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Immune Epitope Database and Analysis Program

2010 Annual IEDB Compendium

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Introduction

The Immune Epitope Database and Analysis Resource (IEDB) is a public repository of immune epitope data and is sponsored by the National Institute for Allergy and Infectious Diseases (NIAID). The IEDB development started in December 2003 and it became available to the public in a beta test phase on 15 February 2006. The IEDB contains data related to antibody and T cell epitopes for humans, non-human primates, rodents, and other animal species. The IEDB also makes available a variety of analytical and epitope prediction tools and resources within its Analysis Resource.

This sixth Annual Compendium of the Immune Epitope Database and Analysis Resource consists of three sections. The first section contains a list of the antibody and T cell epitope information in the database as of 14 January 2011. The second section describes the features of the IEDB 2.5 website. The third section lists the scientific publications in 2010 and 2009 for which the IEDB played a contributory role.

Since the publication of last year's 2009 Annual Compendium, the quantity of data available in the IEDB has increased significantly with the addition of almost 3000 fully curated references. By the end of the year, curation of peptidic epitope data relating to all infectious diseases, including NIAID Category A, B, and C priority pathogens and NIAID Emerging and Re-emerging infectious diseases, allergens, and autoimmune diseases was current through September 2010. Curation of non-peptidic epitopes related to infectious diseases was over 40% complete. Curation of non-peptidic allergen epitopes was 95% completed. Curation of transplant and alloantigen references had been started.

1 Antibody and T Cell Epitopes

Many new references and many new pathogens were added to the IEDB in 2010, as demonstrated in Table 1.1. The table lists the number of distinct B cell and T cell epitopes in the database by source species at the end of 2009 and 2010. Of the 2439 species/strains listed, 420 were added in 2010. It should be noted that the source organism is the species/strain from which the epitopes originate, and may not be from an infecting organism or vaccine target. The curation of MHC binding peptides, cross-reactive epitopes, and autoimmune epitopes has resulted in the appearance of human (*Homo sapiens*) and mouse (*Mus musculus*) epitopes on the list.

In the table below, the leftmost column labeled "New 2010" indicates with an "X" if epitopes for the species/strain was added to the IEDB in 2010. A font color of red is used to highlight the information in the row. The Organism ID matches the NCBI taxonomy ID if it exists. Otherwise it represents an IEDB-assigned identifier, which are eight digit numbers starting with "1000". These are easily distinguishable from the NCBI identifiers which are six digits or less. The columns labeled "B-09", "T-09", "B-10", and "T-10" indicate the cumulative number of distinct B cell and T cell epitopes in the database at the end of 2009 and 2010, respectively. The two rightmost columns display the differences in the B and T cell epitope counts from 2009 to 2010. The changes in B and T cell epitope counts are shown in red. In 2010, the number of B cell epitopes increased by 4660, from 20,653 to 25,313, and the number of T cell epitopes increased by 8,198, from 44,285 to 52,483.

Table 1.1 Summary of B and T cell epitopes contained in the IEDB

NEW-2010	ORGANISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	5755	Acanthamoeba castellanii	14	1	14	2		1
X	2148	Acholeplasma laidlawii				2		2
X	351607	Acidothermus cellulolyticus 11B				1		1
X	470	Acinetobacter baumannii				1		1
X	471	Acinetobacter calcoaceticus				1		1
X	29430	Acinetobacter haemolyticus			1		1	
X	10001530	Acinetobacter haemolyticus strain 57			1		1	
X	10001531	Acinetobacter haemolyticus strain 61			1		1	
X	10001503	Acinetobacter lwoffii F78			1		1	
X	62977	Acinetobacter sp. ADP1				1		1
	228399	Actinobacillus pleuropneumoniae serovar 1 str. 4074	2		2	1		1
X	416269	Actinobacillus pleuropneumoniae serovar 5b str. L20				1		1
	209841	Actinobacillus pleuropneumoniae serovar 7	1		1			
X	272636	Adeno-associated virus				10		10
	10804	Adeno-associated virus - 2	20	8	20	67		59
	202813	Adeno-associated virus - 8			7		7	
	10508	Adenoviridae	3	2	3	2		
	4494	Aegilops markgrafii			1		1	
X	105751	Aeromonas bestiarum				1		1
	645	Aeromonas salmonicida	59		59			
X	29491	Aeromonas salmonicida subsp. salmonicida				1		1
	272557	Aeropyrum pernix K1		2		2		
	117204	African horse sickness virus 3	21		21			
	36421	African horse sickness virus 4	28		28			
	10497	African swine fever virus	1		1			
	85777	Agelas mauritiana			1		1	
	714	Aggregatibacter actinomycetemcomitans			4	1	4	1
X	358	Agrobacterium tumefaciens				1		1
	5039	Ajellomyces dermatitidis		1		1		
	10000828	Ajellomyces dermatitidis ATCC 60636		1		1		
	65690	AK7 murine leukemia virus			1		1	
	11791	AKR (endogenous) murine leukemia virus			7		7	
	11790	AKT8 murine leukemia virus			1		1	
X	10001458	Alcaligenes eutrophus				1		1
X	511	Alcaligenes faecalis				1		1
	512	Alcaligenes sp.			1		1	
	10783	Aleutian mink disease parvovirus (STRAIN G)	3		3			
	28314	Aleutian mink disease virus	1		1			
	172148	Alkhurma hemorrhagic fever virus			4		4	
	5599	Alternaria alternata	5		5			
X	314275	Alteromonas macleodii 'Deep ecotype'				1		1
	261202	Alto Paraguay hantavirus			3		3	
	45218	Amapari virus			3		3	
	4212	Ambrosia artemisiifolia	4	2	4	2		
	4215	Ambrosia artemisiifolia var. elatior	9	12	9	12		
	4214	Ambrosia trifida			6		6	
	86782	Amur virus			34		34	
	170955	Amur virus Solovey/AP63/1999			2		2	
X	4011	Anacardiaceae				4		4
	171929	Anacardium occidentale	27		50		23	
	4615	Ananas comosus	3		3			
	769	Anaplasma centrale			1		1	

NEW 2010	ORGANISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	770	Anaplasma marginale	15	14	15	14		
	10000760	Anaplasma marginale South Idaho		2		2		
	320483	Anaplasma marginale str. Florida	19	59	19	59		
	234826	Anaplasma marginale str. St. Maries	16	1	16	1		
	212042	Anaplasma phagocytophilum HZ	21		21			
X	10001573	Anatid herpesvirus 1 Clone-03			1		1	
	46607	Andes virus	2	44	2	77		33
	10000553	Andes virus CHI-7913	53		53			
	6858	Androctonus australis	9		9			
	70175	Androctonus australis hector	21		23		2	
X	6860	Androctonus mauretanicus mauretanicus			1		1	
X	334426	Angiostrongylus costaricensis			1	1	1	1
	6269	Anisakis simplex	1		8		7	
	7165	Anopheles gambiae		1		1		
	7460	Apis mellifera	14	99	15	99	1	
	7469	Apis mellifera ligustica		1		1		
	4045	Apium graveolens		14		14		
	224324	Aquifex aeolicus VF5		3		3		
	3702	Arabidopsis thaliana		4		4		
	201444	Aracatuba virus		5		5		
	3818	Arachis hypogaea	370	26	377	26	7	
X	308159	Araucaria virus				3		3
	10000980	Arcanobacterium pyogenes Strain 42	4		4			
	224325	Archaeoglobus fulgidus DSM 4304		1		1		
	41118	Arenavirus sp.		2		2		
	3704	Armoracia rusticana	1		3		2	
	6661	Artemia franciscana			2		2	
	4220	Artemisia vulgaris	1	19	1	19		
X	29320	Arthrobacter nicotinovorans			1		1	
X	290399	Arthrobacter sp. FB24				1		1
	6253	Ascaris suum		1		1		
	5085	Aspergillus fumigatus	115	28	115	30		2
X	5061	Aspergillus niger				1		1
X	5067	Aspergillus parasiticus				1		1
	5064	Aspergillus restrictus	1		1			
	287752	Aurantimonas manganoxydans SI85-9A1		1		1		
	4497	Avena nuda		1		1		
	4498	Avena sativa		4		4		
	11861	Avian erythroblastosis virus	1		1			
	172851	Avian hepatitis E virus	10		10			
	11127	Avian infectious bronchitis virus (strain M41)	4	3	4	3		
	231428	Avian infectious bronchitis virus (strain Vic S)	11	8	11	8		
	38171	Avian reovirus strain S1133	2		2			
	195700	Avian rotavirus PO-13	6		6			
X	354	Azotobacter vinelandii				2		2
X	322710	Azotobacter vinelandii DJ				1		1
	5866	Babesia bigemina		3		3		
	5865	Babesia bovis	3	8	3	8		
X	10001459	Babesia bovis Argentina R1A			4		4	
	10000382	Babesia bovis Mexico	1	4	1	4		
	10000383	Babesia bovis Mexico Mo7	1	33	1	33		
	5872	Babesia equi	8		8			
	5868	Babesia microti	2		2			

NEW 2010	ORGANISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	120505	Baboon cytomegalovirus		1		1		
	196403	Baboon endogenous virus		1	1	1		
	11764	Baboon endogenous virus strain M7		1		1		
	1390	Bacillus amyloliquefaciens		2		2		
	1392	Bacillus anthracis	127	188	215	197	88	9
X	486619	Bacillus anthracis str. A0193				1		1
X	592021	Bacillus anthracis str. A0248				1		1
	486623	Bacillus anthracis str. A0389		1		4		3
X	486621	Bacillus anthracis str. A0442				1		1
	486620	Bacillus anthracis str. A0465		5		5		
	486624	Bacillus anthracis str. A0488		2		81		79
X	191218	Bacillus anthracis str. A2012				28		28
X	198094	Bacillus anthracis str. Ames				1		1
	261594	Bacillus anthracis str. 'Ames Ancestor'		1		1		
X	568206	Bacillus anthracis str. CDC 684				2		2
	260799	Bacillus anthracis str. Sterne	5	3	5	3		
	10000291	Bacillus anthracis str. Sterne 34F2		1		1		
X	405536	Bacillus anthracis Tsiankovskii-I				1		1
	1396	Bacillus cereus		1	2	1	4	2
X	572264	Bacillus cereus 03BB102				1		1
	451709	Bacillus cereus 03BB108		8		8		
	405533	Bacillus cereus AH1134		3		3		
	222523	Bacillus cereus ATCC 10987		1		1		
	226900	Bacillus cereus ATCC 14579		1		2		1
X	526977	Bacillus cereus ATCC 4342				1		1
	405532	Bacillus cereus B4264		1		1		
	269801	Bacillus cereus G9241		3		34		31
	405531	Bacillus cereus G9842		1		1		
	451708	Bacillus cereus H3081.97		11		12		1
	451707	Bacillus cereus NVH0597-99		7		7		
	405917	Bacillus cereus W		1		1		
	441769	Bacillus coahuilensis m4-4		1		1		
	315749	Bacillus cytotoxicus NVH 391-98		2		2		
	1467	Bacillus lentus		1		1		
	1402	Bacillus licheniformis		9		10		1
	1404	Bacillus megaterium		1		1		
	1423	Bacillus subtilis	2	1	2	4		3
X	224308	Bacillus subtilis subsp. subtilis str. 168				1		1
X	535026	Bacillus subtilis subsp. subtilis str. NCIB 3610				1		1
	1428	Bacillus thuringiensis	5		5			
	339854	Bacillus thuringiensis serovar israelensis ATCC 35646		2		6		4
	29339	Bacillus thuringiensis serovar kurstaki	3		3			
	1435	Bacillus thuringiensis serovar san diego		2		2		
X	527026	Bacillus thuringiensis serovar sotto str. T04001				1		1
X	527024	Bacillus thuringiensis serovar tochigiensis BGSC 4Y1				1		1
	412694	Bacillus thuringiensis str. Al Hakam		2		9		7
	315730	Bacillus weihenstephanensis KBAB4		3		4		1
	2	Bacteria	2		6		4	
	12040	Barley yellow dwarf virus-PAV	1		1			
X	283166	Bartonella henselae str. Houston-1				1		1
	37962	Bayou virus	1		1			
	12260	Bean pod mottle virus	2		2			
	31715	Bean-pod mottle virus (strain Kentucky G7)	9	1	9	1		

NEW 2010	ORGANISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	31721	Beet necrotic yellow vein virus	10		10			
	12161	Beet yellows virus	5		5			
	3645	Bertholletia excelsa	7	24	7	24		
X	161934	Beta vulgaris			3		3	
	3505	Betula pendula	36	182	36	203		21
	10629	BK polyomavirus		45		52		7
	65743	Blackcurrant reversion virus	2		2			
	6973	Blattella germanica	5	15	5	15		
	40697	Blomia tropicalis	18		18			
	40051	Bluetongue virus	7		7			
	10904	Bluetongue virus (serotype 1 / isolate Australia)	7		7			
	10900	Bluetongue virus (serotype 10 / American isolate)	3		3			
	33717	Bluetongue virus (serotype 13 / isolate USA)	2		2			
	33718	Bluetongue virus (serotype 17 / isolate USA)	4		4			
	35327	Bluetongue virus 1	3		3			
	10906	Bluetongue virus 10	3		3			
	35329	Bluetongue virus 11	6		6			
	35330	Bluetongue virus 13	1		1			
	10903	Bluetongue virus 17	4		4			
	94967	Bluetongue virus 4	2		2			
	388634	Bombyx mandarina nuclear polyhedrosis virus			1		1	
	271108	Bombyx mori NPV	2		2			
	360910	Bordetella avium 197N			1		1	
	518	Bordetella bronchiseptica			1		1	
	520	Bordetella pertussis	221	44	224	45	3	1
	257313	Bordetella pertussis Tohama I	1	11	1	11		
	12455	Borna disease virus	8	4	8	4		
X	10000518	Borna disease virus Giessen strain He/80				6		6
	29518	Borrelia afzelii			1		1	
	390236	Borrelia afzelii PKo	2		2			
	139	Borrelia burgdorferi	39	38	39	39		1
	498740	Borrelia burgdorferi 64b			2		2	
	224326	Borrelia burgdorferi B31	66	10	66	10		
	10001091	Borrelia burgdorferi BEP4	1		1			
	10000675	Borrelia burgdorferi CA12			6		6	
	64895	Borrelia burgdorferi group	3		3			
	521007	Borrelia burgdorferi N40	4	3	4	3		
	445985	Borrelia burgdorferi ZS7	2	45	2	45		
	412419	Borrelia duttonii Ly	7		7			
	29519	Borrelia garinii	2	29	2	29		
	10000530	Borrelia garinii IP90	3		3			
	9913	Bos taurus	847	238	1026	362	179	124
	8722	Bothrops asper	1		1			
	11128	Bovine coronavirus	7		7			
	12064	Bovine enterovirus		13		13		
	12065	Bovine enterovirus strain VG-5-27	6		6			
	10000472	Bovine ephemeral fever virus BB7721	2		2			
	10320	Bovine herpesvirus 1	4	35	4	36		1
	10000404	Bovine herpesvirus 1 Lam	1		1			
	263683	Bovine herpesvirus 5 strain TX89	2		2			
	79889	Bovine herpesvirus type 1.1	1		1			
	10323	Bovine herpesvirus type 1.1 (strain Cooper)	8	18	8	18		
	10324	Bovine herpesvirus type 1.1 (strain P8-2)	1		1			

NEW 2010	ORGANISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	11901	Bovine leukemia virus	32	29	32	29		
	10562	Bovine papillomavirus - 4	10	3	10	3		
	10559	Bovine papillomavirus type 1	1		1			
	11215	Bovine parainfluenza virus 3		1		1		
	11246	Bovine respiratory syncytial virus	2	1	2	1		
	31611	Bovine respiratory syncytial virus (strain 391-2)	3		3			
	11249	Bovine respiratory syncytial virus (strain RB94)	3		3			
	82823	Bovine respiratory syncytial virus strain lelystad	1		1			
	82824	Bovine respiratory syncytial virus strain snook		75		75		
	10927	Bovine rotavirus	7	1	7	1		
	36439	Bovine rotavirus strain NCDV/G6	1		1			
	10933	Bovine rotavirus strain RF	12	7	12	7		
	10934	Bovine rotavirus strain UK/G6		1		1		
	11099	Bovine viral diarrhea virus 1	1	11	1	11		
	11100	Bovine viral diarrhea virus 1-NADL	3	3	3	3		
	54315	Bovine viral diarrhea virus 2	1		1			
	82470	Bovine viral diarrhea virus strain Oregon C24V	5		5			
	158474	Bovine viral diarrhea virus strain Trangie Y546	1		1			
X	288000	Bradyrhizobium sp. BTAl1					1	1
	3707	Brassica juncea	9		9			
	235	Brucella abortus	4	29	4	29		
X	520450	Brucella abortus bv. 2 str. 86/8/59					1	1
X	575591	Brucella abortus NCTC 8038					1	1
X	430066	Brucella abortus S19					13	13
X	641140	Brucella abortus str. 2308 A					1	1
	10001424	Brucella abortus W99	1		1			
X	483179	Brucella canis ATCC 23365					6	6
	29459	Brucella melitensis	2	32	2	63		31
	359391	Brucella melitensis biovar Abortus 2308		30		31		1
	224914	Brucella melitensis bv. 1 str. 16M	6	2	6	6		4
	520464	Brucella melitensis bv. 1 str. Rev.1	1		1			
	236	Brucella ovis	2		2	1		1
	444178	Brucella ovis ATCC 25840		1		5		4
	29461	Brucella suis		7		7		
	204722	Brucella suis 1330		26		28		2
	470137	Brucella suis ATCC 23445		1		3		2
	6279	Brugia malayi		1	2	3	2	2
	89462	Bubalus bubalis		1		1		
	32605	Buffalopox virus		2		2		
	8616	Bungarus multicinctus	20	3	20	3		
	339670	Burkholderia ambifaria AMMD		3		4		1
X	331271	Burkholderia cenocepacia AU 1054					4	4
	350702	Burkholderia cenocepacia PC184		1		1		
	292	Burkholderia cepacia		1	3	1	3	
X	134537	Burkholderia fungorum				1		1
	13373	Burkholderia mallei		45		45		
	243160	Burkholderia mallei ATCC 23344		36		38		2
X	334802	Burkholderia mallei FMH				1		1
X	320390	Burkholderia mallei GB8 horse 4				1		1
X	334803	Burkholderia mallei JHU				2		2
	412022	Burkholderia mallei NCTC 10229		1		9		8
X	320388	Burkholderia mallei SAVP1					24	24
	28450	Burkholderia pseudomallei		1		1		

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X	357348	Burkholderia pseudomallei 1106a				1		1
X	441160	Burkholderia pseudomallei 14				1		1
	320372	Burkholderia pseudomallei 1710b		3		3		
X	425067	Burkholderia pseudomallei 305				1		1
X	360118	Burkholderia pseudomallei 406e				1		1
	320373	Burkholderia pseudomallei 668		1		5		4
X	441158	Burkholderia pseudomallei 9				1		1
X	441159	Burkholderia pseudomallei 91				1		1
X	272560	Burkholderia pseudomallei K96243				5		5
X	320374	Burkholderia pseudomallei S13				1		1
	269483	Burkholderia sp. 383		2		2		
	271848	Burkholderia thailandensis E264		2		2		
X	269482	Burkholderia vietnamiensis G4				9		9
	6239	Caenorhabditis elegans		11		11		
	35305	California encephalitis virus		1		1		
	28873	Camelpox virus		2		2		
	203172	Camelpox virus CMS		26		28		2
	203174	Camelpox virus CP1		1		1		
	203173	Camelpox virus M-96		132		133		1
	306254	Campylobacter coli RM2228		1		3		2
	32019	Campylobacter fetus subsp. fetus		3		3		
	197	Campylobacter jejuni		1	16	1	20	4
	195099	Campylobacter jejuni RM1221			13		13	
	407148	Campylobacter jejuni subsp. jejuni 81116			13		14	1
	354242	Campylobacter jejuni subsp. jejuni 81-176			1		3	2
X	482628	Campylobacter jejuni subsp. jejuni BH-01-0142				1		1
	360112	Campylobacter jejuni subsp. jejuni HB93-13			2		2	
	192222	Campylobacter jejuni subsp. jejuni NCTC 11168		10		10	12	12
X	306263	Campylobacter lari RM2100				1		1
	44088	Canarypox virus			2		6	4
	5476	Candida albicans		85	56	91	66	6 10
	10000335	Candida albicans A-9 (serotype B)		1		1		
	10000337	Candida albicans KIT 1113		1		1		
	10000338	Candida albicans LGH1095 (serotype B)		1		1		
X	237561	Candida albicans SC5314					1	1
	10000339	Candida albicans serotype A		1		1		
	5480	Candida parapsilosis		1		1		
	234267	Candidatus Solibacter usitatus Ellin6076			1		2	1
	292348	Canine calicivirus (strain 48)		2		2		
	11232	Canine distemper virus			1		2	1
	11233	Canine distemper virus strain Onderstepoort		6	19	6	19	
	35258	Canine oral papillomavirus			25		25	
	10788	Canine parvovirus		21	20	21	20	
	246878	Canine parvovirus 2		3		3		
	10790	Canine parvovirus strain CPV-D CORNELL 320		1		1		
	9615	Canis lupus familiaris			61	3	61	3
	11660	Caprine arthritis encephalitis virus		12	1	12	1	
	11662	Caprine arthritis encephalitis virus G63		6		6		
	11661	Caprine arthritis encephalitis virus strain Cork		9		9		
	7957	Carassius auratus		3		3		
	64289	Carey Island virus			2		2	
	10141	Cavia porcellus		15	63	29	72	14 9
	6878	Centruroides noxioides		7		7		

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	13415	Chamaecyparis obtusa	1	62	1	62		
X	266779	Chelativorans sp. BNC1				2		2
	12618	Chicken anemia virus	3		3			
	37124	Chikungunya virus		3		3		
	310542	Chimpanzee adenovirus	2		2			
	7154	Chironomus thummi	2	3	2	3		
	7155	Chironomus thummi thummi	58	27	58	27		
	810	Chlamydia	5		5			
	243161	Chlamydia muridarum Nigg		14		14		
	813	Chlamydia trachomatis	77	43	118	52	41	9
	315277	Chlamydia trachomatis A/HAR-13	6	1	6	1		
	10000858	Chlamydia trachomatis B/Jali-20/OT	2		2			
	10000762	Chlamydia trachomatis B/Tw-5/OT	5		5			
	272561	Chlamydia trachomatis D/UW-3/CX		1		1		
	10001141	Chlamydia trachomatis Serovar A	22	8	22	8		
	10000804	Chlamydia trachomatis Serovar B	42	5	43	5	1	
	10000763	Chlamydia trachomatis Serovar C	16	1	16	1		
	10001085	Chlamydia trachomatis Serovar D	1		1			
	10001121	Chlamydia trachomatis Serovar Da	1		1			
	10000764	Chlamydia trachomatis Serovar E	18	1	18	1		
	10000845	Chlamydia trachomatis Serovar F	3		3			
	10000765	Chlamydia trachomatis Serovar H	7		7			
	10000766	Chlamydia trachomatis Serovar I	10		10			
	10001008	Chlamydia trachomatis Serovar J	6		6			
	10000767	Chlamydia trachomatis serovar K	14		14			
	10000853	Chlamydia trachomatis Serovar L1	21	3	21	3		
	10000768	Chlamydia trachomatis Serovar L2	15	7	15	7		
	10000769	Chlamydia trachomatis Serovar L3	1		2		1	
	204428	Chlamydiae	1		1			
	83555	Chlamydiphila abortus	3		3			
	10000559	Chlamydiphila abortus B-577	11		11			
	83558	Chlamydiphila pneumoniae	52	19	56	21	4	2
	115713	Chlamydiphila pneumoniae CWL029		49		49		
	10000852	Chlamydiphila pneumoniae Kajaani 6		7		7		
	182082	Chlamydiphila pneumoniae TW-183	7		7			
	83554	Chlamydiphila psittaci	71		71	2		2
	9534	Chlorocebus aethiops		3		3		
X	169173	Chocho virus				2		2
	12162	Citrus tristeza virus	1		4		3	
	11096	Classical swine fever virus	3	1	4	1	1	
	358769	Classical swine fever virus - Alfort/187	2		2			
	358805	Classical swine fever virus - Alfort/Tuebingen	5		5			
	11098	Classical swine fever virus - Brescia	4		4			
X	279150	Classical swine fever virus 96TD			2		2	
	10001025	Classical swine fever virus Glentorf		26		26		
X	10001578	Classical swine fever virus LPC/AHRI			2		2	
	10000451	Classical swine fever virus Shimen	16		16			
	36911	Clavispora lusitaniae	1		1			
	214432	Cloning vector pscFvCA-E8VHd		1		1		
	1491	Clostridium botulinum	121	57	123	60	2	3
	36826	Clostridium botulinum A	44		44			
	10000293	Clostridium botulinum A 1	1		1			
	10000294	Clostridium botulinum A 2	2		2			

