH broth with corresponding antibiotic. Mid-log cells were induced with different concentration of D-(+)-xylose 45 (0, 0.001, 0.01, 0.1, 0.5 and 1% wt/vol) and the cell cultures were measured either directly in medium to assess the extent of autofluorescence of the media, on the cell suspension or the suspension buffer (PBS buffer). 20 μ l samples of the cell cultures were diluted 1/10 on PBS buffer, on 96-well plates 50 with flat bottoms. Fluorescence of cell suspensions or media was read on a plate reader. mCherry fluorescence was measured using an excitation wavelength of 558 nm and emission at 612 nm. Absorbance of the resuspended cells was measured at OD 600 nm. A minimum of three inde-55 pendent biological replicates was done for each experiment.

1.8. Activation of CRISPR Array in *S. thermophilus S. thermophilus* LMD-9 and *E. coli* TOP10 both with the plasmid containing the CRISPR array targeting the DNA polymerase III and tetA of *S. thermophilus* were grown 60 overnight in 3 ml cultures supplemented with 30 μ g ml⁻¹ of kanamycin for plasmid maintenance. The next day 96 well deep well plates were inoculated with 500 μ l of 1/100 of overnight culture in fresh TH media, supplemented with 30 ml kanamycin. Mid-log cell cultures were induced with 1% 65 xylose. The killing effect was tested on *S. thermophilus* and *E. coli* alone. For each strain and condition tested a negative

control was kept without xylose. The cells were grown till ~OD 0.5 and next 10-fold serially diluted in TH media and using a 96-well replicator (Mettler Toledo LiquidatorTM 96) 5 μ L volume drops were spotted on TH agar and TH agar supplemented with g l⁻¹ PEA plates. The plates were incubated for 24H at 37° C. and the colony forming units (CFU) were calculated from triplicate measurements.

2. Results

2.1 Growth Condition and Selective Media

We first set out to establish the bacterial strains and cultivation protocol that would support growth for all strains we planned to use for the co-cultivation experiments. We used *S. thermophilus* strain LMD-9 which was able to support a similar growth as *E. coli* in TH broth at 37° C. (FIG. **3**).

Distinguishing the different bacteria from a mixed culture is important in order to determine cell number of the different species. With MacConkey agar is possible to selectively grow *E. coli*, however there is no specific media for selective growth of *S. thermophilus*. PEA agar is a selective medium that is used for the isolation of gram-positive (*S. thermophilus*) from gram-negative (*E. coli*). Additionally, we found that different concentrations of PEA partially inhibit the growth of other gram positives, which allow for selection between the other gram-positive bacteria used in this work (FIG. **4**). 3 g l⁻¹ of PEA proved to selectively grow *S. thermophilus* LMD-9 while limiting growth of *E. coli*.

2.2 Design and Validation of Inducible System

An induction system for *Streptococcus* species was previously developed based on the *Bacillus megaterium* xylose operon (FIG. 5) by creating a heterologous xylose induction cassette (Xyl-S). The xylR and xylA promoters were replaced with *S. mutans*' constitutively expressed gyrA and ldh promoters respectively. This expression cassette for *Streptococcus* species showed differences in sensitivity and expression levels between different species, however the system was not tested in *S. thermophilus* (Xie et al. 2013). Therefore we first set out to validate the xylose induction cassette in *S. thermophilus*.

An alternative version of the induction cassette was constructed by only replacing the xylR promoter with the *S. mutans*' gyrA promoter but left the endogenous *B. megate-rium* xylA promoter intact. During the design of the xylose inducible system we considered both versions of the inducible promoter, the natural P_{XylA} promoter found in *Bacillus megaterium* and a hybrid promoter of the highly conserved promoter P_{tdha} fused with the repressor binding sites of P_{XylA} promoter (FIG. 5). Only a few *Streptococcus* species have been reported to metabolize xylose, and thus the presence of a regulatory machinery to recognize the xylA promoter in the other *Streptococcus* species is not likely. Therefore we constructed both xylose induction systems but only tested the inducibility of mCherry with the $P_{ldha+XylA}$ system.

In order to determine mCherry inducible expression by xylose, mid-log cultures of cells with the plasmid (pBAV1KT5-XylR-mCherry- $P_{idha+XylA}$) were induced with different concentrations of xylose. Six hours after the induction we measured mCherry fluorescence in the cultures, where we observed substantially higher overall expression levels in cells carrying the plasmid (FIG. **6**). It is worth noticing that the system showed a substantial level of basal expression even in the cultures where xylose was not added. This means that the system is 'leaky' and in context of the kill-array this can lead to cell death even before the system is induced with xylose. However, in the subsequent course

of this study we used both versions of the plasmid (pBAV1KT5-XylR-mCherry- $P_{ldha+XylA}$ and pBAV1KT5-XylR-mCherry- P_{xylA}).

2.3 Design of CRISPR/CAS9 Array

In order to determine if the genomic targeting spacers in a CRISPR array can cause death in *S. thermophilus* LMD-9, we inserted the CRISPR array we designed into the two xylose inducible systems previously constructed (pBAV1KT5-XylR-mCherry-P_{*ldha+XylA*} and pBAV1KT5-XylR-mCherry-P_{*ldha+XylA*} and pBAV1KT5-XylR-mCherry-P_{*xylA*}). In these plasmids we replaced mCherry with the gBlock containing the CRISPR array (FIG. 7). The variant with the P_{*ldha+XylA*} promoter was expected to be stronger and have a higher basal activity than the P_{*xylA*} (Xie et al. 2013).

2.4 Inhibition of Bacterial Population Growth Using Endogenous Cas9

After we constructed the plasmids in E. coli, we transformed the plasmids into S. thermophilus. This would allow us to determine if we could cause cell death of a specific 20 bacterial species. Interestingly, bacterial host population size (indicated by growing bacteria and counting colony numbers on agar plates) in S. thermophilus exposed to the plasmid containing the strong $P_{1dh+XylA}$ hybrid promoter was 10-fold less when compared to *S. thermophilus* exposed to the 25 plasmid containing the weak, normal P_{xylA} promoter (FIG. 8; 52 colonies with the strong array expression versus 556 colonies with weak array expression, 10.7-fold difference), the 2 strains having been transformed in parallel using the same batch of electrocompetent S. thermophilus cells. This suggests to us that the plasmid carrying the CRISPR array targeting S. thermophilus genes is able to kill the cells using the endogenous Cas nuclease and RNase III, thereby inhibiting population growth by 10-fold.

We expect that weak array expression in host cells trans- 35 formed by the plasmid comprising the P_{xylA} promoter led to a degree of cell killing, albeit much less than with the strong promoter plasmid. We expect that population growth inhibition that is greater than the observed 10-fold inhibition would be determined if a comparison of the activity of 40 strong array expression was made with S thermophilus that is not exposed to any array-encoding plasmid (such as bacteria directly isolated from gut microbiota). Thus, we believe that array (or single guide RNA) expression in host cells for harnessing endogenous Cas nuclease will be useful 45 for providing effective growth inhibition of target host cells in environmental, medical and other settings mentioned herein. Co-administration of antibiotic may also be useful to enhance the growth inhibition, particularly when one or more antibiotic resistance genes are targeted. 50

3. Discussion and Outlook

In this study we set out to design a CRISPR-array to specifically kill *S. thermophilus* using the endogenous Cas9 system. In order to gain control over the killing signal we sought to apply an inducible system that can be applied in *S. 55 thermophilus*. The xylose inducible XylR system from *B. megaterium* was previously applied in *S. mutans* (Xie, 2013) but not in *S. thermophilus*. In this study we demonstrated the functionality of the xylR induction system using the designed XylR-mCherry-Pldha circuit in *S. thermophilus*. 60 We found 0.1% wt/vol is sufficient to fully induce the XylR system in *S. thermophilus* (FIG. **6**).

In order to observe abundance when co-culturing *S*. *thermophilus* and *E. coli* we established that supplementation of the culture media with 3 g 1^{-1} of PEA, allows for the 65 selective growth of *S. thermophilus* while limiting the growth of *E. coli* (FIG. 4).

A ST1-CRISPR array, targeting the DNA polymerase III subunit alpha and a tetA like gene in the *S. thermophilus* LMD-9 genome, was placed under the xylose inducible promoter (Xie et al. 2013). Targeting these regions should lead to a double strand break and thus limit *S. thermophilus* viability (FIG. 9). Since the engineered array was designed to target *S. thermophilus* genome using the endogenous CRISPR/Cas machinery to process the encoded CRISPR array, the array is expected to have no influence on growth of unrelated strains such as *E. coli*, even similar targets could be found on its genome. This was successfully tested in a mixed bacterial population (simulating aspects of a human microbiota) as discussed in Example 3.

The demonstration of the ability to inhibit host cell growth on a surface is important and desirable in embodiments where the invention is for treating or preventing diseases or conditions mediated or caused by microbiota as disclosed herein in a human or animal subject. Such microbiota are typically in contact with tissue of the subject (eg, gut tissue) and thus we believe that the demonstration of activity to inhibit growth of a microbiota bacterial species (exemplified by *Streptococcus*) on a surface supports this utility.

Example 2: Specific Microbiota Bacterial Population Growth Inhibition in Different Strains

Example 1 demonstrated specific growth inhibition of *Streptococcus thermophilus* LMD-9. Here we demonstrate growth inhibition can also be obtained in a second strain: *Streptococcus thermophilus* DSM 20617. Methods described in Example 1 were, therefore, applied to the latter strain (except that selective media for *S. thermophilus* DSM 20617 was TH media supplemented with 2.5 g 1^{-1} of 2-phenylethanol (PEA)).

Streptococcus thermophilus DSM 20617 transformed with the CRISPR array plasmids were incubated for recovery in liquid media for a period of 3 hours at 37° C. that would allow for expression of kanamycin resistance. After a recovery period, cells were plated in different selection media in presence of 1% xylose in order to induce cell death, and without xylose as a control (FIG. **10**). It is evident that; (1) by xylose induction the growth of *S. thermophilus* can be inhibited (around 10-fold for the 'strong' promoter plasmid versus control), (2) the 'strong' system (pBAV1KT5-XylR-CRISPR-P_{idhA}) results in more growth reduction than the 'weak' system (pBAV1KT5-XylR-CRISPR-P_{xvlA})

Example 3: Selective Bacterial Population Growth Inhibition in a Mixed Consortium of Different Microbiota Species

We next demonstrated selective growth inhibition of a specific bacterial species in a mixed population of three species. We selected species found in gut microbiota of humans and animals (*S thermophilus* DSM 20617(T), *Lactobacillus lactis* and *E coli*). We included two gram-positive species (the *S thermophilus* and *L lactis*) to see if this would affect the ability for selective killing of the former species; furthermore to increase difficulty (and to more closely simulate situations in microbiota) *L lactis* was chosen as this is a phylogenetically-related species to *S thermophilus* (as indicated by high 16s ribosomal RNA sequence identity between the two species). The *S thermophilus* and *L lactis* are both *Firmicutes*. Furthermore, to simulate microbiota, a human commensal gut species (*E coli*) was included.

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1. Materials & Methods

Methods as set out in Example 1 were used (except that selective media was TH media supplemented with 2.5 g l^{-1} of 2-phenylethanol (PEA)).

1.1 Preparation of Electro-Competent L. lactis Cells

Overnight cultures of L. lactis in TH media supplemented with 0.5 M sucrose and 1% glycine were diluted 100-fold in 5 ml of the same media and grown at 30° C. to an OD₆₀₀ between 0.2-0.7 (approximately 2 hours after inoculation). The cells were collected at 7000×g for 5 min at 4° C. and 10 washed three times with 5 ml of ice cold wash buffer (0.5 M sucrose+10% glycerol). After the cells were washed, they were suspended to an OD₆₀₀ of 15-30 in electroporation buffer (0.5 M sucrose, 10% glycerol and 1 mM MgCl₂). The cells in the electroporation buffer were kept at 4° C. until use 15 (within one hour) or aliquot 50 µl in eppendorf tubes, freezing them in liquid nitrogen and stored at -80° C. for later use.

Electroporation conditions for all species were as described in Example 1.

1.2 Activation of CRISPR Array: Consortium Experiments.

S. thermophilus DSM 20617, L. lactis MG1363 and E. coli TOP10 were genetically transformed with the plasmid containing the CRISPR array targeting the DNA polymerase 25 III and tetA of S. thermophilus. After transformation all cells were grown alone and in co-culture for 3 hours at 37° C. allowing for recovery to develop the antibiotic resistance encoded in the plasmid. We decided to use transformation efficiency as a read out of CRISPR-encoded growth inhibi- 30 tion. Therefore, after allowing the cells for recovery the cultures were plated in TH media, TH supplemented with PEA and MacConkey agar all supplemented with Kanamycin, and induced by 1% xylose.

2. Results

2.0 Phylogenetic Distance Between L. lactis, E. Coli and S. thermophilus

The calculated sequence similarity in the 16S rRNAencoding DNA sequence of the S. thermophilus and L. lactis was determined as 83.3%. The following 16S sequences 40 were used: E. coli: AB030918.1, S. thermophilus: AY188354.1, L. lactis: AB030918. The sequences were aligned with needle (http://www.ebi.ac.uk/Tools/psa/emboss_needle/nucleotide.html) with the following parameters: -gapopen 10.0 -gapextend 0.5 -endopen 10.0 -endex- 45 tend 0.5 -aformat3 pair -snucleotide1 -snucleotide2. FIG. 11 shows the maximum-likelihood phylogenetic tree of 16S sequences from S. thermophilus, L. lactis and E. coli.

2.1 Growth Condition and Selective Media

S. thermophilus and L. lactis are commonly used in 50 combination in many fermented foods and voghurt. We chose these strains since they are commonly known to be gut microbes that form an intimate association with the host and previous characterizations of the 16S ribosomal RNA region of S. thermophilus and L. lactis have shown that these 55 organisms are phylogenetically closely related (Ludwig et al., 1995). In parallel we also evaluated the growth of E. coli for our mixed population co-culture experiments, since this organism is also commonly found in gut microbe communities. We first set out to establish the bacterial strains and 60 cultivation protocol that would support growth for all strains we planned to use for the co-cultivation experiments. We found that all strains were able to support growth in TH broth at 37° C. (FIG. 3).

Distinguishing the different bacteria from a mixed culture 65 6. Mercenier, A. (1990). Molecular genetics of Streptococis important in order to determine cell number of the different species. With MacConkey agar is possible to selec-

tively grow E. coli, however there is no specific media for selective growth of S. thermophilus. PEA agar is a selective medium that is used for the isolation of gram-positive (S. thermophilus) from gram-negative (E. coli). Additionally, different concentrations of PEA partially inhibit the growth of the different grams positive species and strains, which allow for selection between the other gram-positive bacteria used in this work. Using 2.5 g 1^{-1} of PEA proved to selectively grow S. thermophilus while limiting growth of L. lactis and E. coli.

All strains were transformed with a plasmid that used the vector backbone of pBAV1KT5 that has a kanamycin selection marker; we found that using media supplemented with 30 ug ml⁻¹ of kanamycin was enough to grow the cells while keeping the plasmid.

2.3 Transformation & Selective Growth Inhibition in a Mixed Population

We transformed S. thermophilus, L. lactis and E. coli with plasmid containing the CRISPR array and cultured them in ²⁰ a consortium of all the bacterial species combined in equal parts, which would allow us to determine if we could cause cell death specifically in S. thermophilus. We transformed all the species with either the pBAV1KT5-XylR-CRISPR- P_{xylA} or pBAV1KT5-XylR-CRISPR-P_{ldha+XylA} plasmid.

FIG. 12 shows the selective S thermophilus growth inhibition in a co-culture of E. coli, L. lactis and S. thermophilus harboring either the pBAV1KT5-XylR-CRISPR- P_{xylA} or the pBAV1KT5-XylR-CRISPR- $P_{ldhA+XylA}$ plasmid. No growth difference is observed between E. coli harboring the pBAV1KT5-XylR-CRISPR-P_{xvl4} or the pBAV1KT5-XylR-CRISPR-P_{ldhA+XylA} plasmid (middle column). However, S. thermophilus (selectively grown on TH agar supplemented with 2.5 gl⁻¹ PEA, last column) shows a decrease in transformation efficiency between the pBAV1KT5-XylR-CRISPR-P_{xylA} (strong) or the pBAV1KT5-XylR-CRISPR-P $_{ldhA+XylA}$ (weak) plasmid as we expected. We thus demonstrated a selective growth inhibition of the target S thermophilus sub-population in the mixed population of cells.

