SUPPLEMENTAL:

Table S1Distribution of reference organisms used sorted by family with list of accession numbers

	Species	
Family	count	Accession List:
unclassified Actinobacteria	1	CP003219, CP000431, AP012204, CP011541, CP015235,
Pseudonocardiales	11	CP000249, CP009110, CP011546, CP012184, CP003119,
Actinomycetales	4	FO082843, CP003876, CP003275, CP011853, F0117623,
•		CP003325, AP012326, AP012327, AP012331, AP012334,
Kineosporiales	1	CP005287, HE804045, CP003696, CP011883, CP003788, CP013747, CP006259, CP007790, CP003924, CP004353,
Nakamurellales	1	CP006272, CP004346, CP008889, CP004351, CP005080,
Frankiales	5	AP012057, CP003170, CP006365, CP011491, CP011545,
Coriobacteriales	4	AP013105, CP006734, CP007699, CP006842, CP006841,
Micromonosporales	8	CP006850, CP006996, CP011786, CP007155, CP007156,
Bifidobacteriales	11	CP013292, CP007595, CP009302, CP011773, CP009211,
Eggerthellales	5	CP011005, CP011311, CP010827, CP011868, CP011489,
		CP012070, CP012171, CP014869, CP012479, CP012697,
Acidimicrobiales	2	LN831026, CP013743, CP014196, CP014475, CP015849,
Rubrobacterales	2	CP012949, CP000518, CP011341, CP008944, CP000509, AE014184, CP009896, CP008953, CP003078, CP000854,
Catenulisporales	1	AP008957, CP012750, AE016958, CP002638, CP000386,
Propionibacteriales	7	CP000750, AE017283, CP000088, CP002385, CP000910,
Corynebacteriales	59	CP000474, CP000454, CP000820, CP002299, CR931997,
Acidothermales	1	AM849034, CT573213, CP000511, CP000481, CP011312,
Streptosporangiales	5	CP010407, CP000667, LN868938, AP009152, CP011340,
Glycomycetales	1	CP000850, CP009215, CP009922, CP007514, CP002786,
		CP011492, CP001630, CP001964, CP002040, CP001819,
Solirubrobacterales	1	CP001778, CP001821, CP013254, AP010968, CP001341, AP009493, CP004370, CP001874, CP001682, CP001854,
Micrococcales	37	CP001738, CP001618, CP001684, CP001706, CP001683,
Geodermatophilales	3	FO203431, CP001686, CP001737, CP001814, CP001700,
Streptomycetales	19	CP001736, CP001726, CP003322, AP012211, AM942444,
		AP012319, AP012322, CP001721, CP001966, CP001631,
		CP001867, CP001802, CP012072, CP007490, CP001992,
		CP001849, CP001606, CP006835, CP006764, CP011112,
		CP011542, CP002666, CP002475, CP002665, CP008802,
		CP001958, CP002162, CP002045, CP001620, CP005929,
		CP002994, CP002801, CP002857, CP002593, FN554889,
		AP011540, CP001829, FN563149, CP002628, CP006936,
		CP003053, CP003169, CP002343, CP012069, CP002810,
		CP002047, CP002280, CP002734, CP015810, CP002917,
		CP012752, AP012333, CP002379, CP009220

Table S2Table of core genes found in the reference set listed by functional category (continued to pg. 6)

Model Classification	Count	Model Names
Amino acid biosynthesis	54	TIGR00032, TIGR00033, TIGR00036, TIGR00069, TIGR00070, TIGR00110, TIGR00112, TIGR00118, TIGR00119, TIGR00120, TIGR00170, TIGR00171, TIGR00191, TIGR00260, TIGR00262, TIGR00263, TIGR00338, TIGR00407, TIGR00465, TIGR00507, TIGR00564, TIGR00652, TIGR00653, TIGR00658, TIGR00674, TIGR00676, TIGR00735, TIGR00761, TIGR00838, TIGR00970, TIGR01027, TIGR01048, TIGR01027, TIGR01048, TIGR01037, TIGR01123, TIGR01127, TIGR01137, TIGR01137, TIGR01137, TIGR01356, TIGR01357, TIGR01356, TIGR01357, TIGR01356, TIGR01356, TIGR01392, TIGR01536, TIGR01366, TIGR01392, TIGR01536, TIGR01855, TIGR01900, TIGR02067, TIGR02082, TIGR03188
Biosynthesis of cofactors, prosthetic groups, and carriers	79	TIGR00018, TIGR00063, TIGR00078, TIGR00083, TIGR00097, TIGR00109, TIGR00114, TIGR00151, TIGR00152, TIGR00154, TIGR00173, TIGR00187, TIGR00190, TIGR00204, TIGR00212, TIGR00216, TIGR00222, TIGR00223, TIGR00243, TIGR00343, TIGR00347, TIGR00379, TIGR00380, TIGR00433, TIGR00453, TIGR00482, TIGR005050, TIGR00555, TIGR00554, TIGR00555, TIGR00551, TIGR00554, TIGR00554, TIGR00562, TIGR00562, TIGR00581, TIGR00612, TIGR00636, TIGR00693, TIGR00708, TIGR00713, TIGR00715, TIGR00751, TIGR01035, TIGR01465, TIGR01469, TIGR01464, TIGR01465, TIGR01469, TIGR01498, TIGR01475, TIGR01496, TIGR01498, TIGR01510, TIGR01513, TIGR01498, TIGR01498, TIGR01994, TIGR01923, TIGR01981, TIGR01994, TIGR02150, TIGR02257, TIGR02352, TIGR02476, TIGR02466, TIGR03160, TIGR03448, TIGR03442, TIGR03447, TIGR03448, TIGR03552, TIGR03699, TIGR03701, TIGR03800