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	10000302	Clostridium botulinum A Kyoto-F	1		1			
X	413999	Clostridium botulinum A str. ATCC 3502				3		3
	441771	Clostridium botulinum A str. Hall	43		44		1	
	10000301	Clostridium botulinum A str. Hall hyper	3		3			
	36827	Clostridium botulinum B	28		28			
	10000295	Clostridium botulinum B 111	2		2			
	10000303	Clostridium botulinum B Lammanea	1		1			
	10000305	Clostridium botulinum B Okra	3		3			
X	508765	Clostridium botulinum B str. Eklund 17B				1		1
	10000297	Clostridium botulinum C 92-13	5		5			
	10000306	Clostridium botulinum C Stockholm	2		2			
	36829	Clostridium botulinum D	2		2			
	10000296	Clostridium botulinum D 1873	2		2			
	36830	Clostridium botulinum E	3	3	3	3		
	10000299	Clostridium botulinum E Beluga	2		2			
X	508767	Clostridium botulinum E3 str. Alaska E43				1		1
	36831	Clostridium botulinum F	2		2			
	10000304	Clostridium botulinum F NCTC 10281	1		1			
	445335	Clostridium botulinum NCTC 2916		2		2		
	445338	Clostridium botulinum str. Iwanei E	6		6			
	1496	Clostridium difficile	15		15			
	10000307	Clostridium difficile BART'S W1	1		1			
X	386415	Clostridium novyi NT				2		2
	29362	Clostridium papyrosolvens	1		1			
	1502	Clostridium perfringens	10		10			
	107819	Clostridium perfringens D	1		1			
	195102	Clostridium perfringens str. 13		1		1		
	1513	Clostridium tetani	76	227	76	231		4
	212717	Clostridium tetani E88		16		16		
X	203119	Clostridium thermocellum ATCC 27405				1		1
	5501	Coccidioides immitis		1		1		
	199306	Coccidioides posadasii		2		2		
	5503	Cochliobolus lunatus	10	10	10	10		
	9014	Colinus virginianus		2	1	2	1	
	8932	Columba livia		7		8		1
X	314285	Congregibacter litoralis KT71				1		1
	6491	Conus geographus	30		30			
	6492	Conus magus	2		2			
	6493	Conus striatus	11		11			
	32614	Convict Creek 107 virus	1	3	1	3		
	13451	Corylus avellana	27	27	27	27		
	1717	Corynebacterium diphtheriae	10	50	10	50		
	152794	Corynebacterium efficiens		5		5		
X	196164	Corynebacterium efficiens YS-314				2		2
	1718	Corynebacterium glutamicum		21		22		1
X	196627	Corynebacterium glutamicum ATCC 13032				7		7
	306537	Corynebacterium jeikeium K411		1		1		
	10703	Corynephage beta	2		2			
	186541	Cote d'Ivoire ebolavirus		1		1		
	128999	Cote d'Ivoire ebolavirus - Cote d'Ivoire (Tai Forest, 1994)		3		3		
	10623	Cottontail rabbit papillomavirus	3	5	3	6		1
	12264	Cowpea mosaic virus		1		1		
	10243	Cowpox virus		19		22		3

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	265872	Cowpox virus (Brighton Red)		1		1		
	10000571	Cowpox virus (Brighton Red) White-pock		1		1		
	777	Coxiella burnetii		32		32		
X	434923	Coxiella burnetii CbuG_Q212				1		1
X	434922	Coxiella burnetii Dugway 5J108-111				6		6
X	360116	Coxiella burnetii 'MSU Goat Q177'				3		3
X	360115	Coxiella burnetii RSA 331				6		6
	360117	Coxiella burnetii RSA 334		14		14		
	227377	Coxiella burnetii RSA 493		1		9		8
X	12066	Coxsackievirus				1		1
	103903	Coxsackievirus B3 (strain Nancy)	13	30	13	30		
	103905	Coxsackievirus B4 (strain E2)	5	41	6	41	1	
	103906	Coxsackievirus B4 (strain JVB / Benschoten / New York/51)		76		76		
	10029	Cricetulus griseus	1		1			
	11593	Crimean-Congo hemorrhagic fever virus		1		1		
	368445	Crocodilepox virus		1		2		1
	8732	Crotalus durissus terrificus	4		4			
X	283643	Cryptococcus neoformans var. neoformans B-3501A				2		2
	10001132	Cryptococcus neoformans var. neoformans Serotype A	1		1			
	3369	Cryptomeria japonica	25	181	39	181	14	
	237895	Cryptosporidium hominis		3		3		
	5807	Cryptosporidium parvum		23		26		3
	353152	Cryptosporidium parvum Iowa II		27		76		49
	220837	Cryptosporidium sp. MNJ-1		1		1		
	12305	Cucumber mosaic virus	1		1			
X	117125	Cucumber mosaic virus (strain Pepo)			1		1	
	3656	Cucumis melo	12		12			
	208899	Cupixi virus		1		1		
X	266264	Cupriavidus metallidurans CH34				2		2
	301964	CY1014 virus		1		1		
	46457	Cycloclasticus oligotrophus		1		1		
	28909	Cynodon dactylon	3	23	3	23		
	10358	Cytomegalovirus		32		32		
X	269798	Cytophaga hutchinsonii ATCC 33406				1		1
X	4509	Dactylis glomerata				2		2
	7955	Danio rerio		3		3		
	4039	Daucus carota		1		1		
	305674	Deerpox virus W-848-83		1		2		1
	243164	Dehalococcoides ethenogenes 195		1		1		
	12637	Dengue virus	1	25	9	47	8	22
	11053	Dengue virus 1	3	46	11	59	8	13
	11059	Dengue virus 1 Nauru/West Pac/1974	1		1			
	33741	Dengue virus 1 Singapore/S275/1990		30		30		
	11060	Dengue virus 2	79	92	81	95	2	3
	31635	Dengue virus 2 16681-PDK53		2		2		
	11064	Dengue virus 2 Jamaica/1409/1983	262	11	262	11		
	11062	Dengue virus 2 Malaysia M2	1		1			
	11066	Dengue virus 2 Puerto Rico/PR159-S1/1969	30		30			
	31634	Dengue virus 2 Thailand/16681/84	9	55	10	55	1	
	11065	Dengue virus 2 Thailand/NGS-C/1944	14	10	15	16	1	6
	11069	Dengue virus 3	2	58	17	60	15	2
	408870	Dengue virus 3 Philippines/H87/1956	4		4	4		4
	11070	Dengue virus 4	5	41	5	41		

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	408871	Dengue virus 4 Dominica/814669/1981	1	5	1	5		
	408688	Dengue virus 4 Thailand/0348/1991		1		1		
	10000965	Dengue virus type 1 FGA/89	1		1			
	10000440	Dengue virus type 1 Hawaii		9		9		
	10000442	Dengue virus type 3 CH53489		6		6		
X	246753	Dengue virus vector p4(Delta30)				2		2
	6954	Dermatophagoides farinae	12	36	12	36		
	6956	Dermatophagoides pteronyssinus	54	104	58	104	4	
X	439235	Desulfatibacillum alkenivorans AK-01				1		1
X	272564	Desulfitobacterium hafniense DCB-2				2		2
X	281689	Desulfovomonas acetoxidans DSM 684				1		1
	11319	Dhori virus (strain Indian/1313/61)	1		1			
X	44689	Dictyostelium discoideum				1		1
	12506	Dobrava-Belgrade virus			13		16	3
	7441	Dolichovespula maculata	11	20	11	20		
	292633	Dragon grouper nervous necrosis virus	20		20			
	7227	Drosophila melanogaster	2	2	3	11	1	9
	12639	Duck hepatitis B virus	189	20	189	20		
	10000466	Duvenhage virus 6			2		2	
X	11021	Eastern equine encephalitis virus				1		1
	11022	Eastern equine encephalitis virus (STRAIN VA33[TEN BROECK])		1		1		
	10000439	Eastern equine encephalitis virus SV	8		8			
	205488	Ebola virus sp.		1		3		2
	6210	Echinococcus granulosus	37		37			
	99586	Echis ocellatus	5		5			
	33758	Echovirus		1		1		
	12643	Ectromelia virus		2		5		3
	944	Ehrlichia canis	7		7			
	945	Ehrlichia chaffeensis	9		13	1	4	1
X	205920	Ehrlichia chaffeensis str. Arkansas			15		15	
	35795	Ehrlichia muris		1		1		
	779	Ehrlichia ruminantium	3		3			
	5801	Eimeria acervulina	1		1			
	5802	Eimeria tenella	6		6			
	35321	El Moro Canyon virus		2		2		
	6035	Encephalitozoon cuniculi		5		5		
	284813	Encephalitozoon cuniculi GB-M1		5		27		22
	12104	Encephalomyocarditis virus		1		1		
X	370354	Entamoeba dispar SAW760				15		15
	5759	Entamoeba histolytica	29	8	29	13		5
	294381	Entamoeba histolytica HM-1:IMSS		46		97		51
	10000352	Entamoeba histolytica YS-27	1		1			
	12340	Enterobacteria phage 933J	2		2			
	10730	Enterobacteria phage 933W	1		1			
X	291401	Enterobacteria phage CP-1639				1		1
	10863	Enterobacteria phage f1		1		1		
	10864	Enterobacteria phage fd	7		7			
	10710	Enterobacteria phage lambda		4	2	6	2	2
	10754	Enterobacteria phage P22	5		5			
	10658	Enterobacteria phage PRD1	2		2			
X	55884	Enterobacteria phage SfV				1		1
	10665	Enterobacteria phage T4	10	20	10	20		
	10760	Enterobacteria phage T7	1		1			

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	543	Enterobacteriaceae	1		2		1	
	12022	Enterobacterio phage MS2	1		1			
	1351	Enterococcus faecalis	1		1			
	1352	Enterococcus faecium	6		6			
X	333849	Enterococcus faecium DO				1		1
	12059	Enterovirus		9		9		
	150846	Enterovirus 5865/sir/000009	2		2			
X	31330	Ephydatia fluviatilis				1		1
	82830	Epstein-barr virus strain ag876	1	1	1	1		
	10326	Equid herpesvirus 1	5		5			
	10000525	Equid herpesvirus 2 16V	1		1			
	10000526	Equid herpesvirus 2 5FN	1		1			
	10000524	Equid herpesvirus 2 691	1		1			
	10000391	Equid herpesvirus 2 ER32	1		1			
	10000527	Equid herpesvirus 2 FIN60	1		1			
	10331	Equid herpesvirus 4	5		5			
	10000405	Equid herpesvirus 4 TH20	2		2			
	82831	Equid herpesvirus type 2 strain 86/87	1		1			
	11047	Equine arteritis virus	2		2			
	11665	Equine infectious anemia virus	91	109	91	109		
	11670	Equine infectious anemia virus (CLONE 1369)		1		1		
	11671	Equine infectious anemia virus (STRAIN WSU5)		16		16		
	10000499	Equine infectious anemia virus PV		2		2		
	10000835	Equine rhinitis A virus 393/76	4		4			
	9796	Equus caballus	2	43	4	44	2	1
	9798	Equus przewalskii		1		1		
	562	Escherichia coli	169	48	176	95	7	47
X	358709	Escherichia coli 101-1				5		5
	10001145	Escherichia coli 1471	2		2			
	10000727	Escherichia coli 180/C3	1		1			
	362663	Escherichia coli 536		2		58		56
	525281	Escherichia coli 83972		2		3		1
X	405955	Escherichia coli APEC O1				7		7
X	481805	Escherichia coli ATCC 8739				1		1
	37762	Escherichia coli B	1	1	1	1		
	10000728	Escherichia coli B B/r CM6		1		1		
X	344601	Escherichia coli B171				11		11
X	340184	Escherichia coli B7A				5		5
X	199310	Escherichia coli CFT073				1		1
X	340186	Escherichia coli E110019				1		1
X	340185	Escherichia coli E22				11		11
X	331111	Escherichia coli E24377A				31		31
	316401	Escherichia coli ETEC H10407	62	31	62	31		
X	340197	Escherichia coli F11				12		12
X	10001528	Escherichia coli F515			1		1	
X	331112	Escherichia coli HS				8		8
	585034	Escherichia coli IAI1	3		3			
X	10001502	Escherichia coli J-5			1		1	
	83333	Escherichia coli K-12	4	7	5	13	1	6
	168807	Escherichia coli O127:H6	1		1			
	83334	Escherichia coli O157:H7	2	103	2	103		
	478005	Escherichia coli O157:H7 str. EC4486		1		1		
	478006	Escherichia coli O157:H7 str. EC4501		2		2		

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	478008	Escherichia coli O157:H7 str. EC869		13		13		
	155864	Escherichia coli O157:H7 str. EDL933		1		19		18
X	386585	Escherichia coli O157:H7 str. Sakai				13		13
	10000733	Escherichia coli O5:K4:H4	1		1			
X	244320	Escherichia coli O55:H7				1		1
	217992	Escherichia coli O6		7		7		
	10000734	Escherichia coli O65:K:-H-	1		1			
X	316385	Escherichia coli str. K-12 substr. DH10B				2		2
X	511145	Escherichia coli str. K-12 substr. MG1655				26		26
X	316407	Escherichia coli str. K-12 substr. W3110				1		1
	364106	Escherichia coli UTI89		2		2		
	2759	Eukaryota	2		2			
	6958	Euroglyphus maynei		10		10		
	420521	Expression vector pNIC-NHT-CF	1		1			
	3617	Fagopyrum esculentum	39		39			
X	62330	Fagopyrum tataricum			4		4	
	6690	Farfantepenaeus aztecus	43		51		8	
	46835	Fasciola gigantica	2		2			
	6192	Fasciola hepatica	24	11	250	11	226	
	11978	Feline calicivirus	4		4			
	11981	Feline calicivirus (STRAIN F9)	2		2			
	11980	Feline calicivirus (STRAIN JAPANESE F4)	6		6			
	11673	Feline immunodeficiency virus		28		28		
	11674	Feline immunodeficiency virus (isolate Petaluma)	2	3	2	3		
	45409	Feline immunodeficiency virus (isolate wo)	3		3			
	36372	Feline immunodeficiency virus (strain UK8)		1		1		
	33734	Feline infectious peritonitis virus (strain 79-1146)	4		4			
	11769	Feline leukemia virus strain A/Glasgow-1	6		6			
	10001130	Feline leukemia virus subtype A	2		2			
	10786	Feline panleukopenia virus		4		4		
	9685	Felis catus	18	48	24	67	6	19
	4606	Festuca arundinacea	1		1			
X	5207	Filobasidiella neoformans				1		1
	156586	Flavobacteria bacterium BBFL7		1		1		
	12110	Foot-and-mouth disease virus	22	2	22	2		
	10000991	Foot-and-mouth disease virus - type A (strain A22 Iraq)	5		5			
	10000989	Foot-and-mouth disease virus - type A (strain A22)	8	6	8	6		
	110195	Foot-and-mouth disease virus - type Asia 1	2	2	7	2	5	
	10000995	Foot-and-mouth disease virus - type Asia 1 Pakistan	1		1			
	12116	Foot-and-mouth disease virus - type C	6	25	6	25		
	12118	Foot-and-mouth disease virus - type O	16	3	16	3		
	10000801	Foot-and-mouth disease virus - type O (O/SKR/2002)	1		1			
	10001118	Foot-and-mouth disease virus - type O (strain HKN/14/82)	2		2			
	10000856	Foot-and-mouth disease virus - type O isolate O/UKG/35/2001		13		13		
	10000992	Foot-and-mouth disease virus - type SAT 1 (Strain Bot 1/68)	1		1			
	10000993	Foot-and-mouth disease virus - type SAT 2 (Strain Ken 3/57)	1		1			
	10001000	Foot-and-mouth disease virus - type SAT 2 (Strain Rho 1/48)	4		4			
	10000994	Foot-and-mouth disease virus - type SAT 3 (Strain Bec 1/65)	1		1			
	12112	Foot-and-mouth disease virus (strain A10-61)	8		8			
	12114	Foot-and-mouth disease virus (strain A12)	21		21			
	12115	Foot-and-mouth disease virus (strain A24 Cruzeiro)	14	3	14	3		
	12113	Foot-and-mouth disease virus (strain A5)	3		3			
	12120	Foot-and-mouth disease virus (strain C1-Santa Pau)	1		1			

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	12117	Foot-and-mouth disease virus (strain C3 Indaial)	2		2			
	73482	Foot-and-mouth disease virus (strain O1)	18	1	18	1		
	10000820	Foot-and-mouth disease virus (strain O1) (O/Taiwan/1/97)	1		1			
	10000964	Foot-and-mouth disease virus (strain O1) (O1 Brugge)	2		2			
	10000516	Foot-and-mouth disease virus (strain O1) (O1 Campos)	18	18	18	18		
	10000513	Foot-and-mouth disease virus (strain O1) (O1 Kaufbeuren)	37	8	37	8		
	10000514	Foot-and-mouth disease virus (strain O1) (O1BFS 1860)	8		8			
	10000515	Foot-and-mouth disease virus (strain O1) (O1BFS)	4		4			
	10000556	Foot-and-mouth disease virus (strain O1) Kaufbeuren	42	11	42	11		
	161727	Foot-and-mouth disease virus A10/Holland	11	14	11	14		
	12121	Foot-and-mouth disease virus C1	1		1			
	10000840	Foot-and-mouth disease virus C1 Brescia It/64	2		2			
	10000836	Foot-and-mouth disease virus C1 CS8	22	24	22	24		
	46290	Foot-and-mouth disease virus C3	3	1	4	1	1	
	10001039	Foot-and-mouth disease virus C3 (strain Resendre-Br/55)	1		1			
	244367	Foot-and-mouth disease virus C-S8c1	12	17	12	17		
	13067	Forficula		2		2		
	31621	Four Corners hantavirus	2	1	2	5	4	
	10261	Fowlpox virus		3		7	4	
	10263	Fowlpox virus isolate HP-438/Munich		1		1		
	263	Francisella tularensis		5		5		
X	351581	Francisella tularensis subsp. holarctica FSC200				1	1	
X	458234	Francisella tularensis subsp. holarctica FTNF002-00				1	1	
	376619	Francisella tularensis subsp. holarctica LVS		68		75	7	
	393011	Francisella tularensis subsp. holarctica OSU18		2		5	3	
	119856	Francisella tularensis subsp. tularensis		5		5		
	430557	Francisella tularensis subsp. tularensis FSC033		26		27	1	
	393115	Francisella tularensis subsp. tularensis FSC198		3		50	47	
	177416	Francisella tularensis subsp. tularensis SCHU S4		90		90		
	418136	Francisella tularensis subsp. tularensis WY96-3418		1		1		
X	298653	Frankia sp. EAN1pec				2	2	
	11795	Friend murine leukemia virus	6	21	6	22	1	
	11797	Friend murine leukemia virus (ISOLATE FB29)		1		1		
X	209882	Fusobacterium nucleatum subsp. vincentii ATCC 49256				1	1	
	8053	Gadus callarias		10		10		
X	7137	Galleria mellonella			1		1	
	9031	Gallus gallus	193	132	228	152	35	20
	11824	Gardner-Arnstein feline leukemia oncovirus B	13		13			
	54290	GB virus C	14		16		2	
	471223	Geobacillus sp. WCH70		2		2		
	1422	Geobacillus stearothermophilus	1		1	1	1	
	5741	Giardia intestinalis		8		8		
X	598745	Giardia intestinalis ATCC 50581				4	4	
	184922	Giardia lamblia ATCC 50803		194		239		45
X	28448	Gluconacetobacter xylinus				1	1	
	3847	Glycine max	71	1	71	1		
X	9593	Gorilla gorilla				1	1	
	9595	Gorilla gorilla gorilla		4		4		
	55951	Grapevine leafroll-associated virus 3	1		1			
	35288	Grapevine virus A	19		19			
	45219	Guanarito virus		835		838		3
	114727	H1N1 subtype	1		3	11	2	11
	10001225	H1N1 subtype Influenza A/Oklahoma/7485/01		5		5		

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	10001315	H1N1 swine influenza virus (A/swine/Korea/S10/2004(H1N1))		3		3		
	10001316	H1N1 swine influenza virus (A/swine/Korea/S175/2004(H1N1))		1		1		
	119210	H3N2 subtype	1	2	1	8		6
	10001157	H3N2 subtype Influenza A virus (A/Hong Kong(H3N2))		15		15		
	102793	H5N1 subtype	2	16	2	16		
	10001317	H9N2 subtype Influenza A virus (A/swine/Korea/S81/2004(H9N2))		1		1		
	10001318	H9N2 subtype Influenza A virus (A/swine/Korea/S83/2004(H9N2))		2		2		
	730	Haemophilus ducreyi		1	2	1	2	
	727	Haemophilus influenzae	17	54	19	61	2	7
X	374927	Haemophilus influenzae 22.1-21					1	1
X	375177	Haemophilus influenzae 3655					1	1
X	521004	Haemophilus influenzae 6P18H1					1	1
	10001055	Haemophilus influenzae 6U	2		2			
	521005	Haemophilus influenzae 7P49H1		1		2		1
	281310	Haemophilus influenzae 86-028NP	1		1			
	10001056	Haemophilus influenzae ATCC 9795	2		2			
	10001053	Haemophilus influenzae MinnA	9		9			
	10000833	Haemophilus influenzae NTHi 1128	6		6			
	10001042	Haemophilus influenzae NTHi 1479	12	4	12	4		
	10000807	Haemophilus influenzae NTHi UC19	2	2	2	2		
	374928	Haemophilus influenzae PittAA		1		2		1
X	374931	Haemophilus influenzae PittGG					1	1
	374932	Haemophilus influenzae PittHH	1		1			
X	262727	Haemophilus influenzae R2846					3	3
X	262728	Haemophilus influenzae R2866					1	1
	375432	Haemophilus influenzae R3021		1		3		2
X	71421	Haemophilus influenzae Rd KW20					1	1
	10000860	Haemophilus influenzae Serotype B	14	9	14	9		
	10001155	Haemophilus influenzae strain 1479	2		2			
X	10001523	Haemophilus influenzae strain I-69 Rd-/b+			2		2	
	10001149	Haemophilus influenzae Subtype 1H	23		23			
	10000861	Haemophilus influenzae Variant d1	18		18			
	10690	Haemophilus phage HP1		1		1		
	205914	Haemophilus somnus 129PT		1		1		
	6454	Haliotis rufescens	1		1			
	10626	Hamster polyomavirus	11		11			
	11599	Hantaan virus	2	41	2	42		1
	11602	Hantaan virus 76-118	11	8	11	18		10
	370830	Hantaan virus Q32		4		4		
	458678	Hantaanvirus CGRn93P8		2		2		
X	11598	Hantavirus					3	3
	74942	Hantavirus CRF355		1		1		
	42358	Hantavirus Monongahela-3		5		5		
	37741	Hantavirus sp.		2		2		
	13557	Hapalemur griseus		1		1		
	4232	Helianthus annuus	18		18			
	32025	Helicobacter hepaticus		1		1		
	235279	Helicobacter hepaticus ATCC 51449	2		2			
	210	Helicobacter pylori	26	4	31	6	5	2
	85962	Helicobacter pylori 26695	13	1	13	3		2
	10000718	Helicobacter pylori J223	2		2			
X	85963	Helicobacter pylori J99					27	27
	570508	Helicobacter pylori P12		2		2		

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	102617	Helicobacter pylori SS1	1		1			
	10000720	Helicobacter pylori UA948	2		2			
	10000721	Helicobacter pylori UA955	1		1			
X	6536	Helix pomatia			1		1	
	63330	Hendra virus	7		7			
	11102	Hepacivirus		3		3		
	12092	Hepatitis A virus	23	9	23	9		
	10407	Hepatitis B virus	210	457	217	458	7	1
	12513	Hepatitis B virus 2		1		1		
	31512	Hepatitis B virus adr/mutant		1		1		
	10409	Hepatitis B virus adr4	1	11	1	11		
	10410	Hepatitis B virus adw/991	2	2	2	2		
	10412	Hepatitis B virus adw/Indonesia/PIDW420		2		2		
	10415	Hepatitis B virus adw/Okinawa/PODW282		4		4		
	45410	Hepatitis B virus adw4/Brazil/isolate w4b		6		6		
	10411	Hepatitis B virus alpha1		15		15		
	489469	Hepatitis B virus ayw/China/Tibet127/2002	1		1			
	391647	Hepatitis B virus ayw2	6		6			
	391646	Hepatitis B virus ayw3	1		1			
	391650	Hepatitis B virus ayw4	1		1			
	10000433	Hepatitis B virus genotype B	1		1			
	10000434	Hepatitis B virus genotype C	3		3			
	10000435	Hepatitis B virus genotype D	1		1			
	10414	Hepatitis B virus LSH/chimpanzee		2		2		
	10001156	Hepatitis B virus subtype AD	3	3	3	3		
	106820	Hepatitis B virus subtype adr	16	25	23	25	7	
	106821	Hepatitis B virus subtype adw	17	60	17	60		
	10408	Hepatitis B virus subtype adw2	71	59	73	60	2	1
	10419	Hepatitis B virus subtype adyw		35		35		
	10000436	Hepatitis B virus subtype AY	7		7			
	10000437	Hepatitis B virus subtype AYR		1		1		
	10418	Hepatitis B virus subtype ayw	57	165	57	165		
	11103	Hepatitis C virus	649	847	656	856	7	9
	11104	Hepatitis C virus (isolate 1)	37	216	37	216		
	356391	Hepatitis C virus (isolate 6a33)		5		5		
	356413	Hepatitis C virus (isolate BEBE1)		10		10		
	11105	Hepatitis C virus (isolate BK)	3	43	3	43		
	333284	Hepatitis C virus (isolate Con1)	3	5	3	5		
	356419	Hepatitis C virus (isolate EUH1480)		4		4		
	329389	Hepatitis C virus (isolate Glasgow)	2	1	2	1		
	11108	Hepatitis C virus (isolate H)	128	81	128	81		
	63746	Hepatitis C virus (isolate H77)	32	174	34	174	2	
	356410	Hepatitis C virus (isolate HC-G9)	2	1	2	1		
	11110	Hepatitis C virus (isolate HCT18)	1		1			
	356416	Hepatitis C virus (isolate HCV-K3a/650)		14		14		
	31644	Hepatitis C virus (isolate HCV-KF)	1	1	1	1		
	356386	Hepatitis C virus (isolate India)		1		1		
	11116	Hepatitis C virus (isolate Japanese)	6	26	6	26		
	356417	Hepatitis C virus (isolate JK049)		5		5		
	356415	Hepatitis C virus (isolate NZL1)	1	1	1	1		
	31645	Hepatitis C virus (isolate Taiwan)	3	19	3	19		
	356421	Hepatitis C virus (isolate Th580)		4		4		
	357355	Hepatitis C virus (isolate Tr Kj)		3		3		