REFERENCES

- 1. Barrangou, R., Fremaux, C., Deveau, H., Richards, M., Patrick Boyaval, Moineau, S., . . . Horvath, P. (2007). CRISPR Provides Acquired Resistance Against Viruses in Prokaryotes. Science, 315 (March), 1709-1712.
- 2. Bryksin, A. V, & Matsumura, I. (2010). Rational design of a plasmid origin that replicates efficiently in both grampositive and gram-negative bacteria. PloS One, 5(10), e13244
- 3. Chan C T Y, Lee J W, Cameron D E, Bashor C J, Collins J J: "Deadman" and "Passcode" microbial kill switches for bacterial containment. Nat Chem Biol 2015, 12(Dec.): 1 - 7
- 4. Horvath, P., Romero, D. A., Cofûté-Monvoisin, A.-C., Richards, M., Deveau, H., Moineau, S., ... Barrangou, R. (2008). Diversity, activity, and evolution of CRISPR loci in Streptococcus thermophilus. Journal of Bacteriology, 190(4), 1401-12.
- 5. Ludwig, E. S., Klipper, R., Magrum L., Wose C., & Stackebrandt, E. (1985). The phylogenetic position of Streptococcus and Enterococcus. Journul of Gencwl Microhiologj., 131, 543-55 1.
- cus thermophilus. FEMS Microbiology Letters, 87(1-2), 61-77. \

- 7. Samaržija, D., Antunac, N., & Havranek, J. (2001). Taxonomy, physiology and growth of *Lactococcus lactis*: a review. *Mljekarstvo*, 51(1), 35-48. Retrieved from
- Sapranauskas, R., Gasiunas, G., Fremaux, C., Barrangou, R., Horvath, P., & Siksnys, V. (2011). The *Streptococcus* 5 *thermophilus* CRISPR/Cas system provides immunity in *Escherichia coli. Nucleic Acids Research*, 39(21), 9275-9282.
- Somkuti, G. A., & Steinberg, D. H. (1988). Genetic transformation of *Streptococcus thermophilus* by elec- 10 troporation. *Biochimie*, 70(4), 579-585
- Sorg, R. A., Kuipers, O. P., & Veening, J.-W. (2014). Gene expression platform for synthetic biology in the human pathogen *Streptococcus pneumoniae*. ACS Synthetic Biology, 4(3), 228-239.
- Suvorov, a. (1988). Transformation of group A streptococci by electroporation. *FEMS Microbiology Letters*, 56(1), 95-99.
- 12. Xie, Z., Qi, F., & Merritt, J. (2013). Development of a tunable wide-range gene induction system useful for the ²⁰ study of streptococcal toxin-antitoxin systems. Applied and Environmental *Microbiology*, 79(20), 6375-84.
- Zhang, X. Z., & Zhang, Y. H. P. (2011). Simple, fast and high-efficiency transformation system for directed evolution of cellulase in *Bacillus subtilis*. *Microbial Biotech*- ²⁵ *nology*, 4(1), 98-105.

Example 4: Altering the Ratio of *Clostridium Dificile* in a Mixed Gut Microbiota Population

Alteration of the ratio of bacteria will be performed according to the present example, which is described by reference to knocking-down *Clostridium dificile* bacteria in a mixed gut microbiota sample. The sample will contain *Bacteroides* and metronidazole (MTZ)-resistant *C dificile* 35 strain 630 sub-populations. Ex vivo the mixed population is combined with a population of carrier bacteria (*Lactobacillus acidophilus* La-14 and/or La-5) that have been engineered to contain CRISPR arrays.

Each CRISPR array is comprised on a plasmid that is 40 compatible with the carrier bacterium and *C dificile* cells. The array is comprised by a *Bacteroides thetaiotamicron* CTnDot transposon that also comprises oriT, an intDOT sequence, a tetQ-rteA-rteB operon, rteC and the operon xis2c-xis2d-orf3-exc. In one experiment, mob and tra oper-45 ons are excluded (instead relying on these supplied by *Bacteroides* cells to which the transposons are transferred in the mixture combined with the carrier bacteria). In another experiment, the mob and tra operons are included in the transposons. 50

Protein translocation across the cytoplasmic membrane is an essential process in all bacteria. The Sec system, comprising at its core an ATPase, SecA, and a membrane channel, SecYEG, is responsible for the majority of this protein transport. A second parallel Sec system has been 55 described in a number of Gram-positive species. This accessory Sec system is characterized by the presence of a second copy of the energizing ATPase, SecA2; where it has been studied, SecA2 is responsible for the translocation of a subset of Sec substrates. In common with many pathogenic 60 Gram-positive species, Clostridium difficile possesses two copies of SecA. Export of the S-layer proteins (SLPs) and an additional cell wall protein (CwpV) is dependent on SecA2. Accumulation of the cytoplasmic precursor of the SLPs SlpA and other cell wall proteins is observed in cells 65 expressing dominant-negative secA1 or secA2 alleles, concomitant with a decrease in the levels of mature SLPs in the

cell wall. Furthermore, expression of either dominant-negative allele or antisense RNA knockdown of SecA1 or SecA2 dramatically impairs growth, indicating that both Sec systems are essential in *C. difficile*.

C. difficile Strain 630 (epidemic type X) has a single circular chromosome with 4,290,252 bp (G+C content=29.06%) and a circular plasmid with 7,881 bp (G+C content=27.9%). The whole genome has been sequenced and found that 11% of the genome consists of mobile genetic elements such as conjugative transposons. These elements provide C. difficile with the genes responsible for its antimicrobial resistance, virulence, host interaction and the production of surface structures. For example, the cdeA gene of C. difficile produces a multidrug efflux pump which was shown to be homologous to known efflux transporters in the multidrug and toxic compound extrusion (MATE) family. The protein facilitates energy-dependent and sodiumcoupled efflux of drugs from cells. In addition, the cme gene in C. difficile has been shown to provide multidrug resistance in other bacteria.

The array comprises a R1-S1-R1' CRISPR unit (spacer flanked by two CRISPR repeats) for targeting a sequence in an essential gene (SecA2) of *C dificile* cells. In another experiment, targeting is to the cdeA gene in the presence of MTZ and optionally one or more other anti-*C dificile* antibiotics. Each spacer (S) comprises a 20mer nucleotide sequence of the SecA or cdeA gene, wherein the sequence comprises a PAM of a *C dificile* strain 630 CRISPR/Cas system that is cognate to the repeat sequences. Each repeat is identical to a *C dificile* strain 630 repeat.

The repeats function with Cas that is endogenous to the *C* dificile cells in the mixed population. The mixed population of bacteria is retrieved as an ex vivo sample from a stool sample of a human patient suffering from *C* dificile infection. The mixed population is mixed with the carrier bacteria in vitro and incubated at 37 degrees centigrade under anaerobic conditions to simulate gut conditions in the presence of tetracycline. It is expected that transposons containing the CRISPR arrays will be transferred to *Bacteroides* and *C* dificile cells in the mixture. Furthermore, it is expected that the target sites in the latter cells will be cut by Cas nuclease action, thus reducing the proportion of *C* dificile in the mixed population (and increasing the ratio of *Bacteroides* versus *C* dificile).

In a follow-on experiment, a drink is produced comprising the carrier bacteria and this is consumed by the human patient once or twice for several consecutive days with or without an ant-acid. The patient is also administered with tetracycline during the treatment period. It is expected that stool analysis will reveal that the proportion of *C dificile* in the stool samples will reduce (and the ratio of *Bacteroides* versus *C dificile* will increase).

Example 5: Vector-Encoded System for Selective Species & Strain Growth Inhibition in A Mixed Bacterial Consortium

In Example 3 we surprisingly established the possibility of harnessing endogenous Cas nuclease activity in host bacteria for selective population growth inhibition in a mixed consortium of different species. We next explored the possibility of instead using vector-encoded Cas activity for selective population growth inhibition in a mixed consortium of different species. We demonstrated selective growth inhibition of a specific bacterial species in a mixed population of three different species, and further including a strain alternative to the target bacteria. We could surprisingly show

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selective growth inhibition of just the target strain of the predetermined target species. Furthermore, the alternative strain was not targeted by the vector-encoded CRISPR/Cas system, which was desirable for establishing the fine specificity of such vector-borne systems in a mixed bacterial ⁵ consortium that mimicked human or animal gut microbiota elements.

We selected species found in gut microbiota of humans and animals (*Bacillus subtilis*, *Lactobacillus lactis* and *E coli*). We included two strains of the human commensal gut ¹⁰ species, *E coli*. We thought it of interest to see if we could distinguish between closely related strains that nevertheless had sequence differences that we could use to target killing in one strain, but not the other. This was of interest as some strains of *E coli* in microbiota are desirable, whereas others may be undesirable (eg, pathogenic to humans or animals) and thus could be targets for Cas modification to knockdown that strain.

1. Material and Methods

1.1. Plasmids and Strains

See Tables 7 and 8. All strains were cultivated in Todd-Hewitt broth (TH) (T1438 Sigma-Aldrich), in aerobic conditions and at 37° C., unless elsewhere indicated. The strains were stored in 25% glycerol at -80° C.

The self-targeting sgRNA-Cas9 complex was tightly regulated by a theophylline riboswitch and the AraC/P_{BAD} expression system respectively. Tight regulation of Cas9 is desired in order to be carried stably in E. coli. The plasmid contained the exogenous Cas9 from Streptococcus pyogenes 30 with a single guide RNA (sgRNA) targeting E. coli's K-12 strains. Therefore K-12 derived strains TOP10 was susceptible to double strand self-cleavage and consequent death when the system was activated. E. coli strains like Nissle don't have the same target sequence therefore they were 35 unaffected by the sgRNA-Cas9 activity. See Tables 9-11 below, which show sequences used in Example 9. We chose a target sequence (ribosomal RNA-encoding sequence) that is conserved in the target cells and present in multiple copies (7 copies), which increased the chances of cutting host cell 40 genomes in multiple places to promote killing using a single gRNA design.

FIG. **13** shows regulators controlling the expression of spCas9 and the self-targeting sgRNA targeting the ribosomal RNA subunit 16s.

1.2. Differential Growth Media

All strains were grown on TH media at 37° C. for 20 hours. Selective media for *B. subtilis* was TH media supplemented with 2.5 g l⁻¹ of 2-phenylethanol (PEA). PEA was added to the media and autoclaved at 121° C. for 15 minutes 50 at 15 psi. Agar plates were prepared by adding 1.5% (wt/vol) agar to the corresponding media.

1.3. Cloning

E. coli (One Shot® ThermoFischer TOP10 Chemically Competent cells) was used in all subcloning procedures. 55 PCR was carried out using PhusionTM polymerase. All PCR products were purified with NucleospinTM Gel and PCR Clean-up by Macherey-NagelTM following the manufacturer's protocol. The purified fragments were digested with restriction enzyme DpnI in 1× FD buffer with 1 μ l enzyme 60 in a total volume of 34 μ l. The digested reaction was again purified with Nucleospin Gel and PCR Clean-up by Macherey-Nagel following the manufacturer's protocol. Gibson assembly was performed in 10 μ l reactions following the manufacturer's protocol (NewEngland Biolab). 65

Plasmid DNA was prepared using Qiagen kits according to the manufacturer's instructions. Modifications for Grampositive strains included growing bacteria in a medium supplemented with 0.5% glycine and lysozyme to facilitate cell lysis.

1.4. Transformation

1.4.1 Electro-Competent E. coli Cells and Transformation

Commercially electrocompetent cells were used for cloning and the experiments (One Shot® ThermoFischer TOP10 electrompetent *E. coli*). Electroporation was done using standard settings: 1800 V, 25 μ F and 200 SI using an Electro Cell Manipulator (BTX Harvard Apparatus ECM630). Following the pulse, 1 ml LB-SOC media was added and the cells were incubated at 37° C. for 1 hour. The transformed cells were plated in LB-agar containing the corresponding antibiotics.

1.5. Activation of sgRNA-Cas9 in *E. coli* and Consortium Experiments.

E. coli TOP10 and Nissle both with the plasmid containing the sgRNA targeting the ribosomal RNA-encoding sequence of K-12 derived strains and the other bacteria were grown overnight in 3 ml of TH broth. The next day the cells were diluted to ~OD 0.5 and next 10-fold serially diluted in TH media and using a 96-well replicator (Mettler Toledo LiquidatorTM 96) 4 μ L volume drops were spotted on TH agar, TH agar with inducers (1% arabinose and 2 mM theophylline), TH agar supplemented with 2.5 g l⁻¹ PEA and MacConkey agar supplemented with 1% maltose. The plates were incubated for 20 h at 37° C. and the colony forming units (CFU) were calculated from triplicate measurements.

2. Results

2.1 Specific Targeting of *E. coli* Strains Using an Exogenous CRISPR-Cas9 System

We first tested if the system could differentiate between two *E. coli* strains by introducing the killing system in both *E. coli* TOP10 and Nissle.

2.1 Targeting of *E. coli* Using an Exogenous CRISPR-Cas9 System in a Mixed Culture

Serial dilutions of overnight cultures were done in duplicate for both *E. coli* strains, *B. subtilis, L. lactis*, and in triplicate for the mixed cultures. All strains were grown at 37° C. for 20 hours in selective plates with and without the inducers. Induction of the system activates the sgRNA-Cas9 targeting K-12 derived strains, while leaving intact the other bacteria.

Distinguishing the different bacteria from a mixed culture is important in order to determine cell numbers of the different species and determine the specific removal of a species. MacConkey agar selectively grows *E. coli*, PEA agar is a selective medium that is used for the isolation of gram-positive (*B. subtilis*) from gram-negative (*E. coli*). Additionally, we found that different concentrations of PEA partially inhibit the growth of other gram positives. 2.5 g l⁻¹ of PEA proved to selectively grow *B. subtilis* while limiting growth of *E. coli* and *L. lactis*.

FIG. 14 shows specific targeting of *E. coli* strain by the inducible, exogenous, vector-borne CRISPR-Cas system. The sgRNA target the genome of K-12 derived *E. coli* strain *E. coli* TOP10, while the other *E. coli* strain tested was unaffected.

FIG. **15** shows spot assay with serial dilutions of individual bacterial species used in this study and mixed culture in TH agar without induction of the CRISPR-Cas9 system.

FIG. 16 shows a spot assay of the dilution 10^3 on different selective media. TH with 2.5 g l⁻¹ PEA is a selective media for B. subtilis alone. MacConkey supplemented with maltose is a selective and differential culture medium for bacteria designed to selectively isolate Gram-negative and enteric 5 bacilli and differentiate them based on maltose fermentation. Therefore TOP10 AmalK mutant makes white colonies on the plates while Nissle makes pink colonies; A is E coli AmalK, B is E coli Nissile, C is B subtilis, D is L lactis, E is mixed culture; the images at MacConkey-/B and E appear 10 pink; the images at MacConkey+/B and E appear pink. FIG. 17 shows selective growth of the bacteria used in this study on different media and selective plates. It can be seen that we clearly, selectively killed the target E coli strain ("E coli" on x-axis in FIG. 17) in the mixed population, whereas the other 15 related strain ("E coli-Nissle") was not similarly killed

Killing of the target strain in the mixed population was 1000-fold in this experiment.

REFERENCES

- Zhang, X. Z., & Zhang, Y. H. P. (2011). Simple, fast and high-efficiency transformation system for directed evolution of cellulase in *Bacillus subtilis*. *Microbial Biotechnology*, 4(1), 98-105. http://doi.org/10.1111/j.1751-7915.2010.00230.x
- Wegmann, U., O'Connell-Motherway, M., Zomer, A., Buist, G., Shearman, C., Canchaya, C., ... Kok, J. (2007). Complete genome sequence of the prototype lactic acid bacterium *Lactococcus lactis* subsp. *cremoris* MG1363. *Journal of Bacteriology*, 189(8), 3256-70. http://doi.org/ 10.1128/JB 0.01768-06

IADLE I	TABLE	1
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Abiotrophia	Acidocella	Actinomyces	Alkalilimnicola	Aquaspirillum
Abiotrophia	Acidocella	Actinomyces bovis	Alkalilimnicola	Aquaspirillum
defectiva	aminolytica	Actinomyces vovis Actinomyces	ehrlichii	polymorphum
Acaricomes	Acidocella facilis	denticolens	Alkaliphilus	Aquaspirillum
Acaricomes	Acidomonas	Actinomyces	Alkaliphilus	putridiconchylium
phytoseiuli	Acidomonas	europaeus	oremlandii	Aquaspirillum
Acetitomaculum	methanolica	Actinomyces	Alkaliphilus	serpens
Acetitomaculum	Acidothermus	georgiae	transvaalensis	Aquimarina
ruminis	Acidothermus	Actinomyces	Allochromatium	Aquimarina
Acetivibrio	cellulolyticus	gerencseriae	Allochromatium	latercula
Acetivibrio	Acidovorax	Actinomyces	vinosum	Arcanobacterium
cellulolyticus	Acidovorax	hordeovulneris	Alloiococcus	Arcanobacterium
Acetivibrio	anthurii	Actinomyces	Alloiococcus otitis	haemolvticum
ethanolgignens	Acidovorax caeni	howellii	Allokutzneria	Arcanobacterium
Acetivibrio	Acidovorax	Actinomyces	Allokutzneria albata	pyogenes
multivorans	cattlevae	hyovaginalis	Altererythrobacter	Archangium
Acetoanaerobium	Acidovorax citrulli	Actinomyces	Altererythrobacter	Archangium
Acetoanaerobium	Acidovorax	israelii	ishigakiensis	gephyra
noterae	defluvii	Actinomyces	Altermonas	Arcobacter
Acetobacter	Acidovorax	johnsonii	Altermonas	Arcobacter butzler
Acetobacter aceti	delafieldii	Actinomyces	haloplanktis	Arcobacter
Acetobacter	Acidovorax facilis	meveri	Altermonas	crvaerophilus
cerevisiae	Acidovorax	Actinomyces	macleodii	Arcobacter
Acetobacter	koniaci	naeslundii	Alvsiella	halophilus
cibinongensis	Acidovorax	Actinomyces neuii	Alysiella crassa	Arcobacter
Acetobacter	temperans	Actinomyces	Alysiella filiformis	nitrofigilis
estunensis	Acidovorax	odontolyticus	Aminobacter	Arcobacter
Acetobacter	valerianellae	Actinomyces oris	Aminobacter	skirrowii
fabarum	Acinetobacter	Actinomyces	aganoensis	Arhodomonas
Acetobacter	Acinetobacter	radingae	Aminobacter	Arhodomonas
ghanensis	baumannii	Actinomyces	aminovorans	aquaeolei
Acetobacter	Acinetobacter	slackii	Aminobacter	Arsenophonus
indonesiensis	baylyi	Actinomyces	niigataensis	Arsenophonus
Acetobacter	Acinetobacter	turicensis	Aminobacterium	nasoniae
lovaniensis	bouvetii	Actinomyces	Aminobacterium	Arthrobacter
Acetobacter	Acinetobacter	viscosus	mobile	Arthrobacter agilis
malorum	calcoaceticus	Actinoplanes	Aminomonas	Arthrobacter albus
Acetobacter	Acinetobacter	Actinoplanes	Aminomonas	Arthrobacter
nitrogenifigens	gerneri	auranticolor	paucivorans	aurescens
Acetobacter oeni	Acinetobacter	Actinoplanes	Ammoniphilus	Arthrobacter
Acetobacter	haemolyticus	brasiliensis	Ammoniphilus	chlorophenolicus
orientalis	Acinetobacter	Actinoplanes	oxalaticus	Arthrobacter
Acetobacter	johnsonii	consettensis	Ammoniphilus	citreus
orleanensis	Acinetobacter junii	Actinoplanes	oxalivorans	Arthrobacter
Acetobacter	Acinetobacter	deccanensis	Amphibacillus	crystallopoietes
pasteurianus	lwoffi	Actinoplanes	Amphibacillus	Arthrobacter
Acetobacter	Acinetobacter	derwentensis	xylanus	cumminsii
pornorurn	parvus	Actinoplanes	Amphritea	Arthrobacter
Acetobacter	Acinetobacter	digitatis	Amphritea balenae	globiformis
senegalensis	radioresistens	Actinoplanes	Amphritea japonica	Arthrobacter
Acetobacter	Acinetobacter	durhamensis	Amycolatopsis	histidinolovorans
xylinus	schindleri	Actinoplanes	Amycolatopsis alba	Arthrobacter ilicis
Acetobacterium	Acinetobacter soli	ferrugineus	Amycolatopsis	Arthrobacter luteu
Acetobacterium	Acinetobacter	Actinoplanes	albidoflavus	Arthrobacter
bakii	tandoii	globisporus	Amycolatopsis	methylotrophus
Acetobacterium	Acinetobacter	Actinoplanes	azurea	Arthrobacter