Model Classification	Count	Model Names
Cell envelope	21	TIGR00031, TIGR00055, TIGR00067, TIGR00179, TIGR00219, TIGR00274, TIGR00445, TIGR00492, TIGR00753, TIGR01072, TIGR01082, TIGR01087, TIGR01099, TIGR01133, TIGR01181, TIGR01207, TIGR01214, TIGR01221, TIGR01695, TIGR03423, TIGR03426
Cellular processes	15	TIGR00065, TIGR00172, TIGR00220, TIGR00647, TIGR00685, TIGR00732, TIGR02210, TIGR02400, TIGR02614, TIGR02673, TIGR03137, TIGR03253, TIGR03445, TIGR03446, TIGR03451
Central intermediary metabolism	16	TIGR00101, TIGR00192, TIGR00193, TIGR00221, TIGR00455, TIGR00642, TIGR00700, TIGR01034, TIGR01135, TIGR01173, TIGR01217, TIGR01455, TIGR01792, TIGR02727, TIGR03383, TIGR03705
DNA metabolism	35	TIGR00042, TIGR00084, TIGR00194, TIGR00228, TIGR00237, TIGR00281, TIGR00362, TIGR00416, TIGR00575, TIGR00577, TIGR00580, TIGR00593, TIGR00595, TIGR00613, TIGR00615, TIGR00628, TIGR00630, TIGR00631, TIGR00634, TIGR00635, TIGR00643, TIGR00663, TIGR00665, TIGR01051, TIGR01059, TIGR01063, TIGR01073, TIGR01083, TIGR01128, TIGR01280, TIGR01389, TIGR01391, TIGR02012, TIGR02168, TIGR02225

Model Classification	Count	Model Names
Energy metabolism	86	TIGR00016, TIGR00021, TIGR00129,
		TIGR00131, TIGR00203, TIGR00209,
		TIGR00218, TIGR00232, TIGR00239,
		TIGR00273, TIGR00330, TIGR00419,
		TIGR00461, TIGR00518, TIGR00527,
		TIGR00528, TIGR00561, TIGR00651,
		TIGR00692, TIGR00720, TIGR00871,
		TIGR00872, TIGR00873, TIGR00876,
		TIGR00936, TIGR00962, TIGR00979,
		TIGR01039, TIGR01060, TIGR01064,
		TIGR01068, TIGR01078, TIGR01131,
		TIGR01132, TIGR01144, TIGR01145,
		TIGR01146, TIGR01163, TIGR01179,
		TIGR01198, TIGR01216, TIGR01224,
		TIGR01225, TIGR01228, TIGR01235,
		TIGR01236, TIGR01255, TIGR01260,
		TIGR01263, TIGR01266, TIGR01292,
		TIGR01311, TIGR01312, TIGR01341,
		TIGR01344, TIGR01515, TIGR01520,
		TIGR01534, TIGR01722, TIGR01771,
		TIGR01798, TIGR01814, TIGR01816,
		TIGR01828, TIGR01915, TIGR01959,
		TIGR01962, TIGR01973, TIGR02022,
		TIGR02152, TIGR02156, TIGR02157,
		TIGR02158, TIGR02159, TIGR02422,
		TIGR02423, TIGR02426, TIGR02427,
		TIGR02866, TIGR02891, TIGR02970,
		TIGR03036, TIGR03181, TIGR03377,
		TIGR04380, TIGR04382
Fatty acid and phospholipid	6	TIGR00189, TIGR00473, TIGR00517,
metabolism		TIGR00560, TIGR01830, TIGR03150
Hypothetical proteins	8	TIGR00247, TIGR00252, TIGR00370,
		TIGR00730, TIGR01777, TIGR02569,
		TIGR03084, TIGR03085
Protein fate	43	TIGR00064, TIGR00077, TIGR00079,
		TIGR00115, TIGR00121, TIGR00214,
		TIGR00357, TIGR00382, TIGR00401,
		TIGR00493, TIGR00500, TIGR00504,
		TIGR00534, TIGR00544, TIGR00546,
		TIGR00556, TIGR00706, TIGR00763,
		TIGR00810, TIGR00945, TIGR00959,
		TIGR00963, TIGR00964, TIGR00966,
		TIGR00967, TIGR01129, TIGR01410,
		TIGR02227, TIGR02348, TIGR02350,
		TIGR02493, TIGR03144, TIGR03346,
		TIGR03534, TIGR03686, TIGR03687,
		TIGR03688, TIGR03689, TIGR03690,
		TIGR03691, TIGR03919, TIGR03920,
		TIGR03921