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	356424	Hepatitis C virus (isolate VN004)		1		1		
	41856	Hepatitis C virus genotype 1		2	42	2	42	
	40271	Hepatitis C virus genotype 2		41	5	41	5	
	356114	Hepatitis C virus genotype 3		114	10	114	10	
	33745	Hepatitis C virus genotype 4			2		2	
	33746	Hepatitis C virus genotype 5			1		1	
	42182	Hepatitis C virus genotype 6			6		6	
	421877	Hepatitis C virus isolate HC-J1		3	26	3	26	
	11113	Hepatitis C virus isolate HC-J6		3	2	3	2	
	11115	Hepatitis C virus isolate HC-J8		2	1	2	1	
	356411	Hepatitis C virus JFH-1		2	1	3	1	1
	31646	Hepatitis C virus subtype 1a		143	312	147	312	4
	10000453	Hepatitis C virus subtype 1a (isolate Gla)		5		5		
	10000455	Hepatitis C virus subtype 1a 1/910			17		17	
	10000457	Hepatitis C virus subtype 1a Chiron Corp.		1	1	1	1	
	31647	Hepatitis C virus subtype 1b		525	157	532	167	7 10
	10000456	Hepatitis C virus subtype 1b AD78		71		71		
	10000968	Hepatitis C virus subtype 1b isolate BE-11		3		3		
	10000460	Hepatitis C virus subtype 1b JK1			13		13	
	31649	Hepatitis C virus subtype 2a		78	21	78	21	
	31650	Hepatitis C virus subtype 2b		100	3	100	3	
	356426	Hepatitis C virus subtype 3a		5	3	5	3	
	42792	Hepatitis C virus subtype 3g		1		1		
	31653	Hepatitis C virus subtype 4a		1	1	1	1	
	31654	Hepatitis C virus subtype 5a		1	1	1	1	
	12475	Hepatitis delta virus		48	7	48	7	
	10000522	Hepatitis delta virus (isolate TW2667)			5		5	
	12461	Hepatitis E virus		131	27	133	27	2
	31767	Hepatitis E virus (strain Burma)		160		160		
	31768	Hepatitis E virus (strain Mexico)		31		31		
	10000519	Hepatitis E virus China Xinjiang		1		1		
	10000520	Hepatitis E virus SAR-55		1	6	1	6	
	39113	Hepatitis GB virus B			4		4	
	28300	Heron hepatitis B virus			1		1	
	10299	Herpes simplex virus (type 1 / strain 17)		44	30	44	30	
	10301	Herpes simplex virus (type 1 / strain Angelotti)			1		1	
	10304	Herpes simplex virus (type 1 / strain F)		17	5	17	5	
	10303	Herpes simplex virus (type 1 / strain HFEM)		2		2		
	10308	Herpes simplex virus (type 1 / strain Patton)		3		3		
	10309	Herpes simplex virus (type 1 / strain SC16)		6	1	6	2	1
	126283	Herpes simplex virus unknown type			1		1	
	10292	Herpesviridae			1		1	
	49011	Hesperocyparis arizonica		1		2		1
	3981	Hevea brasiliensis		164	43	164	43	
X	388799	HIV-1 group O				1		1
	11685	HIV-1 M:B_ARV2/SF2			43		43	
	11706	HIV-1 M:B_HXB2R			12		12	
	11696	HIV-1 M:B_MN		1	21	1	21	
X	327105	HIV-1 O_ANT70				1		1
	11583	HoJo virus			1		1	
	29679	Holcus lanatus		14		14		
	9606	Homo sapiens		1397	2872	4108	5110	2711 2238
X	397342	Horsepox virus					5	5

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	343462	Human adenovirus 11p	7		7			
	28282	Human adenovirus 12	9	2	10	2	1	
X	46921	Human adenovirus 13			1		1	
	28276	Human adenovirus 15	1		1			
	10515	Human adenovirus 2	28	5	28	5		
	45659	Human adenovirus 3	17	1	20	1	3	
	28284	Human adenovirus 40	2	1	2	1		
X	46941	Human adenovirus 46			1		1	
	28285	Human adenovirus 5	21	116	22	154	1	38
	31545	Human adenovirus 8	1		1			
	10001392	Human adenovirus B strain Harbin04B	5		5			
	129951	Human adenovirus C	1	3	1	3		
	130309	Human adenovirus F	1		1			
	11137	Human coronavirus 229E			2		2	
	31631	Human coronavirus OC43			1		1	
	12067	Human coxsackievirus A9	7		30		23	
	12071	Human coxsackievirus B1			12			
	12072	Human coxsackievirus B3	7	11	7	14		3
	10001213	Human coxsackievirus B3 (strain RK)	1		1			
	12073	Human coxsackievirus B4	5	49	14	50	9	1
	11827	Human endogenous retrovirus	2	1	2	1		
X	64382	Human Endogenous Retrovirus IDDMK1.2-22			3		3	
X	45617	Human endogenous retrovirus K			12		12	
	39054	Human enterovirus 71	2		4		2	
X	138948	Human enterovirus A			1	1	1	1
	138950	Human enterovirus C	6	3	6	3		
	208726	Human hepatitis A virus			4			
	12098	Human hepatitis A virus Hu/Australia/HM175/1976	86	62	86	62		
	10298	Human herpesvirus 1	139	69	140	78	1	9
	10000394	Human herpesvirus 1 103/65	1		1			
	10000396	Human herpesvirus 1 McIntyre			1		1	
	10000398	Human herpesvirus 1 NS	1		1			
	10306	Human herpesvirus 1 strain KOS	17	4	17	4		
	10310	Human herpesvirus 2	88	109	90	139	2	30
	10312	Human herpesvirus 2 strain 186			1		1	
	10313	Human herpesvirus 2 strain 333			4		4	
	10315	Human herpesvirus 2 strain HG52			33		33	
	10335	Human herpesvirus 3	4	78	6	79	2	1
	10000406	Human herpesvirus 3 H-551	26	15	26	15		
	10338	Human herpesvirus 3 strain Dumas			28		28	
	10376	Human herpesvirus 4	80	321	117	350	37	29
	10377	Human herpesvirus 4 (strain B95-8)	62	234	65	242	3	8
	10000420	Human herpesvirus 4 BL74			1		1	
	10000421	Human herpesvirus 4 CKL			10		10	
	10000424	Human herpesvirus 4 GD1			1		1	
	36352	Human herpesvirus 4 type 1	3	11	4	12	1	1
	12509	Human herpesvirus 4 type 2	1	1	2	1	1	
	10000427	Human herpesvirus 4 type A			5		5	
	10359	Human herpesvirus 5	76	341	80	465	4	124
	10360	Human herpesvirus 5 strain AD169	27	302	27	316		14
	10363	Human herpesvirus 5 strain Towne	4	23	4	24		1
	10368	Human herpesvirus 6	2	1	2	2		1
	10369	Human herpesvirus 6 (strain GS)	2		2			

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	10370	Human herpesvirus 6 (strain Uganda-1102)	1	2	1	2		
	36351	Human herpesvirus 6 strain Z29	1		1			
	32604	Human herpesvirus 6B	1	1	1	2		1
	10000535	Human herpesvirus 6B HST	1		1			
	10372	Human herpesvirus 7	2		2	1		1
	57278	Human herpesvirus 7 strain JI		1		1		
	37296	Human herpesvirus 8	26	127	26	127		
	12721	Human immunodeficiency virus	1	14	1	16		2
	11676	Human immunodeficiency virus 1	7	191	8	276	1	85
	10000500	Human immunodeficiency virus 1 IIIB		2		2		
	11709	Human immunodeficiency virus 2		6		6		
X	11682	Human immunodeficiency virus type 1 (BH5 ISOLATE)				1		1
	11686	Human immunodeficiency virus type 1 (BRU ISOLATE)	1	3	1	3		
	11687	Human immunodeficiency virus type 1 (CDC-451 ISOLATE)		3		3		
	11679	Human immunodeficiency virus type 1 (CLONE 12)		6		6		
	11689	Human immunodeficiency virus type 1 (ELI ISOLATE)		4		6		2
	362651	Human immunodeficiency virus type 1 (isolate YU2)		21		21		
	11694	Human immunodeficiency virus type 1 (JH3 ISOLATE)		3		3		
	11688	Human immunodeficiency virus type 1 (JRCSF ISOLATE)		35		35		
	11697	Human immunodeficiency virus type 1 (MAL ISOLATE)		6		7		1
	11704	Human immunodeficiency virus type 1 (MFA ISOLATE)		2		2		
	11695	Human immunodeficiency virus type 1 (NDK ISOLATE)		8		8		
	11698	Human immunodeficiency virus type 1 (NEW YORK-5 ISOLATE)		1		1		
	11699	Human immunodeficiency virus type 1 (OYI ISOLATE)		3		3		
	11701	Human immunodeficiency virus type 1 (RF/HAT ISOLATE)	1	19	1	19		
	11691	Human immunodeficiency virus type 1 (SF162 ISOLATE)		1		1		
	11690	Human immunodeficiency virus type 1 (SF33 ISOLATE)		2		2		
	11703	Human immunodeficiency virus type 1 (STRAIN UGANDAN / ISOLATE U455)			14		14	
	31678	Human immunodeficiency virus type 1 (WMJ1 isolate)		15		15		
	11705	Human immunodeficiency virus type 1 (WMJ2 ISOLATE)		1		1		
	11683	Human immunodeficiency virus type 1 (Z2/CDC-Z34 ISOLATE)		1		1		
	11678	Human immunodeficiency virus type 1 BH10		16		19		3
	11714	Human immunodeficiency virus type 2 (ISOLATE BEN)		7		7		
	11715	Human immunodeficiency virus type 2 (ISOLATE CAM2)		1		1		
	11713	Human immunodeficiency virus type 2 (ISOLATE D194)		1		1		
	11716	Human immunodeficiency virus type 2 (ISOLATE D205,7)		1		1		
	11717	Human immunodeficiency virus type 2 (ISOLATE GHANA-1)		5		5		
	73484	Human immunodeficiency virus type 2 (isolate KR)		2		2		
	11718	Human immunodeficiency virus type 2 (ISOLATE SBLISY)		2		2		
	162145	Human metapneumovirus	7	22	7	22		
	10566	Human papillomavirus	4	2	4	2		
	334203	Human papillomavirus 1		41	1	42	1	1
	337042	Human papillomavirus 18	9	4	9	4		
	337043	Human papillomavirus 2		6		6		
	10580	Human papillomavirus type 11	4	32	4	33		1
	10573	Human papillomavirus type 13	1		1	1		1
	333760	Human papillomavirus type 16		232	306	236	324	4 18
	333761	Human papillomavirus type 18	18	75	18	76		1
	10583	Human papillomavirus type 1a	7	7	7	7		
	333751	Human papillomavirus type 2		2		2		
	333762	Human papillomavirus type 26		1		1		
	37112	Human papillomavirus type 29		1		1		

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	10614	Human papillomavirus type 3		3		3		
	10585	Human papillomavirus type 31		2	6	7	6	5
	333763	Human papillomavirus type 32		1		1	1	
	10586	Human papillomavirus type 33		5	8	5	8	
	10587	Human papillomavirus type 35			1		1	
	10588	Human papillomavirus type 39			1		1	
	10617	Human papillomavirus type 4			4		4	
	10615	Human papillomavirus type 40		1		1	1	1
	10592	Human papillomavirus type 44			1		1	
	10593	Human papillomavirus type 45			5		5	
	10595	Human papillomavirus type 51			1		1	
	10618	Human papillomavirus type 52			4		4	
	333765	Human papillomavirus type 53			1		1	
	10596	Human papillomavirus type 56			3		3	
	10598	Human papillomavirus type 58			4		32	28
	37115	Human papillomavirus type 59			1		1	
	31552	Human papillomavirus type 6		9	3	9	3	
	45240	Human papillomavirus type 68			1		1	
	10600	Human papillomavirus type 6b		11	24	11	24	
	10620	Human papillomavirus type 7		1	1	1	3	2
	51033	Human papillomavirus type 73			1		1	
	36412	Human parainfluenza 1 virus (strains A1426 / 86-315 / 62M-753)			1		1	
	11217	Human parainfluenza 3 virus (strain NIH 47885)		8		8		
	11212	Human parainfluenza virus 2		10		10		
	11214	Human parainfluenza virus 2 (strain Toshiba)		3		3		
	11216	Human parainfluenza virus 3		2		2		
	12063	Human parechovirus 1		5		5		
	10798	Human parvovirus B19		90	44	93	44	3
	10000438	Human parvovirus B19 genotype 1		1		1		
	12080	Human poliovirus 1		6	1	6	1	
	12081	Human poliovirus 1 Mahoney		47	20	47	20	
	12082	Human poliovirus 1 strain Sabin		7		7		
	10001028	Human poliovirus 2 (strain MEF-1)		1		1		
	10001040	Human poliovirus 2 (strain Sabin)		2	7	2	7	
	12086	Human poliovirus 3		3	2	3	2	
	10001086	Human poliovirus 3 (strain Sabin)		36	1	36	1	
	11250	Human respiratory syncytial virus		15	31	15	32	1
	11255	Human respiratory syncytial virus (strain RSB6190)		6		6		
	11256	Human respiratory syncytial virus (strain RSB6256)		10		10		
	11252	Human respiratory syncytial virus (strain RSB642)		1		1		
	11251	Human respiratory syncytial virus (subgroup B / strain 18537)		2	1	2	1	
	208893	Human respiratory syncytial virus A		1	3	1	3	
	10000960	Human respiratory syncytial virus A Mon/3/88		6		6		
	11260	Human respiratory syncytial virus A strain Long		120	7	120	16	9
X	10001456	Human respiratory syncytial virus A strain RGH				1		1
	11259	Human respiratory syncytial virus A2		48	107	50	108	2
X	79692	Human respiratory syncytial virus B1				1		1
	410078	Human respiratory syncytial virus S2		1		1		
	12131	Human rhinovirus 14		12		12		
	12134	Human rhinovirus 1A			19		19	
	12130	Human rhinovirus 2		11		11		
	10000987	Human rhinovirus 2 Vienna		5		5		
	147711	Human rhinovirus A		1		1		

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	10950	Human rotavirus (SEROTYPE 2 / STRAIN DS1)	1		1			
	31569	Human rotavirus (serotype 2 / strain RV-5)	1		1			
	10941	Human rotavirus A	13	2	13	2		
	10960	Human rotavirus G4 strain St. Thomas 3	6	1	6	1		
	374507	Human rotavirus G9 isolate F45	7		7			
	10001215	Human rotavirus G9 WI61	1		1			
	408599	Human rotavirus G9P[8]	1		1			
X	94432	Human rotavirus MP409			1	1	1	1
	10952	Human rotavirus strain KU	9		9			
	10957	Human rotavirus strain P	1		1	4		4
	10958	Human rotavirus strain RRV	1		1			
	10962	Human rotavirus strain WA	10	7	10	7		
	11927	Human T-cell lymphotropic virus type 1 (Caribbean isolate)		5		5		
	10001004	Human T-cell lymphotropic virus type 1 (Caribbean isolate) (Strain HS35)	1		1			
	11928	Human T-cell lymphotropic virus type 1 (isolate MT-2)	1		1			
	11926	Human T-cell lymphotropic virus type 1 (strain ATK)	30	118	30	118		
	39015	Human T-cell lymphotropic virus type 1 (african isolate)		2		2		
	11908	Human T-lymphotropic virus 1	214	162	223	162	9	
	11909	Human T-lymphotropic virus 2	63	2	63	2		
	77644	IncQ plasmid pIE1120		1		1		
	11120	Infectious bronchitis virus	5		5			
	10000825	Infectious bronchitis virus Avian strain D207	11		11			
X	10001431	Infectious bronchitis virus CK/CH/LDL/97I			1		1	
X	633557	Infectious bronchitis virus CK/CH/LHLJ/04V			1		1	
	10995	Infectious bursal disease virus	11		11			
	10997	Infectious bursal disease virus 002-73/AUS	1		1			
	11290	Infectious hematopoietic necrosis virus	8		8			
X	11002	Infectious pancreatic necrosis virus			1		1	
	11320	Influenza A virus	29	248	39	262	10	14
X	229051	Influenza A virus (A/81/HO)				3		3
	387139	Influenza A virus (A/Aichi/2/1968(H3N2))	5	10	5	10		
X	203129	Influenza A virus (A/Aichi/2/1994(H3N2))				1		1
	553345	Influenza A virus (A/Aichi/72/2007(H3N2))	1		1			
	385576	Influenza A virus (A/Alaska/6/1977(H3N2))		1		1		
	383602	Influenza A virus (A/Anas acuta/Primorje/695/1976(H2N3))		10		10		
	384498	Influenza A virus (A/Ann Arbor/6/1960(H2N2))	1	180	1	180		
	62446	Influenza A virus (A/Argentina/3779/94(H3N2))		1		1		
	385630	Influenza A virus (A/Bangkok/1/1979(H3N2))	1	4	1	4		
X	223931	Influenza A virus (A/Bangkok/10/1983(H1N1))				1		1
X	336238	Influenza A virus (A/Bar-headed Goose/Qinghai/61/05(H5N1))				8		8
	384500	Influenza A virus (A/Beijing/11/1956(H1N1))		22		22		
X	518922	Influenza A virus (A/Beijing/262/1995(H1N1))			1		1	
	380950	Influenza A virus (A/Beijing/32/1992(H3N2))		54		54		
X	648856	Influenza A virus (A/Beijing/501/2009(H1N1))			1		1	
	304883	Influenza A virus (A/Belfast/53582/2004(H3N?))		2		2		
	282811	Influenza A virus (A/Bilthoven/4791/81(H3N2))		3		3		
	88776	Influenza A virus (A/Brevig Mission/1/1918(H1N1))	1		1			
	385587	Influenza A virus (A/budgerigar/Hokkaido/1/1977(H4N6))		4		4		
X	641501	Influenza A virus (A/California/04/2009(H1N1))				32		32
	352963	Influenza A virus (A/California/7/2004(H3N2))		1		1		
	387191	Influenza A virus (A/camel/Mongolia/1982(H1N1))		8		8		
	205946	Influenza A virus (A/Chicken/Beijing/1/95(H9N2))		1		1		

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	404137	Influenza A virus (A/chicken/Beijing/nl/2002(H9N2))		1		1		
	36418	Influenza A virus (A/chicken/Brescia/1902(H7N7))		1		1		
	342137	Influenza A virus (A/chicken/CA/375TR/2002(H6N2))		1		1		
X	196462	Influenza A virus (A/chicken/California/1002a/00(H6N2))				1		1
	11384	Influenza A virus (A/chicken/FPV/Weybridge(H7N7))		1		1		
X	329593	Influenza A virus (A/chicken/Fujian/25/00(H9N2))				5		5
	197579	Influenza A virus (A/Chicken/Guangdong/11/97(H9N2))		1		1		
X	407418	Influenza A virus (A/chicken/Guiyang/2147/2005(H5N1))				1		1
	435359	Influenza A virus (A/chicken/Hebei/706/2005(H5N1))		1		1		
	357205	Influenza A virus (A/chicken/Hong Kong/3123.1/2002(H5N1))		1		1		
	97348	Influenza A virus (A/Chicken/Hong Kong/G23/97(H9N2))		1		1		
	404150	Influenza A virus (A/chicken/Hubei/wk/1997(H5N1))		2		2		
	404155	Influenza A virus (A/chicken/Hunan/fg/2004(H5N1))		2		2		
	447133	Influenza A virus (A/chicken/Japan/1925(H7N7))	1		1			
	404163	Influenza A virus (A/chicken/Jilin/hg/2002(H5N1))		1		1		
X	300750	Influenza A virus (A/chicken/Korea/S1/2003(H9N2))				1		1
	342221	Influenza A virus (A/chicken/Korea/S4/2003(H9N2))		1		1		
	298713	Influenza A virus (A/chicken/Korea/SNU0091/00(H9N2))		1		1		
	298716	Influenza A virus (A/chicken/Korea/SNU1035C/00(H9N2))		1		1		
X	680739	Influenza A virus (A/chicken/Mexico/37821-771/1996(H5N2))				1		1
	270490	Influenza A virus (A/chicken/Nakorn-Patom/Thailand/CU-K3/2004(H5N1))		1		1		
	215862	Influenza A virus (A/Chicken/Nanchang/2-220/2001(H3N6))		1		1		
	385617	Influenza A virus (A/chicken/Pennsylvania/1370/1983(H5N2))	1		1			
	392810	Influenza A virus (A/chicken/Rostock/8/1934(H7N1))	2	4	2	4		
X	402527	Influenza A virus (A/chicken/Scotland/1959(H5N1))				1		1
	197585	Influenza A virus (A/chicken/Shandong/6/96(H9N2))	1		1			
	264510	Influenza A virus (A/chicken/Thailand/LV1NF/2004(H5N1))		1		1		
	393548	Influenza A virus (A/chicken/Tula/Russia/Oct-5/2005(H5N1))		1		1		
	299730	Influenza A virus (A/chicken/Viet Nam/AG-010/2004(H5N1))		1		1		
	380836	Influenza A virus (A/chicken/Vietnam/5/2003(H5N1))		1		1		
	365129	Influenza A virus (A/chicken/Vietnam/DT171/2004(H5N1))		1		1		
	380985	Influenza A virus (A/Chile/1/1983(H1N1))		1		1		
	62541	Influenza A virus (A/Christchurch/2/1988(H3N2))		1		1		
X	514193	Influenza A virus (A/CHU/2-524/2005(H3N2))				2		2
	284168	Influenza A virus (A/Ck/HK/2133.1/2003(H5N1))		1		1		
	354235	Influenza A virus (A/Ck/HK/CSW161/2003(H9N2))		1		1		
	284195	Influenza A virus (A/Ck/YN/115/2004(H5N1))		1		1		
	107493	Influenza A virus (A/Cordoba/3278/96(H3N2))		1		1		
X	320306	Influenza A virus (A/crested eagle/Belgium/01/2004(H5N1))				2		2
	291592	Influenza A virus (A/crow/Kyoto/53/2004(H5N1))	1		1			
	385583	Influenza A virus (A/Denver/1957(H1N1))		1		2		1
X	88297	Influenza A virus (A/duck/Bavaria/1/1977 (H1N1))				2		2
	383550	Influenza A virus (A/duck/England/1/1956(H11N6))		1		1		
	274829	Influenza A virus (A/duck/Fujian/17/2001(H5N1))		1		1		
	365078	Influenza A virus (A/duck/Guangxi/1586/2004(H5N1))		1		1		
X	365080	Influenza A virus (A/duck/Guangxi/1793/2004(H5N1))				1		1
	395841	Influenza A virus (A/duck/Hokkaido/1130/01(H1N1))		1		1		
	249372	Influenza A virus (A/duck/Hokkaido/13/00(H9N2))	2		2			
	387207	Influenza A virus (A/duck/Hokkaido/8/1980(H3N8))		6		6		
X	183783	Influenza A virus (A/duck/Hong Kong/3096/99(H6N2))				1		1
	176672	Influenza A virus (A/Duck/Hong Kong/ww461/2000(H5N1))		1		1		
	384946	Influenza A virus (A/duck/Hungary/1/1970(H6N2))		1		1		