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TABLE 1-continued

	EXAMPLE BACTERI	A	
Acinetobacter	Actinoplanes	coloradensis	Arthrobacter
towneri	italicus	Amycolatopsis	nicotianae
Acinetobacter	Actinoplanes	lurida	Arthrobacter
ursingii	liguriensis	Amycolatopsis	nicotinovorans
Acinetobacter	Actinoplanes	mediterranei	Arthrobacter
venetianus	lobatus	Amycolatopsis	oxydans
Acrocarpospora	Actinoplanes	rifamycinica	Arthrobacter
Acrocarpospora	missouriensis	Amycolatopsis	pascens
corrugata	Actinoplanes	rubida	Arthrobacter
Acrocarpospora	palleronii	Amycolatopsis	phenanthrenivorans
macrocephala	Actinoplanes	sulphurea Amycolatopsis	Arthrobacter polychromogenes
Acrocarpospora pleiomorpha	philippinensis Actinoplanes	tolypomycina	Atrhrobacter
Actibacter	rectilineatus	Anabaena	protophormiae
Actibacter	Actinoplanes	Anabaena cylindrica	Arthrobacter
sediminis	regularis	Anabaena flosaquae	psychrolactophilus
Actinoalloteichus	Actinoplanes	Anabaena variabilis	Arthrobacter
Actinoalloteichus	teichomyceticus	Anaeroarcus	ramosus
cyanogriseus	Actinoplanes	Anaeroarcus	Arthrobacter
Actinoalloteichus	utahensis	burkinensis	sulfonivorans
hymeniacidonis	Actinopolyspora	Anaerobaculum	Arthrobacter
Actinoalloteichus	Actinopolyspora	Anaerobaculum	sulfureus
spitiensis	halophila	mobile	Arthrobacter
Actinobaccillus	Actinopolyspora	Anaerobiospirillum	uratoxydans
Actinobacillus	mortivallis	Anaerobiospirillum	Arthrobacter
capsulatus	Actinosynnema	succiniciproducens	ureafaciens
Actinobacillus	Actinosynnema	Anaerobiospirillum	Arthrobacter
delphinicola	mirum	thomasii	viscosus
Actinobacillus	Actinotalea	Anaerococcus	Arthrobacter
hominis	Actinotalea	Anaerococcus	woluwensis
Actinobacillus	fermentans	hydrogenalis	Asaia
indolicus	Aerococcus	Anaerococcus	Asaia bogorensis
Actinobacillus	Aerococcus	lactolyticus	Asanoa
lignieresii Actinobacillus	sanguinicola Aerococcus urinae	Anaerococcus	Asanoa ferruginea Asticcacaulis
minor	Aerococcus urinue Aerococcus	prevotii Anaerococcus	Asticcacaulis
Actinobacillus	urinaeequi	tetradius	biprosthecium
muris	Aerococcus	Anaerococcus	Asticcacaulis
Actinobacillus	urinaehominis	vaginalis	excentricus
pleuropneumoniae	Aerococcus	Anaerofustis	Atopobacter
Actinobacillus	viridans	Anaerofustis	Atopobacter
porcinus	Aeromicrobium	stercorihominis	phocae
Actinobacillus	Aeromicrobium	Anaeromusa	Atopobium
rossii	erythreum	Anaeromusa	Atopobium fossor
Actinobacillus	Aeromonas	acidaminophila	Atopobium
scotiae	Aeromonas	Anaeromyxobacter	minutum
Actinobacillus	allos a ccharophila	Anaeromyxobacter	Atopobium
seminis	Aeromonas	dehalogenans	parvulum
Actinobacillus	bestiarum	Anaerorhabdus	Atopobium rimae
succinogenes	Aeromonas caviae	Anaerorhabdus	Atopobium vaginae
Actinobaccillus	Aeromonas	furcosa	Aureobacterium
suis Actinohacillus	encheleia	Anaerosinus Anaerosinus	Aureobacterium
11011100 000101110	Aeromonas	11111111100000000	barkeri
Actinghamlum	enteropelogenes	glycerini Apagrovincula	Aurobacterium
Actinobaculum Actinobaculum	Aeromonas eucrenophila	Anaerovirgula Anaerovirgula	Aurobacterium liquefaciens
massiliense	Aeromonas	multivorans	Avibacterium
Actinobaculum	ichthiosmia	Ancalomicrobium	Avibacterium avium
schaalii	Aeromonas	Ancalomicrobium	Avibacterium
Actinobaculum	jandaei	adetum	gallinarum
suis	Aeromonas media	Ancylobacter	Avibacterium
Actinomyces	Aeromonas	Ancylobacter	paragallinarum
urinale	popoffii	aquaticus	Avibacterium
Actinocatenispora	Aeromonas sobria	Aneurinibacillus	volantium
Actinocatenispora	Aeromonas veronii	Aneurinibacillus	Azoarcus
rupis	Agrobacterium	aneurinilyticus	Azoarcus indigens
Actinocatenispora	Agrobacterium	Aneurinibacillus	Azoarcus
thailandica	gelatinovorum	migulanus	tolulyticus
Actinocatenispora	Agrococcus	Aneurinibacillus	Azoarcus
sera	Agrococcus	thermoaerophilus	toluvorans
Actinocorallia	citreus	Angiococcus	Azohydromonas
Actinocorallia	Agrococcus	Angiococcus	Azohydromonas
aurantiaca	jenensis	disciformis	australica
Actinocorallia	Agromonas	Angulomicrobium	Azohydromonas
aurea	Agromonas	Angulomicrobium	lata
Actinocorallia	oligotrophica	tetraedrale	Azomonas
cavernae	Agromyces	Anoxybacillus	Azomonas avilis

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dehalogenans Acetobacterium fimetarium Acetobacterium malicum Acetobacterium paludosum Acetobacterium tundrae Acetobacterium wieringae Acetobacterium woodii Acetofilamentum Acetofilamentum rigidum Acetohalobium Acetohalobium arabaticum Acetomicrobium Acetomicrobium faecale Acetomicrobium flavidum Acetonema Acetonema longum Acetothermus Acetothermus paucivorans Acholeplasma Acholeplasma axanthum Acholeplasma brassicae Acholeplasma cavigenitalium Acholeplasma equifetale Acholeplasma granularum Acholeplasma hippikon Acholeplasma laidlawii Acholeplasma modicum Acholeplasma morum Acholeplasma multilocale Acholeplasma oculi Acholeplasma palmae Acholeplasma parvum Acholeplasma pleciae Acholeplasma vituli Achromobacter Achromobacter denitrificans Achromobacter insolitus Achromobacter piechaudii Achromobacter ruhlandii Achromobacter spanius Acidaminobacter Acidaminobacter hydrogenoformans Acidaminococcus Acidaminococcus fermentans

cavernae

Actinocorallia

Agromyces

Agromyces

Anoxybacillus

Anoxybacillus

Azomonas agilis

Azomonas insignis

Acetobacterium

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TABLE 1-continued

Bacteroides eggerthii Bacteroides fragilis Bacteroides galacturonicus Bacteroides helcogenes

Bifidobacterium angulatum Bifidobacterium animalis Bifidobacterium asteroides Bifidobacterium bifidum

Borrelia carolinensis Borrelia coriaceae Borrelia garinii Borrelia japonica Bosea Bosea minatitlanensis

alba Brevundimonas aurantiaca Brevundimonas diminuta Brevundimonasintermedia Brevundimonas

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TABLE 1-continued

EXAMPLE BACTERIA

Bacteroides ovatus Bacteroides pectinophilus Bacteroides pyogenes Bacteroides salyersiae Bacteroides stercoris Bacteroides suis Bacteroides tectus Bacteroides thetaiotaomicron Bacteroides uniformis Bacteroides ureolvticus **Bacteroides** vulgatus Balnearium Balnearium lithotrophicum Balneatrix Balneatrix alpica Balneola Balneola vulgaris Barnesiella Barnesiella viscericola Bartonella Bartonella alsatica Bartonella bacilliformis Bartonella clarridgeiae Bartonella doshiae Bartonella elizabethae Bartonella grahamii Bartonella henselae Bartonella rochalimae Bartonella vinsonii Bavariicoccus Bavariicoccus seileri Bdellovibrio Bdellovibrio bacteriovorus Bdellovibrio exovorus Beggiatoa Beggiatoa alba Beijerinckia Beijerinckia derxii Beijerinckia fluminensis Beijerinckia indica Beijerinckia mobilis Belliella Belliella baltica Bellilinea Bellilinea caldifistulae Belnapia Belnapia moabensis

Bifidobacterium boum Bifidobacterium breve Bifidobacterium catenulatum Bifidobacterium choerinum Bifidobacterium coryneforme Bifidobacterium cuniculi Bifidobacterium dentium Bifidobacterium gallicum Bifidobacterium gallinarum Bifidobacterium indicum Bifidobacterium longum Bifidobacterium magnumBifidobacterium mervcicum Bifidobacterium minimum **Bifidohacterium** pseudocatenulatum Bifidobacterium pseudolongum Bifidobacterium pullorum Bifidohacterium ruminantium Bifidobacterium saeculare Bifidobacterium subtile Bifidobacterium thermophilumBilophila Bilophila wadsworthia Biostraticola Biostraticola tofi Bizionia Bizionia argentinensis Blastobacter Blastobacter capsulatus Blastobacter denitrificans Blastococcus Blastococcus aggregatus Blastococcus saxobsidens Blastochloris Blastochloris viridis Blastomonas Blastomonas natatoria Blastopirellula Blastopirellula marina Blautia Blautia coccoides Blautia hansenii Blautia producta Blautia wexlerae Bogoriella Bogoriella caseilvtica Bordetella Bordetella avium Bordetella bronchiseptica Bordetella hinzii Bordetella holmesii Bordetella parapertussis Bordetella pertussis

Bosea thiooxidans Brachybacterium Brachybacterium alimentarium Brachybacterium faecium Brachybacterium paraconglomeratum Brachybacterium rhamnosum Brachybacterium tyrofermentans Brachyspira Brachyspira alvinipulli Brachyspira hyodysenteriae Brachyspira innocens Brachyspira murdochii Brachyspira pilosicoli Bradyrhizobium Bradvrhizobium canariense Bradyrhizobium elkanii Bradyrhizobium japonicum Bradyrhizobium liaoningense Brenneria Brenneria alni Brenneria nigrifluens Brenneria quercina Brenneria quercina Brenneria salicis Brevibacillus Brevibacillus agri Brevibacillus borstelensis Brevibacillus brevis Brevibacillus centrosporus Brevibacillus choshinensis Brevibacillus invocatus Brevibacillus laterosporus Brevibacillus parabrevis Brevibacillus reuszeri Brevibacterium Brevibacterium abidum **Brevihacterium** album Brevibacterium aurantiacum Brevibacterium celere Brevibacterium epidermidis Brevibacterium frigoritolerans Brevibacterium halotolerans Brevibacterium iodinum Brevibacterium linens Brevibacterium lyticum

subvibrioides Brevundimonas vancanneytii Brevundimonas variabilis Brevundimonas vesicularis Brochothrix Brochothrix campestris Brochothrix thermosphacta Brucella Brucella canis Brucella neotomae Bryobacter Bryobacter aggregatus Burkholderia Burkholderia ambifaria Burkholderia andropogonis Burkholderia anthina Burkholderia caledonica Burkholderia caryophylli Burkholderia cenocepacia Burkholderia cepacia Burkholderia cocovenenans Burkholderia dolosa Burkholderia fungorum Burkholderia glathei Burkholderia glumae Burkholderia graminis Burkholderia kururiensis Burkholderia multivorans Burkholderia phenazinium . Burkholderia plantarii , Burkholderia pyrrocinia Burkholderia silvatlantica Burkholderia stabilis Burkholderia thailandensis Burkholderia tropica Burkholderia unamae Burkholderia vietnamiensis Buttiauxella Buttiauxella agrestis Buttiauxella brennerae Buttiauxella ferragutiae Buttiauxella gaviniae