Model Classification	Count	Model Names
Protein synthesis	115	Ribosomal_L10, Ribosomal_L23,
		Ribosomal_S14, Ribosomal_S8,
		Ribosomal_S9, TIGR00001, TIGR00008,
		TIGR00009, TIGR00011, TIGR00012,
		TIGR00019, TIGR00020, TIGR00029,
		TIGR00038, TIGR00043, TIGR00048,
		TIGR00057, TIGR00059, TIGR00060,
		TIGR00061, TIGR00062, TIGR00071,
		TIGR00086, TIGR00088, TIGR00090,
		TIGR00091, TIGR00095, TIGR00096,
		TIGR00105, TIGR00116, TIGR00132,
		TIGR00133, TIGR00135, TIGR00138,
		TIGR00150, TIGR00157, TIGR00158,
		TIGR00165, TIGR00166, TIGR00168,
		TIGR00174, TIGR00186, TIGR00233,
		TIGR00234, TIGR00256, TIGR00344,
		TIGR00392, TIGR00396, TIGR00398,
		TIGR00409, TIGR00414, TIGR00418,
		TIGR00420, TIGR00422, TIGR00430,
		TIGR00431, TIGR00435, TIGR00436,
		TIGR00442, TIGR00447, TIGR00456,
		TIGR00459, TIGR00460, TIGR00464,
		TIGR00467, TIGR00468, TIGR00472,
		TIGR00479, TIGR00484, TIGR00485,
		TIGR00487, TIGR00496, TIGR00499,
		TIGR00563, TIGR00731, TIGR00755,
		TIGR00855, TIGR00952, TIGR00981,
		TIGR01009, TIGR01011, TIGR01017,
		TIGR01021, TIGR01022, TIGR01023,
		TIGR01024, TIGR01029, TIGR01030,
		TIGR01031, TIGR01032, TIGR01044,
		TIGR01049, TIGR01050, TIGR01066,
		TIGR01067, TIGR01071, TIGR01079,
		TIGR01125, TIGR01164, TIGR01169,
		TIGR01171, TIGR01308, TIGR01575,
		TIGR01632, TIGR02692, TIGR02729,
		TIGR03594, TIGR03631, TIGR03632,
		TIGR03635, TIGR03654, TIGR03704,
		TIGR03723, TIGR03725, TIGR03953
Purines, pyrimidines,	43	TIGR00017, TIGR00041, TIGR00081,
nucleosides, and nucleotides		TIGR00126, TIGR00184, TIGR00302,
		TIGR00336, TIGR00337, TIGR00355,
		TIGR00639, TIGR00655, TIGR00670,
		TIGR00877, TIGR00878, TIGR00884,
		TIGR00928, TIGR01036, TIGR01090,
		TIGR01091, TIGR01134, TIGR01161,
		TIGR01162, TIGR01203, TIGR01302,
		TIGR01354, TIGR01368, TIGR01369,
		TIGR01694, TIGR01704, TIGR01736,
		TIGR01737, TIGR01744, TIGR02075,
		TIGR02127, TIGR02170, TIGR02274,
		TIGR02487, TIGR02491, TIGR02504,
		TIGR02506, TIGR02961, TIGR03263,
		TIGR03284

Model Classification	Count	Model Names
Regulatory functions	9	TIGR00242, TIGR00244, TIGR00331, TIGR00498, TIGR01394, TIGR01529, TIGR01693, TIGR01950, TIGR03968
Transcription	19	TIGR00082, TIGR00188, TIGR00690, TIGR00766, TIGR00767, TIGR00922, TIGR01388, TIGR01951, TIGR01953, TIGR01966, TIGR02013, TIGR02027, TIGR02191, TIGR02258, TIGR02273, TIGR02386, TIGR02393, TIGR02949, TIGR03988
Transport and binding proteins	20	TIGR00383, TIGR00400, TIGR00750, TIGR00754, TIGR00773, TIGR00972, TIGR00974, TIGR00975, TIGR01104, TIGR01256, TIGR02135, TIGR02138, TIGR02275, TIGR03409, TIGR03410, TIGR03770, TIGR03771, TIGR03772, TIGR03851, TIGR04520
Unclassified	64	TIGR00177, TIGR00180, TIGR00369, TIGR00759, TIGR01350, TIGR01412, TIGR01417, TIGR01428, TIGR01430, TIGR01581, TIGR01698, TIGR01701, TIGR01751, TIGR01788, TIGR01885, TIGR01919, TIGR02133, TIGR02188, TIGR02200, TIGR02234, TIGR02278, TIGR02288, TIGR02349, TIGR02412, TIGR02631, TIGR02753, TIGR02857, TIGR02927, TIGR02947, TIGR02952, TIGR02960, TIGR03003, TIGR03005, TIGR03081, TIGR03083, TIGR03086, TIGR03089, TIGR03178, TIGR03180, TIGR03356, TIGR03449, TIGR03450, TIGR03465, TIGR03459, TIGR03464, TIGR03465, TIGR03467, TIGR03535, TIGR03539, TIGR03625, TIGR03664, TIGR03815, TIGR03817, TIGR03819, TIGR03843, Cpn10, GrpE, Methyltransf_5, PGK, SHMT, TIGR03869, TIGR03873, TIGR03936, TIGR03997
Unknown function	31	TIGR00044, TIGR00092, TIGR00103, TIGR00149, TIGR00164, TIGR00196, TIGR00250, TIGR00257, TIGR00368, TIGR00481, TIGR00486, TIGR00494, TIGR00724, TIGR00726, TIGR00762, TIGR00977, TIGR01303, TIGR01304, TIGR01393, TIGR01448, TIGR01490, TIGR01764, TIGR01967, TIGR01970, TIGR01976, TIGR03156, TIGR03816, TIGR03941, TIGR03954, TIGR03960, TIGR04047