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	370813	Influenza A virus (A/duck/IT/701/2005(H10N7))		1		1		
	311169	Influenza A virus (A/duck/Korea/S10/03(H3N2))		1		1		
	441962	Influenza A virus (A/duck/Mongolia/47/2001(H7N1))		3		3		
	353253	Influenza A virus (A/duck/Novosibirsk/56/2005(H5N1))		2	6	2	6	
	210656	Influenza A virus (A/Duck/Shantou/1881/00(H9N2))		1		1		
	210659	Influenza A virus (A/Duck/Shantou/2088/01(H9N2))		1		1		
	210663	Influenza A virus (A/Duck/Shantou/2134/00(H9N2))		1		1		
	210655	Influenza A virus (A/Duck/Shantou/2144/00(H9N2))		1		1		
	385580	Influenza A virus (A/duck/Ukraine/1/1963(H3N8))	3		3			
	380842	Influenza A virus (A/duck/Vietnam/15/2003(H5N1))		1		1		
	11375	Influenza A virus (A/Dunedin/4/73(H3N2))	2		2			
	279797	Influenza A virus (A/egret/Hong Kong/757.3/2002(H5N1))		3		3		
	192549	Influenza A virus (A/Egypt/96/2002(H1N2))		1		1		
	380210	Influenza A virus (A/England/333/1980(H1N1))	1	1	1	1		
	198059	Influenza A virus (A/England/627/01(H1N2))		2		2		
	573797	Influenza A virus (A/England/654/2007(H1N1))		1		1		
	387147	Influenza A virus (A/England/878/1969(H3N2))	1	2	1	2		
	137578	Influenza A virus (A/England/939/69 x A/PR/8/34)		2		2		
X	135661	Influenza A virus (A/environment/Hong Kong/437-10/99 (H5N1))				1		1
X	680789	Influenza A virus (A/environment/Viet Nam/1203/2004(H5N1))			1		1	
X	475493	Influenza A virus (A/equine/California/8560/2002(H3N8))				2		2
	336794	Influenza A virus (A/equine/Massachusetts/213/2003(H3N8))		1		1		
	387223	Influenza A virus (A/equine/Miami/1963(H3N8))	5		5			
	336800	Influenza A virus (A/equine/New York/1/1999(H3N8))		3		3		
X	225089	Influenza A virus (A/equine/NewMarket/D64/79(H3N8))				1		1
	207401	Influenza A virus (A/finch/Canada/NS1301/2001(H3N8))		2		2		
	367120	Influenza A virus (A/Fort Monmouth/1/1947(H1N1))	1		1			
	380282	Influenza A virus (A/Fort Monmouth/1/1947-mouse adapted(H1N1))		6		24		18
	260806	Influenza A virus (A/fowl/Dutch/27(H7N7))	1		1			
	107558	Influenza A virus (A/France/75/97(H3N2))		1		1		
X	354228	Influenza A virus (A/Gf/HK/SSP607/2003(H9N2))				1		1
X	93838	Influenza A virus (A/Goose/Guangdong/1/96(H5N1))				3		3
	165512	Influenza A virus (A/goose/Guangdong/3/1997(H5N1))	3		3	1		1
	182781	Influenza A virus (A/Goose/Hong Kong/385.3/2000(H5N1))		1		1		
X	680799	Influenza A virus (A/goose/Hong Kong/437-6/1999(H5N1))				1		1
X	680803	Influenza A virus (A/goose/Hong Kong/739.2/2002(H5N1))				1		1
	380330	Influenza A virus (A/goose/Hong Kong/8/1976(H1N1))		2		2		
X	402464	Influenza A virus (A/gray teal/Australia/2/1979(H4N4))				1		1
X	387239	Influenza A virus (A/gull/Delaware/475/1986(H2N2))				1		1
	384499	Influenza A virus (A/gull/Maryland/704/1977(H13N6))		1		1		
	387156	Influenza A virus (A/Harbin/1/1988(H1N2))		4		11		7
X	303316	Influenza A virus (A/Hatay/2004/(H5N1))				3		3
	462695	Influenza A virus (A/Hiroshima/52/2005(H3))	2		2			
X	223947	Influenza A virus (A/Hokkaido/20/89(H3N2))				2		2
	506350	Influenza A virus (A/Hong Kong/1/1968(H3N2))		2	1	2	1	
	130760	Influenza A virus (A/Hong Kong/1073/99(H9N2))	1		1			
X	130761	Influenza A virus (A/Hong Kong/1074/99(H9N2))				1		1
	164326	Influenza A virus (A/Hong Kong/1131/1998(H1N1))		1		1		
	130763	Influenza A virus (A/Hong Kong/156/97(H5N1))	1	2	1	2		
X	231851	Influenza A virus (A/Hong Kong/1774/99(H3N2))				1		1
	317652	Influenza A virus (A/Hong Kong/2/68(H3N2))		3		3		
	432070	Influenza A virus (A/Hong Kong/213/2003(H5N1))		4		4		

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X	155218	Influenza A virus (A/Hong Kong/482/97(H5N1))				1		1
	88104	Influenza A virus (A/Hong Kong/483/1997(H5N1))	1		1			
	680692	Influenza A virus (A/Hong Kong/485/1997(H5N1))		1		1		
	155222	Influenza A virus (A/Hong Kong/488/97(H5N1))		1		1		
	155225	Influenza A virus (A/Hong Kong/507/97(H5N1))		5		5		
	155226	Influenza A virus (A/Hong Kong/514/97(H5N1))		2		2		
	155227	Influenza A virus (A/Hong Kong/516/97(H5N1))		2		2		
X	88103	Influenza A virus (A/Hong Kong/532/1997(H5N1))				1		1
X	499286	Influenza A virus (A/Hong Kong/CUHK13527/2003(H3N2))				1		1
	400788	Influenza A virus (A/Indonesia/5/2005(H5N1))	1	1	1	1		
	400828	Influenza A virus (A/Indonesia/CDC669/2006(H5N1))	1		1			
	400830	Influenza A virus (A/Indonesia/CDC699/2006(H5N1))	2		2			
	402194	Influenza A virus (A/Istanbul/CTF/10/2004(H3N2))		1		1		
	387161	Influenza A virus (A/Japan/305/1957(H2N2))	2	27	2	28		1
	307819	Influenza A virus (A/Jiangsu/38/2004(H3N2))		1		1		
	203126	Influenza A virus (A/Kamata/14/91(H3N2))	1		1			
X	154540	Influenza A virus (A/Kayano/57(H2N2))				1		1
	384495	Influenza A virus (A/Kiev/59/1979(H1N1))		1		1		
	327310	Influenza A virus (A/Kinmen/645/04(H3N2))		1		1		
X	225088	Influenza A virus (A/Kitakyushu/159/93(H3N2))				1		1
	488241	Influenza A virus (A/Korea/426/1968(H2N2))		1		1		
	436004	Influenza A virus (A/kunke/1/71(H3N2))		1		1		
	393557	Influenza A virus (A/Leningrad/1954/1(H1N1))		2		4		2
	393697	Influenza A virus (A/Liaoning/25/2000(H1N1))		1		1		
	62559	Influenza A virus (A/Los Angeles/1987(H3N2))	5		5			
	95888	Influenza A virus (A/mallard duck/PA/10218/84(H5N2))	8		8			
	282124	Influenza A virus (A/mallard/Alberta/127/00(H3N8))		1		1		
	352520	Influenza A virus (A/mallard/Alberta/35/1976(H1N1))		2		2		
	185794	Influenza A virus (A/mallard/NZ/1/97(H5N2))		1		1		
	383586	Influenza A virus (A/Memphis/1/1971(H3N2))	19	11	19	11		
X	378139	Influenza A virus (A/Memphis/1/1978(H1N1))				5		5
X	383577	Influenza A virus (A/Memphis/1/1990(H3N2))				1		1
	79695	Influenza A virus (A/Memphis/1/71H-A/Bellamy/42N)		1		1		
	385640	Influenza A virus (A/Memphis/102/1972(H3N2))		7		8		1
	228928	Influenza A virus (A/Memphis/31/98(H3N2))	6		6			
	383571	Influenza A virus (A/Memphis/6/1986(H3N2))	2		4		2	
	480019	Influenza A virus (A/Moscow/10/1999(H3N2))		2		2		
	334590	Influenza A virus (A/Moscow/343/2003(H3N2))		1		1		
	389259	Influenza A virus (A/Nagasaki/N03/2005(H3N2))		1		1		
	62488	Influenza A virus (A/Nanchang/58/1993(H3N2))		1		1		
	132884	Influenza A virus (A/Netherlands/458/98 (H3N2))		1		1		
	132841	Influenza A virus (A/Netherlands/785e/90 (H3N2))		1		1		
	381512	Influenza A virus (A/New Caledonia/20/1999(H1N1))	2	51	3	127	1	76
	311695	Influenza A virus (A/New York/124/2001(H3N2))		1		1		
	335358	Influenza A virus (A/New York/232/2004(H3N2))		102		102		
X	327205	Influenza A virus (A/New York/348/2003(H1N1))				56		56
	335313	Influenza A virus (A/New York/364/2004(H3N2))		1		1		
X	335333	Influenza A virus (A/New York/384/2005(H3N2))				61		61
	335337	Influenza A virus (A/New York/388/2005(H3N2))		14		14		
X	342508	Influenza A virus (A/New York/444/2001(H1N1))				14		14
X	364132	Influenza A virus (A/New York/504/1998(H3N2))				1		1
	62564	Influenza A virus (A/New_York/15/94(H3N2))		1		1		
	62496	Influenza A virus (A/New_York/17/94(H3N2))		1		1		

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X	370128	Influenza A virus (A/Northern Territory/60/1968(H3N2))				1		1
	384505	Influenza A virus (A/nt/60/1968(H3N2))		61		66		5
	155917	Influenza A virus (A/NWS/33HA-A/tern/Australia/G70C/75NA(H1N9))		1		1		
	382820	Influenza A virus (A/NWS/G70C(H1N9))		1		1		
	62503	Influenza A virus (A/Ohio/3/95(H3N2))			1		1	
	428486	Influenza A virus (A/Oklahoma/309/06(H3N2))			3		3	
	223935	Influenza A virus (A/Okuda/57(H2N2))		1	9	1	9	
	381513	Influenza A virus (A/Panama/2007/1999(H3N2))		10	8	10	10	
X	137885	Influenza A virus (A/parakeet/Chiba/1/97(H9N2))					3	3
X	137886	Influenza A virus (A/parakeet/Narita/92A/98(H9N2))					2	2
X	11448	Influenza A virus (A/parrot/Ulster/73(H7N1))					1	1
	382825	Influenza A virus (A/Philippines/2/1982(H3N2))		1		1		
	385624	Influenza A virus (A/Port Chalmers/1/1973(H3N2))		7	1	7	1	
	211044	Influenza A virus (A/Puerto Rico/8/1934(H1N1))		21	355	22	366	1 11
	183764	Influenza A virus (A/Puerto Rico/8/34/Mount Sinai(H1N1))		7	85	7	87	
	216877	Influenza A virus (A/quail/Hong Kong/FY298/00(H9N2))			4		4	
X	183778	Influenza A virus (A/quail/Hong Kong/SF550/00(H6N1))					2	2
	197589	Influenza A virus (A/Quail/Shanghai/8/96(H9N2))			1		1	
	417758	Influenza A virus (A/quail/Shantou/1475/2004(H9N2))		1		1		
X	411167	Influenza A virus (A/Queensland/11/2001(H3N2))					1	1
X	286239	Influenza A virus (A/red knot/Delaware/2561/87(H10N5))					1	1
	382828	Influenza A virus (A/RI/5/-1957(H2N2))			1		1	
	221012	Influenza A virus (A/Rio/6/69(H3N2))			1		1	
	142948	Influenza A virus (A/ruddy turnstone/Delaware/67/1998(H12N4))			1		1	
X	385616	Influenza A virus (A/ruddy turnstone/NJ/60/1985(N9))					1	1
	192720	Influenza A virus (A/Saudi Arabia/2581/2001(H1N2))			1		1	
	192535	Influenza A virus (A/Saudi Arabia/7971/2000(H1N1))			1		1	
	384493	Influenza A virus (A/seal/Mass/1/1980(H7N7))		1	1	1	1	
	62512	Influenza A virus (A/Shangdong/5/94(H3N2))			1		1	
	380948	Influenza A virus (A/Shangdong/9/1993(H3N2))		13		13		
X	383568	Influenza A virus (A/Shanghai/11/1987(H3N2))					1	1
	383567	Influenza A virus (A/Shanghai/16/1989(H3N2))			1		1	
	342207	Influenza A virus (A/shorebird/Delaware/141/2002(H9N9))			1		1	
	342210	Influenza A virus (A/shorebird/Delaware/275/2001(H9N7))			1		1	
	342211	Influenza A virus (A/shorebird/Delaware/277/2000(H9N7))			1		1	
	62575	Influenza A virus (A/Sichuan/2/1987(H3N2))			3		4	1
	382781	Influenza A virus (A/Singapore/1/1957(H2N2))		1		1		
	59375	Influenza A virus (A/South Carolina/1/1918(H1N1))		2		3		1
	177131	Influenza A virus (A/SW/KS/13481-T/00(H1N2))			1		1	
	177128	Influenza A virus (A/SW/MN/12883/00(H1N2))			2		2	
	177188	Influenza A virus (A/SW/MN/3227/00(H1N2))			1		1	
	382842	Influenza A virus (A/swine/29/1937(H1N1))			2		2	
	336207	Influenza A virus (A/swine/Bakum/1832/00(H1N2))			3		3	
	336205	Influenza A virus (A/swine/Bakum/5/95(H1N1))			2		2	
	256045	Influenza A virus (A/swine/Chai/77-10/2001(H3N1))			1		1	
X	136474	Influenza A virus (A/Swine/Colorado/23619/99(H3N2))					1	1
	169169	Influenza A virus (A/swine/Cotes d'Armor/1482/1999(H1N1))			1		1	
	158306	Influenza A virus (A/swine/Cotes d'Armor/800/00(H1N2))			1		1	
X	522740	Influenza A virus (A/swine/Hannover/1/1981(H1N1))					6	6
	145306	Influenza A virus (A/swine/Hong Kong/10/98(H9N2))		5		5		
	11498	Influenza A virus (A/swine/Hong Kong/126/82(H3N2))			1		1	
X	384483	Influenza A virus (A/swine/Hong Kong/127/1982(H3N2))					1	1

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	173473	Influenza A virus (A/swine/Hong Kong/2106/98(H9N2))		2		2		
	380332	Influenza A virus (A/swine/Hong Kong/273/1994(H1N1))		1		1		
	173477	Influenza A virus (A/swine/Hong Kong/3297/98(H9N2))		3		3		
X	380217	Influenza A virus (A/swine/Hong Kong/4/1976(H3N2))				1		1
	253682	Influenza A virus (A/swine/Hong Kong/5190/99(H3N2))		3		3		
	384484	Influenza A virus (A/swine/Hong Kong/81/1978(H3N2))		8		11		3
	145307	Influenza A virus (A/swine/Hong Kong/9/98(H9N2))	2		2			
	253691	Influenza A virus (A/swine/Hong Kong/9840/01(H3N2))		1		1		
	306061	Influenza A virus (A/swine/IDT/Bakum1832/2000(H1N2))		1		1		
	384487	Influenza A virus (A/Swine/Indiana/1726/1988(H1N1))		1		2		1
X	128982	Influenza A virus (A/Swine/Indiana/9K035/99 (H1N2))				2		2
X	161502	Influenza A virus (A/swine/Italy/1509-6/97(H1N1))				12		12
X	164042	Influenza A virus (A/Swine/Italy/1513-1/98(H1N1))				1		1
X	169166	Influenza A virus (A/swine/Italy/1521/98(H1N2))				2		2
	170342	Influenza A virus (A/swine/Italy/1654-1/99(H1N2))		1		1		
	383561	Influenza A virus (A/swine/Italy/1850/1977(H3N2))		1		1		
X	383531	Influenza A virus (A/swine/Italy/2/1979(H1N1))				3		3
	147095	Influenza A virus (A/Swine/Italy/25823/94(H3N2))		2		2		
	383566	Influenza A virus (A/swine/Italy/425/1976(H1N1))		1		1		
	383564	Influenza A virus (A/swine/Italy/547/1985(H1N1))		3		3		
	383558	Influenza A virus (A/swine/Italy/809/1989(H3N2))		1		1		
X	538555	Influenza A virus (A/swine/Korea/Hongsong2/2004(H1N2))				2		2
	300748	Influenza A virus (A/swine/Korea/S190/2004(H9N2))		3		3		
X	187164	Influenza A virus (A/Swine/Minnesota/55551/2000(H1N2))				1		1
	426738	Influenza A virus (A/swine/Miyagi/5/2003(H1N2))		2		2		
	186460	Influenza A virus (A/swine/New Jersey/11/1976(H1N1))	1		1			
X	515581	Influenza A virus (A/swine/Ohio/24366/2007(H1N1))				1		1
X	83206	Influenza A virus (A/swine/Scotland/410440/94(H1N2))				3		3
	385644	Influenza A virus (A/swine/Ukkel/1/1984(H3N2))		1		1		
	82372	Influenza A virus (A/Sydney/05/97-like(H3N2))		1		1		
X	380213	Influenza A virus (A/Taiwan/1/1986(H1N1))				1		1
	371082	Influenza A virus (A/Taiwan/2/2006(H1N1))		8		8		
	395842	Influenza A virus (A/tern/Australia/1/04(H2N5))		1		1		
	405050	Influenza A virus (A/tern/Australia/1402/2004(H2N5))		1		1		
	384509	Influenza A virus (A/tern/Australia/G70C/1975(H11N9))	10		10			
	444318	Influenza A virus (A/Texas/1/1977(H3N2))	2	13	2	13		
X	270945	Influenza A virus (A/Thailand/4(SP-528)/2004(H5N1))				20		20
	427773	Influenza A virus (A/Thailand/SP83/2004(H5N1))	1		1			
	270484	Influenza A virus (A/tiger/Thailand/CU-LV/2004(H5N1))		2		2		
	322564	Influenza A virus (A/tiger/Thailand/CU-T4/04(H5N1))		6		6		
	322566	Influenza A virus (A/tiger/Thailand/CU-T6/04(H5N1))		1		1		
X	225071	Influenza A virus (A/Tokyo/1566/98(H3N2))				1		1
	342396	Influenza A virus (A/turkey/Canada-Ontario/NS-01839-1/05(H3))		1		1		
X	380285	Influenza A virus (A/turkey/Ireland/1378/1983(H5N8))				1		1
	402489	Influenza A virus (A/turkey/Minnesota/1200/1980(H7N3))		1		1		
X	383603	Influenza A virus (A/turkey/Minnesota/833/1980(H4N2))				1		1
	342197	Influenza A virus (A/turkey/MN/511/78(H9N2))		2		2		
	342145	Influenza A virus (A/turkey/MN/735/79(H6N2))		2		2		
	165420	Influenza A virus (A/Turkey/MO/24093/99(H1N2))		5		5		
	533026	Influenza A virus (A/turkey/Ohio/313053/2004(H3N2))		1		3		2
	293054	Influenza A virus (A/turkey/Ontario/6213/66(H5N?))	1		1			
	380301	Influenza A virus (A/turkey/Ontario/7732/1966(H5N9))	5	6	5	6		
X	380299	Influenza A virus (A/turkey/Wisconsin/1968(H5N9))				1		1

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	385599	Influenza A virus (A/udorn/1972(H3N2))		1		1		
	381517	Influenza A virus (A/Udorn/307/1972(H3N2))		4		4		
X	392811	Influenza A virus (A/Udorn/8/1972(H3N2))			1		1	
	62596	Influenza A virus (A/USSR/26/1985(H3N2))		1		1		
	381516	Influenza A virus (A/USSR/90/1977(H1N1))	13	1	13	1		
	370127	Influenza A virus (A/Victoria/1968(H3N2))		1		1		
	392809	Influenza A virus (A/Victoria/3/1975(H3N2))	43	8	43	8		
	284217	Influenza A virus (A/Viet Nam/1194/2004(H5N1))	1	4	1	4		
	284218	Influenza A virus (A/Viet Nam/1203/2004(H5N1))	43	50	44	57	1	7
	299390	Influenza A virus (A/Viet Nam/HN/2004(H5N1))		6		6		
X	644788	Influenza A virus (A/Vietnam/1194/2004(H5N1))			1		1	
	357208	Influenza A virus (A/Vietnam/CL26/2004(H5N1))		72		72		
	382832	Influenza A virus (A/VM113-V1(H1N1))	1		1			
X	383121	Influenza A virus (A/Waikato/7/2000(H3N2))				2		2
	191090	Influenza A virus (A/Weiss/1943(H1N1))		1		1		
X	383152	Influenza A virus (A/Wellington/22/2001(H3N2))				1		1
X	383231	Influenza A virus (A/Wellington/8/2004(H3N2))				1		1
	11484	Influenza A virus (A/whale/Maine/1/84(H13N9))	3		3			
	344598	Influenza A virus (A/whooper swan/Mongolia/6/05(H5N1))		1		1		
	381518	Influenza A virus (A/Wilson-Smith/1933(H1N1))		108		109		1
	461772	Influenza A virus (A/Wisconsin/43/2006(H3N2))	1		1			
	380978	Influenza A virus (A/Wisconsin/4755/1994(H1N1))		6		6		
	382835	Influenza A virus (A/WSN/1933(H1N1))	4	2	4	3		1
	63106	Influenza A virus (A/Wuhan/359/1995(H3N2))	5		5			
	480024	Influenza A virus (A/Wyoming/03/2003(H3N2))		5		5		
X	432094	Influenza A virus (A/Wyoming/3/2003-X-147(H3N2))				1		1
	132504	Influenza A virus (A/X-31(H3N2))	77	151	77	174		23
	380905	Influenza A virus (A/X-47(H3N2))	3	2	3	2		
	327311	Influenza A virus (A/Yilan/515/03(H3N2))		1		1		
X	255681	Influenza A virus (A/Yokohama/22/2002(H1N2))				1		1
	11408	Influenza A virus (STRAIN A/EQUINE/NEW MARKET/76)		1		1		
X	55532	Influenza A virus (strain A/SW/Quebec/1192/1986)				1		1
X	168278	Influenza A virus (swine/Finistere/127/99(H3N2))				4		4
	41857	Influenza A virus H3N2	1	15	1	39		24
	10000865	Influenza A virus H3N2 (A/Kiev/301/94)		3		3		
	10000550	Influenza A virus H3N2 (A/Netherlands/9/03 (H3N2))			1		1	
	10000474	Influenza A virus H3N2 (A/Resvir-9 (H3N2))		10		10		
	415166	Influenza A virus(A/swine/Henan/2/2004(H9N2))		1		1		
	415167	Influenza A virus(A/swine/Henan/3/2004(H9N2))		2		2		
	415169	Influenza A virus(A/swine/Henan/5/2004(H9N2))		1		1		
	415172	Influenza A virus(A/swine/Henan/8/2004(H9N2))		2		2		
	415176	Influenza A virus(A/swine/Shandong/fNY/2003(H9N2))		3		3		
	415177	Influenza A virus(A/swine/Shandong/fZC/2003(H9N2))		7		7		
	11520	Influenza B virus	2	6	2	6		
	11521	Influenza B virus (B/Ann Arbor/1/1986)		3		3		
	107404	Influenza B virus (B/Beijing/184/93)		1		1		
	107406	Influenza B virus (B/Chiba/447/98)			1		1	
	206203	Influenza B virus (B/Hong Kong/330/2001)		16		16		
	224964	Influenza B virus (B/Johannesburg/5/99)	1		1			
	184816	Influenza B virus (B/Kadoma/122/99)	2		2			
	256080	Influenza B virus (B/Kobe/1/2003)	1		1			
	504664	Influenza B virus (B/Kobe/113/2005)	1		1			
	504666	Influenza B virus (B/Kobe/115/2005-T1)	1		1			

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	503386	Influenza B virus (B/Kobe/3/2004)	1		1			
	504662	Influenza B virus (B/Kobe/39/2005)	1		1			
	504660	Influenza B virus (B/Kobe/67/2005)	1		1			
	171425	Influenza B virus (B/Kobe/87/2001)	1		1			
	107412	Influenza B virus (B/Lee/40)	5	3	5	3		
	11541	Influenza B virus (B/Oregon/5/80)	16		16			
	150127	Influenza B virus (B/Osaka/983/97-V3)	1		1			
	335812	Influenza B virus (B/Shanghai/361/2002)	1		1			
	107418	Influenza B virus (B/Victoria/2/87)	2		2	1	1	
	11531	Influenza B virus (STRAIN B/HONG KONG/8/73)		1		1		
	11532	Influenza B virus (STRAIN B/HT/84)		1		1		
	11552	Influenza C virus		1		1		
	11553	Influenza C virus (C/Ann Arbor/1/50)	9		9			
	197911	Influenzavirus A		6		6		
	9725	Inia geoffrensis		1		1		
X	55635	Inula helenium				1		1
	42097	Isla Vista virus	1	2	1	2		
	261204	Itapua hantavirus		10		10		
	11072	Japanese encephalitis virus	19	13	19	18		5
	10000444	Japanese encephalitis virus CH2195LA	1		1			
	10000445	Japanese encephalitis virus JaOH0566	1		1			
	11075	Japanese encephalitis virus strain JAOARS982	4	4	4	4		
	11076	Japanese encephalitis virus strain Nakayama	2	2	2	2		
	11073	Japanese encephalitis virus strain SA-14	1	1	1	1		
	10001335	Japanese encephalitis virus Vellore P20778	1		1	1	1	
	10632	JC polyomavirus	1	5	1	5		
	51240	Juglans regia		27		27		
	11619	Junin virus	3	867	3	871		4
	13101	Juniperus ashei	13		13			
	453927	Juniperus formosana	1		1			
	430511	Juquitiba virus		2		2		
X	42894	Khabarovsk virus				1		1
	10638	Kilham polyomavirus		1		1		
	573	Klebsiella pneumoniae	16	3	21	6	5	3
X	484021	Klebsiella pneumoniae NTUH-K2044				1		1
	11077	Kunjin virus		1	3	1	3	
	11078	Kunjin virus (STRAIN MRM61C)		2		2		
	11577	La Crosse virus	2	1	2	1		
	11578	La Crosse virus L74		1		2		1
	8753	Lachesis muta muta	21		21			
	300015	Lactate dehydrogenase elevating virus C	1		1			
	300016	Lactate dehydrogenase elevating virus Plagemann	1		1			
	11048	Lactate dehydrogenase-elevating virus	14		14			
X	1585	Lactobacillus delbrueckii subsp. bulgaricus			1		1	
X	1358	Lactococcus lactis				2		2
	61172	Laguna Negra virus		9		11		2
	11269	Lake Victoria marburgvirus		18		70		52
	33727	Lake Victoria marburgvirus - Musoke	4	81	4	81		
	33728	Lake Victoria marburgvirus - Popp	1	14	1	14		
	378809	Lake Victoria marburgvirus - Ravn		32		32		
	11085	Langat virus		1		1		
	11620	Lassa virus	4	626	4	633		7
	11621	Lassa virus GA391		43		45		2