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TABLE 1-continued

	EXAMPLE BACTERIA					
Bet den Bet Bet	0	detella petrii detella trematum	Brevibacterium mcbrellneri Brevibacterium otitidis Brevibacterium oxydans Brevibacterium paucivorans Brevibacterium stationis	Buttiauxella izardii Buttiauxella noackiae Buttiauxella warmboldiae Butyrivibrio fibrisolvens Butyrivibrio fibrisolvens Butyrivibrio hungatei Butyrivibrio proteoclasticus		
Bacillus						
B. acidiceler	B. aminovorans	B. glucanolyticus	B. taeanensis	B. lautus		
B. acidicola B. acidiproducens	B. amylolyticus B. andreesenii	B. gordonae B. gottheilii	B. tequilensis B. thermantarcticus	B. lehensis B. lentimorbus		
B. acidocaldarius	B. aneurinilyticus	B. graminis	B. thermoaerophilus	B. lentus		
B. acidoterrestris	B. anthracis	B. halmapalus	B. thermoamylovorans	B. licheniformis		
B. aeolius	B. aquimaris	B. haloalkaliphilus	B. thermocatenulatus	B. ligniniphilus		
B. aerius	B. arenosi	B. halochares	B. thermocloacae	B. litoralis		
B. aerophilus	B. arseniciselenati		B. thermocopriae	B. locisalis		
B. agaradhaerens B. agri	B. arsenicus B. aurantiacus	B. halodurans B. halophilus	B. thermodenitrificans B. thermoglucosidasius	B. luciferensis B. luteolus		
B. aidingensis	B. arvi	B. halosaccharovorans		B. luteus		
B. akibai	B. aryabhattai	B. hemicellulosilyticus		B. macauensis		
B. alcalophilus	B. asahii	B. hemicentroti	B. thermophilus	B. macerans		
B. algicola	B. atrophaeus	B. herbersteinensis	B. thermoruber	B. macquariensis		
B. alginolyticus B. alkalidiazotrophicus	B. axarquiensis B. azotofixans	B. horikoshii B. horneckiae	B. thermosphaericus B. thiaminolyticus	B. macyae B. malacitensis		
3. alkalinitrilicus	B. azotoformans	B. horti	B. thioparans	B. mannanilyticus		
3. alkalisediminis	B. badius	B. huizhouensis	B. thuringiensis	B. marisflavi		
B. alkalitelluris	B. barbaricus	B. humi	B. tianshenii	B. marismortui		
B. altitudinis	B. bataviensis	B. hwajinpoensis	B. trypoxylicola	B. marmarensis		
B. alveayuensis B. alvei	B. beijingensis B. benzoevorans	B. idriensis B. indicus	B. tusciae B. validus	B. massiliensis B. megaterium		
B. amyloliquefaciens	B. beringensis	B. infantis	B. vallismortis	B. mesonae		
B.a. subsp. amyloliquefac	ciens B. berkeleyi	B. infernus	B. vedderi	B. methanolicus		
B.a. subsp. plantarum	B. beveridgei	B. insolitus	B. velezensis	B. methylotrophicu:		
B. dipsosauri B. drentensis	B. bogoriensis B. boroniphilus	B. invictae B. iranensis	B. vietnamensis B. vireti	B. migulanus B. mojavensis		
3. edaphicus	B. borstelensis	B. isabeliae	B. vulcani	B. mojuvensis B. mucilaginosus		
B. ehimensis	B. brevis Migula	B. isronensis	B. wakoensis	B. muralis		
3. eiseniae	B. butanolivorans	B. jeotgali	B. weihenstephanensis	B. murimartini		
3. enclensis	B. canaveralius	B. kaustophilus	B. xiamenensis	B. mycoides		
B. endophyticus	B. carboniphilus B. cecembensis	B. kobensis B. kochii	B. xiaoxiensis	B. naganoensis B. nanhaiensis		
3. endoradicis 3. farraginis	B. cellulosilyticus	B. kokeshiiformis	B. zhanjiangensis B. peoriae	B. nanhaiisediminis		
3. fastidiosus	B. centrosporus	B. koreensis	B. persepolensis	B. nealsonii		
3. fengqiuensis	B. cereus	B. korlensis	B. persicus	B. neidei		
3. firmus	B. chagannorensis		B. pervagus	B. neizhouensis		
3. flexus 3. foraminis	B. chitinolyticus B. chondroitinus	B. krulwichiae B. laevolacticus	B. plakortidis B. pocheonensis	B. niabensis B. niacini		
3. fordii	B. choshinensis	B. larvae	B. polygoni	B. novalis		
3. formosus	B. chungangensis	B. laterosporus	B. polymyxa	B. oceanisediminis		
3. fortis	B. cibi	B. salexigens	B. popilliae	B. odysseyi		
3. fumarioli	B. circulans	B. saliphilus	B. pseudalcalophilus	B. okhensis		
3. funiculus 3. fusiformis	B. clarkii B. clausii	B. schlegelii B. sediminis	B. pseudofirmus B. pseudomycoides	B. okuhidensis B. oleronius		
3. galactophilus	B. coagulans	B. selenatarsenatis	B. psychrodurans	B. oryzaecorticis		
B. galactosidilyticus	B. coahuilensis	B. selenitireducens	B. psychrophilus	B. oshimensis		
B. galliciensis	B. cohnii	B. seohaeanensis	B. psychrosaccharolyticus	B. pabuli		
B. gelatini B. cibronii	B. composti	B. shacheensis	B. psychrotolerans	B. pakistanensis		
B. gibsonii B. ginsengi	B. curdlanolyticus B. cycloheptanicus		B. pulvifaciens B. pumilus	B. pallidus B. pallidus		
5. ginsengi B. ginsengihumi	B. cyclonepiunicus B. cytotoxicus	B. silvestris	B. purgationiresistens	B. panacisoli		
B. ginsengisoli	B. daliensis	B. simplex	B. pycnus	B. panaciterrae		
B. globisporus	B. decisifrondis	B. siralis	B. qingdaonensis	B. pantothenticus		
eg, B.g. subsp.	B. decolorationis	B. smithii B. soli	B. qingshengii	B. parabrevis		
<i>Globisporus</i> ; or B.g. subsp.	B. deserti	B. soli B. solimangrovi	B. reuszeri B. rhizosphaerae	B. paraflexus B. pasteurii		
Marinus)		B. solisalsi	В. rnizospnaerae B. rigui	B. patagoniensis		
		B. songklensis	B. ruris	P		
		B. sonorensis	B. safensis			
		B. sphaericus	B. salarius			
		R snorothermodurans				

B. sphaericus B. sporothermodurans B. stearothermophilus

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TABLE 1-continued

	TABL	E 1-continued		
	EXAMI	PLE BACTERIA		
B. stratosphericus B. subterraneus B. subtilis (eg, B.s. subsp. Inaquosorum; or B.s. subsp. Spizizeni; or B.s. subsp. Subtilis)				
Caenimonas Caenimonas koreensis Caldalkalibacillus Caldalkalibacillus uzonensis Caldanaerobacter Caldanaerobacter Subterraneus Caldanaerobius Caldanaerobius Caldanaerobius caldanaerobius zeae Caldanaerobius zeae Caldanaerovirga acetigignens Caldicellulosiruptor Caldicellulosiruptor kristjanssonii Caldicellulosiruptor owensensis	Campylobacter Campylobacter concisus Campylobacter curvus Campylobacter curvus Campylobacter gracilis Campylobacter helveticus Campylobacter hominis Campylobacter hyointestinalis Campylobacter jejuni Campylobacter mucosalis Campylobacter rectus Campylobacter showae Campylobacter showae Campylobacter sputorum Campylobacter sputorum Campylobacter sputorum Campylobacter sputorum Campylobacter upsaliensis Capnocytophaga connocytophaga granulosa Capnocytophaga funcosalis Capnocytophaga cannocytophaga	Cardiobacterium Cardiobacterium hominis Carnimonas Carnimonas Carnobacterium Carnobacterium alterfinditum Carnobacterium divergens Carnobacterium funditum Carnobacterium gallinarum Carnobacterium maltaromaticum Carnobacterium mobile Carnobacterium wiridans Carnobacterium viridans Carnobacterium taltaromaticum Carnobacterium viridans Caryophanon latum Caryophanon tenue Catellatospora citrea Catellatospora methionotrophica Catenococcus Catenococcus Catenococcus	Catenuloplanes Catenuloplanes atrovinosus Catenuloplanes castaneus Catenuloplanes crispus Catenuloplanes indicus Catenuloplanes indicus Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes nepalensis Catenuloplanes Corynebacterium Corynebacterium flavescens Corynebacterium variabile	Curtobacterium Curtobacterium albidum Curtobacterium citreus

Clostridium

Clostridium absonum, Clostridium aceticum, Clostridium acetireducens, Clostridium acetobutylicum, Clostridium acidisoli, Clostridium aciditolerans, Clostridium acidurici, Clostridium aerotolerans, Clostridium aestuarii, Clostridium akagii, Clostridium aldenense, Clostridium aldrichii, Clostridium algidicarni, Clostridium algidixylanolyticum, Clostridium algifaecis, Clostridium algoriphilum, Clostridium alkalicellulosi, Clostridium aminophilum, Clostridium aminovalericum, Clostridium amygdalinum, Clostridium amylolyticum, Clostridium arbusti, Clostridium arcticum, Clostridium argentinense, Clostridium asparagiforme, Clostridium aurantibutyricum, Clostridium autoethanogenum, Clostridium baratii, Clostridium barkeri, Clostridium bartlettii, Clostridium beijerinckii, Clostridium bifermentans, Clostridium bolteae, Clostridium bornimense, Clostridium botulinum, Clostridium bowmanii, Clostridium bryantii, Clostridium butyricum, Clostridium cadaveris, Clostridium caenicola, Clostridium caminithermale, Clostridium carboxidivorans, Clostridium carnis, Clostridium cavendishii, Clostridium celatum, Clostridium celerecrescens, Clostridium cellobioparum, Clostridium cellulofermentans, Clostridium cellulolyticum, Clostridium cellulosi, Clostridium cellulovorans, Clostridium chartatabidum, Clostridium chauvoei, Clostridium chromiireducens, Clostridium citroniae, Clostridium clariflavum, Clostridium clostridioforme, Clostridium coccoides, Clostridium cochlearium, Clostridium colletant, Clostridium colicanis, Clostridium colinum, Clostridium collagenovorans, Clostridium cylindrosporum, Clostridium difficile, Clostridium diolis, Clostridium disporicum, Clostridium drakei, Clostridium durum, Clostridium estertheticum, Clostridium estertheticum estertheticum, Clostridium estertheticum laramiense, Clostridium fallax, Clostridium felsineum, Clostridium fervidum, Clostridium fimetarium, Clostridium formicaceticum, Clostridium frigidicarnis, Clostridium frigoris, Clostridium ganghwense, Clostridium gasigenes, Clostridium ghonii, Clostridium glycolicum, Clostridium glycyrrhizinilyticum, Clostridium grantii, Clostridium haemolyticum, Clostridium halophilum, Clostridium hastiforme, Clostridium hathewayi, Clostridium herbivorans, Clostridium hiranonis, Clostridium histolyticum, Clostridium homopropionicum, Clostridium huakuii, Clostridium hungatei, Clostridium hydrogeniformans,

sputigena

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TABLE 1-continued

EXAMPLE BACTERIA

Clostridium hydroxybenzoicum, Clostridium hylemonae, Clostridium jejuense, Clostridium indolis, Clostridium innocuum, Clostridium intestinale, Clostridium irregulare, Clostridium isatidis, Clostridium josui, Clostridium kluyveri, Clostridium lactatifermentans, Clostridium lacusfryxellense, Clostridium laramiense, Clostridium lavalense, Clostridium lentocellum, Clostridium lentoputrescens, Clostridium leptum, Clostridium limosum, Clostridium litorale, Clostridium lituseburense, Clostridium ljungdahlii, Clostridium lortetii, Clostridium lundense, Clostridium magnum, Clostridium malenominatum, Clostridium mangenotii, Clostridium mayombei, Clostridium methoxybenzovorans, Clostridium methylpentosum, Clostridium neopropionicum, Clostridium nexile, Clostridium nitrophenolicum, Clostridium novyi, Clostridium oceanicum, Clostridium orbiscindens, Clostridium oroticum, Clostridium oxalicum, Clostridium papyrosolvens, Clostridium paradoxum, Clostridium paraperfringens (Alias: C. welchii), Clostridium paraputrificum, Clostridium pascui, Clostridium pasteurianum, Clostridium peptidivorans, Clostridium perenne, Clostridium perfringens, Clostridium pfennigii, Clostridium phytofermentans, Clostridium piliforme, Clostridium polysaccharolyticum, Clostridium populeti, Clostridium propionicum, Clostridium proteoclasticum, Clostridium proteolyticum, Clostridium psychrophilum, Clostridium puniceum, Clostridium purinilyticum, Clostridium putrefaciens, Clostridium putrificum, Clostridium quercicolum, Clostridium quinii, Clostridium ramosum, Clostridium rectum, Clostridium roseum, Clostridium saccharobutylicum, Clostridium saccharogumia, Clostridium saccharolyticum, Clostridium saccharoperbutylacetonicum, Clostridium sardiniense, Clostridium sartagoforme, Clostridium scatologenes, Clostridium schirmacherense, Clostridium scindens, Clostridium septicum, Clostridium sordellii, Clostridium sphenoides, Clostridium spiroforme, Clostridium sporogenes, Clostridium sporosphaeroides, Clostridium stercorarium, Clostridium stercorarium leptospartum, Clostridium stercorarium stercorarium. Clostridium stercorarium thermolacticum. Clostridium sticklandii, Clostridium straminisolvens, Clostridium subterminale, Clostridium sufflavum, Clostridium sulfidigenes, Clostridium symbiosum, Clostridium tagluense, Clostridium tepidiprofundi, Clostridium termitidis, Clostridium tertium, Clostridium tetani, Clostridium tetanomorphum, Clostridium thermaceticum, Clostridium thermautotrophicum, Clostridium thermoalcaliphilum, Clostridium thermobutyricum, Clostridium thermocellum, Clostridium thermocopriae, Clostridium thermohydrosulfuricum, Clostridium thermolacticum, Clostridium thermopalmarium, Clostridium thermopapyrolyticum, Clostridium thermosaccharolyticum, Clostridium thermosuccinogenes, Clostridium thermosulfurigenes, Clostridium thiosulfatireducens, Clostridium tyrobutyricum, Clostridium uliginosum, Clostridium ultunense, Clostridium villosum, Clostridium vincentii, Clostridium viride, Clostridium xylanolyticum, Clostridium xylanovorans

Dactylosporangium Dactylosporangium aurantiacum Dactylosporangium fulvum Dactylosporangium matsuzakiense Dactylosporangium roseum Dactylosporangium thailandense Dactylosporangium winaceum	Deinococcus Deinococcus aerius Deinococcus apachensis Deinococcus aquaticus Deinococcus aquatilis Deinococcus caeni Deinococcus radiodurans Deinococcus radiophilus	Delftia Delftia acidovorans Desulfovibrio Desulfovibrio desulfuricans Diplococcus Diplococcus pneumoniae	Echinicola Echinicola pacifica Echinicola vietnamensis
Enterobacter E. aerogenes E. amnigenus E. agglomerans E. arachidis E. asburiae E. cancerogenous E. cloacae E. cowanii E. dissolvens E. gergoviae E. helveticus E. helveticus E. hormaechei E. intermedius	Enterobacter kobei E. ludwigii E. mori E. nimipressuralis E. oryzae E. pulveris E. pyrinus E. radicincitans E. taylorae E. turicensis E. sakazakii Enterobacter soli Enterococcus factores faccalis Enterococcus faccalis Enterococcus faccalis Enterococcus faccalis Enterococcus faccium Erwinia hapontici Escherichia Escherichia coli	Faecalibacterium Faecalibacterium prausnitzii Fangia Fangia hongkongensis Fastidiosipila sanguinis Fusobacterium rusobacterium nucleatum	Flavobacterium antarcticum Flavobacterium aquatile Flavobacterium aquatile Flavobacterium balustinum Flavobacterium croceum Flavobacterium cucumis Flavobacterium daejeonense Flavobacterium defluvii Flavobacterium defluvii Flavobacterium degerlachei Flavobacterium denitrificans Flavobacterium filum Flavobacterium filum Flavobacterium filevense Flavobacterium filevense Flavobacterium filevense Flavobacterium filevense Flavobacterium filevense Flavobacterium filevense