Figure S3Core gene comparison to the Database of Essential Genes (DEG) sorted by function. Red columns indicate hits not found in the database, compared to all core gene classifications shown in blue

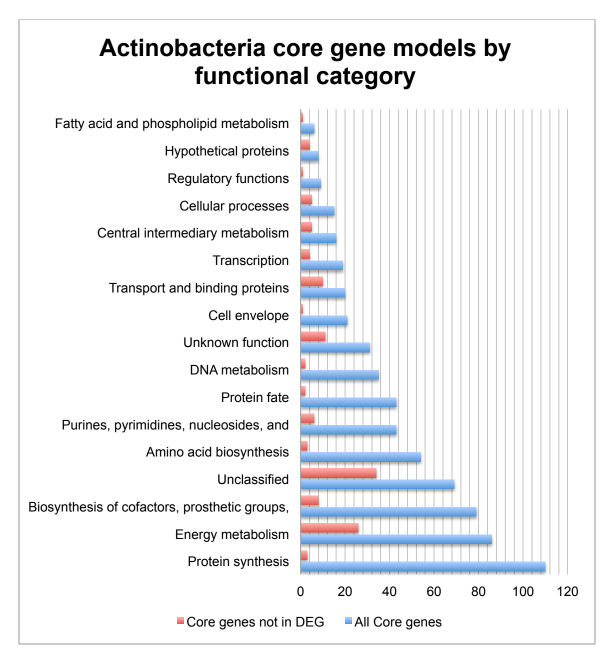


Figure S4Comparison of counts for all proteins in the reference set sorted by function. Red columns indicate distribution of all core gene hits and blue indicate distribution of all TIGRFAM model hits

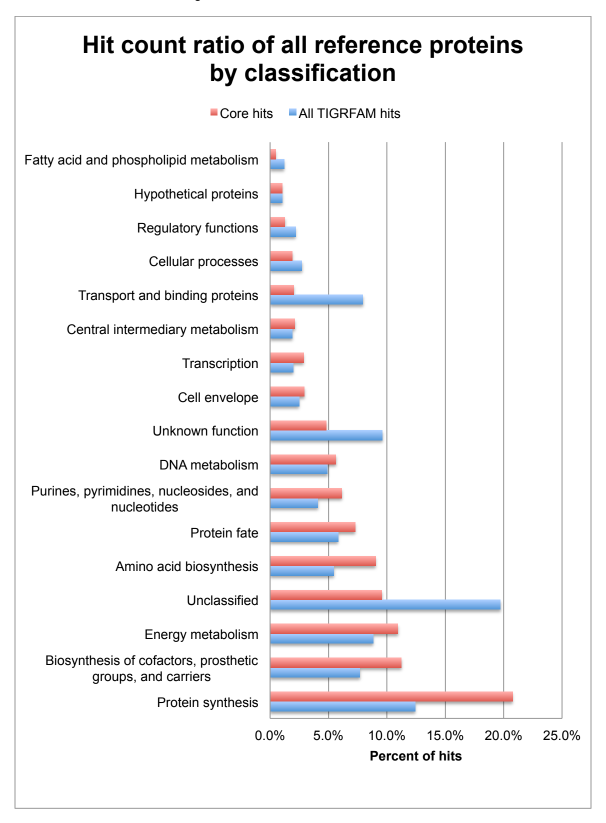


Figure S5 Histograms of essentiality measures. (a) Median values of pairwise dN/dS calculations for all alignments. (b) Reference ubiquity by species represents presence in all genomes. If the majority of organisms in a genus have a gene it is counted and these counts are shown relative to total genera in the set. (c) Single copy ratio shows number of occurrences a gene is found only once divided by total genomes that have one or more copies