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	11622	Lassa virus Josiah		372		376		4
	6925	Latrodectus tredecimguttatus		1		1		
X	85223	Laurus nobilis				1		1
	69245	Lechiguanas virus		1		5		4
X	446	Legionella pneumophila				1		1
X	272624	Legionella pneumophila subsp. pneumophila str. Philadelphia 1				1		1
X	218308	Lehmannia valentiana				1		1
	5658	Leishmania	1		1			
	5667	Leishmania aethiopica		16		16		
	5659	Leishmania amazonensis			2		2	
	5660	Leishmania braziliensis		6		6		
	44271	Leishmania chagasi		2		2		
	5661	Leishmania donovani		9	38	9	38	
	99875	Leishmania donovani donovani		1		1		
	10000341	Leishmania donovani donovani 1S2D		1		1		
	5671	Leishmania infantum		93	1	93	1	
	10000345	Leishmania infantum LEM 75			14		14	
	5664	Leishmania major		13	117	13	119	2
	347515	Leishmania major strain Friedlin			14		14	
	5665	Leishmania mexicana		1		1	1	1
	5679	Leishmania panamensis		3	6	3	6	
	5682	Leishmania pifanoi			20		20	
	11049	Lelystad virus		24		26	12	12
	36936	Lepidoglyphus destructor		5	10	5	10	
X	173	Leptospira interrogans				4	4	4
X	44275	Leptospira interrogans serovar Copenhageni				2		2
	267671	Leptospira interrogans serovar Copenhageni str. Fiocruz L1-130		11		11		
X	57678	Leptospira interrogans serovar Lai				2	2	2
	189518	Leptospira interrogans serovar Lai str. 56601		1		1		
	10000814	Leptospira interrogans serovar Lai str. HY-1		1		1		
X	44276	Leptospira interrogans serovar Pomona					3	3
	10000847	Leptospira sp. Akiyami A AUT10		1		1		
	1642	Listeria innocua		2	3	2	3	
X	1638	Listeria ivanovii				1	1	1
	1639	Listeria monocytogenes		18	75	18	75	
	393133	Listeria monocytogenes 10403S			4		4	
	10000308	Listeria monocytogenes ATCC 35967			1		1	
	10000309	Listeria monocytogenes ATCC 43251			2		2	
	169963	Listeria monocytogenes EGD-e		5	45	5	56	11
X	393128	Listeria monocytogenes F6900					1	1
X	393117	Listeria monocytogenes FSL J1-194					1	1
	393124	Listeria monocytogenes FSL N3-165			2		2	
X	393126	Listeria monocytogenes FSL R2-561					1	1
X	552536	Listeria monocytogenes HCC23					2	2
X	401650	Listeria monocytogenes HPB2262					3	3
X	393130	Listeria monocytogenes J0161					4	4
	267409	Listeria monocytogenes serotype 1/2a str. F6854			2		3	1
X	568819	Listeria monocytogenes serotype 4b str. CLIP 80459					4	4
	265669	Listeria monocytogenes serotype 4b str. F2365			3		6	3
	267410	Listeria monocytogenes serotype 4b str. H7858			19		23	4
X	386043	Listeria welshimeri serovar 6b str. SLCC5334					2	2
	55601	Listonella anguillarum		1		1		
	6299	Litomosoides carinii		3		3		

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	217686	Little cherry virus 1		1		1		
	4522	Lolium perenne	36	78	36	81		3
	36386	Louping ill virus (strain 31)	1		1			
X	58218	Loxosceles intermedia			1		1	
	11623	Lymphocytic choriomeningitis virus	2	339	2	348		9
	11624	Lymphocytic choriomeningitis virus (strain Armstrong)		1145		1145		
	10000496	Lymphocytic choriomeningitis virus (strain Armstrong) (clone 3)		1		1		
	10000487	Lymphocytic choriomeningitis virus (strain Armstrong) (clone 4)	1		1			
	10000497	Lymphocytic choriomeningitis virus (strain Armstrong) (clone 5)	1		1			
	10000488	Lymphocytic choriomeningitis virus (strain Armstrong) (clone 53b)		45		45		
	11625	Lymphocytic choriomeningitis virus (strain Pasteur)		5		5		
	11626	Lymphocytic choriomeningitis virus (strain Traub)		3		3		
	11627	Lymphocytic choriomeningitis virus (strain WE)		70		70		
	10000494	Lymphocytic choriomeningitis virus (strain WE) variant 8.7		1		1		
	10000495	Lymphocytic choriomeningitis virus (strain WE) WE CL1.2		1		1		
	10000490	Lymphocytic choriomeningitis virus A22.2b		1		1		
	10000491	Lymphocytic choriomeningitis virus Docile		2		2		
	9541	Macaca fascicularis	1	3	1	3		
	9544	Macaca mulatta		1		1		
	10373	Macaca mulatta cytomegalovirus		44		44		
X	9548	Macaca radiata			14	1	14	1
	10325	Macacine herpesvirus 1	1		1			
	47929	Macacine herpesvirus 3		68		68		
	45455	Macacine herpesvirus 4		18		18		
	11628	Machupo virus		778		784		6
X	272627	Magnetospirillum magnetotacticum MS-1				3		3
	76777	Malassezia sympodialis	1		1			
	3750	Malus x domestica	27	1	27	1		
	40674	Mammalia	3	1	3	1		
	351073	Mammalian orthoreovirus		1		1		
	7130	Manduca sexta		3		3		
	29780	Mangifera indica	1		1			
	45201	Mannheimia haemolytica serotype 1		82		82		
	221988	Mannheimia succiniciproducens MBEL55E		2		2		
	186537	Marburgvirus		2		2		
	270374	Marinobacter sp. ELB17		1		1		
X	38020	marmosets			2		2	
	11234	Measles virus	54	36	63	37	9	1
	10000462	Measles virus CAM/RB		1		1		
	11235	Measles virus strain Edmonston	167	203	167	203		
	70146	Measles virus strain Edmonston-B		13		13		
X	70149	Measles virus strain Edmonston-Zagreb			1		1	
	11236	Measles virus strain Halle		34		34		
	132487	Measles virus strain Schwarz	1	1	1	1		
X	9103	Meleagris gallopavo			2		2	
X	152219	Menangle virus			2		2	
	12107	Mengo virus	5	13	5	13		
	10036	Mesocricetus auratus	31		31			
	243232	Methanocaldococcus jannaschii DSM 2661		1		1		
	187420	Methanothermobacter thermautrophicus str. Delta H		1		1		
X	420662	Methylibium petroleiphilum PM1				1		1
	10793	Mink enteritis virus strain Abashiri	1		1			
	467144	Modified Vaccinia Ankara virus		54		54		

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	11801	Moloney murine leukemia virus		5		5		
	10244	Monkeypox virus		5		5		
	264732	Moarella thermoacetica ATCC 39073		1		1		
	300180	Mopeia Lassa reassortant 29		1		3		2
	11629	Mopeia virus		12		12		
	480	Moraxella catarrhalis	9		10		1	
X	10001556	Moraxella catarrhalis 26404			1		1	
	11757	Mouse mammary tumor virus		10		10		
	11161	Mumps virus	3		3			
	11169	Mumps virus (STRAIN KILHAM)		1		1		
	11173	Mumps virus (STRAIN SBL-1)	3		3			
	10366	Murid herpesvirus 1		41		41		
	10000411	Murid herpesvirus 1 deltaMS94.5		2		2		
	10000412	Murid herpesvirus 1 Isolate G4		1		1		
	10000413	Murid herpesvirus 1 Isolate K6		1		1		
	33708	Murid herpesvirus 4	1	31	1	52		21
	10000536	Murid herpesvirus 4 G2.4		3		3		
	10000537	Murid herpesvirus 4 WUMS		2		2		
X	10530	Murine adenovirus 1			1	2	1	2
	69156	Murine cytomegalovirus (strain K181)	2	11	2	11		
	10367	Murine cytomegalovirus (strain Smith)		30		30		
	10001396	Murine cytomegalovirus (strain Smith) MW97.01		5		5		
	35275	Murine endogenous retrovirus		11		11		
	11138	Murine hepatitis virus	3	15	3	15		
	12760	Murine hepatitis virus strain 4		6		6		
	11142	Murine hepatitis virus strain A59	26	3	26	3		
	11144	Murine hepatitis virus strain JHM	8	14	8	14		
	11786	Murine leukemia virus		7		7		
	10001207	Murine leukemia virus LP-BM5		1		1		
	223997	Murine norovirus 1	1		2		1	
	10634	Murine polyomavirus	1	25	2	25	1	
	10636	Murine polyomavirus strain A2		1		3		2
	28327	Murine rotavirus		3		3		
	70865	Murine rotavirus EDIM		27		27		
	11812	Murine sarcoma virus 3611	4		4			
	44561	Murine type C retrovirus		1		1		
	11079	Murray Valley encephalitis virus	12	6	12	7		1
	301478	Murray valley encephalitis virus (strain MVE-1-51)	11	10	11	10		
	10090	Mus musculus	245	932	437	1824	192	892
	10000000	Mus musculus BALB/c		2		2		
	10092	Mus musculus domesticus				1		1
	10095	Mus sp.		1		1		
	4641	Musa acuminata	50	1	50	1		
	214697	Musa acuminata AAA Group		1		1		
	1763	Mycobacterium	2	11	2	15		4
	1764	Mycobacterium avium	5	12	6	12	1	
	243243	Mycobacterium avium 104		10		10		
	10000313	Mycobacterium avium serovar 1	1		1			
	10000314	Mycobacterium avium serovar 2	1		1			
	10000316	Mycobacterium avium serovar 25	1		1			
	10000317	Mycobacterium avium serovar 26	1		1			
	10000318	Mycobacterium avium serovar 4	2		2			
	10000319	Mycobacterium avium serovar 8	1	1	1	1		

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	10000320	Mycobacterium avium serovar 9	1		1			
	1770	Mycobacterium avium subsp. paratuberculosis		3	2	4	2	1
X	262316	Mycobacterium avium subsp. paratuberculosis K-10			2		2	
	10000328	Mycobacterium avium subsp. paratuberculosis Strain Ben		1		1		
	1765	Mycobacterium bovis	44	122	47	125	3	3
	233413	Mycobacterium bovis AF2122/97		49		49		
	10000322	Mycobacterium bovis AN5	9	28	9	28		
	33892	Mycobacterium bovis BCG	11	136	11	147		11
	410289	Mycobacterium bovis BCG str. Pasteur 1173P2	1	15	1	88		73
	10000323	Mycobacterium bovis T/91/1378		6		6		
	670516	Mycobacterium chelonae group	1		1			
	1766	Mycobacterium fortuitum		1		1		
	144549	Mycobacterium fortuitum subsp. fortuitum		1		1		
	10000331	Mycobacterium gastri W471	1		1			
X	350054	Mycobacterium gilvum PYR-GCK				4		4
X	1778	Mycobacterium gordonaee			1		1	
	1767	Mycobacterium intracellulare	1	1	1	1		
	1768	Mycobacterium kansasii	9	8	9	8		
	557599	Mycobacterium kansasii ATCC 12478	1		1			
	10000325	Mycobacterium kansasii Subspecies IV		3		3		
	10000326	Mycobacterium kansasii Subspecies V		2		2		
	1769	Mycobacterium leprae	116	357	117	363	1	6
	272631	Mycobacterium leprae TN		6		6		
X	525368	Mycobacterium parascrofulaceum ATCC BAA-614				1		1
	43304	Mycobacterium peregrinum	2		2			
	1783	Mycobacterium scrofulaceum	4		4			
	1772	Mycobacterium smegmatis		3		3		
	1785	Mycobacterium sp.		2		2		
	164757	Mycobacterium sp. JLS		1		1		
	189918	Mycobacterium sp. KMS		1		1		
	1773	Mycobacterium tuberculosis	449	993	475	1043	26	50
	10000329	Mycobacterium tuberculosis 103	1		1			
	348776	Mycobacterium tuberculosis C		13		67		54
	83331	Mycobacterium tuberculosis CDC1551		16		22		6
	10000330	Mycobacterium tuberculosis Erdman	15	19	15	19		
	336982	Mycobacterium tuberculosis F11		9		13		4
	419947	Mycobacterium tuberculosis H37Ra		10		56		46
	83332	Mycobacterium tuberculosis H37Rv	23	377	53	419	30	42
	395095	Mycobacterium tuberculosis str. Haarlem		82		88		6
	182785	Mycobacterium tuberculosis subsp. tuberculosis		1		1		
	1809	Mycobacterium ulcerans		7		7		
	362242	Mycobacterium ulcerans Agy99		1		1		
	347257	Mycoplasma agalactiae PG2	3		3			
	28903	Mycoplasma bovis	21		21			
	10001075	Mycoplasma gallisepticum strain S6	1		1			
	2097	Mycoplasma genitalium		1		1		
	243273	Mycoplasma genitalium G37	7		7			
	2099	Mycoplasma hyopneumoniae	1		2		1	
X	295358	Mycoplasma hyopneumoniae 232			21		21	
X	754503	Mycoplasma hyopneumoniae 7422			1		1	
X	262722	Mycoplasma hyopneumoniae 7448			1		1	
	28227	Mycoplasma penetrans	34		34			
	2104	Mycoplasma pneumoniae	9	2	9	2		

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	10000332	Mycoplasma pneumoniae FH	1		1			
	272634	Mycoplasma pneumoniae M129	9		9			
	13618	Myrmecia pilosula	2		2			
X	10273	Myxoma virus				3		3
	8656	Naja atra	7		7			
	35670	Naja naja	3		3			
	8654	Naja nigricollis	3	2	3	2		
	8657	Naja oxiana	4		4			
	8658	Naja pallida	1	1	1	1		
X	51031	Necator americanus			2		2	
	367400	Neembucu hantavirus		3		4		1
	485	Neisseria gonorrhoeae	19	10	21	11	2	1
X	10001558	Neisseria gonorrhoeae 15253			1		1	
X	10000863	Neisseria gonorrhoeae 4505			1		1	
	528354	Neisseria gonorrhoeae MS11	8		8			
	10001077	Neisseria gonorrhoeae SU89	1		1			
	10001078	Neisseria gonorrhoeae SU96	3		3			
	487	Neisseria meningitidis	38	18	39	22	1	4
	272831	Neisseria meningitidis FAM18	1		1			
X	10001560	Neisseria meningitidis M982B			2		2	
	122586	Neisseria meningitidis MC58	12		12			
	65699	Neisseria meningitidis serogroup A	1		1			
	10001051	Neisseria meningitidis serogroup A Strain 8659	1		1			
	491	Neisseria meningitidis serogroup B	36	1	36	1		
	10000843	Neisseria meningitidis serogroup B H44/76	73	22	74	22	1	
	10001003	Neisseria meningitidis serogroup B CU385	2		2			
	10001050	Neisseria meningitidis serogroup B Strain 7967	1		1			
	10000979	Neisseria meningitidis serogroup B Strain 8047	4		4			
	10000972	Neisseria meningitidis serogroup B Strain B16.B6	1		1			
	10001162	Neisseria meningitidis serogroup B Strain M1239	2		2			
	10001049	Neisseria meningitidis serogroup B Strain S3446	1		1			
	10001044	Neisseria meningitidis serogroup C MC51	1		1			
	29176	Neospora caninum	2	2	2	2		
	452646	Neovison vison		1		1		
	5141	Neurospora crassa	4		6	3	2	3
	11176	Newcastle disease virus	11		14	3	3	3
	11177	Newcastle disease virus (STRAIN AUSTRALIA-VICTORIA/32)	4		4			
	10001138	Newcastle disease virus (strain Eaves)	1		1			
	10001032	Newcastle disease virus (strain La Sota)	1		1			
	10001140	Newcastle disease virus (strain WA2116)	1		1			
	351071	Newcastle disease virus AF2240	1		1			
	11178	Newcastle disease virus strain Beaudette C/45	12		12			
	11180	Newcastle disease virus strain D26/76	4		4			
	11186	Newcastle disease virus strain Queensland/66	4		4			
	4097	Nicotiana tabacum	2		2			
	121791	Nipah virus	5		5			
	314278	Nitrococcus mobilis Nb-231		1		1		
X	142786	Norovirus			2	1	2	1
	122928	Norovirus genogroup 1	1		1			
X	10001484	Norovirus genogroup 1 GI.12			1		1	
	10001116	Norovirus genogroup 1 isolates 96-908	4		4			
	122929	Norovirus genogroup 2	2		2			
	10000829	Norovirus genogroup 2 Mexico type strain 36	3		3			

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	10000560	Norovirus genogroup 3 Bo/Jena/1980/DE	1		1			
X	10001485	Norovirus genogroup 3 GIII.1			1		1	
X	647514	Norovirus genogroup GI.1			1		1	
X	647515	Norovirus genogroup GI.2			1		1	
X	647519	Norovirus genogroup GI.5			1		1	
X	647521	Norovirus genogroup GI.7			1		1	
X	499191	Norovirus genogroup GII.1			1		1	
X	520963	Norovirus genogroup GII.11			1		1	
X	520964	Norovirus genogroup GII.18			1		1	
X	490039	Norovirus genogroup GII.2			1		1	
	489821	Norovirus genogroup GII.4	1		1			
X	508775	Norovirus genogroup GII.9			1		1	
X	588533	Norovirus Hu/GII/GZ-1/2008/CHN			1		1	
	150080	Norovirus isolates	1		1			
X	63737	Nostoc punctiforme PCC 73102					1	1
	8663	Notechis scutatus	2		2			
	8996	Numida meleagris	1		1			
	4146	Olea europaea	65	14	65	14		
	42764	Oliveros virus			1		1	
	6282	Onchocerca volvulus	3	38	3	39		1
	69247	Oran virus			2		2	
	9733	Orcinus orca			1		1	
	784	Orientia tsutsugamushi	86		86			
	10000761	Orientia tsutsugamushi Karp	1		1			
	357244	Orientia tsutsugamushi str. Boryong	2	1	2	2		1
X	334380	Orientia tsutsugamushi str. Ikeda					1	1
	9986	Oryctolagus cuniculus	1	6	15	7	14	1
	39947	Oryza sativa Japonica Group	5	1	5	1		
	28869	Ovine respiratory syncytial virus	1		1			
	9940	Ovis aries	74	2	80	3	6	1
	8667	Oxyuranaus scutellatus scutellatus	3		3			
	9598	Pan troglodytes	6	13	6	17		4
	188763	Panine herpesvirus 2			2		2	
	3469	Papaver somniferum	1		1			
	61183	Papio sp.	1		1			
	121759	Paracoccidioides brasiliensis	4	14	4	14		
	10000566	Paracoccidioides brasiliensis B339			8		8	
	502780	Paracoccidioides brasiliensis Pb18	1		1			
	266	Paracoccus denitrificans	1		1			
X	318586	Paracoccus denitrificans PD1222					2	2
	11207	Parainfluenza virus 5		1	1	1	1	
	45222	Parana virus			1		1	
	33127	Parietaria judaica	61	1	61	2		1
	13187	Parietaria officinalis	2		2			
	747	Pasteurella multocida			10		10	
X	272843	Pasteurella multocida subsp. multocida str. Pm70					4	4
	10000759	Pasteurella multocida X-73	2		2			
	1254	Pediococcus acidilactici	1		1			
	133894	Penaeus	2		2			
	5076	Penicillium chrysogenum	45		45			
	69248	Pergamino virus			1		1	
	6978	Periplaneta americana	9		9			
	31604	Peste-des-petits-ruminants virus	9	1	9	1		

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	10001020	Peste-des-petits-ruminants virus (strain Nigeria 75/1)		1		1		
	3885	Phaseolus vulgaris		1	3	1	3	
X	9057	Phasianus colchicus colchicus			1		1	
	15957	Phleum pratense	49	72	49	717		645
	9742	Phocoena phocoena		1		1		
X	314292	Photobacterium angustum S14				1		1
X	659	Photobacterium phosphoreum				1		1
	72539	Physalis mottle virus	2		2			
	9755	Physeter catodon	1	27	1	28		1
	11630	Pichinde virus		7		8		1
	3352	Pinus taeda	4		4			
X	60876	Pixuna virus				1		1
	2439	Plasmid ColB2	1		1			
	2465	Plasmid F	1		1			
	141833	Plasmid piPO2T		2		2		
	28399	Plasmid R100		1		1		
	5820	Plasmodium	4	5	4	5		
	5821	Plasmodium berghei	14	44	14	45		1
	10000356	Plasmodium berghei NK65	2	2	2	2		
	5823	Plasmodium berghei str. ANKA	6		6			
	5824	Plasmodium brasilianum	2		2			
	5825	Plasmodium chabaudi	4	4	4	4		
	10000357	Plasmodium chabaudi adami DS	2	37	2	37		
	5827	Plasmodium cynomolgi	1		2		1	
	5833	Plasmodium falciparum	564	693	573	695	9	2
	10000358	Plasmodium falciparum 366		1		1		
	36329	Plasmodium falciparum 3D7	120	197	142	199	22	2
	57266	Plasmodium falciparum 7G8	40	50	40	50		
	10000363	Plasmodium falciparum Brazil-608	1	1	1	1		
	5835	Plasmodium falciparum CAMP/Malaysia	36	3	36	3		
	5836	Plasmodium falciparum CDC/Honduras	1		1			
	10000364	Plasmodium falciparum Clone PNG3		1		1		
	5837	Plasmodium falciparum FC27/Papua New Guinea	40	45	40	45		
	10000366	Plasmodium falciparum FCB-2	1		1			
	5838	Plasmodium falciparum FCR-3/Gambia	8		8			
	10000371	Plasmodium falciparum Indochina I/CDC	32		32			
	5848	Plasmodium falciparum isolate WELLCOME	26	53	26	53		
	10000373	Plasmodium falciparum ItG2G1		1		1		
	5839	Plasmodium falciparum K1	15	19	18	19	3	
	5840	Plasmodium falciparum LE5		2		2		
	5841	Plasmodium falciparum Mad20/Papua New Guinea	8	47	8	47		
	5843	Plasmodium falciparum NF54	14	33	14	35		2
	5842	Plasmodium falciparum NF7/Ghana	5		5			
	57270	Plasmodium falciparum Palo Alto/Uganda	14		14			
	5834	Plasmodium falciparum RO-33	8	1	8	1		
	10000375	Plasmodium falciparum RO71	1	1	1	1		
	5846	Plasmodium falciparum T4/Thailand	25		25			
	10000816	Plasmodium falciparum T9/96	1	1	1	1		
	10000376	Plasmodium falciparum UF-5	2		2			
	5857	Plasmodium fragile		4		4		
	5850	Plasmodium knowlesi	2	7	2	7		
	5851	Plasmodium knowlesi strain H	7	6	7	6		
	5852	Plasmodium knowlesi strain Nuri		3		3		

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	5858	Plasmodium malariae	1		1			
	5854	Plasmodium reichenowi		3		3		
	35085	Plasmodium simiovale	1		1			
	5859	Plasmodium simium	2		2			
	31272	Plasmodium sp.	1		1			
	5855	Plasmodium vivax	57	143	57	148		5
	31273	Plasmodium vivax (strain Belem)	6	34	6	34		
	10000378	Plasmodium vivax NK		2		2		
	126793	Plasmodium vivax Sal-1	3	1	3	1		
	10000862	Plasmodium vivax VK247	1		1			
	27990	Plasmodium vivax-like sp.	2		2			
	5861	Plasmodium yoelii	9	33	10	33	1	
	73239	Plasmodium yoelii yoelii	22	26	22	28		2
	10000555	Plasmodium yoelii yoelii 265BY		1		1		
	352914	Plasmodium yoelii yoelii str. 17XNL	6	1	6	1		
	10000381	Plasmodium yoelii yoelii YM	1	5	1	5		
X	103448	Pleistophora sp. LS				1		1
X	703	Plesiomonas shigelloides				1		1
	12211	Plum pox virus	1		1			
	12213	Plum pox virus (isolate NAT)		1		1		
	10001100	Plum pox virus (strain W)	6		6			
	4754	Pneumocystis carinii	2		2	1		1
	263815	Pneumocystis murina	3		3			
	270473	Pneumonia virus of mice J3666		6		6		
	4545	Poa pratensis	34	18	34	18		
X	4479	Poaceae				1		1
X	365044	Paromomonas naphthalenivorans CJ2				1		1
	12088	Poliovirus type 3 (strains P3/LEON/37 AND P3/LEON 12A[1]B)	4	15	4	15		
	80956	Pomacentrus moluccensis		1		1		
X	9601	Pongo abelii				1		1
	9600	Pongo pygmaeus		2		2		
	46221	Porcine circovirus		49		49		
	133704	Porcine circovirus 1	5		5			
	85708	Porcine circovirus 2	13		15		2	
	10001239	Porcine circovirus strain ISU31	7		7			
	194958	Porcine endogenous retrovirus A	2		2			
	28295	Porcine epidemic diarrhea virus	1		1			
	229032	Porcine epidemic diarrhea virus (strain CV777)	4		4			
	10796	Porcine parvovirus	37		37			
	28344	Porcine reproductive and respiratory syndrome virus	62	3	62	3		
X	10001586	Porcine reproductive and respiratory syndrome virus 07V063			3		3	
X	10001587	Porcine reproductive and respiratory syndrome virus 08V204			2		2	
	10000969	Porcine respiratory and reproductive syndrome virus 111/92	9		9			
	10000529	Porcine respiratory and reproductive syndrome virus CH-1a	6		6			
	10000859	Porcine respiratory and reproductive syndrome virus JA142	1		1			
	10000864	Porcine respiratory and reproductive syndrome virus MD-001	1		1			
	10000888	Porcine respiratory and reproductive syndrome virus Olot/91	3		3			
	10919	Porcine rotavirus (STRAIN YM)	2	1	2	1		
	101350	Porcine rotavirus strain A253	2		2			
	10917	Porcine rotavirus strain Gottfried	2		2			
	53179	Porcine rubulavirus	3		3			
	11150	Porcine transmissible gastroenteritis coronavirus strain FS772/70	2		2			
	11151	Porcine transmissible gastroenteritis coronavirus strain Purdue	14		14			