TABLE 1-continued

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		EXAMPLE BACT	EDIA		
		EAAMITLE DACT	EKIA	1	Flavobacterium
					okeanokoites
Gaetbulibacter	Haemoph		Ideonella		ibacter
Gaetbulibacter saemankumensis	Haemoph aegyptius		Ideonella azotifigens		nibacter ophelis
Gallibacterium	aegypnus Haemoph		Idiomarina		ubacter
Gallibacterium anatis			Idiomarina		allicola
Gallicola	Haemoph		abyssalis		ubacter
Gallicola barnesae	Haemoph		Idiomarina		osus
Garciella	gallinarii		baltica	Jan	iibacter
Garciella	Haemoph		Idiomarina	mei	lonis
nitratireducens	haemolyti		fontislapidosi		ubacter
Geobacillus	Haemoph		Idiomarina	teri	
Geobacillus	influenzae		loihiensis		naschia
thermoglucosidasius Geobacillus	Haemoph paracunio		Idiomarina ramblicola		maschia tauzzna
stearothermophilus	Haemoph		Idiomarina		taugens maschia
Geobacter	parahaen		seosinensis		golandensis
Geobacter	Haemoph		Idiomarina		maschia
bemidjiensis	parainflu		zobellii		angensis
Geobacter brememis	Haemoph	ilus	Ignatzschineria		maschia
Geobacter chapellei		haemolyticus	Ignatzschineria	rub	
Geobacter grbiciae	Haemoph	nilus	larvae		thinobacterium
Geobacter	parasuis	1	Ignavigranum		nthinobacterium
hydrogenophilus Cachactan laulani	Haemoph		Ignavigranum		uricidamnosum uthinobacterium
Geobacter lovleyi Geobacter	<i>pittmania</i> Hafnia	ie .	<i>ruoffiae</i> Ilumatobacter		ithinobacterium dum
metallireducens	Hafnia al	luei	Ilumatobacter	Jeji	
Geobacter pelophilus	~		fluminis	Jeji	
Geobacter pickeringi			Ilyobacter		lidilutea
Geobacter	ganghwei	nsis	Nyobacter	Jeo	tgalibacillus
sulfurreducens	Halalkali	bacillus	delafieldii	Jeo	tgalibacillus
Geodermatophilus	Halalkali	bacillus	Ilyobacter		nentarius
Geodermatophilus	halophilu	10	insuetus	Ieo	tgalicoccus
obscurus	Helicobad	cter	Ilyobacter	Jeo	tgalicoccus
Gluconacetobacter	Helicobad Helicobad	cter	Ilyobacter polytropus	Jeo	
Gluconacetobacter Gluconacetobacter	Helicobad	cter	Ilyobacter polytropus Ilyobacter	Jeo	tgalicoccus
Gluconacetobacter Gluconacetobacter xylinus	Helicobad Helicobad	cter	Ilyobacter polytropus	Jeo	tgalicoccus
Gluconacetobacter <i>Gluconacetobacter</i> <i>xylinus</i> Gordonia	Helicobad Helicobad	cter	Ilyobacter polytropus Ilyobacter	Jeo	tgalicoccus
Gluconacetobacter Gluconacetobacter xylinus	Helicobad Helicobad	cter	Ilyobacter polytropus Ilyobacter	Jeo	tgalicoccus
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta	Helicobad Helicobad	cter	Ilyobacter polytropus Ilyobacter	Jeo hal	tgalicoccus
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia Kaistia adipata	Helicobad Helicobad pylori Labedella Labedella	cter cter Listeria ivanovii L. marthii	Ilyobacter polytropus Ilyobacter tartaricus Microc Microc	Jeo hal	rgalicoccus otolerans Nesterenkonia Nesterenkonia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia Kaistia adipata Kaistia soli	Helicobad Helicobad pylori Labedella gwakjiensis	cter cter Listeria ivanovii L. marthii L. monocytogenes	Ilyobacter polytropus Ilyobacter tartaricus Microc Microc luteus	Jeo hal coccus coccus	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia
Gluconacetobacter Gluconacetobacter xylinus Gordonia gordonia rubripertincta Kaistia Kaistia adipata Kaistia soli Kangiella	Helicobad Helicobad pylori Labedella gwakjiensis Labrenzia	cter <i>Listeria ivanovii</i> <i>L. marthii</i> <i>L. monocytogenes</i> <i>L. newyorkensis</i>	Ilyobacter polytropus Ilyobacter tartaricus Microc Microc luteus Microc	Jeo hal coccus coccus	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia Kaistia adipata Kaistia soli Kangiella Kangiella	Helicobad Helicobad pylori Labedella gwakjiensis Labrenzia Labrenzia	cter Listeria ivanovii L. marthii L. monocytogenes L. newyorkensis L. riparia	Ilyobacter polytropus Ilyobacter tartaricus Microo <i>Microo</i> <i>luteus</i> <i>Microo</i> <i>luteus</i>	Jeo hal	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia Kaistia adipata Kaistia soli Kangiella Kangiella aquimarina	Helicobad Helicobad pylori Labedella gwakjiensis Labrenzia aggregata	cter <i>Listeria ivanovii</i> <i>L. marthii</i> <i>L. monocytogenes</i> <i>L. newyorkgenis</i> <i>L. riparia</i> <i>L. riparia</i>	Ilyobacter polytropus Ilyobacter tartaricus Microc Iuteus Microc luteus Microc luteus Microc	Jeo hal coccus coccus coccus ella	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia argentinensis
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella kangiella koreensis	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Labrenzia aggregata Labrenzia alba	cter <i>Listeria ivanovii</i> <i>L. marthii</i> <i>L. monocytogenes</i> <i>L. newyorkensis</i> <i>L. riparia</i> <i>L. rocourtiae</i> <i>L. seeligeri</i>	Ilyobacter polytropus Ilyobacter tartaricus Microc Iuteus Microc Iuteus Moraxu Moraxu	Jeo hal voccus voccus eccus ella ella bovis	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia argentinensis Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia Kaistia adipata Kaistia soli Kangiella aquimarina Kangiella koreensis Kerstersia	Helicobad Helicobad pylori Labedella gwakjiensis Labrenzia aggregata	cter <i>Listeria ivanovii</i> <i>L. marthii</i> <i>L. monocytogenes</i> <i>L. newyorkgenis</i> <i>L. riparia</i> <i>L. riparia</i>	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxu moraxu moraxu Moraxu	Jeo hal coccus coccus coccus ella ella ella bovis ella	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia argentinensis
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia aggregata Labrenzia alba Labrenzia alba	cter <i>Listeria ivanovii</i> <i>L. marthii</i> <i>L. monocytogenes</i> <i>L. newyorkensis</i> <i>L. riparia</i> <i>L. rocourtiae</i> <i>L. seeligeri</i> <i>L. weihenstephaner</i>	Ilyobacter polytropus Ilyobacter tartaricus Microc Iuteus Microc Iuteus Morax Morax Morax noniiqi	Jeo hal coccus coccus coccus ella ella bovis ella effaciens	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia argentinensis Nocardia corallina
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella Kangiella kerstersia Kerstersia Kioniella	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Labrenzia Labrenzia algaregata Labrenzia alba Labrenzia alexandrii	cter <i>Listeria ivanovii</i> <i>L. marthii</i> <i>L. monocytogenes</i> <i>L. newyorkensis</i> <i>L. riparia</i> <i>L. rocourtiae</i> <i>L. seeligeri</i> <i>L. weikenstephanen</i> <i>L. weishimeri</i>	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxu moraxu moraxu Moraxu	Jeo hal coccus coccus coccus ella ella bovis ella uefaciens ella	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Klebsiella	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Aggregata Labrenzia alba Labrenzia alexandrii Labrenzia marina Labrys Labrys	cter cter Listeria ivanovii L. marthii L. monocytogenes L. newyorkensis L. riparia L. rocourtiae L. seeligeri L. seeligeri L. seeligeri L. weihenstephanen L. stonella Listonella anguillarum	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxa Moraxa nonliqu Moraxa siss Moraxa nonliqu Moraxa Noraxa	Jeo hal	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Aquimarina Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Klebsiella	Helicobaa Helicobaa pylori	cter <i>Listeria ivanovii</i> <i>L. marthii</i> <i>L. monocytogenes</i> <i>L. riparia</i> <i>L. rocourtiae</i> <i>L. seeligeri</i> <i>L. weihenstephanet</i> <i>L. weihenstephanet</i> <i>L. weishimeri</i> <i>Listonella</i> <i>Listonella</i> <i>Listonella</i> <i>anguillarum</i> Macrococcus	Ilyobacter polytropus Ilyobacter tartaricus Microcc luteus Microc luteus Moraxx moraxx monliqu Moraxx nonliqu Moraxx Nakam Nakam	Jeo hal	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia adipata Kaistia soli Kangiella Kangiella aquimarina Kangiella koreensis Kerstersia gyiorum Kiloniella Kiloniella laminariae Klebsiella K. granulomatis K. oxytoca	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Aggregata Labrenzia aggregata Labrenzia alba Labrenzia alexandrii Labrenzia marina Labrys Labrys methylaminiphilus Labrys	cter cter Listeria ivanovii L. marthii L. monocytogenes L. niparia L. rocourtiae L. seeligeri L. seeligeri L. weihenstephanei L. weihstimeri Listonella Listonella Listonella Macrococcus Macrococcus	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc lylae Moraxa monitqu Moraxa nonitqu Nakam Multipo	Jeo hal	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Adigiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella Kiloniella laminariae Klebsiella K. granulomatis K. goxytoca K. pneumoniae	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia aggregata Labrenzia alba Labrenzia alba Labrenzia marina Labrenzia marina Labrys Labrys methylaminiphilus Labrys miyagiensis	cter cter Listeria ivanovii L. marthii L. monocytogenes L. riparia L. rocourtiae L. seeligeri L. weikenstephanen L. weikenstephanen L. weishimeri Listonella Listonella Listonella anguillarum Macrococcus Macrococcus	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc lylae Moraxu nonliqu Moraxu osloen: Nakam Nakam multipo Nannou	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia aoli Kangiella Aquimarina Kangiella koreensis Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella ks. granulomatis K. granulomatis K. speumoniae K. terrigena	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Labrenzia alexandrii Labrenzia marina Labrys Labrys Labrys methylaminiphilus Labrys miyagiensis Labrys monachus	cter cter Listeria ivanovii L. marthii L. monocytogenes L. newyorkensis L. rocourtiae L. seeligeri L. weihenstephanen L. weihenstephanen L. seeligeri L. weihenstephanen L. stonella Listonella Listonella anguillarum Macrococcus bovicus Marinobacter	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxa Moraxa nonliqu Moraxa Moraxa Noraxa Noraxa Nakam Nakam Multipo Nannon Nannon Nannon	Jeo hall coccus coccus coccus ella bovis ella uefaciens ella sis uurella urella urella urella urella cystis cystis	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia adipata Kangiella aquimarina Kangiella koreensis Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kilobsiella K. granulomatis K. oxytoca K. terrigena K. terrigena K. variicola	Helicobaa Helicobaa pylori	cter cter Listeria ivanovii L. marthii L. monocytogenes L. newyorkensis L. riparia L. rocourtiae L. seeligeri L. weihenstephaneri L. seeligeri L. weikshimeri Listonella anguillarum Macrococcus Macrococcus bovicus Marinobacter Marinobacter	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxa Moraxa nonliqu Moraxa sis Moraxa nonliqu Moraxa soloen: Nakam Nakam Nakam Nakam Nakam Nakam Nakam Nakam Nakam Nakam Nakam Nakam	Jeo hall voccus voccus voccus ella bovis ella ella covis ella sis ella sis ella urella urella urella urella urella vritia cystis cystis	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella Kiloniella laminariae Kiloniella K. granulomatis K. oxytoca K. preumoniae K. terrigena K. variicola	Helicobaa Helicobaa pylori Labedella Labedella Labenzia Labrenzia aggregata Labrenzia alba Labrenzia alba Labrenzia marina Labrys Labrys Labrys methylaminiphilus Labrys miyagiensis Labrys miyagiensis Labrys babrys monachus Labrys okinawensis	cter cter Listeria ivanovii L. marthii L. monocytogenes L. niparia L. rocourtiae L. seeligeri L. seeligeri L. weihenstephanei L. seeligeri L. weihstimeri Listonella Listonella Listonella Macrococcus Macrococcus Macrococcus Marinobacter Algicola	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc lylae Moraxa monilqu Moraxa osloem. Nakam Nakam Nakam Nakam Nakam Nakam Nakam	Jeo hal	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella aquimarina Kangiella koreensis Kerstersia guimarina Kangiella koreensis Kerstersia Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella koreansis K. granulomatis K. oxytoca K. preumoniae K. tertigena K. variicola K. luyvera ascorbata	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia aggregata Labrenzia alba Labrenzia alba Labrenzia marina Labrys Labrys methylaminiphilus Labrys miyagiensis Labrys miyagiensis Labrys okinawensis Labrys	cter cter Listeria ivanovii L. marthii L. monocytogenes L. niparia L. rocourtiae L. seeligeri L. seeligeri L. weihenstephanen L. seeligeri L. weishimeri Listonella Listonella Listonella anguillarum Macrococcus Macrococcus bovicus Marinobacter Agrinobacter Marinobacter	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc lylae Moraxu Moraxu nonliqu Moraxu osloen: Nakam multipu Nannov Nannov pusilla Natran Natran	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella koreensis Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella koreensis Kerstersia gyiorum Kiloniella koreensis Kerstersia gyiorum Kiloniella koreensis Kerstersia gyiorum Kiloniella koreensis Kiloniella koreensis Koreensis Koreensis Koreensis Koreensis Koreensis Kiloniella koreensis Koreensis Kiloniella koreensis Kiloniella koreensis Kiloniell	Helicobaa Helicobaa pylori Labedella Labedella Labenzia Labrenzia aggregata Labrenzia alba Labrenzia alba Labrenzia marina Labrys Labrys Labrys methylaminiphilus Labrys miyagiensis Labrys miyagiensis Labrys babrys monachus Labrys okinawensis	cter cter Listeria ivanovii L. marthii L. monocytogenes L. niparia L. rocourtiae L. seeligeri L. seeligeri L. weihenstephanei L. seeligeri L. weihstimeri Listonella Listonella Listonella Macrococcus Macrococcus Macrococcus Marinobacter Algicola	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Microc luteus Moraxu Moraxu nonliqu Moraxu osloens Nakam Moraxu Nakam Matran Natran Natran Natran	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia adipata Kangiella aquimarina Kangiella koreensis Kerstersia gyiorum Kiloniella koreensis Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella k. granulomatis K. spreumoniae K. terrigena K. variicola Kluyvera Kluyvera ascorbata Koccuria	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Labrenzia alexandrii Labrenzia aalba Labrenzia alexandrii Labrenzia marina Labrys Labrys Labrys methylaminiphilus Labrys miyagiensis Labrys miyagiensis Labrys okinawensis Labrys portucalensis	cter cter	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Microc luteus Moraxu Moraxu nonliqu Moraxu osloens Nakam Moraxu Nakam Matran Natran Natran Natran	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella kiloniella K. granulomatis K. syrioca K. pneumoniae K. tariicola K. variicola Kluyvera Kluyvera Kluyvera Kluyvera ascorbata Kocuria roasea Kocuria roasea Kocuria varians	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Labrenzia aggregata Labrenzia alba Labrenzia alba Labrenzia marina Labrys Labrys methylaminiphilus Labrys miyagiensis Labrys miyagiensis Labrys motachus Labrys portucalensis Labrys portucalensis Lacobacillus [see below] Laceyella	cter cter	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc lylae Morax Morax nonliqu Morax osloen: Nakam multipo Nanno Nakam multipo Nanno Natran Natran thermo Natran thermo Natran thermo Natran	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella kiloniella K. granulomatis K. syrioca K. pneumoniae K. tariicola K. variicola Kluyvera Kluyvera Kluyvera Kluyvera ascorbata Kocuria roasea Kocuria roasea Kocuria varians	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia aggregata Labrenzia alba Labrenzia alba Labrenzia marina Labrenzia marina Labrys Labrys methylaminiphilus Labrys miyagiensis Labrys okinawensis Labrys okinawensis Labrys portucalensis Lactobacillus [see below] Laceyella Laceyella putida	cter cter cter Listeria ivanovii L. marthii L. monocytogenes L. newyorkensis L. riparia L. rocourtiae L. seeligeri L. weishimeri Listonella digeri Listonella Listonella anguillarum Macrococcus bovicus Marinobacter Marinobacter algicola Marinobacter bryozoorum Marinobacter flavimaris Meiothermus	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxu Moraxu nonliqu Moraxu osloen: Nakam multipa Nanno Narmo pusilla Natran Natran Natran thermo Narva	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella kiloniella K. granulomatis K. syrioca K. pneumoniae K. tariicola K. variicola Kluyvera Kluyvera Kluyvera Kluyvera Kluyvera ascorbata Kocuria roasea Kocuria roasea Kocuria varians	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Labrenzia alexandrii Labrenzia alba Labrenzia alexandrii Labrenzia marina Labrys Labrys Labrys methylaminiphilus Labrys miyagiensis Labrys miyagiensis Labrys okinawensis Labrys portucalensis Lacopella Laceyella Laceyella putida Lechevalieria	cter cter cter Listeria ivanovii L. marthii L. monocytogenes L. newyorkensis L. riparia L. rocourtiae L. seeligeri L. weikenstephanet L. seeligeri L. weikenstephanet L. seeligeri L. weikenstephanet Listonella Listonella Listonella Listonella Macrococcus Macrococcus Macrococcus Marinobacter Marinobacter algicola Marinobacter flavimaris Meiothermus Reiothermus ruber	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxa Moraxa Moraxa nonliqu Moraxa Moraxa Nosaxi Nakam Nakam Nakam Nakam Nakam Nakam Nakam Natran Natran thermo Natran truepen Naxiba alkalita	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella kiloniella K. granulomatis K. syrioca K. pneumoniae K. tariicola K. variicola Kluyvera Kluyvera Kluyvera Kluyvera Kluyvera ascorbata Kocuria roasea Kocuria roasea Kocuria varians	Helicobaa Helicobaa pylori	cter cter	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxa Moraxa monliqu Moraxa nonliqu Moraxa osloen: Nakam multipo Nannoo Nannoo pusilla Natran thermo, Natran trueper Naxiba alkaliti Neisse:	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella kiloniella K. granulomatis K. syrioca K. pneumoniae K. tariicola K. variicola Kluyvera Kluyvera Kluyvera Kluyvera Kluyvera ascorbata Kocuria roasea Kocuria roasea Kocuria varians	Helicobaa Helicobaa pylori	cter cter cter Listeria ivanovii L. marthii L. monocytogenes L. niparia L. rocourtiae L. seeligeri L. weishimeri Listonella Listonella Listonella Listonella Anguillarum Macrococcus bovicus Marinobacter Marinobacter Marinobacter algicola Marinobacter flavimaris Meiothermus Meiothermus Methylophilus Methylophilus	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc lylae Morax Morax Morax nonliqu Morax osloen; Nakam multipo Nanno Nakam multipo Nanno Natran Natran thermo Natran trueper Naxiba Natse Natran tureper Naxiba Natran	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
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Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella kiloniella K. granulomatis K. syrioca K. pneumoniae K. tariicola K. variicola Kluyvera Kluyvera Kluyvera Kluyvera Kluyvera ascorbata Kocuria roasea Kocuria roasea Kocuria varians	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Labrenzia alexandrii Labrenzia alba Labrenzia alba Labrenzia alba Labrenzia marina Labrys Labrys Labrys Labrys miyagiensis Labrys miyagiensis Labrys okinawensis Labrys portucalensis Lactobacillus [see below]	cter cter cter Listeria ivanovii L. marthii L. monocytogenes L. newyorkensis L. riparia L. rocourtiae L. seeligeri L. weikenstephanen L. weishimeri Listonella Listonella Listonella Listonella anguillarum Macrococcus bovicus Marinobacter Marinobacter Marinobacter bryozoorum Marinobacter flavimaris Meiothermus ruber Methylophilus Methylophilus Meitrophus	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxu Moraxu nonliqu Moraxi Moraxi Moraxi Nakam Moraxi Nakam Natam Natam Natam Natam Natran thermo Nata Nata Nata Nata Nata Nata Nata Nat	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae K. syraiulomatis K. syraiulomatis K. variicola Kluyvera Kluyvera Kluyvera Kluyvera Kluyvera Kocuria roasea Kocuria roasea Kocuria varians	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Labrenzia Labrenzia alba Labrenzia alba Labrenzia alba Labrenzia marina Labrys Labrys methylaminiphilus Labrys methylaminiphilus Labrys miyagiensis Labrys motachus Labrys motachus Labrys motachus Labrys bainawensis Labrys okinawensis Lacobacillus [see below] Laceyella Lechevalieria Lechevalieria Legionella [see below] Listeria	cter cter cter Listeria ivanovii L. marthii L. monocytogenes L. newyorkensis L. riparia L. rocourtiae L. seeligeri L. weihenstephanen L. seeligeri L. weihenstephanen L. seeligeri L. weihenstephanen L. seeligeri L. seeligeri L. seeligeri L. seeligeri Macrococcus Macrococcus Macrococcus Macrococcus Marinobacter Marinobacter flavimaris Meiothermus Meiothermus Meiothermus muber Methylophilus Methylophilus Microbacterium Microbacterium	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxa Moraxa Moraxa Moraxa Moraxa Moraxa Moraxa Moraxa Nakam multipo Nannon Nakam Nakam nultipo Nannon Nannon pusilla Natram thermo Natram trueper Naxiba alkalita Neisser Neissee Neissee Neissee Neissee Neissee Neissee Neissee	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia	Helicobaa Helicobaa pylori Labedella Labedella Labedella Laberazia Labrenzia Labrenzia aggregata Labrenzia alba Labrenzia alba Labrenzia marina Labrys Labrys miyagiensis Labrys miyagiensis Labrys miyagiensis Labrys mothylaminiphilus Labrys miyagiensis Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys portucalensis Labrys portucalensis Lacobacillus [see below] Laceyella Lucchevalieria aerocolonigenes Legionella [see below] Listeria Listeria Laquatica	cter cter cter Listeria ivanovii L. marthii L. monocytogenes L. riparia L. rocourtiae L. seeligeri L. weikenstephanei L. seeligeri L. weishimeri Listonella anguillarum Macrococcus Macrococcus Marinobacter Marinobacter Marinobacter Marinobacter darinobacter flavimaris Meiothermus Meiothermus Meiothermus Methylophilus Methylophilus methylotrophus Microbacterium Microbacterium Microbacterium	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc lylae Moraxa Moraxa nonliqu Moraxa osloem Nakam Nakam Nakam Nakam Nakam Nakam Natran Natran Natran thermo Natran trueper Naxiba Natran tureper Naxiba Natran Natran tureper Naxiba Natran Natran tureper Naxiba Natran Natran tureper Naxiba Natran Natran tureper Naxiba Natran Natran tureper Naxiba Natran Natran tureper Naxiba Natran Natran tureper Naxiba Natse Neisser Neisser denity Neisser gonorr	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella laminariae Kiloniella laminariae Kiloniella laminariae Kiloniella kiloniella K. granulomatis K. syrioca K. pneumoniae K. tariicola K. variicola Kluyvera Kluyvera Kluyvera Kluyvera Kluyvera ascorbata Kocuria roasea Kocuria roasea Kocuria varians	Helicobaa Helicobaa pylori Labedella Labedella gwakjiensis Labrenzia Labrenzia Labrenzia alba Labrenzia alba Labrenzia alba Labrenzia marina Labrys Labrys methylaminiphilus Labrys methylaminiphilus Labrys miyagiensis Labrys motachus Labrys motachus Labrys motachus Labrys bainawensis Labrys okinawensis Lacobacillus [see below] Laceyella Lechevalieria Lechevalieria Legionella [see below] Listeria	cter cter cter Listeria ivanovii L. marthii L. monocytogenes L. newyorkensis L. riparia L. rocourtiae L. seeligeri L. weihenstephanen L. seeligeri L. weihenstephanen L. seeligeri L. weihenstephanen L. seeligeri L. seeligeri L. seeligeri L. seeligeri Macrococcus Macrococcus Macrococcus Macrococcus Marinobacter Marinobacter flavimaris Meiothermus Meiothermus Meiothermus muber Methylophilus Methylophilus Microbacterium Microbacterium	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxa Moraxa Moraxa Moraxa Moraxa Moraxa Moraxa Moraxa Nakam multipo Nannon Nakam Nakam nultipo Nannon Nannon pusilla Natram thermo Natram trueper Naxiba alkalita Neisser Neissee Neissee Neissee Neissee Neissee Neissee Neissee	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia
Gluconacetobacter Gluconacetobacter xylinus Gordonia Gordonia rubripertincta Kaistia adipata Kaistia soli Kangiella Kangiella Kangiella koreensis Kerstersia Kerstersia gyiorum Kiloniella lawinariae Kiloniella laminariae Kiloniella laminariae Kilobiella K. granulomatis K. syrtoca K. peneumoniae K. tertigena K. varitoola Kluyvera Kluyvera Kluyvera Kluyvera Kluyvera Kluyvera sacorbata Kocuria roasea Kocuria varians Kurthia	Helicobaa Helicobaa pylori Labedella Labedella Labedella gwakjiensis Labrenzia Aggregata Labrenzia alba Labrenzia alba Labrenzia marina Labrys Labrys miyagiensis Labrys miyagiensis Labrys miyagiensis Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys mothylaminiphilus Labrys portucalensis Labrys portucalensis Labrys portucalensis Lacoyella Laceyella Laceyella Laceyella Laceyella Laceyella Laceinia Lagionella [see below] Listeria L. aquatica L. booriae	cter cter cter Listeria ivanovii L. marthii L. monocytogenes L. niparia L. rocourtiae L. seeligeri L. weihenstephaner L. seeligeri L. weishimeri Listonella anguillarum Macrococcus bovicus Marinobacter Marinobacter Marinobacter algicola Marinobacter flavimaris Meiothermus Meiothermus Meiothermus ruber Methylophilus Methylophilus methylotrophus Microbacterium Microbacterium	Ilyobacter polytropus Ilyobacter tartaricus Microc luteus Microc luteus Moraxu Moraxu nonliqu Moraxu nonliqu Moraxu osloen: Nakam multipa Nanno Nam multipa Natran Natran thermo, Natse Natse the See Neissee gonorr, Neissee Neissee See See See See See See See See See	Jeo hall	rgalicoccus otolerans Nesterenkonia Nesterenkonia holobia Nocardia Nocardia Nocardia corallina Nocardia corallina Nocardia