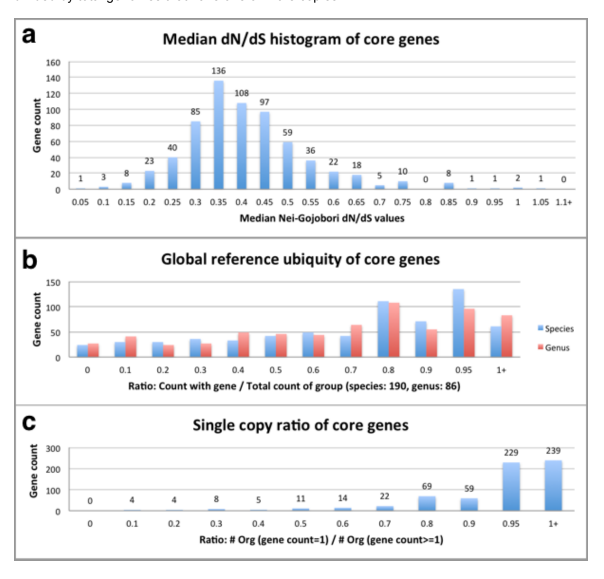


Figure S6Median and mode distributions of all bootstrap support values of reference trees using all nodes and terminal nodes respectively

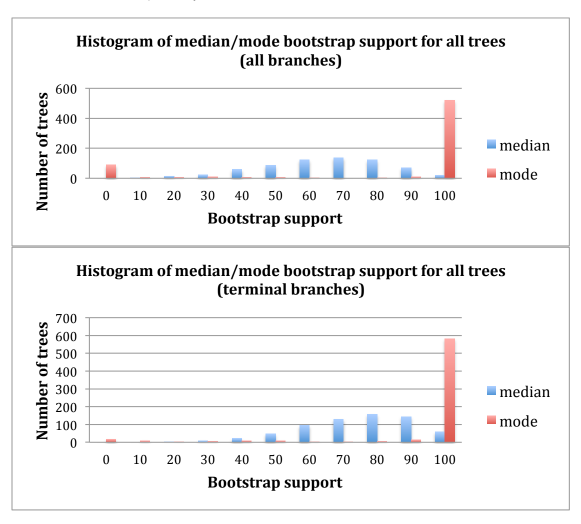


Figure S7

(a) Functional classification of all MIBiG cluster core gene hits. Inner ring shows hits with exploration mode enabled while outer ring is default filtered core hits. Actual counts are presented in parenthesis (b) Distribution of hit counts for clusters where blue, red and purple columns represent exploration core, resistance, and default core model hits respectively

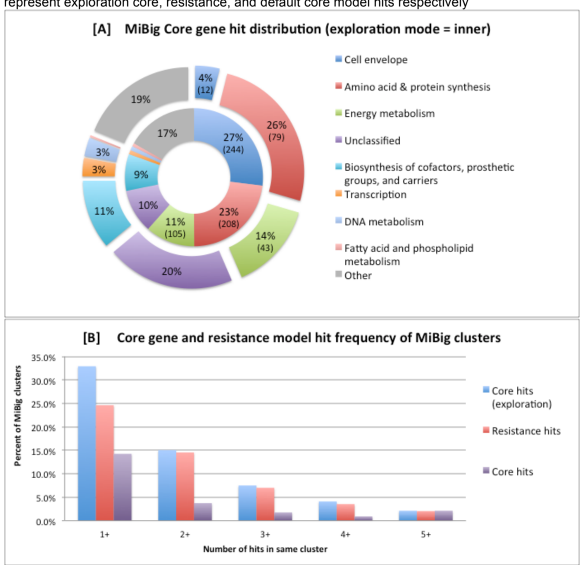


Table S8Detection frequency statistics from complete actinobacterial genomes using combined 189 reference and 11 positive controls run through the ARTS pipeline.

Hit type	Avg	Std	min	max
Core	489	79	271	653
Duplicate	27	23	0	96
BGC prox.	27	24	0	140
Phylogeny	125	88	7	422
Two +	16	16	0	83
Three +	2	4	0	22

Percent of c				
Hit type	Avg	Std	min	max
Duplicate	5.07%	3.85%	0.00%	16.03%
BGC prox.	5.03%	4.34%	0.00%	25.78%
Phylogeny	25.74%	17.53%	1.68%	70.69%
Two +	3.32%	3.00%	0.00%	13.74%
Three +	0.35%	0.65%	0.00%	3.69%

Table S9

Positive examples of genomes with known self-resistance mechanisms analyzed with ARTS exploration mode. Hits to ARTS criteria are shown as; D: Duplication, B: BGC proximity, P: Phylogeny, R: Resistance model. Rows in grey indicate non-actinobacterial genomes where phylogeny criteria do not apply. Tan rows indicate resistance that is not within a BGC. Notes marked with stars are explained in the bottom row