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	837	Porphyromonas gingivalis	72	18	72	18		
	10001079	Porphyromonas gingivalis 381	71	55	71	55		
	10001081	Porphyromonas gingivalis HG66	2		2			
	10001223	Porphyromonas gingivalis OMZ 409	23		23			
	242619	Porphyromonas gingivalis W83	1		2		1	
	37128	Potato mop-top virus	8		8			
	12216	Potato virus Y	1		1			
	10001152	Potato virus Y strain D	6		6			
X	11083	Powassan virus				2		2
X	1752	Propionibacterium freudenreichii subsp. shermanii				1		1
	11603	Prospect Hill virus	1	2	1	2		
X	583	Proteus			1		1	
	584	Proteus mirabilis	1		6		5	
	10001006	Proteus mirabilis CFT322	1		1			
X	529507	Proteus mirabilis HI4320				1		1
X	10001498	Proteus mirabilis O23				1		1
X	10001500	Proteus mirabilis O43				1		1
X	10001499	Proteus mirabilis O6				1		1
X	102862	Proteus penneri				1		1
X	10001501	Proteus penneri ATCC 33519				1		1
X	10001489	Proteus vulgaris Strain 5/43				1		1
	88086	Protobothrops elegans	7		7			
	300559	PRRSV VR2332	18		18			
	36596	Prunus armeniaca	4		4			
	42229	Prunus avium	3	1	3	1		
	3758	Prunus domestica	4		4			
	3760	Prunus persica	15	43	17	49	2	6
	8671	Pseudechis porphyriacus	3		3			
X	342610	Pseudoalteromonas atlantica T6c					1	1
	287	Pseudomonas aeruginosa	58	48	65	50	7	2
X	350704	Pseudomonas aeruginosa C3719					2	2
	10001057	Pseudomonas aeruginosa CD4	1		1			
X	10000822	Pseudomonas aeruginosa Immunotype 3				1		1
	10000723	Pseudomonas aeruginosa Immunotype 4	4		4			
	10001058	Pseudomonas aeruginosa K122-4	1		1			
	10000817	Pseudomonas aeruginosa KB7	2		2			
	10000818	Pseudomonas aeruginosa P1	1		1			
X	388272	Pseudomonas aeruginosa PACS2					4	4
	10000725	Pseudomonas aeruginosa PAK	7	3	7	3		
	10000815	Pseudomonas aeruginosa PAO	2	2	2	2		
	208964	Pseudomonas aeruginosa PAO1	1		1	1		1
X	10001519	Pseudomonas aeruginosa serotype O11				1		1
	208963	Pseudomonas aeruginosa UCBPP-PA14			16		20	4
	237609	Pseudomonas alkylphenolia			1		1	
	294	Pseudomonas fluorescens			4		4	
	10879	Pseudomonas phage phi6	7		7			
	303	Pseudomonas putida	1	7	1	8		1
	351746	Pseudomonas putida F1			2		4	2
	76869	Pseudomonas putida GB-1			1		1	
	306	Pseudomonas sp.			1		1	
	74138	Pseudomonas sp. DJ-12			1		1	
	71238	Pseudomonas sp. G-179			1		1	
	159091	Pseudomonas sp. KIE171			1		1	

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X	91465	Pseudomonas sp. MIS38					1	1
	150396	Pseudomonas sp. MT-1		1		1		
	264730	Pseudomonas syringae pv. phaseolicola 1448A		1		1		
	29442	Pseudomonas tolaasii		1		1		
	11604	Puumala virus	30	13	30	15		2
X	11605	Puumala virus (STRAIN HALLNAS B1)					1	1
	10000485	Puumala virus (STRAIN HALLNAS B1) Vranica/Hallnas	4		4			
	39002	Puumala virus (strain sotkamo/v-2969/81)	196		196			
	10001238	Puumala virus (strain Umea/hu)	1		1			
	10000483	Puumala virus CG18-20	5		5			
	10000484	Puumala virus Kazan	25	7	25	7		
	272844	Pyrococcus abyssi GE5		1		1		
	10000507	Rabbit hemorrhagic disease virus Olot/89	1		1			
	122291	Rabbit oral papillomavirus	2		2			
X	32606	Rabbitpox virus					2	2
	11292	Rabies virus	6	13	6	13		
	11293	Rabies virus AVO1		4		4		
	10000467	Rabies virus CVS	14		14			
	11294	Rabies virus CVS-11	1		1			
	11295	Rabies virus ERA	9	23	9	23		
	10000470	Rabies virus Flury LEP		1		1		
	11296	Rabies virus HEP-FLURY	7	1	7	1		
	11298	Rabies virus Nishigahara RCEH	1		1			
	37132	Rabies virus Ontario fox	3		3			
	10000471	Rabies virus RC-HL	4		4			
	103929	Rabies virus strain Pasteur vaccin	1	1	1	1		
	10114	Rattus	2	1	2	1		
	10116	Rattus norvegicus	60	230	169	377	109	147
X	10001412	Rattus norvegicus Wistar-Furth					7	7
X	48483	Reclinomonas americana					1	1
	1646	Renibacterium salmoninarum	12		12			
X	10891	Reovirus sp.					2	2
	12814	Respiratory syncytial virus	3	6	3	6		
	186539	Reston ebolavirus		3		6		3
	129003	Reston ebolavirus - Reston	2	27	2	27		
X	386032	Reston ebolavirus - Reston (1989)					1	1
	103930	Rhesus cytomegalovirus strain 68-1		5		5		
	10969	Rhesus rotavirus	8	1	8	1		
	60189	Rhipicephalus decoloratus	4		5		1	
X	1063	Rhodobacter sphaeroides					1	1
	43767	Rhodococcus equi	15		18		3	
	1833	Rhodococcus erythropolis		1		1		
X	1828	Rhodococcus fascians					1	1
X	316056	Rhodopseudomonas palustris BisB18					1	1
	160061	Ricinus		1		1		
	3988	Ricinus communis	7	3	9	3	2	
	35788	Rickettsia africae		1		1		
X	293614	Rickettsia akari str. Hartford					1	1
	787	Rickettsia australis		1		1		
	391896	Rickettsia bellii OSU 85-389		3		3		
	336407	Rickettsia bellii RML369-C		4		7		3
	788	Rickettsia canadensis		1		1		
	781	Rickettsia conorii		13		13		

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	272944	Rickettsia conorii str. Malish 7		5		5		
	42862	Rickettsia felis		1		1		
	315456	Rickettsia felis URRWXCal2		1		6		5
	35791	Rickettsia massiliae		1		1		
X	416276	Rickettsia massiliae MTU5				1		1
	782	Rickettsia prowazekii		20		21		1
	272947	Rickettsia prowazekii str. Madrid E		2		5		3
	783	Rickettsia rickettsii		1		1		
	452659	Rickettsia rickettsii str. Iowa		1		1		
X	392021	Rickettsia rickettsii str. 'Sheila Smith'				1		1
	35793	Rickettsia sibirica		4		4		
	272951	Rickettsia sibirica 246		8		15		7
	785	Rickettsia typhi		3		3		
	257363	Rickettsia typhi str. Wilmington		1		5		4
	11588	Rift Valley fever virus	4	2	4	2		
	11589	Rift valley fever virus (STRAIN ZH-548 M12)		2		2		
	10000482	Rift Valley fever virus ZH501	2		2			
	11241	Rinderpest virus	1	2	1	2		
	11243	Rinderpest virus (strain L)	6		6			
	36409	Rinderpest virus (strain RBOK)	33	5	33	5		
	10000465	Rinderpest virus LATC	3		3			
	46920	Rio Mamore virus	1	1	1	1		
	37207	Rio Segundo virus	1		1			
X	313596	Robiginitalea biformata HTCC2501				1		1
	1	root	1	1	1	5		4
	314262	Roseobacter sp. MED193		1		1		
	11029	Ross River virus		1		1		
	11032	Ross river virus (STRAIN T48)	3		3			
	10912	Rotavirus		1		1		
	10930	Rotavirus A bovine/B223/G10	2		2			
	72132	Rotavirus G1	1	1	1	1		
X	73036	Rotavirus G3				1		1
	10001216	Rotavirus G3 strain RV-3	1		1			
	11886	Rous sarcoma virus	4	4	4	4		
	11041	Rubella virus	77	92	77	96		4
	11043	Rubella virus (strain M33)	3	11	3	11		
	11045	Rubella virus (strain THERIEN)	20	35	20	35		
	11044	Rubella virus (vaccine strain RA27/3)	2	6	2	6		
	159479	Saaremaa virus		3		9		6
	45709	Sabia virus		874		874		
	64284	Saboya virus		1		1		
	4932	Saccharomyces cerevisiae	15	5	16	9	1	4
	545124	Saccharomyces cerevisiae AWRI1631		2		2		
	285006	Saccharomyces cerevisiae RM11-1a		2		2		
	10381	Saimiriine herpesvirus 2		3		3		
	8030	Salmo salar	1		1			
	590	Salmonella	1		3		2	
	28901	Salmonella enterica		51		52		1
	59203	Salmonella enterica subsp. arizonae		1		1		
	41514	Salmonella enterica subsp. arizonae serovar 62:z4,z23:--		1		4		3
	119912	Salmonella enterica subsp. enterica serovar Choleraesuis		1		2		1
	321314	Salmonella enterica subsp. enterica serovar Choleraesuis str. SC-B67		2		5		3

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	98360	Salmonella enterica subsp. enterica serovar Dublin		2		2		
	149539	Salmonella enterica subsp. enterica serovar Enteritidis		1		2		1
	29477	Salmonella enterica subsp. enterica serovar Essen	1		1			
X	10001569	Salmonella enterica subsp. enterica serovar Gallinarum/pullorum str. 11			1		1	
X	10001570	Salmonella enterica subsp. enterica serovar Gallinarum/pullorum str. 77			1		1	
X	286783	Salmonella enterica subsp. enterica serovar Indiana				1		1
X	10000554	Salmonella enterica subsp. enterica serovar Minnesota R595			2		2	
	596	Salmonella enterica subsp. enterica serovar Muenchen	1		1			
	54388	Salmonella enterica subsp. enterica serovar Paratyphi A		1		1		
X	554290	Salmonella enterica subsp. enterica serovar Paratyphi A str. AKU_12601				2		2
	295319	Salmonella enterica subsp. enterica serovar Paratyphi A str. ATCC 9150		3		3		
X	272994	Salmonella enterica subsp. enterica serovar Paratyphi B str. SPB7				5		5
	605	Salmonella enterica subsp. enterica serovar Pullorum		1		1		
	90370	Salmonella enterica subsp. enterica serovar Typhi	9	22	9	111		89
X	497977	Salmonella enterica subsp. enterica serovar Typhi str. 404ty				1		1
	220341	Salmonella enterica subsp. enterica serovar Typhi str. CT18		2		6		4
	209261	Salmonella enterica subsp. enterica serovar Typhi str. Ty2		1		1		
	90371	Salmonella enterica subsp. enterica serovar Typhimurium	9	64	12	66	3	2
	99287	Salmonella enterica subsp. enterica serovar Typhimurium str. LT2		23		30		7
	48409	Salmonella enterica subsp. enterica serovar Virchow		1		1		
X	59205	Salmonella enterica subsp. houtenae				1		1
X	59202	Salmonella enterica subsp. salamae				1		1
X	10001511	Salmonella 'group A'			1		1	
	599	Salmonella sp.	1		1			
	72590	Salmonella sp. 'group B'			4		4	
X	10001526	Salmonella thompson C1 strain IS40			1		1	
	10000739	Salmonella typhi 620Ty	1		1			
	10000740	Salmonella typhi Ty21a		6		6		
	10000738	Salmonella typhimurium PL5 (O9,12)	1		1			
	10000742	Salmonella typhimurium SH 4809	1		3		2	
	10000743	Salmonella typhimurium SL3261		4		4		
	358400	Sangassou virus		6		6		
	11607	Sapporo rat virus	1	4	1	4		
	82659	Sapporo virus-Manchester		1		1		
	7386	Sarcophaga peregrina		1		1		
	227859	SARS coronavirus	141	226	144	235	3	9
	228407	SARS coronavirus BJ01	25	28	27	28	2	
	228415	SARS coronavirus CUHK-W1	17		17			
	229992	SARS coronavirus Frankfurt 1	6		6	26		26
	627442	SARS coronavirus P2	1		1			
	227984	SARS coronavirus Tor2	171	1965	171	1965		
	228330	SARS coronavirus Urbani	22	42	22	51		9
	6184	Schistosoma bovis	2		2			
	6182	Schistosoma japonicum	36	12	36	12		
	6183	Schistosoma mansoni	41	44	41	44		
	10000385	Schistosoma mansoni Puerto Rico	17	7	17	7		
X	5334	Schizophyllum commune				1		1
	4550	Secale cereale	51		51			
	11033	Semliki forest virus	35	5	35	5		
	11191	Sendai virus	1	7	1	9		2

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	11194	Sendai virus (strain Enders)	7	11	7	11		
	11196	Sendai virus (strain Harris)	2		2			
	302272	Sendai virus (strain Ohita)	1		1			
	11198	Sendai virus (Z)		1		1		
	11608	Seoul virus		1		2		1
	12557	Seoul virus 80-39		1		1		
	11610	Seoul virus SR11	1		1			
	44026	Sepik virus		1		1		
	8570	Serpentes	4		4			
	615	Serratia marcescens		1		3		2
	399741	Serratia proteamaculans 568		1		1		
	4182	Sesamum indicum	11		11			
X	325240	Shewanella baltica OS155				2		2
X	319224	Shewanella putrefaciens CN-32				1		1
X	351745	Shewanella sp. W3-18-1				1		1
	621	Shigella boydii		1		2		1
X	344609	Shigella boydii CDC 3083-94				17		17
	300268	Shigella boydii Sb227		2		4		2
	622	Shigella dysenteriae	5	2	5	2		
X	358708	Shigella dysenteriae 1012				1		1
X	300267	Shigella dysenteriae Sd197				4		4
	10000748	Shigella dysenteriae serotype 1	2		3		1	
	10000749	Shigella dysenteriae serotype 1 114Sd	1		1			
	623	Shigella flexneri	71	10	72	10	1	
X	591020	Shigella flexneri 2002017				1		1
	42897	Shigella flexneri 2a	26		31		5	
X	198215	Shigella flexneri 2a str. 2457T				2		2
X	198214	Shigella flexneri 2a str. 301				3		3
	10000750	Shigella flexneri 2b	1		1			
	424717	Shigella flexneri 3a	2		2			
	373384	Shigella flexneri 5 str. 8401		2		19		17
	424718	Shigella flexneri 5a	2		2			
	10000752	Shigella flexneri 5b	1		1			
	10000754	Shigella flexneri X	1		1			
	424720	Shigella flexneri Y	6		6			
	624	Shigella sonnei		1		1		
	300269	Shigella sonnei Ss046		3		9		6
	92652	Shrimp white spot syndrome virus	3		3			
	11723	Simian immunodeficiency virus		154		154		
	11711	Simian immunodeficiency virus - mac		1		1		
	10000501	Simian immunodeficiency virus - mac - mac 239		279		279		
	10000502	Simian immunodeficiency virus - mac - mac 32H		7		7		
	10000503	Simian immunodeficiency virus - mac - mac BK28		3		3		
	10000504	Simian immunodeficiency virus - mac - mac F965		1		1		
	31682	Simian immunodeficiency virus - mac1A11		8		8		
X	224206	Simian immunodeficiency virus - mon				1		1
	10000506	Simian immunodeficiency virus - sm - sm PT573		1		1		
	31683	Simian immunodeficiency virus - stm		3		3		
	11737	Simian immunodeficiency virus (F236/SMH4 ISOLATE) (SOOTY MANGABEY)		7		7		
	11735	Simian immunodeficiency virus (K6W ISOLATE)		72		72		
	11736	Simian immunodeficiency virus (K78 ISOLATE)		13		13		
	11733	Simian immunodeficiency virus (MM142-83 ISOLATE)		50		50		

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	11734	Simian immunodeficiency virus (MM251 ISOLATE)		1		1		
	160753	Simian immunodeficiency virus 17E-Fr		2		2		
	11942	Simian retrovirus 1	1		1			
	10923	Simian rotavirus A/SA11	8	4	8	5		1
	37137	Simian rotavirus A/SA11-both	20	2	20	2		
	10633	Simian virus 40	13	40	14	42	1	2
	31608	Simian virus 5 (isolate canine/CPI+)	1		1			
	57667	Simian-Human immunodeficiency virus		1		1		
	37705	Sin Nombre virus	1	4	1	12		8
	10000544	Sin Nombre virus NM H10	4	8	4	8		
	3728	Sinapis alba	2		2			
	11034	Sindbis virus	9		9			
	366394	Sinorhizobium medicae WSM419		1		1		
	382	Sinorhizobium meliloti		1		1		
X	266834	Sinorhizobium meliloti 1021				2		2
	254355	Small ruminant lentivirus	1		1			
	10001307	Small ruminant lentivirus strain It-561		1		1		
	10001308	Small ruminant lentivirus strain It-Pi1		1		1		
	11780	Snyder-Theilen feline sarcoma virus		1		1		
	28375	Soil-borne wheat mosaic virus		1		1		
	4081	Solanum lycopersicum	2		3		1	
	11307	Sonchus yellow net virus		1		1		
X	286542	Soochong virus-2				1		1
	11984	Southampton virus	1		1			
X	58024	Spermatophyta				1		1
X	185949	Sphingomonas aurantiaca				1		1
	2133	Spiroplasma citri		1		1		
	240426	Squirrel poxvirus		1		1		
	11080	St. Louis encephalitis virus	2		2			
	11081	St. Louis encephalitis virus (strain MS1-7)	7		7			
	1279	Staphylococcus	1		1			
	1280	Staphylococcus aureus	85	33	86	35	1	2
	93062	Staphylococcus aureus subsp. aureus COL	17	1	17	1		
	282458	Staphylococcus aureus subsp. aureus MRSA252	40		40			
X	158879	Staphylococcus aureus subsp. aureus N315				4		4
X	93061	Staphylococcus aureus subsp. aureus NCTC 8325				1		1
	1301	Streptococcus	2		2			
	1311	Streptococcus agalactiae	1		1			
X	208435	Streptococcus agalactiae 2603V/R				1		1
X	211110	Streptococcus agalactiae NEM316				1		1
	1317	Streptococcus downei	2	1	2	1		
	119602	Streptococcus dysgalactiae subsp. equisimilis	21		21			
	1336	Streptococcus equi	17		22	14	5	14
	1309	Streptococcus mutans	137	87	140	87	3	
	10000974	Streptococcus mutans GS-5		14		14		
	10000773	Streptococcus mutans MT 8148	39	11	40	12	1	1
	10000813	Streptococcus mutans SJ	1	1	1	1		
	1303	Streptococcus oralis	12		12			
	1313	Streptococcus pneumoniae	9	2	11	3	2	1
	373153	Streptococcus pneumoniae D39		1		2		1
X	170187	Streptococcus pneumoniae TIGR4				2		2
	1314	Streptococcus pyogenes	93	91	105	103	12	12
	10000775	Streptococcus pyogenes 156	1		1			

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	10000776	Streptococcus pyogenes 88/25	1		1			
	10000777	Streptococcus pyogenes 88/30	1		1			
	10000778	Streptococcus pyogenes 88/544	1		1			
	10000779	Streptococcus pyogenes 90/85	1		1			
	10000781	Streptococcus pyogenes BSA10	3		3			
	160490	Streptococcus pyogenes M1 GAS	12		12	1		1
	286636	Streptococcus pyogenes MGAS10394		1		1		
	10000782	Streptococcus pyogenes NS1	1		1			
	10000783	Streptococcus pyogenes NS14	1		1			
	10000784	Streptococcus pyogenes NS27	1		1			
	10000785	Streptococcus pyogenes NS5	1		1			
	10000786	Streptococcus pyogenes serotype M11	2		2			
	342023	Streptococcus pyogenes serotype M12	3	1	3	1		
	10000780	Streptococcus pyogenes serotype M12 A374	1		1			
	10000787	Streptococcus pyogenes serotype M13	1		1			
	301451	Streptococcus pyogenes serotype M18	1		1			
	404330	Streptococcus pyogenes serotype M2	2		2			
	10000788	Streptococcus pyogenes serotype M22	1		1			
	10000789	Streptococcus pyogenes serotype M24	6	2	6	2		
	301448	Streptococcus pyogenes serotype M3	1		1			
	10000542	Streptococcus pyogenes serotype M3 D58	2		2			
	10000790	Streptococcus pyogenes serotype M30	1		1			
	404331	Streptococcus pyogenes serotype M4	2		2			
	10000791	Streptococcus pyogenes serotype M41	1		1			
	301452	Streptococcus pyogenes serotype M49	1		1			
	301449	Streptococcus pyogenes serotype M5	53	53	56	57	3	4
	10000792	Streptococcus pyogenes serotype M52	1		1			
	10000793	Streptococcus pyogenes serotype M54	1		1			
	10000794	Streptococcus pyogenes serotype M55	1		1			
	10000795	Streptococcus pyogenes serotype M57	1		1			
	301450	Streptococcus pyogenes serotype M6	9	7	9	7		
	10000824	Streptococcus pyogenes serotype M6 strain D471	4		4			
	10000796	Streptococcus pyogenes serotype M60	1		1			
	10000797	Streptococcus pyogenes serotype M75	1		1			
	10000798	Streptococcus pyogenes serotype M8	1		1			
	410069	Streptococcus pyogenes serotype M80	1		1			
	160491	Streptococcus pyogenes str. Manfredo	3	17	3	34		17
X	1305	Streptococcus sanguinis					1	1
X	10001439	Streptococcus sanguinis strain BD113-20					6	6
	246202	Streptococcus sobrinus 6715	4	4	4	4		
	36470	Streptococcus sp. 'group A'	11		11			
	1307	Streptococcus suis	1		1			
	1349	Streptococcus uberis	5		5			
X	1902	Streptomyces coelicolor					2	2
X	100226	Streptomyces coelicolor A3(2)					1	1
X	1946	Streptomyces kasugaensis					1	1
	80859	Streptomyces ribosidificus		1		1		
	465541	Streptomyces sp. Mg1		1		1		
	35297	Striped Jack nervous necrosis virus	1		1			
X	7726	Styela plicata					6	6
	186540	Sudan ebolavirus		261		261		
	128948	Sudan ebolavirus - Boniface (1976)		1		1		
	128949	Sudan ebolavirus - Maleo (1979)		30		30		

NEW 2010	ORGANISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	386033	Sudan ebolavirus - Uganda (2000)		6		6		
	10345	Suid herpesvirus 1	2	2	2	2		
	10349	Suid herpesvirus 1 (strain NIA-3)	4		4			
	33703	Suid herpesvirus 1 strain Kaplan	3		3			
	9823	Sus scrofa	7	5	55	26	48	21
	9825	Sus scrofa domesticus		3		4		1
X	10000187	Sus scrofa Landrace X Large White				1		1
	12075	Swine vesicular disease virus	1		1			
	12076	Swine vesicular disease virus (STRAIN H/3 '76)	5		5			
	12077	Swine vesicular disease virus (STRAIN UKG/27/72)	3		3			
	10000809	Swine vesicular disease virus ITL/1/66	1		1			
	10000842	Swine vesicular disease virus NET/1/92	1		1			
	10001002	Swine vesicular disease virus SPA/1/93	16		16			
	10276	Swinepox virus		1		1		
	32049	Synechococcus sp. PCC 7002		1		1		
	1148	Synechocystis sp. PCC 6803		1		2		1
	32630	synthetic construct		3		6		3
X	215169	Synthetic plasmid pMOL98					1	1
	11631	Tacaribe virus		105		106		1
	31616	Tacaribe virus strain V7		1		1		
	6207	Taenia crassiceps	7	1	11	1	4	
	10000802	Taenia crassiceps Strain ORF	3	3	3	3		
	6206	Taenia saginata	5		5			
	6204	Taenia solium	8	4	8	4		
	127999	Tanacetum parthenium		1		1		
	425088	Tanganya virus		1		1		
X	28871	Taterapox virus					2	2
	99883	Tetraodon nigroviridis		1		1		
	5874	Theileria annulata		1		1		
	5875	Theileria parva	28		28			
	333668	Theileria parva strain Muguga	14	15	14	15		
	10000848	Theileria sergenti Type B1		1		1		
	10000849	Theileria sergenti Type B2		2		2		
	10000850	Theileria sergenti Type C		5		5		
	10000851	Theileria sergenti Type I		1		1		
	12124	Theiler's encephalomyelitis virus		16		18		2
	12125	Theiler's encephalomyelitis virus (STRAIN BEAN 8386)	16	52	16	52		
	12126	Theiler's encephalomyelitis virus (STRAIN DA)	3	2	3	2		
	10000855	Theiler's murine encephalomyelitis virus (strain BeAn 8386) (variant M2)					1	1
	204711	Theilovirus		5		9		4
	360549	Themiste hennahi	1		1			
	10479	Thermoproteus tenax virus 1	1		1			
	271	Thermus aquaticus	1		1			
X	274	Thermus thermophilus					1	1
	13286	Theromyzon tessulatum	1		1			
	11084	Tick-borne encephalitis virus	14	1	14	2		1
	11087	Tick-borne encephalitis virus (STRAIN SOFJIN)	6		6			
	10000449	Tick-borne encephalitis virus (WESTERN SUBTYPE) - Neudoerfl	2		10		8	
	6887	Tityus serrulatus	92		92			
	223337	Tobacco leaf curl Zimbabwe virus		1		1		
	12242	Tobacco mosaic virus	28	9	36	13	8	4
	12246	Tobacco mosaic virus (strain Dahlemense)	2		2			