TABLE 1-continued

EXAMPLE BACTERIA				
	L. grandensis L. grayi L. innocua	Microbacteriun oxydans	n Neisseria sub, Neptunomona Neptunomona japonica	s
Lactobacillus				
	L. catenaformis	L. mali	L. parakefiri	L. sakei
J	L. ceti	L. manihotivoran	1	
1	L. coleohominis L. collinoides	L. mindensis L. mucosae	L. paraplantarum L. pentosus	L. sanfranciscensis L. satsumensis
	L. composti	L. murinus	L. periosus L. perolens	L. saisumensis L. secaliphilus
0	L. concavus	L. nagelii	L. plantarum	L. sharpeae
	L. corvniformis	L. namurensis	L. pontis	L. siliginis
	L. crispatus	L. nantensis	L. protectus	L. spicheri
L. amylophilus	L. crustorum	L. oligofermentar	ıs L. psittaci	L. suebicus
· ·	L. curvatus	L. oris	L. rennini	L. thailandensis
L. amylovorus	L. delbrueckii	L. panis	L. reuteri	L. ultunensis
L. animalis	subsp. bulgaricus	L. pantheris	L. rhamnosus	L. vaccinostercus
	L. delbrueckii	L. parabrevis	L. rimae	L. vaginalis
L. apodemi L. aviarius	subsp. delbrueckii L. delbrueckii	L. parabuchneri L. paracasei	L. rogosae L. rossiae	L. versmoldensis L. vini
L. bifermentans	subsp. lactis	L. paracollinoide		L. vitulinus
L. brevis	L. dextrinicus	L. parafarraginis		L. zeae
	L. diolivorans	L. homohiochii	L. jensenii	L. zymae
	L. equi	L. iners	L. johnsonii	L. gastricus
L. casei	L. equigenerosi	L. ingluviei	L. kalixensis	L. ghanensis
	L. farraginis	L. intestinalis	L. kefiranofaciens	L. graminis
	L. farciminis	L. fuchuensis	L. kefiri	L. hammesii
L. leichmannii L. lindneri	L. fermentum	L. gallinarum L. gasseri	L. kimchii L. helveticus	L. hamsteri L. harbinensis
	L. fornicalis L. fructivorans	L. gasseri	L. hilgardii	L. havakitensis L. havakitensis
L. matejermentans	L. frumenti		L. migurun	L. huyuhitehsis
Legionella				
Legionella		onella	Candidatus Legionella	Legionella
adelaidensis		courtii	jeonii	quinlivanii
Legionella anisa		onella	Legionella	Legionella
Legionella boli audou si s		lenensis	jordanis Lociou alla	rowbothamii Lasianalla
beliardensis Legionella		onella Inskii	Legionella lansingensis	Legionella rubrilucens
birminghamensis		onella	Legionella	Legionella
Legionella	dum		londiniensis	sainthelensi
bozemanae		onella erythra	Legionella	Legionella
Legionella brunens		onella	longbeachae	santicrucis
Legionella		eldensis	Legionella lytica	Legionella
busanensis	0	onella fallonii	Legionella	shakespearei
Legionella cardiaco	0	onella feeleii	maceachernii	Legionella
Legionella cherrii Legionella		onella tiana	Legionella	spiritensis Lagionalla
Legionella cincinnatiensis	gees Lagi	onella	massiliensis Legionella	Legionella steelei
Legionella		mospecies	micdadei	Legionella
clemsonensis		onella	Legionella	steigerwaltii
Legionella	gorn		monrovica	Legionella
donaldsonii		onella	Legionella	taurinensis
	grati		moravica	Legionella
		onella	Legionella	tucsonensis
		lensis	nagasakiensis	Legionella
	Legi hack	onella eliae	Legionella	tunisiensis Legionella
		enae onella	nautarum Legionella	Legionella wadsworthii
	0	etisoli	norrlandica	Legionella
		onella	Legionella	waltersii
		lensis	oakridgensis	Legionella
		onella	Legionella	worsleiensis
	jame	stowniensis	parisiensis	Legionella
			Legionella	yabuuchiae
			pittsburghensis Lociouslla	
			Legionella preumophila	
			pneumophila Legionella	
			quateirensis	
Oceanibulbus	Paen	ibacillus	Prevotella	Quadrisphaera
Oceanibulbus			Prevotella	Quadrisphaera
	thia	ninolyticus	albensis	granulorum
indolifex				
Oceanicaulis	Pant	bea .	Prevotella amnii	Quatrionicoccus
	Pant Pant	oea .	Prevotella amnii Prevotella bergensis	Quatrionicoccus Quatrionicoccus australiensis

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TABLE 1-continued

EXAMPLE BACTERIA

Oceanicola	Paracoccus	Prevotella bivia	Quinella
Oceanicola batsensis	Paracoccus	Prevotella brevis	Quinella ovalis
Oceanicola	alcaliphilus	Prevotella	Ralstonia
granulosus	Paucimonas	bryantii	Ralstonia
Oceanicola	Paucimonas	Prevotella buccae	eutropha
nanhaiensis	lemoignei	Prevotella	Ralstonia
Oceanimonas	Pectobacterium	buccalis	insidiosa
Oceanimonas	Pectobacterium	Prevotella copri	Ralstonia
baumannii	aroidearum	Prevotella	mannitolilytica
Oceaniserpentilla	Pectobacterium	dentalis	Ralstonia
		Prevotella	pickettii
Oceaniserpentilla	atrosepticum		1
haliotis	Pectobacterium	denticola	Ralstonia
Oceanisphaera	betavasculorum	Prevotella disiens	pseudosolanacearum
Oceanisphaera	Pectobacterium	Prevotella	Ralstonia syzygii
donghaensis	cacticida	histicola	Ralstonia
Oceanisphaera	Pectobacterium	Prevotella	solanacearum
litoralis	carnegieana	intermedia	Ramlibacter
Oceanithermus			
	Pectobacterium	Prevotella	Ramlibacter
Oceanithermus	carotovorum	maculosa	henchirensis
desulfurans	Pectobacterium	Prevotella	Ramlibacter
Oceanithermus	chrysanthemi	marshii	tataouinensis
profundus	Pectobacterium	Prevotella	Raoultella
Oceanobacillus	cypripedii	melaninogenica	Raoultella
	· · · ·		
Oceanobacillus caeni	Pectobacterium	Prevotella micans	ornithinolytica
Oceanospirillum	rhapontici	Prevotella	Raoultella
Oceanospirillum	Pectobacterium	multiformis	planticola
linum	wasabiae	Prevotella	Raoultella
	Planococcus	nigrescens	terrigena
	Planococcus	Prevotella oralis	Rathayibacter
	citreus	Prevotella oris	Rathayibacter
	Planomicrobium	Prevotella	caricis
	Planomicrobium	oulorum	Rathayibacter
	okeanokoites	Prevotella pallens	festucae
	Plesiomonas	Prevotella salivae	Rathayibacter
	Plesiomonas	Prevotella	iranicus
	shigelloides	stercorea	Rathayibacter
	Proteus	Prevotella	rathayi
	Proteus vulgaris	tannerae	Rathayibacter
		Prevotella	toxicus
		timonensis	Rathayibacter
		Prevotella	tritici
		veroralis	Rhodobacter
		Providencia	Rhodobacter
		Providencia	sphaeroides
		stuartii	Ruegeria
		Pseudomonas	Ruegeria
		Pseudomonas	gelatinovorans
		aeruginosa	genuintororuno
		0	
		Pseudomonas	
		alcaligenes	
		Pseudomonas	
		anguillispetica	
		Pseudomonas	
		fluorescens	
		Pseudoalteromonas	
		haloplanktis	
		Pseudomonas	
		mendocina	
		Pseudomonas	
		pseudoalcaligenes	
		Pseudomonas	
		putida	
		Pseudomonas	
		tutzeri	
		Pseudomonas	
		syringae	
		Psychrobacter	
		Psychrobacter	
		· ·	
		faecalis	
		Psychrobacter	
		phenylpyruvicus	

Saccharococcus Saccharococcus thermophilus Saccharomonospora Saccharomonospora azurea

Sagittula *Sagittula stellata* Salegentibacter Salegentibacter salegens Salimicrobium

Sanguibacter Sanguibacter keddieii Sanguibacter suarezii Saprospira

Stenotrophomonas Stenotrophomonas maltophilia Streptococcus [also see below] Streptomyces