Product	Resistance gene	Organism	Gene Accession (ref)	ARTS hits	Criteria hits (>2, >3)	BGCs (total, core hit, res hit)	Genes (core, total)
Novobiocin	duplicated gyrB	Streptomyces niveus NCIMB 11891	WP_03123 2360 (1)	D,B, R,*P	46 , 13	30 , 24 , 8	577 , 7815
Clorobiocin	duplicated gyrB	Streptomyces roseochromogenes DS 12.976	AAN65247 (2)	D , B , R, *P	71 , 20	43 , 24 , 13	590 , 9055
Albicidin	pentapeptide repeat protein for GyrB	Xanthomonas albilineans GPE PC73	CBA16025	B,R	2, 0	8,7,3	434 , 3208
Streptolydigin	mutated <i>rpoB</i>	Streptomyces lydicus NRRL2433	AAQ19729 (4)	R	42 , 3	35 , 17 , 13	576 , 8518
Rifamycin	mutated <i>rpoB</i>	Amycolatopsis mediterranei S699	AAS07760 (5)	R	49 , 5	30 , 16 , 8	564 , 9575
Rifampicin	duplicated rpoB	Nocardia farcinica IFM 10152	BAD59497. 1_(6)	D	21 , 3	17, 13, 7	550 , 5946
Thiocillin	duplicated ribosomal L11	Bacillus cereus ATCC 14579	AAP11944, AAP11947 (7)	D,B	6,0	10,7,2	449 , 5255
Erythromycin	duplicated 23S rRNA methyltransferase	Saccharopolyspora erythraea NRRL23338	WP_00995 0391 (8)	**D, B,P, R	83 , 17	36 , 16 , 13	653 , 7198
Agrocin 84	duplicated Leu- tRNA synthase	Agrobacterium radiobacter K84	ACM31456 (9)	D	4,0	10,7,0	475 , 6684
Thiolactomycin	duplicated FabB/F	Salinispora pacifica DSM 45543	ALJ49913 (10)	**D, B,P	61 , 20	25 , 20 , 9	641 , 4784
Salinospora- mide A	duplicated beta- proteasome subunit	Salinispora tropica CNB-440	ABP53490 (11)	D,B, P,R	30 , 7	19 , 15 , 7	531 , 4536
Vancomycin	Peptidoglycan remodeling	Amycolatopsis orientalis DSM 40040	CCD33128, CCD33129, CCD33130 (12)	B, R ***	50 , 8	39 , 22 , 17	563 , 8194
Cephamycin	duplicated beta- lactamase	Streptomyces clavuligerus ATCC 27064	AAF86620 (13)	B, D, R	26 , 3	45 , 20 , 15	546 , 7730
GE2270	duplicated elongation factor	Planobispora rosea ATCC 53733 y seen; ** Other hits four	AGY49599, AGY49600 (14)	D,B, P	34 , 6	26 , 15 , 9	549 , 8176

Notes: * Intra-genus phylogeny hits only seen; ** Other hits found using the advanced noise cutoff "E1" (90% model noise cutoff); *** Only *vanX* gene is detected

Table S10

ARTS results of example BGCs from the MIBiG database that contain known self-resistance mechanisms in their sequence. "Core" and "Res." Indicate a match to a gene for core and resistance model respectively. Where both models match the same protein, a "+" is indicated. Notes indicated with a star are listed in the row below