NEW 2010	ORGANISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	10001210	Tobacco mosaic virus (strain Ni568)	1	1				
	10001209	Tobacco mosaic virus (strain PM5)	1	1				
	12243	Tobacco mosaic virus (<i>vulgare</i>)	1	1				
	12146	Tomato bushy stunt virus (STRAIN BS-3)	3	3				
	83192	Topografov virus	1	1				
	7787	Torpedo californica	18	137	61	119	61	
	7788	Torpedo marmorata		3	3	3	3	
X	68887	Torque teno virus			5		5	
	5811	Toxoplasma gondii	12	29	12	38		9
	10000353	Toxoplasma gondii 76K	5	5	5	5		
	10000354	Toxoplasma gondii BK	1	1	1	1		
X	507601	Toxoplasma gondii GT1				1		1
X	508771	Toxoplasma gondii ME49				1		1
	383379	Toxoplasma gondii RH	5	11	5	19		8
	11149	Transmissible gastroenteritis virus	9	9				
	10001154	Transmissible gastroenteritis virus MAD88		1		1		
	158	Treponema denticola		1		1		
	160	Treponema pallidum	5	1	5	2		1
	10001021	Treponema pallidum subsp. pallidum (strain Chicago)	18	19		1		
	243276	Treponema pallidum subsp. pallidum str. Nichols	204	9	204	9		
	6334	Trichinella spiralis		1	2	1	2	
X	203124	Trichodesmium erythraeum IMS101				2		2
	5722	Trichomonas vaginalis	2	2				
	5551	Trichophyton rubrum	2	27	2	27		
	3677	Trichosanthes kirilowii	1		1			
	5554	Trichosporon cutaneum		1				
X	9337	Trichosurus vulpecula			45		45	
	88087	Trimeresurus flavoviridis	3	3				
	4565	Triticum aestivum	273	239	284	244	11	5
	279889	Triticum aestivum var. arduini		4		4		
	4567	Triticum turgidum subsp. durum		1		1		
	5691	Trypanosoma brucei	4	6	4	6		
	31286	Trypanosoma brucei rhodesiense		9		9		
	5693	Trypanosoma cruzi	113	321	113	323		2
	10000347	Trypanosoma cruzi Dm28c	1		1			
	353153	Trypanosoma cruzi strain CL Brener	2		2			
	37133	Tula virus	1	3	1	9		6
X	37347	Tupaia belangeri				1		1
	63673	Turbo cornutus	2		2			
	10001158	unclassified Poxviridae Mule deer poxvirus		1		1		
	11309	unidentified influenza virus		12		12		
X	237631	Ustilago maydis 521				1		1
	10245	Vaccinia virus		298		415		117
	332193	Vaccinia Virus Acambis 3000 MVA		2		2		
	126794	Vaccinia virus Ankara		17		21		4
	10001027	Vaccinia virus Connaught	1		1			
	10249	Vaccinia virus Copenhagen	31	472	31	511		39
X	502057	Vaccinia virus GLV-1h68				1		1
	10251	Vaccinia virus IHD-J	1		1			
	10248	Vaccinia virus LC16M8		4		4		
	31531	Vaccinia virus L-IPV		2		8		6
	10000388	Vaccinia virus NYCBH - Dryvax		29		29		
	10253	Vaccinia virus Tian Tan		3		15		12

NEW 2010	ORGANISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	10254	Vaccinia virus WR	25	5306	25	5333		27
X	10247	Vaccinia virus WR 65-16				4		4
	12870	Variola major virus		11		16		5
	10000390	Variola major virus India-1967		1		1		
	10255	Variola virus		176		223		47
X	11036	Venezuelan equine encephalitis virus				1		1
	36382	Venezuelan equine encephalitis virus (strain 3880)		1		1		
	11037	Venezuelan equine encephalitis virus (strain TC-83)	21		22		1	
	11038	Venezuelan equine encephalitis virus (strain Trinidad donkey)	15		15			
	7742	Vertebrata	8		8			
	11277	Vesicular stomatitis Indiana virus	6	7	6	7		
	11278	Vesicular stomatitis Indiana virus strain Glasgow		1		1		
	11279	Vesicular stomatitis Indiana virus strain Mudd-Summers	1		1			
	11285	Vesicular stomatitis Indiana virus strain San Juan		1		1		
	11280	Vesicular stomatitis New Jersey virus		1		1		
	11276	Vesicular stomatitis virus		6		6		
	7444	Vespa basalis	1		1			
	7453	Vespula maculifrons	1		1			
	7454	Vespula vulgaris		36		36		
	314288	Vibrio alginolyticus 12G01		3		3		
	666	Vibrio cholerae	33	81	34	86	1	5
X	412966	Vibrio cholerae 1587				1		1
X	412614	Vibrio cholerae 2740-80				28		28
	44104	Vibrio cholerae 569B	37		37			
X	404974	Vibrio cholerae AM-19226				1		1
X	417400	Vibrio cholerae B33				1		1
X	412967	Vibrio cholerae MAK 757				5		5
X	345072	Vibrio cholerae MO10				1		1
X	417399	Vibrio cholerae NCTC 8457				1		1
	127906	Vibrio cholerae O1	8		9		1	
	686	Vibrio cholerae O1 biovar El Tor	3	1	3	1		
X	243277	Vibrio cholerae O1 biovar El Tor str. N16961				1		1
	10000567	Vibrio cholerae O1 serotype Inaba	4		4			
	10000568	Vibrio cholerae O1 serotype Ogawa	5		5			
	345073	Vibrio cholerae O395		3		4		1
X	345074	Vibrio cholerae RC385				1		1
	345075	Vibrio cholerae V51		1		1		
	345076	Vibrio cholerae V52		24		31		7
	670	Vibrio parahaemolyticus		57		57		
X	563773	Vibrio parahaemolyticus AN-5034				3		3
	419109	Vibrio parahaemolyticus AQ3810		23		31		8
X	627611	Vibrio parahaemolyticus K5030				12		12
	223926	Vibrio parahaemolyticus RIMD 2210633	2	8	2	16		8
X	70203	Vibrio phage fs1				1		1
X	150340	Vibrio sp. Ex25				9		9
X	314291	Vibrio splendidus 12B01				2		2
X	575788	Vibrio splendidus LGP32				1		1
	672	Vibrio vulnificus	1	64	1	66		2
	216895	Vibrio vulnificus CMCP6		35		49		14
	196600	Vibrio vulnificus YJ016		24		55		31
	8704	Vipera ammodytes	1		1			
	11288	Viral hemorrhagic septicemia virus (STRAIN 07-71)	11	1	11	1		
	3972	Viscum album	17		17			

NEW 2010	ORGANISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	11743	Visna lentivirus (strain 1514 / clone LV1-1KS1)	3		4		1	
	11742	Visna lentivirus (strain 1514)	1		1			
	11741	Visna/Maedi virus	5	7	5	7		
	12750	Visna/maedi virus EV1	1		1			
	36374	Visna/maedi virus EV1 KV1772	2		2			
	29760	Vitis vinifera	1		1			
	74537	Vladivostok virus	1		1			
X	9627	<i>Vulpes vulpes</i>			1		1	
	148360	Watermelon mosaic virus 2 (isolate USA)	2		2			
	43141	Watermelon silver mottle virus	3		3			
	11082	West Nile virus	16	332	17	339	1	7
	10000447	West Nile virus 3000.0259	2		2			
	10000971	West Nile virus NY-99	7	19	11	19	4	
	449278	West Nile virus SPU116/89			1		1	
	10001047	West Nile virus strain 2741	32		32			
	307044	West Nile virus strain 385-99	4	12	4	12		
	406269	West Nile virus strain PTRoxo			12		12	
X	11039	Western equine encephalomyelitis virus					1	1
	46919	Whitewater Arroyo virus			344		344	
	4963	Williopsis saturnus var. mrankii	1	1	1	1		
	66077	Wolbachia sp. wMel			1		1	
	35269	Woodchuck hepatitis virus	5	2	5	2		
	10430	Woodchuck hepatitis virus 1	3		3			
	341946	Woodchuck hepatitis virus 2		4		4		
	10433	Woodchuck hepatitis virus 8			24		34	10
	280	Xanthobacter autotrophicus			1		1	
X	78245	Xanthobacter autotrophicus Py2					1	1
	8364	Xenopus (Silurana) tropicalis			1		1	
	8355	Xenopus laevis			2		3	1
X	132475	Yaba-like disease virus					3	3
	11089	Yellow fever virus			219		235	16
	11090	Yellow fever virus 17D			121		145	24
	617102	Yellow fever virus 17D/Tiantan	4		4			
	31641	Yellow fever virus 1899/81			12		21	9
	407141	Yellow fever virus isolate Ethiopia/Couma/1961			1		1	
	407134	Yellow fever virus strain Ghana/Asibi/1927			5		5	
X	349968	<i>Yersinia bercovieri</i> ATCC 43970					5	5
	630	<i>Yersinia enterocolitica</i>	3	17	5	23	2	6
X	34054	<i>Yersinia enterocolitica</i> (type O:8)					1	1
	393305	<i>Yersinia enterocolitica</i> subsp. <i>enterocolitica</i> 8081			12		15	3
X	349966	<i>Yersinia frederiksenii</i> ATCC 33641					1	1
X	28152	<i>Yersinia kristensenii</i>					1	1
X	349967	<i>Yersinia mollaretii</i> ATCC 43969					3	3
	632	<i>Yersinia pestis</i>	14	125	15	134	1	9
	10000757	<i>Yersinia pestis</i> 195/P	5		5			
X	349746	<i>Yersinia pestis</i> Angola					3	3
	360102	<i>Yersinia pestis</i> Antiqua			8		29	21
	412420	<i>Yersinia pestis</i> CA88-4125			21		21	
	214092	<i>Yersinia pestis</i> CO92			4	1	19	15
X	375450	<i>Yersinia pestis</i> FV-1					2	2
	187410	<i>Yersinia pestis</i> KIM 10			2		8	6
	10000756	<i>Yersinia pestis</i> KIM 5			21		21	
	377628	<i>Yersinia pestis</i> Nepal516			7		110	103

NEW 2010	ORGAN- ISM ID	SPECIES/STRAIN	B-09	T-09	B-10	T-10	Δ B	Δ T
	386656	<i>Yersinia pestis</i> Pestoides F		1		1		
	633	<i>Yersinia pseudotuberculosis</i>		1		1		
X	349747	<i>Yersinia pseudotuberculosis</i> IP 31758				7		7
X	10001504	<i>Yokenella regensburgei</i> PCM 2476			1		1	
X	10001505	<i>Yokenella regensburgei</i> PCM 2477			1		1	
	186538	Zaire ebolavirus	778		811		33	
	129000	Zaire ebolavirus - Eckron (Zaire, 1976)		12		12		
	128947	Zaire ebolavirus - Gabon (1994-1997)		18		18		
	128952	Zaire ebolavirus - Mayinga (Zaire, 1976)	13	14	13	14		
	128951	Zaire ebolavirus - Zaire (1995)		1		1		
	4577	<i>Zea mays</i>	2		2			
	34245	<i>Zinnia violacea</i>		1		1		
	157914	<i>Ziziphus mauritiana</i>	4		4			

2 Website Features

IEDB 2.4 was released on 8 July 2010 and IEDB 2.5 was released on 29 October 2010. In addition, new versions of the Analysis Resource were introduced on June 24 (Version 2.3) and November 5 (version 2.4). The features introduced since the last Annual Compendium include:

- Epitope Structure section on home page search has been redesigned to utilize radio boxes and the Molecule Finder.
- Molecule Finder has been updated to include tree navigation/selection functionality incorporating peptidic and non-peptidic hierarchies.
- Auto-complete feature has been added to home page search for all Finder input fields.
- Allergen tree in Source Organism Finder has been updated to display only nodes referenced by curated data (eliminating empty searches). Additionally, all nodes in the Allergen tree are now selectable.
- Links have been added to PATRIC using NCBI tax IDS.
- Landing page has been created for each NCBI tax ID.
- Related References tab has been added to the Reference Information page.
- ChEBI ID Search has been added to the Identifier Search page.
- XML links have been added to assays with structure data on assay information pages.
- Database Export page has been split into separate pages for current and archived versions.
- New Clustering feature added to Epitope List pages.
- Submission PDFs are now viewable on the Reference Information page (if they exist).
- Query Tutorial added which is accessible from home page and Support links.
- Assay Type labels changed throughout site to use new “Method/Technique” terminology.
- Messages added to Molecule Finder to indicate Molecule Finder is loading and searches are suppressed until Molecule Finder is completely loaded.
- Date headers removed from News section.
- NetMHCpan integrated into the MHC class I binding prediction tool, allowing the prediction for over 1,650 alleles including
- NetMHCIIpan integrated into the MHC class II binding prediction tool, allowing the prediction for over 500 HLA-DR alleles
- MHC class I and class II binding prediction interfaces have been overhauled to allow the simultaneous prediction for multiple alleles and to simplify use
- Combined processing predictor has been updated to reflect changes to class I tool

The IEDB website functionality can be divided into five categories—Browse, Search, Tools, Support, and More IEDB, which largely correspond to the pull-down menus at the top of the home page. The subsections that follow describe the website features within these categories.

The reader will find it helpful to keep in mind how data are stored in the IEDB. Each item contained in the IEDB consists of a reference (article or submission) containing information about one or more epitopes and associated binding or response information. The same epitope can exist in multiple references.

2.1 Home Page

The IEDB Home Page is the default screen displayed when users enter the IEDB system. Besides providing a general description of the IEDB project, the home page displays system level status and notification of scheduled updates or maintenance. As one can see in Figure 2.1, the page is divided into three columns that contain a basic search capability, introductory information, Summary Metrics of the data, available Resources, and project-related News. As users browse the IEDB system, they can return to the home page anytime by clicking *Home* on the far left of the main menu bar.

Summary Metrics are displayed in the center column of the screen. These numbers are intended to be a gauge of the volume of data available in the system. The Summary Metrics provide the number of epitopes (peptidic and non-peptidic), assays (T cell, B cell, MHC ligand elution, and MHC binding), epitope source organisms, restricting MHC alleles, and references. Clicking on the “See all Metrics” brings up a Summary Metrics page that shows the positive, negative, and total (all) count for epitopes and assays, and the counts for epitope source organism, host organism, restricting MHC allele, and references. Further detail is revealed when the user clicks on the counts.

The screenshot shows the IEDB 2.5 Home page with a blue header bar containing the logo and the text "IMMUNE EPITOPE DATABASE AND ANALYSIS RESOURCE". Below the header is a navigation bar with links for Home, Browse, Search, Tools, Support, More IEDB, and a Keyword Search bar with a "Search" button. The main content area is divided into three columns:

- Search:** Contains sections for "Epitope Structure" (radio buttons for Any, Linear Peptide, Discontinuous Peptide, Non-Peptidic; input fields for Linear Peptide and Non-Peptidic, and "Molecule Finder" buttons), "Epitope Source" (input fields for Source Organism and Source Antigen, and "Organism Finder" and "Molecule Finder" buttons), and "Immune Recognition Context" (checkboxes for B Cell Response, T Cell Response, MHC Binding, MHC Ligand Elution, and a "Host Organism" dropdown with "Organism Finder" button). There is also a "Search" and "Clear" button, and a link to "Help With Common Queries".
- Welcome!**: A red-bordered box contains the text "Swine Flu Analysis: The IEDB has performed an analysis of swine flu epitopes. Click [here](#) for details." Below this is a section about the IEDB's analysis of swine flu epitopes, mentioning antibody and T cell epitopes for humans, non-human primates, rodents, and other animal species. It notes curation of peptidic epitope data for infectious diseases, including NIAID Category A, B, and C priority pathogens, and emerging and re-emerging infectious diseases, allergens, and autoimmune diseases current through June 2010.
- Summary Metric**: A table showing the count of various metrics:

Summary Metric	Count
Peptidic Epitopes	79778
Non-Peptidic Epitopes	789
T Cell Assays	154709
B Cell Assays	118588
MHC Ligand Elution Assays	2665
MHC Binding Assays	201123
Epitope Source Organisms	2631
Restricting MHC Alleles	579
References	11833

[See all Metrics](#)
- Resources**: A section titled "Resources" with a "Resources" icon. It says "We have provided a variety of resources to analyze our data and enhance your IEDB experience:" followed by a list of links:
 - T Cell Epitope Prediction
 - B Cell Epitope Prediction
 - Epitope Analysis Tools
 - Database Export
 - IEDB Ontology
 - Data Field Descriptions
- News**: A section titled "News" with a "News" icon. It includes a "Newsletters" section with links to November 2010, August 2010, June 2010, March 2010, October 2009, and July 2009 newsletters, and a "more" link. It also includes sections for "Publications", "Upcoming Events", "Meta-Analyses", "Compendia", and "Release Notes".

At the bottom of the page are links for "Provide Feedback", "Help Request", and "Solutions Center", and a note that it is supported by a contract from the National Institute of Allergy and Infectious Diseases. The page is last updated on February 07, 2011.

Figure 2.1 IEDB 2.5 Home page

The rightmost column has links to a variety of resources, including epitope prediction and analysis tools, database export, the IEDB ontology, and documentation describing data fields. These resources can also be accessed via the pull-down menus on the page. The News panel shows six categories – Newsletters, Publications, Upcoming Events, Meta-analyses, Compendia, and Release Notes. Each category can be expanded to show its linkable content, as shown in the figure for Newsletters.

2.2 Browse

The IEDB allows users to browse for records in three different ways – by MHC allele, by source species, and by the new 3D structure.

2.2.1 Browse Records by MHC Allele

All users can find records associated with a specific MHC allele by browsing records by allele.

The screenshot shows a hierarchical tree structure for browsing MHC alleles. The root node is 'MHC Alleles' (108 structure(s)*). It branches into 'Class I' (4614 structure(s)*), which further branches into 'Mamu-A' and 'Mamu-B'. 'Mamu-A' has sub-nodes for A*01 through A*11. 'Mamu-B' has sub-nodes for B*01 through B*17. Other nodes include 'Class II' (108 structure(s)*), 'sheep (Ovis aries)' (8 structure(s)*), and 'cattle (Bos taurus)' (354 structure(s)*). A note at the bottom states: '* Indicates the number of distinct molecular structures tested in the context of the selected group of MHC molecules.'

- + bonobo (*Pan paniscus*) (1 structure(s)*)
- + cattle (*Bos taurus*) (354 structure(s)*)
- + chicken (*Gallus gallus*) (37 structure(s)*)
- + chimpanzee (*Pan troglodytes*) (705 structure(s)*)
- + cotton-top tamarin (*Saguinus oedipus*) (2 structure(s)*)
- + dog (*Canis lupus familiaris*) (27 structure(s)*)
- + gorilla (*Gorilla gorilla*) (15 structure(s)*)
- + horse (*Equus caballus*) (75 structure(s)*)
- + human (*Homo sapiens*) (41459 structure(s)*)
- + mouse (*Mus musculus*) (15493 structure(s)*)
- + pig (*Sus scrofa*) (25 structure(s)*)
- + rat (*Rattus norvegicus*) (114 structure(s)*)
- rhesus macaque (*Macaca mulatta*) (4721 structure(s)*)
 - Class I (4614 structure(s)*)
 - Mamu-A*01 (1972 structure(s)*)
 - Mamu-A*02 (595 structure(s)*)
 - Mamu-A*08 (1 structure(s)*)
 - Mamu-A*11 (913 structure(s)*)
 - Mamu-B*01 (630 structure(s)*)
 - Mamu-B*03 (32 structure(s)*)
 - Mamu-B*04 (9 structure(s)*)
 - Mamu-B*08 (10 structure(s)*)
 - Mamu-B*17 (1360 structure(s)*)
 - + Class II (108 structure(s)*)
- + sheep (*Ovis aries*) (8 structure(s)*)

* Indicates the number of distinct molecular structures tested in the context of the selected group of MHC molecules.

Figure 2.2 Browse by MHC Allele interface

To browse records by allele, the user selects *Browse by MHC Allele* under the *Browse* heading on the main menu. The interface for the *Browse by MHC Allele* is a tree structure that makes it easy for users to find and investigate information on specific MHC alleles. As Figure 2.2 shows, the tree structure expands (and collapses) so users can drill down on species, MHC type, and allele to find the number of records in the IEDB for their MHC allele of interest. This number serves as a link that will display the records associated with the selected allele.

2.2.2 Browse Records by Source Organism

Users can find records associated with a specific epitope source species by browsing records by source organism. To do this, the user accesses the *Browse by Source Organism* page via the *Browse* pull-down menu. The interface for the *Browse by*

Source Organism is a tree structure that expands (and collapses) so users can drill down on species as Figure 2.3 shows. Each entry in the tree indicates the number of distinct molecular structures tested from the selected group of source organisms. This number serves as a link that will display the records associated with the selected source organism. As one mouses over an entry, a descriptive balloon appears. As seen in the figure, “biota” appears to describe “cellular organisms”. The user can also search for names. For example, a search for “human”, as shown in Figure 2.4, generates a table of responses for the term and corresponding synonyms. If the user clicks on [Details], Source Organism Information is displayed. The page for the first result, *Homo sapiens*, is displayed in Figure 2.5, and includes the NCBI taxonomy identifier numbers and information on epitope structure, relevant literature and submission references, source antigens, and assays. If the user clicks on [Highlight] the tree expands and highlights the location. In the case of “*Homo sapiens*”, the tree expands to over 20 levels. Clicking on the number structures displays the same Source Organism Information as seen in Figure 2.5.

Browse By Source Organism

Search

- Species
 - Viruses (63586 structure(s)*)
 - unclassified phages (3 structure(s)*)
 - dsDNA viruses, no RNA stage (24272 structure(s)*)
 - dsRNA viruses (305 structure(s)*)
 - ssRNA viruses (32651 structure(s)*)
 - Deltavirus (247 structure(s)*)
 - ssDNA viruses (604 structure(s)*)
 - Retro-transcribing viruses (5507 structure(s)*)
 - other sequences (13 structure(s)*)
 - artificial sequences (7 structure(s)*)
 - plasmids (6 structure(s)*)
 - cellular organisms (43672 structure(s)*)
 - Bacteria (17037 structure(s)*)
 - Archaea (6 structure(s)*)
 - Eukaryota (26631 structure(s)*)

* Indicates the number of distinct molecular structures tested from the selected group of source organisms.

Figure 2.3 Browse by Source Organism interface

Browse By Source Organism

Search Results

438 item(s) found, displaying 1 to 5 (Click the column headers to adjust the sorting)

« previous
1
2
3
4
5
6
7
8
9
...
87
88
next »
Go To »
1

Organism Name	Matched Value
Homo sapiens (ID:9606) [Details] [Highlight]	Genbank common name: human
Human respiratory syncytial virus (ID:11250) [Details] [Highlight]	Synonym: human RSV
Human rotavirus A (ID:10941) [Details] [Highlight]	Scientific name: Human rotavirus A
Rotavirus G4 (ID:35336) [Details] [Highlight]	Synonym: Human rotavirus 4
Rotavirus G9 (ID:42567) [Details] [Highlight]	Synonym: Human rotavirus 9

438 item(s) found, displaying 1 to 5

« previous
1
2
3
4
5
6
7
8
9
...
87
88
next »
Go To »
1

+ Species

Figure 2.4 Using the search feature on the Browse by Source Organism page

Source Organism Information

Source Organism

Source Organism: Homo sapiens
 Source NCBI Taxonomy ID: [9606](#)
 Parent NCBI Taxonomy ID: [9605](#)

Structure (5805) Reference (1138) Source Antigen (878) MHC Binding (7701) B Cell Assay (4422) T Cell Assay (6175) MHC Ligand Elution (392)

5805 item(s) found, displaying 1 to 25 (Click the column headers to adjust the sorting)

« previous
1
2
3
4
5
6
7
8
9
...
232
233
next »
Go To »
1

Export all results: ([compact](#) | [full](#))

Epitope ID ↑	Structure	Source Antigen	Source Organism
10	AAAAAIFVI	MHC class I related protein A	Homo sapiens
11	AAAAALDKKQRNFDFKILA	Myosin-7	Homo sapiens
129	AAEYWNSQKEVLER	HLA class II histocompatibility antigen, DQ	Homo sapiens
155	AAGIGILTIV	Melanoma antigen recognized by T-cells 1	Homo sapiens
156	AAGIGILTIVI	Melanoma antigen recognized by T-cells 1	Homo sapiens
157	AAGIGILTIVLGVL	Melanoma antigen recognized by T-cells 1	Homo sapiens
309	AAMMAEELKKEQDTSAHL	Myosin-7	Homo sapiens
316	AANDPIFVV	L-dopachrome tautomerase precursor	Homo sapiens
363	AAPLILSR	Prostate-specific antigen precursor	Homo sapiens
368	AAPPVAPA	Cellular tumor antigen p53	Homo sapiens

Figure 2.5 Source Organism Information page

2.2.3 Browse by 3D Structure

All users can find records associated with a curated 3D structure by using *Browse by 3D Structure* found under the *Browse* heading on the main menu. The interface for the Browse by 3D Structure is a tree structure that makes it easy for users to find and investigate information on B cell, T cell, and MHC binding structures. As seen in Figure 2.6, the tree structure expands (and collapses) so users can drill down on species, MHC type, and/or allele to find the number of records in the IEDB for their structure of interest. This entry serves as a link that will display the records associated with the selected structure.

Browse By 3D Structure

The B Cell, T Cell, and MHC Binding trees below are organized by the organism that is the source of the antibody, T Cell, and MHC molecule, respectively.