Tatlockia Tatlockia maceachernii Tatlockia micdadei Tenacibaculum

TABLE 1-continued

	1	EXAMPLE BACTER	IA	
Saccharomonospora cyanea	Salimicrobium album	Saprospira grandis	Streptomyces achromogenes	Tenacibaculum amylolyticum
Saccharomonospora	Salinibacter	Sarcina	Streptomyces	Tenacibaculum
viridis	Salinibacter ruber	Sarcina maxima	cesalbus	discolor
Saccharophagus	Salinicoccus	Sarcina ventriculi	Streptomyces	Tenacibaculum
Saccharophagus docradans	Salinicoccus	Sebaldella Sebaldella	cescaepitosus Strontomucos	gallaicum Tanaaibaaulum
<i>degradans</i> Saccharopolyspora	alkaliphilus Salinicoccus	Sebaldella termitidis	Streptomyces cesdiastaticus	Tenacibaculum lutimaris
Saccharopolyspora	hispanicus	Serratia	Streptomyces	Tenacibaculum
ervthraea	Salinicoccus roseus	Serratia fonticola	cesexfoliatus	mesophilum
Saccharopolyspora	Salinispora	Serratia	Streptomyces	Tenacibaculum
gregorii	Salinispora	marcescens	fimbriatus	skagerrakense
Saccharopolyspora	arenicola	Sphaerotilus	Streptomyces	Tepidanaerobacter
hirsuta	Salinispora tropica	Sphaerotilus	fradiae	Tepidanaerobacter
Saccharopolyspora	Salinivibrio	natans	Streptomyces	syntrophicus
hordei Saccharopolyspora	Salinivibrio costicola	Sphingobacterium Sphingobacterium	fulvissimus Streptomyces	Tepidibacter <i>Tepidibacter</i>
rectivirgula	Salmonella	multivorum	griseoruber	formicigenes
Saccharopolyspora	Salmonella bongori	Staphylococcus	Streptomyces	Tepidibacter
spinosa	Salmonella enterica	[see below]	griseus	thalassicus
Ŝaccharopolyspora	Salmonella		Streptomyces	Thermus
taberi	subterranea		lavendulae	Thermus
Saccharothrix	Salmonella typhi		Streptomyces	aquaticus
Saccharothrix			phaeochromogenes	Thermus
australiensis Saacharothrix			Streptomyces thormodiastations	filiformis Thormus
Saccharothrix coeruleofusca			thermodiastaticus Streptomyces	Thermus thermophilus
coeruieojusca Saccharothrix			tubercidicus	inermopnius
espanaensis			moor cratcad	
Saccharothrix				
longispora				
Saccharothrix mutabilis				
Saccharothrix syringae				
Saccharothrix tangerinus				
Saccharothrix texasensis				
Staphylococcus				
S. arlettae	S. equorum	S. micro	oti	S. schleiferi
S. agnetis	S. felis	S. musc		S. sciuri
S. aureus	S. fleurettii	S. nepa		S. simiae
S. auricularis	S. gallinarum	S. paste	21181	
	0 I I			S. simulans
S. capitis	S. haemolyticus	S. petra	ısii	S. stepanovicii
S. caprae	S. hominis	S. petra S. pette	ısii nkoferi	S. stepanovicii S. succinus
S. caprae S. carnosus	S. hominis S. hyicus	S. petra S. petre S. piscij	ısii nkoferi fermentans	S. stepanovicii S. succinus S. vitulinus
S. caprae S. carnosus S. caseolyticus	S. hominis	S. petra S. petra S. piscij S. pseud	ısii nkoferi fermentans dintermedius	S. stepanovicii S. succinus S. vitulinus S. warneri
S. caprae S. carnosus	S. hominis S. hyicus S. intermedius	S. petra S. petra S. piscij S. pseud	nsii nkoferi fermentans dintermedius dolugdunensis	S. stepanovicii S. succinus S. vitulinus
S. caprae S. carnosus S. caseolyticus S. chromogenes	S. hominis S. hyicus S. intermedius S. kloosii	S. petra S. petre S. piscij S. pseud S. pseud S. pseud	isii nkoferi fermentans dintermedius dolugdumensis ereri	S. stepanovicii S. succinus S. vitulinus S. warneri
S. caprae S. carnosus S. caseolyticus S. chromogenes S. cohnii	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leetus S. lugdunensis	S. petra S. petra S. pisci S. pseu S. pseu S. pulve S. rostri	isii nkoferi fermentans dintermedius dolugdumensis ereri	S. stepanovicii S. succinus S. vitulinus S. warneri
S. caprae S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. delphini S. devriesei	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leetus S. lugdunensis S. lutrae	S. petra S. petra S. pisci, S. pseu S. pseu S. pulve S. rostr S. sacci	isii nkoferi fermentans dintermedius dolugdumensis ereri i	S. stepanovicii S. succinus S. vitulinus S. warneri
S. caprae S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condiimenti S. delphini	S. hominis S. hyicus S. intermedius S. leei S. leetus S. lugdunensis S. lutrae S. lyticans	S. petra S. petra S. pisci, S. pseu S. pseu S. pulve S. rostr S. sacci	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus	S. stepanovicii S. succinus S. vitulinus S. warneri
S. caprae S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. delphini	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leetus S. lugdunensis S. lutrae	S. petra S. petra S. pisci, S. pseu S. pseu S. pulve S. rostr S. sacci	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus	S. stepanovicii S. succinus S. vitulinus S. warneri
S. caprae S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. delphini S. devriesei S. epidermidis	S. hominis S. hyicus S. intermedius S. leei S. leetus S. lugdunensis S. lutrae S. lyticans	S. petra S. petra S. pisci, S. pseu S. pseu S. pulve S. rostr S. sacci	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus	S. stepanovicii S. succinus S. vitulinus S. warneri
S. caprae S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. delphini	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leitus S. lugdunensis S. lutrae S. lyticans S. massiliensis	S. petra S. petra S. pisci, S. pseu S. pseu S. pulve S. rostr S. sacci S. sapro	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus	S. stepanovicii S. succinus S. vitulinus S. warneri
S. caprae S. carnosus S. careolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus agalactiae	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. lentus S. lugdunensis S. lutrae S. lyticans S. massiliensis Streptococ infantarius	S. petra S. petra S. pisci, S. pseu S. polve S. rostr S. sacci S. sapre	isii nkoferi fermentans dintermedius dolugdumensis ereri i harolyticus ophyticus Streptococcus orisratti	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus
S. caprae S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus agalactiae Streptococcus anginos	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. lentus S. lugdunensis S. lutrae S. lyticans S. massiliensis Streptococ infantarius S. treptococ	S. petra S. petra S. pisci, S. pseu S. pulve S. rostr S. sacci S. sapre	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus orisratti Streptococcus	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus
S. caprae S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus agalactiae Streptococcus anginos Streptococcus anginos	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. lentus S. lugdunensis S. lutrae S. lutrae S. lyticans S. massiliensis S. treptococ infantarius sus Streptococ Streptococ	S. petra S. petra S. pisci, S. pseu S. pseu S. pseu S. pseu S. pseu S. pseu S. sacci S. sacci S. sapre	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus orisratti Streptococcus parasanguinis	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis
S. caprae S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus agalactiae Streptococcus anginos Streptococcus bovis Streptococcus bovis	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. lentus S. lugdunensis S. lutrae S. lyticans S. lyticans S. massiliensis sus Streptococ Streptococ Streptococ infantarius	S. petra S. petra S. pisci, S. pseu S. pseu S. pulve S. rostr S. sacci S. sapro cus iniae cus iniae	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus phyticus Streptococcus orisratti Streptococcus parasanguinis Streptococcus	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus
S. caprae S. carnosus S. carnosus S. chromogenes S. cohnii S. condimenti S. delphini S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus Streptococcus anginos Streptococcus canis Streptococcus canis Streptococcus canis	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leet S. lutrae S. lutrae S. lyticans S. massiliensis Streptococ Streptococ Streptococ infantarius Streptococ intermediu Streptococ	S. petra S. petra S. pisci, S. pseu S. pseu S. polve S. rostr S. sacci S. sapro cus tus iniae cus iniae cus iniae	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus orisratti Streptococcus parasanguinis Streptococcus perosis	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sanguinis
S. caprae S. carnosus S. careolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus agalactiae Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus anginos	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leei S. lugdunensis S. lugtae S. lyticans S. hyticans S. massiliensis Streptococ infantarius Streptococ intermediu Streptococ lactarius	S. petra S. petra S. pisci, S. pseu S. pseu S. pulve S. rostr S. sacci S. sapre	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus orisratti Streptococcus parasanguinis Streptococcus peroris Streptococcus Streptococcus	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus
S. caprae S. carnosus S. caroosus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus Streptococcus anginos Streptococcus anginos Streptococcus canis Streptococcus canis	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. lentus S. lugdunensis S. lutrae S. lyticans S. lyticans S. massiliensis Streptococ infantarius Streptococ intermediu Streptococ lactarius Streptococ	s S. petra S. petra S. pisci, S. pseu S. pulve S. rostr S. sacci S. sacci S. sapre	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus parasanguinis Streptococcus peroris Streptococcus peroris Streptococcus peroris	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus sobrinus
S. caprae S. carnosus S. caroosus S. chromogenes S. cohnii S. condimenti S. delphini S. delphini S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus Streptococcus anginos Streptococcus anginos Streptococcus dovis Streptococcus canis Streptococcus canis Streptococcus downei Streptococcus downei Streptococcus downei	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. lentus S. lugdunensis S. lutrae S. lyticans S. lyticans S. massiliensis Streptococ infantarius Streptococ intermediu Streptococ lactarius Streptococ lactarius Streptococ lactarius	cus iniae cus	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus ophyticus Streptococcus orisratti Streptococcus parasanguinis Streptococcus peroris Streptococcus peroris Streptococcus peneumoniae Streptococcus	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus
S. caprae S. carnosus S. caroosus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus Streptococcus anginos Streptococcus anginos Streptococcus canis Streptococcus canis	S. hominis S. hyicus S. hyicus S. kloosii S. leei S. leentus S. lugtane S. lyticans S. lyticans S. massiliensis Streptococ infantarius Streptococ intermediu Streptococ lactarius Streptococ lactarius Streptococ futermediu	S. petra S. petra S. pisci, S. pseu S. pseu S. postu S. rostr S. sacci S. sapro cus triae cus s cus cus s cus cus s cus cus s	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus parasanguinis Streptococcus peroris Streptococcus peroris Streptococcus peroris	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus suis Streptococcus suis Streptococcus
S. caprae S. carnosus S. carnosus S. chromogenes S. chromogenes S. conhii S. condimenti S. delphini S. delphini S. devriesei S. epidermidis Streptococcus agalactiae Streptococcus anginos Streptococcus canis Streptococcus canis Streptococcus canis Streptococcus canis Streptococcus downei Streptococcus downei Streptococcus downei	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leei S. lugdunensis S. lugtae S. lyticans S. lyticans S. massiliensis Streptococ infantarius Streptococ intermediu Streptococ lactarius Streptococ streptococ streptococ	S. petra S. petra S. pisci, S. pseu S. pseu S. postu S. postu S. sacci S. sacci S. sapro cus iniae cus iniae cus iniae cus iniae cus iniae cus iniae cus iniae	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus parasanguinis Streptococcus peroris Streptococcus peroris Streptococcus pneumoniae Streptococcus pneumoniae	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus suis Streptococcus suis Streptococcus suis
S. caprae S. carnosus S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus Streptococcus anginos Streptococcus dovis Streptococcus dovis Streptococcus dovis Streptococcus dovis Streptococcus dovis Streptococcus dovis Streptococcus dovie Streptococcus dovnei Streptococcus dovnei	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leei S. lugdunensis S. lugtae S. lyticans S. lyticans S. massiliensis Streptococ infantarius Streptococ intermediu Streptococ lactarius Streptococ streptococ streptococ	s S. petra S. petra S. pisci, S. pseu S. pseu S. pulve S. rostr S. sacci S. sacci S. sapre	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus orisratti Streptococcus parasanguinis Streptococcus peroris Streptococcus perotos Streptococcus penumoniae Streptococcus pseudopneumoniae Streptococcus	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus suis Streptococcus suis Streptococcus suis Streptococcus suis Streptococcus suis Streptococcus suis Streptococcus
S. caprae S. carnosus S. carnosus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus Streptococcus anginos Streptococcus anginos Streptococcus dovis Streptococcus dovis Streptococcus dovis Streptococcus canis Streptococcus dovnei Streptococcus dovnei Streptococcus dovnei Streptococcus dovnei Streptococcus equines Streptococcus equines Streptococcus equines	S. hominis S. hyicus S. hyicus S. kloosii S. leei S. leei S. leutus S. lugtae S. lyticans S. hyticans S. massiliensis Streptococ intermediu Streptococ lactarius Streptococ milleri Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ	cus mitis cus oralis cus oralis cus soralis cus cus cus cus cus cus cus cus cus cu	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus ophyticus Streptococcus parasanguinis Streptococcus peroris Streptococcus pneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus suis Streptococcus suis Streptococcus suis Streptococcus uberis Streptococcus vestibularis Streptococcus vestibularis
S. caprae S. carnosus S. carnosus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus Streptococcus anginos Streptococcus anginos Streptococcus dovis Streptococcus dovis Streptococcus dovis Streptococcus canis Streptococcus dovnei Streptococcus dovnei Streptococcus dovnei Streptococcus dovnei Streptococcus equines Streptococcus equines Streptococcus equines	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. lentus S. lutrae S. lutrae S. lyticans S. massiliensis Streptococ infantarius Streptococ intermediu Streptococ lactarius Streptococ streptococ intermediu Streptococ attreptococ streptococ intermediu Streptococ streptococ streptococ attrention Streptococ streptococ milleri Streptococ streptococ streptococ streptococ	Cus mitis cus oralis cus oralis cus	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus parasanguinis Streptococcus peroris Streptococcus prevoris Streptococcus prevoris Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus progenes Streptococcus ratti	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus suis Streptococcus suis Streptococcus uberis Streptococcus vestibularis Streptococcus vestibularis Streptococcus vestibularis Streptococcus vestibularis Streptococcus vestibularis Streptococcus vestibularis
S. caprae S. carnosus S. careolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus Streptococcus anginos Streptococcus anginos Streptococcus dovis Streptococcus dovis Streptococcus dovis Streptococcus canis Streptococcus canis Streptococcus dovnei Streptococcus dovnei Streptococcus dovnei Streptococcus equines Streptococcus equines Streptococcus equines	S. hominis S. hyicus S. hyicus S. kloosii S. leei S. leei S. leutus S. lugtae S. lyticans S. hyticans S. massiliensis Streptococ intermediu Streptococ lactarius Streptococ milleri Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ	Cus mitis cus oralis cus oralis cus	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus ophyticus Streptococcus parasanguinis Streptococcus peroris Streptococcus pneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus suis Streptococcus suis Streptococcus suis Streptococcus uberis Streptococcus vestibularis Streptococcus vestibularis
S. caprae S. carnosus S. caroosus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus dovis Streptococcus facadis Streptococcus facadis Streptococcus facadis	S. hominis S. hyicus S. hyicus S. kloosii S. leei S. leei S. leutus S. lugtae S. lyticans S. hyticans S. massiliensis Streptococ intermediu Streptococ lactarius Streptococ milleri Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ	Cus mitis cus oralis cus oralis cus	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus Streptococcus parasanguinis Streptococcus peroris Streptococcus prevoris Streptococcus prevoris Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus progenes Streptococcus ratti	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus suis Streptococcus suis Streptococcus suis Streptococcus uberis Streptococcus vestibularis Streptococcus vestibularis Streptococcus vestibularis Streptococcus vestibularis Streptococcus viridans
S. carnosus S. carnosus S. carnosus S. chromogenes S. chromogenes S. conhii S. delphini S. delphini S. devrisei S. epidermidis S. epidermidis Streptococcus agalactiae Streptococcus anginos Streptococcus canis Streptococcus canis Streptococcus canis Streptococcus downei Streptococcus downei Streptococcus downei Streptococcus downei Streptococcus downei Streptococcus dis Streptococcus dis Streptococcus dis Streptococcus dis Streptococcus dis Streptococcus dis Streptococcus dis Streptococcus dis Streptococcus faecalis Streptococcus ferus	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leei S. leutus S. lugdunensis S. lutrae S. lyticans S. lyticans S. massiliensis Streptococ intermediu Streptococ lactarius Streptococ atterneoiu Streptococ Streptococ intermediu Streptococ	cus mitis cus oralis cus	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus ophyticus Streptococcus parasanguinis Streptococcus peroris Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus psogenes Streptococcus ratti Streptococcus salivariu	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus suis Streptococcus suis Streptococcus uberis Streptococcus vueris Streptococcus vueris Streptococcus vueris Streptococcus vueris Streptococcus vueris Streptococcus vueris Streptococcus vueris Streptococcus vueris Streptococcus vueris Streptococcus vueris Streptococcus viridans Streptococcus viridans
S. caprae S. carnosus S. carnosus S. caseolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus Streptococcus anginos Streptococcus canis Streptococcus canis Streptococcus downei Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus ferus	S. hominis S. hyicus S. hyicus S. kloosii S. leei S. leei S. leutus S. lugtaensis S. lugtaensis S. lugtaensis S. lugtaensis S. hassiliensis Streptococ intermediu Streptococ lactarius Streptococ intermediu Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ Streptococ	s S. petra S. petra S. pisci, S. pisci, S. pseu S. pseu S. pulve S. rostr S. sacci S. sapro cus iniae cus iniae cus mitis cus mitis cus oralis cus oralis	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus ophyticus Streptococcus parasanguinis Streptococcus pneumoniae Streptococcus pneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus psogenes Streptococcus progenes Streptococcus ratti Streptococcus ratti Streptococcus ratti Streptococcus	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus suberis Streptococcus suberis Streptococcus suberis Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus
S. caprae S. carnosus S. carnosus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus agalactiae Streptococcus anginos Streptococcus anginos Streptococcus downei Streptococcus downei Streptococcus downei Streptococcus downei Streptococcus downei Streptococcus gaunes Streptococcus equines Streptococcus equines Streptococcus equines Streptococcus equines Streptococcus faecalis Streptococcus ferus	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leei S. lugdunensis S. lutrae S. lyticans S. massiliensis S. massiliensis S. massiliensis S. treptococ infantarius Streptococ lactarios Streptococ lactarios Streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ	s S. petra S. petra S. pisci, S. pisci, S. pseu S. pseu S. pulve S. rostr. S. sacro S. sapro cus cus iniae cus cus iniae cus cus cus cus cus cus cus cus cus cus	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus ophyticus Streptococcus parasanguinis Streptococcus parasanguinis Streptococcus protecoccus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus streptococcus streptococcus streptococcus streptococcus salivariu Virgibacillus halodenitrificans Virgibacillus	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus suberis Streptococcus suis Streptococcus suis Streptococcus vestibularis
S. caprae S. carnosus S. caroosus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus downei Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leei S. lentus S. lugdunensis S. lutrae S. lyticans S. lyticans S. massiliensis Streptococ intermediu Streptococ intermediu Streptococ actarius Streptococ streptococ intermediu Streptococ streptococ Streptococ	s S. petra S. petra S. pisci, S. pisci, S. pseu S. pseu S. pulve S. rostri S. sacci S. sapro cus iniae cus iniae cus mitis cus mitis cus oralis cus oralis cus vibrio aerogenes Vibrio albensis	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus ophyticus Streptococcus parasanguinis Streptococcus parasanguinis Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus psteptococcus streptococcus streptococcus streptococcus streptococcus streptococcus salivariu Virgibacillus Virgibacillus halodenitrificans Virgibacillus pantothenticus	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus sobrinus Streptococcus sus sus Streptococcus sus sus Streptococcus uberis Streptococcus vestibularis Streptococcus vestibularis Streptococcus vestibularis Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans Streptococcus viridans
S. caprae S. carnosus S. careolyticus S. chromogenes S. cohnii S. condimenti S. delphini S. devriesei S. epidermidis Streptococcus Streptococcus anginos Streptococcus anginos Streptococcus anginos Streptococcus downei Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis Streptococcus faecalis	S. hominis S. hyicus S. intermedius S. kloosii S. leei S. leei S. lugdunensis S. lutrae S. lyticans S. massiliensis S. massiliensis S. massiliensis S. treptococ infantarius Streptococ lactarios Streptococ lactarios Streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ streptococ	s S. petra S. petra S. pisci, S. pisci, S. pseu S. pseu S. pulve S. rostr. S. sacro S. sapro cus cus iniae cus cus iniae cus cus cus cus cus cus cus cus cus cus	isii nkoferi fermentans dintermedius dolugdunensis ereri i harolyticus ophyticus ophyticus Streptococcus parasanguinis Streptococcus parasanguinis Streptococcus protecoccus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus pseudopneumoniae Streptococcus streptococcus streptococcus streptococcus streptococcus salivariu Virgibacillus halodenitrificans Virgibacillus	S. stepanovicii S. succinus S. vitulinus S. warneri S. xylosus Streptococcus thermophilus Streptococcus sanguinis Streptococcus suberis Streptococcus suis Streptococcus suis Streptococcus vestibularis