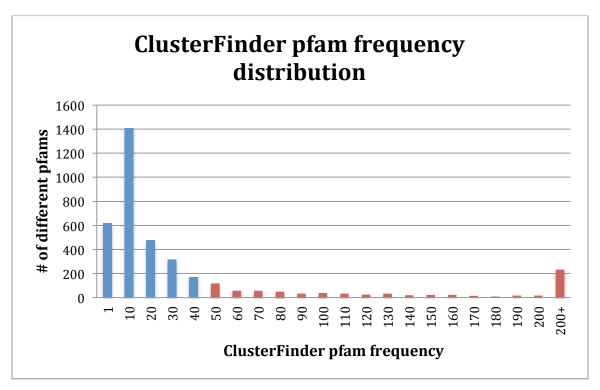
			Accession		
Product	Resistance gene	Organism	(ref)	BGC ID	ARTS Hit
Fiouuct	Resistance gene	Streptomyces sp. DSM	AKC91855	DGC ID	ANISTIIC
Griselimycin	Copy of dnaN	40835	(15)	BGC0001414	Core + Res.
Griseinnychi	copy of analy	Streptomyces	AAO47226	BGC0001414	Core i ites.
Coumermycin	Copy of gyrB	rishiriensis DSM 40489	(16)	BGC0000833	Core + Res.
, , , , , , , , , , , , , , , , , , , ,		Streptomyces niveus	(- /		
Novobiocin	Copy of gyrB	NCIMB 9219	AFI47646 (1)	BGC0000834	Core + Res.
	pentapeptide repeat	Xanthomonas			
Albicidin	protein for GyrB	albilineans GPE PC73	CBA16025 (3)	BGC0001088	Res.
	pentapeptide repeat		AKP45389		
Cystobactamide	protein for GyrB	Cystobacter sp. Cbv34	(17)	BGC0001413	Res.
D.C		Amycolatopsis		500000105	***
Rifamycin	mutated rpoB	mediterranei	AAS07760 (5)	BGC0000136	***Core + Res
Vatalianista	Copy of DNA	Streptomyces sp. TP- A2060	ADZ13541	DCC0000466	Ni a lata
Yatakemycin	glycosylase Copy of DNA	A2060	(18) ABY83174	BGC0000466	No hit
Azinomycin	glycosylase	Streptomyces sahachiroi	(19)	BGC0000960	No hit
, anomy cm	two copies of	Streptomyces	(13)	200000000	140 IIIC
	Initiation factor	achromogenes subsp.	CAI94679,		
Rubradirin	****	rubradiris	CAI94684 (20)	BGC0000141	Core,Core
	two copies of	7407441115	C/ 113 100 1 (20)	200000112	55.5/55.5
	Ribosomal protein L11	Bacillus cereus ATCC	AAP11944,		
Thiocillin	****	14579	AAP11947 (7)	BGC0000612	Core,Core
			AGY49599,		
	two copies of	Planobispora rosea	AGY49600		
GE2270	elongation factor	ATCC 53733	(14)	BGC0001155	Core,Core
	duplicated 23S rRNA	Saccharopolyspora	WP_00995039		
Erythromycin	methyltransferase	erythraea NRRL2338	1 (8)	BGC0000055	Res.
			AAC69328,		
	duplicated 23S rRNA	Streptomyces	AAC69327		
Pikromycin	methyltransferase	venezuelae ATCC 15439	(21)	BGC0000094	Res.
		Streptomyces	AAG32067,		
	duplicated 23S rRNA	viridochromogenes	AAG32066		
Avilamycin	methyltransferase	Tue57	(22)	BGC0000026	No hit
		Pseudomonas			
	duplicated Ile-tRNA	fluorescens NCIMB	AAM12927	BCC0000403	
Mupirocin	synthetase duplicated Thr-tRNA	10586	(23) CAE45679	BGC0000182	Core
Borrelidin	synthetase	Streptomyces parvulus	(24)	PCC0000031	Coro
Borrellalli	duplicated Trp-tRNA	Streptomyces parvaius Streptomyces griseus	(24)	BGC0000031	Core
Indolmycin	synthase	ATCC12648	AJT38681 (25)	BGC0001206	Res.
domyon	duplicated beta-	Salinospora tropica	ABP53490	200001200	11001
Salinosporamide A	proteasome subunit	CNB-440	(11)	BGC0001041	Core + Res.
p		Streptomyces	, ,		
	duplicated beta-	hygroscopicus ATCC	AHB38505		
Eponemycin	proteasome subuint	53709	(26)	BGC0000345	Core + Res.
			CCD33128,		
			CCD33129,		
	Peptidoglycan	Amycolatopsis orientalis	CCD33130		
Vancomycin	remodeling	HCCB 10007	(12)	BGC0000455	Res.
		Streptomyces			
	duplicated beta-	clavuligerus ATCC	AAF86620		
Cephamycin	lactamase	27064	(13)	BGC0000319	Res.
District.	1 11 1 5 1 5 /5	Streptomyces platensis	A 054 274 0 (27)	DCC0001175	*Core
Platencin	duplicated FabB/F	MA7339	ACS13710 (27)	BGC0001156	**No hit

Thiolactomycin	duplicated FabB/F	Salinispora pacifica DSM 45543	ALJ49913 (28)	BGC0001237	*Core **No hit
Thiotetroamide	two copies of FabB/F	Streptomyces afghaniensis NRRL5621	ALJ49924, ALJ49919 (28)	BGC0001236	*Core,*Core **No hit
Kalimantacin	duplicated Fabl	Pseudomonas fluorescens BCCM_ID9359	ADD82948 (29)	BGC0001099	No hit
Andrimid	One copy of acetyl- CoA carboxyltransferase	Pantoea agglomerans	AAO39114 (30)	BGC0000956	Res.

^{*} Search used exploration mode with relaxed cutoff: 90% Noise cutoff (E1)

Figure S11

Distribution of pfam frequencies taken from the biosynthetic pfam frequency file in ClusterFinder. These were frequencies defined from a manually curated set of 732 known BGCs. Here we have highlighted those that are most consistently found to remove higher confidence biosynthetic functions from the core gene list



SUPPLEMENTAL METHODS

Core gene and known resistance searches

Initial steps of the analyses consist of searching for BGCs, essential genes, and known resistance models. Submitted sequences that do not have BGC annotations are first processed with antiSMASH v3 using default settings to identify BGCs. The annotated Genbank is then parsed using Biopython (31) to identify all protein coding sequences, rRNAs, and cluster annotations. DNA sequences and protein translations are written to FASTA files for Hidden Markov Model (HMM) searches using HMMER (32) v3. These models include those for known

^{**} No hits when using default search mode

^{***} Using the provided MIBiG cluster annotation by running though "Existing Antismash Job" section

^{****} Putative resistance, in vitro experiments not shown

resistance factors and core genes, as determined from previous analysis detailed in the reference set section. HMM domain results are parsed and the best model hit for a gene is extracted if it passes 50% coverage length thresholds of both model and gene. This value was chosen to allow for missing domains and incomplete sequences while reducing fragmented hits. Resistance models are based on Resfam (33) which include known resistance factors from several databases (34–36). Domain of Unknown Function (DUF) models are from the Pfam (37) database and are used to highlight potential novel chemistry in a cluster. Custom submitted models are appended to corresponding core gene and known resistance models then searches are performed using model specific trusted cutoffs by default.