TABLE 1-continued

EXAMPLE BACTERIA				
Undibacterium pigrum Ureaplasma Ureaplasma Ureaplasma Ureibacillus composti Ureibacillus composti Ureibacillus terrenus Ureibacillus terrenus Ureibacillus thermophilus thermophilus thermosphaericus	Vagococcus lutrae Vagococcus salmoninarum Variovorax boronicumulans Variovorax dokdonensis Variovorax paradoxus Variovorax soli Veillonella atypica Veillonella dispar Veillonella caviae Veillonella criceti Veillonella criceti Veillonella parvula Veillonella parvula Veillonella ratti Veillonella ratti Veillonella ratti Veillonella ratti Veillonella ratti Veinenivibrio stagnispumantis Verninephrobacter eiseniae Verrucomicrobium Verrucomicrobium spinosum	EXAMPLE BACTERIA Vibrio campbellii Vibrio cholerae Vibrio cincinnatiensis Vibrio coralliilyticus Vibrio diazotrophicus Vibrio fluvialis Vibrio halioticoli Vibrio ichthyoenteri Vibrio mediterranei Vibrio metischnikovii Vibrio mytili Vibrio mytili Vibrio natriegens Vibrio navarrensis Vibrio nigripulchritudo Vibrio ordalii Vibrio ordalii Vibrio ordentiis Vibrio pectenicida Vibrio penaeicida Vibrio pshionii Vibrio shilonii Vibrio shilonii Vibrio tubiashii Vibrio tubiashii Vibrio vulnificus	Weissella confusa Weissella halotolerans Weissella kellenica Weissella kandleri Weissella minor Weissella minor Weissella paramesenteroides Weissella thailandensis Weissella thailandensis Weissella thailandensis Weissella thailandensis Weissella thailandensis Weissella thailandensis Weissella thailandensis Williamsia marianensis Williamsia serinedens Winogradskyella Winogradskyella Winogradskyella Winogradskyella Winogradskyella Winogradskyella Wolbachia persica Wolbachia persica Zobellia Zobellia Zobellia Zobellia Zobellia Zobellia Zobellia Zoogloea ramigera Zoogloea resiniphila	Xanthobacter flavus Xanthobacter viscosus Xanthobacter viscosus Xanthomonas albilineans Xanthomonas alfalfae Xanthomonas arboricola Xanthomonas arboricola Xanthomonas campestris Xanthomonas citri Xanthomonas citri Xanthomonas cucurbitae Xanthomonas fragariae Xanthomonas fragariae Xanthomonas fragariae Xanthomonas fragariae Xanthomonas gardneri Xanthomonas paseoli Xanthomonas perforans Xanthomonas phaseoli Xanthomonas pisi Xanth
Xenophilus Xenophilus azovorans Xenorhabdus beddingii Xenorhabdus bovienii Xenorhabdus bovienii Xenorhabdus doucetiae Xenorhabdus griffiniae Xenorhabdus hominickii Xenorhabdus hominickii Xenorhabdus koppenhoeferi Xenorhabdus nematophila Xenorhabdus poinarii Xylanibacter Xylanibacter oryzae	Yangia Yangia pacifica Yaniella Yaniella flava Yaniella flava Yaniella halotolerans Yeosuana Yeosuana aromativorans Yersinia Yersinia aldovae Yersinia enterocolitica Yersinia enterocolitica Yersinia entomophaga Yersinia frederiksenii Yersinia intermedia Yersinia kristensenii	Yersinia mollaretii Yersinia pastis Yersinia pestis Yersinia pestis Yersinia ruckeri Yersinia ruckeri Yokenella Yokenella Yokenella Yonghaparkia Yonghaparkia Alkaliphila Zavarzinia Zavarzinia compransoris	Zooshikella Zooshikella ganghwensis Zunongwangia Zunongwangia profunda Zymobacter Zymobacter palmae Zymomonas Zymomonas Zymophilus Zymophilus paucivorans Zymophilus raffinosivorans	Zobellella Zobellella denitrificans Zobellella taiwanensis Zeaxanthinibacter enoshimensis Zhihengliuella Zhihengliuella Zhihengliuella halotolerans Xylanibacterium Xylanibacterium ulmi

Yersinia kristensenii

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TABLE 2

MEDICAMENTS

ACE inhibitors with calcium channel blocking agents ACE inhibitors with thiazides adamantane antivirals adrenal cortical steroids adrenal corticosteroid inhibitors adrenergic bronchodilators agents for hypertensive emergencies agents for pulmonary hypertension aldosterone receptor antagonists alkylating agents allergenics alpha-glucosidase inhibitors alternative medicines amebicides aminoglycosides aminopenicillins aminosalicvlates AMPA receptor antagonists amylin analogs analgesic combinations analgesics androgens and anabolic steroids angiotensin converting enzyme inhibitors angiotensin II inhibitors with calcium channel blockers angiotensin II inhibitors with agents thiazides angiotensin receptor blockers angiotensin receptor blockers and neprilysin inhibitors anorectal preparations anorexiants antacids anthelmintics agents anti-angiogenic ophthalmic agents anti-CTLA-4 monoclonal antibodies agents anti-infectives Anti-PD-1 monoclonal antibodies antiadrenergic agents (central) with thiazides antiadrenergic agents (peripheral) with thiazides antiadrenergic agents, centrally acting antiadrenergic agents, peripherally acting antiandrogens antianginal agents antiarrhythmic agents antiasthmatic combinations antibiotics/antineoplastics anticholinergic antiemetics anticholinergic antiparkinson agents anticholinergic bronchodilators anticholinergic chronotropic agents anticholinergics/ antispasmodics agents anticoagulant reversal agents anticoagulants anticonvulsants antidepressants antidiabetic agents antidiabetic combinations antidiarrheals antidiuretic hormones

decongestants respiratory agents dermatological agents sex hormones diagnostic topical agents radiopharmaceuticals uncategorized diarylquinolines agents dibenzazepine vaginal agents anticonvulsants mitotic inhibitors digestive enzymes monoamine oxidase dipeptidyl peptidase 4 inhibitors mouth and throat inhibitors diuretics products mTOR inhibitors dopaminergic antiparkinsonism agents mucolytics drugs used in alcohol multikinase dependence inhibitors echinocandins muscle relaxants EGFR inhibitors mydriatics estrogen receptor narcotic analgesic antagonists combinations estrogens narcotic analgesics nasal anti-infectives expectorants factor Xa inhibitors nasal antihistamines fatty acid derivative and decongestants nasal lubricants and anticonvulsants fibric acid derivatives irrigations nasal preparations first generation cephalosporins nasal steroids natural penicillins fourth generation cephalosporins neprilysin inhibitors functional bowel neuraminidase disorder agents inhibitors gallstone solubilizing neuromuscular blocking agents gamma-aminobutyric neuronal potassium acid analogs channel openers gamma-aminobutyric next generation acid reuptake inhibitors cephalosporins gastrointestinal agents nicotinic acid general anesthetics derivatives genitourinary tract NK1 receptor antagonists GI stimulants NNRTIS glucocorticoids non-cardioselective glucose elevating beta blockers non-iodinated glycopeptide antibiotics contrast media glycoprotein platelet non-ionic iodinated inhibitors contrast media glycylcyclines non-sulfonvlureas gonadotropin releasing nonsteroidal antihormones inflammatory gonadotropin-releasing agents NS5A inhibitors hormone antagonists gonadotropins nucleoside reverse group I antiarrhythmics transcriptase inhibitors (NRTIs) group II antiarrhythmics nutraceutical group III products antiarrhythmics nutritional products group IV ophthalmic antiarrhythmics anesthetics ophthalmic antigroup V antiarrhythmics infectives ophthalmic antigrowth hormone receptor blockers inflammatory growth hormones agents ophthalmic guanylate cyclase-C antihistamines and agonists H. pylori eradication decongestants ophthalmic H2 antagonists diagnostic agents ophthalmic hedgehog pathway glaucoma agents inhibitors hematopoietic stem cell ophthalmic mobilizer lubricants and heparin antagonists irrigations ophthalmic heparins HER2 inhibitors preparations

otic agents renin inhibitors products salicylates cephalosporins selective selective inhibitors serotoninnorepinephrine serotoninergic neuroenteric modulators sex hormone combinations sex hormones skeletal muscle skeletal muscle relaxants agents spermicides statins sterile irrigating solutions streptomyces derivatives succinimide anticonvulsants sulfonamides sulfonvlureas stimulants tetracyclic antidepressants tetracyclines therapeutic thioxanthenes third generation cephalosporins thrombolytics thyroid drugs tocolvtic agents topical agents agents

respiratory agents respiratory inhalant rifamycin derivatives sclerosing agents second generation selective estrogen receptor modulators immunosuppressants phosphodiesterase-4 selective serotonin reuptake inhibitors reuptake inhibitors SGLT-2 inhibitors relaxant combinations smoking cessation somatostatin and somatostatin analogs synthetic ovulation radiopharmaceuticals therapeutic vaccines thiazide diuretics thiazolidinediones thrombin inhibitors TNF alfa inhibitors topical acne agents topical anesthetics topical anti-infectives topical anti-rosacea topical antibiotics topical antifungals topical antihistamines topical antineoplastics topical antipsoriatics topical antivirals topical astringents

TABLE 2-continued

MEDICAMENTS

antidotes herbal products antiemetic/antivertigo agents histone deacetylase antifungals inhibitors antigonadotropic agents hormones antigout agents hormones antineoplastics antihistamines antihyperlipidemic agents hydantoin antihyperlipidemic anticonvulsants combinations hydrazide derivatives antihypertensive combinations immune globulins antihyperuricemic agents immunologic agents antimalarial agents immunostimulants antimalarial combinations immunosuppressive antimalarial quinolines agents antimetabolites impotence agents in vivo diagnostic antimigraine agents antineoplastic detoxifying biologicals incretin mimetics agents antineoplastic interferons inhaled anti-infectives antineoplastics inhaled corticosteroids antiparkinson agents inotropic agents antiplatelet agents insulin antipseudomonal penicillins insulin-like growth antipsoriatics factor integrase strand transfer antipsychotics antirheumatics inhibitor antiseptic and germicides interferons interleukin inhibitors antithyroid agents antitoxins and antivenins interleukins antituberculosis agents intravenous nutritional products antituberculosis combinations antitussives iodinated contrast antiviral agents media antiviral boosters ionic iodinated contrast antiviral combinations media antiviral interferons iron products anxiolytics, sedatives, and ketolides hypnotics laxatives aromatase inhibitors leprostatics atypical antipsychotics leukotriene modifiers azole antifungals lincomycin derivatives bacterial vaccines local injectable barbiturate anticonvulsants anesthetics barbiturates local injectable BCR-ABL tyrosine kinase anesthetics with inhibitors corticosteroids loop diuretics benzodiazepine lung surfactants anticonvulsants benzodiazepines lymphatic staining beta blockers with calcium agents channel blockers lysosomal enzymes beta blockers with thiazides macrolide derivatives beta-adrenergic blocking macrolides magnetic resonance agents beta-lactamase inhibitors imaging contrast media bile acid sequestrants mast cell stabilizers biologicals medical gas bisphosphonates meglitinides bone morphogenetic proteins metabolic agents bone resorption inhibitors methylxanthines mineralocorticoids bronchodilator combinations bronchodilators minerals and electrolytes calcimimetics calcineurin inhibitors agents calcitonin analgesics calcium channel blocking antibiotics anticonvulsants agents carbamate anticonvulsants antidepressants carbapenems antidiabetic agents carbonic anhydrase inhibitors antiemetics antifungals antihyperlipidemic agents

antihypertensive

combinations

antimalarials

antineoplastics antiparkinson agents

ophthalmic steroids ophthalmic steroids with anti-infectives ophthalmic surgical agents oral nutritional supplements immunostimulants immunosuppressants otic anesthetics otic anti-infectives otic preparations otic steroids otic steroids with anti-infectives oxazolidinedione anticonvulsants oxazolidinone antibiotics parathyroid hormone and analogs PARP inhibitors PCSK9 inhibitors penicillinase resistant penicillins penicillins peripheral opioid receptor antagonists peripheral opioid receptor mixed agonists/antagonists peripheral vasodilators peripherally acting antiobesity agents phenothiazine antiemetics phenothiazine antipsychotics phenylpiperazine antidepressants phosphate binders plasma expanders platelet aggregation inhibitors platelet-stimulating agents polyenes potassium sparing diuretics with thiazides potassium-sparing diuretics probiotics progesterone receptor modulators progestins prolactin inhibitors prostaglandin D2 antagonists protease inhibitors protease-activated receptor-1 antagonists proteasome inhibitors proton pump inhibitors psoralens psychotherapeutic agents psychotherapeutic combinations purine nucleosides pyrrolidine anticonvulsants

topical debriding agents topical depigmenting agents topical emollients topical keratolytics topical non-steroidal anti-inflammatories topical photochemotherapeutics topical rubefacient topical steroids topical steroids with anti-infectives triazine anticonvulsants tricyclic antidepressants trifunctional monoclonal antibodies ultrasound contrast media upper respiratory combinations urea anticonvulsants urea cycle disorder agents urinary anti-infectives urinary antispasmodics urinary pH modifiers uterotonic agents vaccine combinations vaginal anti-infectives vaginal preparations vasodilators vasopressin antagonists vasopressors VEGF/VEGFR inhibitors viral vaccines viscosupplementation agents vitamin and mineral combinations vitamins 5-alpha-reductase inhibitors 5-aminosalicylates 5HT3 receptor antagonists chloride channel activators cholesterol absorption inhibitors cholinergic agonists cholinergic muscle stimulants cholinesterase inhibitors CNS stimulants coagulation modifiers colony stimulating factors contraceptives corticotropin coumarins and indandiones cox-2 inhibitors

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TABLE 2-continued

MEDICAMENTS			
antipsychotic agents antituberculosis agei antivirals anxiolytics, sedative and hypnotics bone resorption inhibitors cardiovascular agent central nervous syst agents coagulation modifie: diagnostic dyes diuretics genitourinary tract agents GI agents hormones metabolic agents ophthalmic agents	nts radiocontrast agents radiologic adjuncts s radiologic agents radiologic conjugating agents radiopharmaceuticals ts recombinant em human erythropoietins		

SEQUENCE LISTING

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Gln Asn Gln Leu Tyr Asn Glu Leu Asn Leu Gly Arg Arg Glu Glu Tyr 20 25 30
Asp Val Leu Asp Lys Arg Arg Gly Arg Asp Pro Glu Met Gly Gly Lys 35 40 45
Pro Arg Arg Lys Asn Pro Gln Glu Gly Leu Tyr Asn Glu Leu Gln Lys 50 55 60
Asp Lys Met Ala Glu Ala Tyr Ser Glu Ile Gly Met Lys Gly Glu Arg 65 70 75 80
Arg Arg Gly Lys Gly His Asp Gly Leu Tyr Gln Gly Leu Ser Thr Ala 85 90 95

100

Thr Lys Asp Thr Tyr Asp Ala Leu His Met Gln Ala Leu Pro Pro Arg

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-continued

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What is claimed is:

1. A method for enhancing efficacy of an immunotherapy of a cancer selected from the group consisting of melanoma, lung cancer, and colon cancer in a human or animal patient, wherein the method comprises increasing the relative proportion of a sub-population of bacteria comprising *Akkermansia* in a gut microbiota of the patient by administering a bacterial transplant to the patient, wherein an effective amount of the immunotherapy is administered to the patient, wherein the immunotherapy is selected from the group consisting of an immune checkpoint therapy, an adoptive cell therapy, a CAR-T (chimeric antigen receptor T) cell 65 therapy, and a TIL (Tumor-infiltrating lymphocyte) therapy, wherein the bacterial transplant comprises *Akkermansia*, and

wherein increasing the relative proportion of the sub-population of bacteria modulates immune cells in the patient, whereby the efficacy of the immunotherapy is enhanced for treatment of the cancer in the patient.

2. The method of claim 1, wherein the immunotherapy is an immune checkpoint therapy comprising administration of an immune checkpoint modulator.

3. The method of claim **2**, wherein the immune checkpoint modulator is a CTLA4 (Cytotoxic T-Lymphocyte Associated Protein 4) inhibitor.

4. The method of claim 3, wherein the immune checkpoint modulator is an antibody.

5. The method of claim 4, wherein the immune checkpoint modulator is ipilimumab or tremelimumab.

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6. The method of claim 1, wherein the bacterial transplant comprises *Akkermansia municiphila*.

7. The method of claim 1, wherein increasing the relative proportion of the sub-population of bacteria modulates immune cells selected from CD8+ cells, tumor infiltrating 5 lymphocytes (TILs), CD4+ cells, Treg cells and memory cells and Th17 cells.

8. The method of claim **1**, wherein the immune cells comprise CD8+ cells.

9. The method of claim **8**, wherein the CD8+ cells are 10 CD8+ cells of the patient.

10. The method of claim **1**, wherein the immune cells are upregulated or expanded in the patient.

11. The method of claim 1, wherein the immune cells comprise memory cells selected from the group consisting 15 of central memory T-cells (TCM), effector memory T-cells (TEM), stem cell memory cells (TSCM), and memory effector T-cells (Teff).

12. The method of claim **1**, wherein the immune cells comprise memory cells selected from the group consisting ²⁰ of CD45RO+CD62L+ and CD25+ CD45RA- CD45RO+ CD127+ cells.

13. The method of claim **1**, wherein the immune cells comprise administered cells via adoptive cell therapy.

14. The method of claim **1**, wherein the method further 25 comprises administering a purine nucleoside.

15. The method of claim **1**, wherein the cancer is a colon cancer.

16. The method of claim **1**, wherein the cancer is melanoma or lung cancer. 30

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