Reference set and core gene detection

The current *Actinobacteria* reference set is comprised of complete genomes from 189 species representing 22 different families that are available through NCBI's RefSeq (38) database (Supplemental S1). Essential genes are inferred by a comparative genomics approach where ubiquitous "core genes" are those consistently found in reference organisms as detected using HMMER and Hidden Markov Models (HMMs) from the TIGRFAM (39) protein family database; In addition, predefined core genes from the TIGRFAM v15 "bacterial core gene set" (GenProp0799) are included. All TIGRFAM homologous proteins with emphasis on conserved function, "equivologs", and their hypothetical and domain variants are used for essential gene analysis. After HMM detection, counts for genes are recorded in a gene matrix consisting of all reference genomes. Family specific core genes are then defined as genes present in greater than 95% of genomes relative to each family based on the count matrix. Families with less than 10 genomes were combined and a lowered ubiquity threshold of 90% is used instead to account for variations due to more distant relations.

For accelerated gene tree creation, all core gene sequences are extracted into corresponding multi-record FASTA files and, where applicable, out-group sequences added using model matches from various sequences of Proteobacteria, which harbored many non-actinobacterial genomes with core gene matches. Each core gene protein FASTA file is then aligned with MAFFT (40) followed by a codon alignment with Pal2nal (41). Trimmed copies of each codon alignment are made using TrimAl (42) with the maximum likelihood optimized "automated1" setting. RaxML (43) is used to build each tree with 100 bootstrap replicates using GTRGAMMAI model selection. Pairwise selection (dN/dS) values were calculated for each alignment using the yn00 tool from the PAML(44) package and the median of all Nei-Gojobori dN/dS values were logged to the model metadata. Metadata for functional classification are taken from model descriptions and associated main categories in TIGRFAM Roles. Additional statistics such as global ubiquity and how often a gene appears as a single copy are also recorded using the matrix of gene counts in the reference.

Core gene filtering and exploration mode

To reduce false positives while retaining the ability to search for more potential targets, a second search mode is provided as an option. The default search filters core genes for unlikely targets whereas "exploration mode" omits this step. Core genes are filtered for transport and regulatory functions by terms found in the descriptions and additional biosynthetic genes are removed if the protein sequences from the corresponding HMM seed alignments yield positive hits for high frequency BGC Pfams. High frequency biosynthetic Pfams are determined using ClusterFinder (45) Pfam frequency data where those above a frequency of 50 were used. This threshold was conservatively chosen based on the histogram of different Pfams at each frequency (Supplemental S11).

Duplication screening

Duplication is determined comparatively using the sum of median and standard deviation of corresponding sequence counts in the reference set. Genes with counts greater than this baseline are recorded along with the bit-score, scaffold location, and reference count statistics provided for manual review.

BGC proximity screening

BGC proximity is calculated by finding all core gene locations that intersect with BGC boundaries on the same scaffold. Visualizations are appended to the antiSMASH generated graphics and colored by criteria to quickly identify the type of proximity hits. Results from DUF and resistance model hits are appended to cluster annotations in a similar manner where hits for both resistance and core models are marked as "CoreRes" indicating a potential known target (a core model that is also in the known resistance set).

Phylogenetic screening

Queries of genomes, which are part of the reference phyla, will be screened for genes subject to HGT. Input sequences that do not have enough core genes, or sequences not part of the reference phyla will fail this screen or produce inaccurate results so it is advised to switch this screen off when using these inputs. The screening for HGT involves making sequence alignments to build many gene trees from which tree reconciliation with species tree is used to infer HGT for each gene. Alignments are created by adding the extracted nucleotide core sequences to the pre-trimmed reference codon alignments using the add method in MAFFT (mafft --add). Appended alignments are stored for user export and trimmed copies are made using the automated1 method in TrimAl. Trimmed alignments and existing reference trees are used with the Evolutionary Placement Algorithm (46)(EPA) option available in RAxML to produce gene trees with the guery sequences added. The species tree is then inferred from a coalescent of multi-locus gene trees using ASTRAL(47). The set of gene trees used are all single copy genes present in every reference and query organism with dN/dS values < 1; 16S rRNA sequences are also included if present. Each gene tree is then compared to the species tree to delineate incongruences due to duplications, transfers, and loss; this is determined using the ranger-dtl-U(48) tool with default cost values. All transfers involving the guery organism are then parsed and sorted and an additional filtering of intra-genus transfers is applied if genus names match with reference.

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