-  B Cell (142 structure(s)*)
 -  Camelus dromedarius (3 structure(s)*)
 -  Cricetinae gen. sp. (2 structure(s)*)
 -  Cricetulus griseus (1 structure(s)*)
 -  Ginglymostoma cirratum (1 structure(s)*)
 -  Homo sapiens (18 structure(s)*)
 -  Lama glama (1 structure(s)*)
 -  Mus musculus (116 structure(s)*)
 -  unidentified (1 structure(s)*)
-  T Cell (28 structure(s)*)
 -  Homo sapiens (16 structure(s)*)
 -  Class I (12 structure(s)*)
 -  Class II (4 structure(s)*)
 -  DR (4 structure(s)*)
 -  HLA-DR1 (3 structure(s)*)
 -  HLA-DR4 (1 structure(s)*)
 -  HLA-DRA*0101/DRB5*0101 (1 structure(s)*)
 -  Mus musculus (12 structure(s)*)
-  MHC Binding (158 structure(s)*)
 -  Gallus gallus (2 structure(s)*)
 -  Homo sapiens (94 structure(s)*)
 -  Mus musculus (60 structure(s)*)
 -  Rattus norvegicus (3 structure(s)*)

* Indicates the number of distinct molecular structures.

Figure 2.6 Browse by 3D Structure interface

2.3 Query

In addition to the Browse functions, there are several other ways users can find information in the IEDB. A keyword search is available on the menu bar on each page. A family of advanced query-by-example searches can be accessed on the Search pull-down menu and consist of “B Cell Search”, “T Cell Search”, “MHC Binding Search”, “Ligand Elution Search”, “Epitope Search”, and “Identifier Search”. This group of queries will be referred to as Detailed Searches in this document. A simplified search that is especially designed for the needs of most

immunologists appears on the home page. This document will refer to this particular query as the Simple Search. All of these methods return results in a common format, as described in Section 2.3.4. These methods are elaborated in the subsections below.

2.3.1 Perform a Keyword Search

The keyword search allows users to locate records in the database using a keyword, identifier, or sequence. Wild card characters '*' and '?' can be used in the search field. The '*' character will match zero or more characters and the '?' character will match exactly one character. Additionally, the operators 'AND', 'OR', and 'NOT' can be utilized. A logical 'OR' is the default option. These three logical operators must be uppercase in order to be identified as operators instead of search terms. In all other cases, the keyword search is case insensitive, so "ABC" is the same as "abc". Regular expressions cannot be used in the search field.

To perform a search, users enter criteria such as a keyword, identifier, or sequence into the text box in the upper right corner of each page next to the "Search" button (Figure 2.1). They then click the Search button and view the Search Result Summary, which is described in Section 2.3.4.

In addition to using the search function from the menu bar, users can submit searches via a properly constructed URL. The example below will perform a search using the URL for records that contain 'dengue'. This would return the equivalent of typing 'dengue' in the search option on the menu bar and clicking the Search button.

http://www.iedb.org/counts_keyword.php?keyword_search_query_term=dengue

2.3.2 Perform a Home Page Search

The search feature available on the left-hand portion of the home page (Figure 2.7) was originally designed to simplify the search process for many commonly asked queries by immunologists. It allows greater control in the search than available in the keyword search and is less involved than the Advanced queries. The ability to specify peptide sequences and the types of searching was enhanced in 2010. Previously the user specified the epitope structure with a pull-down menu. This has been replaced with radio button selections for *Any*, *Linear Peptide*, *Discontinuous Peptide*, and *Non-Peptidic*. The user can prescribe a linear peptide sequence and the type of search to be performed on it. When specifying a linear sequence, a window appears in which the user must enter the sequence and press "Apply". The user can prescribe the type of search desired - an exact match to the entered sequence, a substring search where epitopes are found that contain the sequence within its overall structure, or a homologous peptides at sequence at identity levels of 70%, 80%, or 90% as determined by BLAST. The user can specify a non-peptidic structure with the Molecule Finder. This Molecule Finder is a new feature and differs from the Molecule Finders available on the Advanced Query pages.

Introduced in 2010 is an autocomplete feature implemented for the Molecule, Organism, and Allele Finders. For example, as one types "hem" in the Source Antigen field in the Epitope Source section, several choices start to appear in a list below the text field, including "hemolysin", "hemK protein", and "hemagglutinin". The user can click on a selection from the list or can continue typing to further narrow down choices. Likewise, a user may type "human"

Search

Epitope Structure

Any
 Linear Peptide: Exact Matches
 Discontinuous Peptide
 Non-Peptidic: Molecule Finder

Epitope Source

Source Organism: Organism Finder

 Source Antigen: Molecule Finder

Immune Recognition Context

B Cell Response
 T Cell Response
 MHC Binding
 MHC Ligand Elution



Host Organism: Organism Finder

 MHC Restriction: Allele Finder

 MHC Class:

[◆ Help With Common Queries?](#)

Figure 2.7 Home page Simple Search

finder has tree structures for non-peptidic molecules and peptidic molecules. They are described further in Section 2.3.5.4. The user can decide whether to include B cell responses, T cell responses, MHC binding, and/or MHC ligand elution results in the search (at least one must be checked). The host organism, the MHC restriction, and the MHC class can also be specified with the help of the organism finder and the allele finder (Section 2.3.5.1). The fields using finders will allow multiple selections as search criteria. In these cases the selections are treated as a set. Records will be considered a match if they include at least one of the selected values in the set. The search is executed by selecting the Search button and query results can be viewed on the Search Result Summary page, which is described in Section 2.3.4.

Finally, a link to “Help with Common Queries” was added to the bottom of the Search panel. This links to a series of 16 questions that have been useful in preparing the epitope meta-analyses conducted and published by LIAI. More information about the Common Queries is available in Section 2.5.5.

2.3.3 Advanced query

The Advanced queries are based on a standard Query by Example (QBE) approach, which is a method of forming queries where a user can enter conditions for each data field they want

or “homo” to select “homo sapiens” in the Host Organism field. The list of matches includes scientific names and synonyms. These actions enable users to quickly specify their search parameters without using the finders. The autocomplete applies wherever the input field has “Enter Search” in grey. Use of the autocomplete circumvents the need to use the finders.

If the user decides not to use the autocomplete feature for a search field, they will need to use the finders. The epitope source can be prescribed for the source organism and source antigen by using the organism finder (Section 2.3.5.5) and molecule finder (Section 2.3.5.4), respectively. The molecule finder for the epitope source is again different from the non-peptidic molecule mentioned above and the molecule finders available on the Advanced Query pages. This

included in the query. The Advanced queries allow users to define example criteria for each field in the system. As there are over 300 fields, the Advanced queries are both powerful and comprehensive.

The fully expanded input screen for the Epitope Search is shown in Figure 2.8. The user can search on several reference criteria, such as author name, article title, the IEDB Reference ID, keywords in the reference abstract, year, author affiliation, and reference type. References are either published literature articles that have been curated by the IEDB curation staff or data submitted directly by researchers. Users can also specify search criteria for the epitope, such as epitope type (e.g. peptide from protein, carbohydrate), source molecule and organism, and various reference details. Further explanation of the search terms can be found in the Curation Manual. This query type returns T cell, B cell, MHC binding, and MHC ligand elution data.

Epitope Search

Reference

Author
 Title

Reference Details

Reference ID
 Abstract
 Affiliations
 Date (Year) Equal To
 Reference Type Journal Article Submission Any

Epitope

Type
 Source Molecule Molecule Finder
 Source Organism Organism Finder

Epitope Reference Details

Epitope Structure Defines
 Evidence Code for Epitope Source Antigen
 Epitope Name
 Reference Start Position Equal To
 Reference End Position Equal To
 Reference Region
 Comments
 Location of Data in Reference

Epitope Related Object

Related Object
 Type
 Source Molecule Molecule Finder
 Source Organism Organism Finder

Buttons

Figure 2.8 Epitope Search input screen

There are five other Advanced queries that can be used to select search criteria for references, epitopes, and assays. These five are B Cell Search, T Cell Search, MHC Binding Search, MHC Ligand Elution Search, and Identifier Search. As an example, the B Cell Search input screen is shown in Figure 2.9. As is the case for all four assay search input screens, the Reference and Epitope criteria fields are listed first, followed by the fields specific for each assay type.

B Cell Search

Reference

Epitope

Type

Source Molecule

Source Organism

Epitope Reference Details

Epitope Related Object

Immunization

Host Organism

Host Details

1st In Vivo Process may or may not MUST MUST NOT be present in search results.

1st Immunogen may or may not MUST MUST NOT be present in search results.

2nd In Vivo Process may or may not MUST MUST NOT be present in search results.

2nd Immunogen may or may not MUST MUST NOT be present in search results.

Immunization Comments

Adoptive Transfer may or may not MUST MUST NOT be present in search results.

B Cell Assay

Qualitative Measurement

Assay Type

Measurement Details

Assayed Antibody

Antigen

3D Structure of Complex

Assay Reference Details

Figure 2.9 B Cell Search input screen

The B cell response assay category captures B cell-mediated immunity information and describes antibody responses related to the epitope/antigen. In a B cell response, B lymphocytes (a type of

IEDB Identifiers

Epitope ID

Reference ID

Submission ID

Assay ID

External Identifiers

PubMed ID

PDB ID

ChEBI ID CHEBI:

Search **Clear**

Figure 2.10 Identifier Search input screen

white blood cell) produce proteins called antibodies that bind to antigens. Antibodies are present on the surface on B lymphocytes and are also secreted. Once an antibody binds an antigen, the bound antigen molecule can be engulfed by phagocytes and broken into fragments.

The T Cell Response assay category captures T cell-mediated immunity information where the MHC molecule/antigen complex is recognized by T cells in the context of presentation by an antigen presenting cell.

The MHC Binding assay category captures details relating to the in vitro interaction of the epitope with specific MHC molecules along with available Epitope-MHC complex structure details. In other words, assays in this category assess the epitope's binding capacity to the MHC molecule.

The MHC Ligand Elution assay category captures data related to epitopes that are naturally processed and presented on the surface of an antigen presenting cell. The MHC Ligand Elution category differs from the MHC Binding category in that for the former, antigen bound to the MHC molecule on the cell surface has been taken up and processed internally for presentation by the antigen presenting cell, whereas for the latter, antigen in solution is bound to MHC molecules on the surface of the antigen presenting cells.

The Identifier Search input screen is shown in Figure 2.10. If user know the epitope, reference, submission, or assay identifier they are seeking, they can enter that information into one of the four IEDB Identifiers fields. Users can also look for information based on the PubMed ID of a reference or the PDB ID of a structure by using the External Identifiers fields. In 2010, a new field was added so users can search by ChEBI ID for non-peptidic structures.

2.3.4 Search Results Summary

The Search Results Summary page displays the results of all the different queries. An example is displayed in Figure 2.11, which was generated with the Home page Simple Search for the linear sequence substring "AAA" for human and mouse host organisms. The Search Results Summary allows the user to quickly assess the results of their search and drill down the level of detail that is of interest. As a convenience, the display of Search Parameters was added to the Search Result Summary page, along with the ability to revise the search criteria. Clicking on the "Revise Search" button will take the user back to the original search form, either the home page for the Simple Search or to the appropriate Detailed Search form.

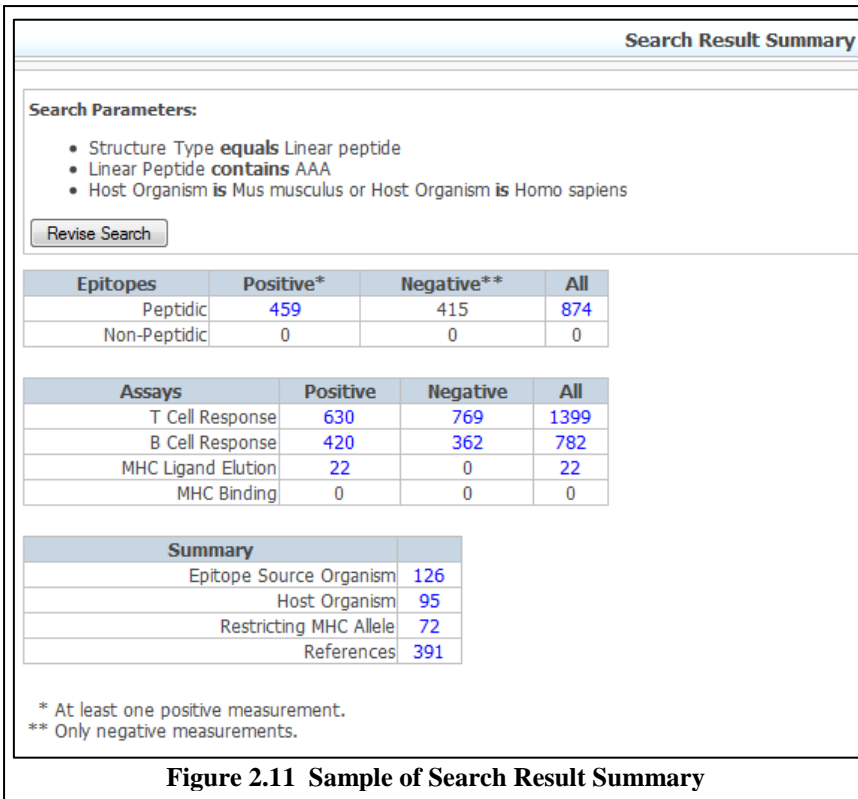


Figure 2.11 Sample of Search Result Summary

the number of corresponding epitope, source antigen, and source species.

As Figure 2.12 and subsequent figures show, comma separated values (CSV) file exports are available on-demand from all list pages and list tabs throughout the site, except for the patent list, which is only provided in PDF format. The CSV file exports are generated in two formats, a compact format that only contains the data presented in the list the user is viewing, and a full record format that contains all of the data associated with the records in the list the user is viewing. On some results pages, only the full format is available. The CSV format can be easily manipulated using a spreadsheet program, such as Microsoft Excel, or edited with a word processing program, such as Microsoft Word or Notepad.

One of the major new features introduced in 2010 is the ability to cluster query results by the sequence identity of the epitope structure. Clustering has been implemented to simplify query results, especially since the amount of data returned in many queries has grown as the amount of data in the IEDB has grown. As an example, if the user would query for all Vaccinia virus epitopes they would get a list of over 10,000 records. If they then hit the ‘Cluster’ button, the epitopes would be grouped according to their sequence identity and the user would end up with a ‘cleaned up’ view of the results.

An algorithm for grouping the sequences has been developed and tested with several IEDB data sets. It groups related sequences based on sequence identity and allows the user to specify the sequence identity threshold from 70%-100%. As an example, at 85% identity, 7,512 Vaccinia virus epitopes form 2,528 clusters. This drastically reduces the number of results displayed to the user and should result in more efficient navigation. In terms of sorting the clusters of

The number peptidic and non-peptidic epitopes are displayed in the topmost table on the Result Summary. Because the IEDB captures both positive and negative assay responses, some “epitopes” lack a positive binding value and therefore are not actually epitopes; the number of these molecular structures is indicated in the Negative column of the table. The user can click on the number in the Positive and All columns to view all the results for that category, as seen in Figure 2.12 for the positive peptidic epitopes. The columns displayed include

epitopes so that the most relevant ones are displayed first, bioinformaticians at LIAI have developed a metric called the ‘evidence score’. This metric takes into account the number of references, number of positive assays, and total number of assays for each epitope. It is used initially to seed the clusters and is summed over all of the epitopes in the cluster to get the ‘cluster score’ (CScore). Clusters are then sorted by the CScore in a descending order by default.

Figure 2.12 shows the initial results page without any clustering. The user can select from four levels of sequence identity for the clustering, 70, 80, 90, or 100%, with 80% being the default (see Figure). Clustering is initiated when the user clicks the *Cluster* button. Figure 2.13 shows clustering by 80% sequence identity. A description of the epitope clustering algorithm and additional screen shots of the clustered results can be found in the IEDB Solutions Center under the *Tutorials and Reference Materials* folder (<http://iedb.zendesk.com/entries/306422-iedb-epitope-clustering-help>). Help can also be reached by clicking on the ‘?’ icon to the right of the % Identify button.

The second table in Figure 2.11 summarizes the positive and negative assay responses for the four general assay types. The user can drill down on any of these values to see the corresponding epitopes. The detail page for the positive T Cell Responses of this search is displayed in Figure 2.14.

The third table lists the number of epitope source organisms, host organisms, restricting MHC alleles, and references resulting from the input search criteria. As before, the user can drill down on the values in the table to find more detail related to each category, as seen in Figure 2.15, Figure 2.16, Figure 2.17, and Figure 2.18.

Epitope					
Cluster	80% Identity	?			
459 item(s) found, displaying 1 to 25 (Click the column headers to adjust the sorting)					
« previous 1 2 3 4 5 6 7 8 9 ... 18 19 next » Go To » 1					
Export all results: [x] (full)					
Epitope ID ↑	Structure	Source Antigen	Source Organism		
10	AAAAAIFVI	MHC class I related protein A	Homo sapiens		
11	AAAALDKKQRNFDKILA	MYH7 protein (1 more)	Homo sapiens		
19	AAAAGLAGLVFPQPPAPIAV	CONSERVED HYPOTHETICAL ALANINE RICH PROTEIN (1 more)	Mycobacterium tuberculosis (1 more)		
33	AAAGAVVGGLGGYMLG	Major prion protein precursor	Mus musculus		
34	AAAGDK	B13 antigen	Trypanosoma cruzi		
35	AAAGFASKTPANQAISMIDG	Phosphate-binding protein ptsS 1 precursor	Mycobacterium tuberculosis		
36	AAAGFMVLQDINCFRPHGVSAAQEKISFGKSSQCREAVGT	Glycoprotein 4	Lelystad virus		
46	AAALEQLLGQTADVA	hypothetical protein ML1057	Mycobacterium leprae (1 more)		
49	AAALPGKCGV	Non-specific lipid-transfer protein 1 (2 more)	Prunus persica		
52	AAANTSDSQKE				
54	AAAQHGHMHGS				
56	AAARTTS	envelope glycoprotein 2	Hepatitis C virus		
59	AAASAIQG	6 kDa early secretory antigenic target	Mycobacterium tuberculosis		
60	AAASAIQGNVTIHSL	6 kDa early secretory antigenic target	Mycobacterium tuberculosis		
84	AADELVGGPPVEASAAL	POSSIBLE phiRv2 PROPHAGE PROTEIN	Mycobacterium tuberculosis		
159	AAGKATTEEQKLIEDINVGFKAAVAAAASVPAA	Pollen allergen Phl p 5b precursor	Phleum pratense		
160	AAGKATTEEQKLIEDINVGFKAAVAAAASVPAA	Pollen allergen Phl p 5b precursor	Phleum pratense		
424	AASARYPNVTIAAAA				
477	AAVAAAASVPAADKFKTFE	Pollen allergen Phl p 5b precursor	Phleum pratense		
707	ADLGDFENSAAAETGVGVIKSIA	Toxin coregulated pilin precursor (1 more)	Vibrio cholerae (1 more)		
805	AEAAAPVVESKWRAL	polyprotein	Hepatitis C virus		
1125	AEPQAEAPAAAASSEQ	brain abundant, membrane attached signal protein 1	Mus musculus		
1266	AFASRGNHVSPTHYVPESDAAR	polyprotein	Hepatitis C virus		
2156	AKAVAAWTLKAAA				
2192	AKFVAAWTLKAAA				

Figure 2.12 Screen generated by drilling down on the positive epitope hyperlink in the Search Result Summary

Epitope Clusters											
Cluster	80% Identity										
Expand/Collapse All											
281 clusters identified at 80% sequence identity, displaying 1 to 25											
« previous 1 2 3 4 5 6 7 8 9 10 11 12 next » <input type="text"/> Go To » 1											
Export all results: [X] (full)											
Cluster	CScore	Count	Epitope	EScore	Source Antigen	Source Organism					
[+] MTEQQWNFAGIEAAASAIQGNVTI	3396	10	MTEQQWNFAGIEAAASAIQG (And More)	1436	6 kDa early secretory antigenic target	Mycobacterium tuberculosis (AND 2 more)					
[+] WNPKSPKPKTNLKHVAGAAAAGAVVGLGGYMLG	2374	11	WNPKSPKPKTNLKHVAGAAAAGAVVGLGGY (And More)	674	Major prion protein precursor	Mus musculus (AND 2 more)					
[+] AFASRGNHVSPTHYVPESDAARVTAILSSL	2199	17	SPTHYVPESDAAR (And More)	290	Genome polyprotein (AND 3 more)	Hepatitis C virus subtype 1a (AND 4 more)					
[+] GDKPPPFQAAAGDKPLF	1821	14	KPPPFQAAAGDKPP (And More)	233	B13 antigen	Trypanosoma cruzi					
[+] WNFAGIEAAASAIQGNVTIHSLLDEGKQS	1511	8	IEAAASAIQGNVTIHSLL (And More)	383	6 kDa early secretory antigenic target	Mycobacterium tuberculosis					
[+] GDKPSLFGQAAAGDKLSLF	1477	11	KPSLFGQAAAGDKLS (And More)	233	B13 antigen	Trypanosoma cruzi					
[+] LDEGKQSLTKLAAAWGGSGSEAYQGVQ	1391	6	EGKQSLTKLAAAWGGSGSEA (And More)	495	6 kDa early secretory antigenic target	Mycobacterium tuberculosis (AND 1 more)					
[-] AAGKATTEEQKLIEDINVGFKAAVAAAASVPAA	1354	4	INVGFKAAVAAAASV	858	Group V allergen Phl p 5.0203 precursor	Phleum pratense					
			AAGKATTEEQKLIEDINVGFKAAVAAA	122	Pollen allergen Phl p 5b precursor	Phleum pratense					
			AAGKATTEEQKLIEDINVGFKAAVAAAASVPAA	155	Pollen allergen Phl p 5b precursor	Phleum pratense					
			MIEKINVGFKAAVAAAAGGP	219	Pollen allergen KBG 60 precursor	Poa pratensis					
[+] CMKKDDQIAAAMVLRGMAKDQFALKD	1301	8	MKKDDQIAAAMVLRGMAKDQFALK (And More)	369	vls recombination cassette Vls10 (AND 4 more)	Borrelia garinii IP90 (AND 3 more)					
[+] MALWMRLPLLALLALWGPDPAAAFVNQHLCGS	1270	7	ALWGPDPAAA (And More)	488	Insulin precursor (AND 2 more)	Homo sapiens					
[+] SLKKGAAALGIGTDSVI	1260	3	KKGAAALGIGTDSVI (And More)	738	Glutamate decarboxylase 2 (AND 1 more)	Homo sapiens					

Figure 2.13 Query results for clustering at 80% sequence identity

T Cell Response Assays															
630 item(s) found, displaying 1 to 25 (Click the column headers to adjust the sorting)															
	« previous	1	2	3	4	5	6	7	8	9	... 25	26	next »	Go To »	1
Export all results: [compact] [full]															
T Cell ID ↑	Reference	Epitope	Host	Immunization	Assay Antigen	Antigen Epitope Relation	MHC Restriction	Assay Description							
2061	Yolanda López-Vidal; Arch Med Res 2004	MTEQQWNFAGIEAAAS 6 kDa early secretory antigenic target (1-16) Mycobacterium tuberculosis	Homo sapiens	Infectious disease via exposure to Mycobacterium tuberculosis (Source Organism)	MTEQQWNFAGIEAAAS 6 kDa early secretory antigenic target (1-16) Mycobacterium tuberculosis	Epitope		3H-thymidine cell proliferation Positive							
2064	Yolanda López-Vidal; Arch Med Res 2004	MTEQQWNFAGIEAAAS 6 kDa early secretory antigenic target (1-16) Mycobacterium tuberculosis	Homo sapiens	Infectious disease via exposure to Mycobacterium tuberculosis (Source Organism) followed by restimulation in vitro	MTEQQWNFAGIEAAAS 6 kDa early secretory antigenic target (1-16) Mycobacterium tuberculosis	Epitope		ELISA cytokine release IL-4 Positive							
2139	Yolanda López-Vidal; Arch Med Res 2004	AAASAIQGNVTSIHL 6 kDa early secretory antigenic target (13-28) Mycobacterium tuberculosis	Homo sapiens	Infectious disease via exposure to Mycobacterium tuberculosis (Source Organism)	AAASAIQGNVTSIHL 6 kDa early secretory antigenic target (13-28) Mycobacterium tuberculosis	Epitope		3H-thymidine cell proliferation Positive							
2142	Yolanda López-Vidal; Arch Med Res 2004	AAASAIQGNVTSIHL 6 kDa early secretory antigenic target (13-28) Mycobacterium tuberculosis	Homo sapiens	Infectious disease via exposure to Mycobacterium tuberculosis (Source Organism) followed by restimulation in vitro	AAASAIQGNVTSIHL 6 kDa early secretory antigenic target (13-28) Mycobacterium tuberculosis	Epitope		ELISA cytokine release IL-4 Positive							
2153	Yolanda López-Vidal; Arch Med Res 2004	LDEGKQSLTKLAAWG 6 kDa early secretory antigenic target (29-44) Mycobacterium tuberculosis	Homo sapiens	Infectious disease via exposure to Mycobacterium tuberculosis (Source Organism)	LDEGKQSLTKLAAWG 6 kDa early secretory antigenic target (29-44) Mycobacterium tuberculosis	Epitope		3H-thymidine cell proliferation Positive							
2156	Yolanda López-Vidal; Arch Med Res 2004	LDEGKQSLTKLAAWG 6 kDa early secretory antigenic target (29-44) Mycobacterium tuberculosis	Homo sapiens	Infectious disease via exposure to Mycobacterium tuberculosis (Source Organism) followed by restimulation in vitro	LDEGKQSLTKLAAWG 6 kDa early secretory antigenic target (29-44) Mycobacterium tuberculosis	Epitope		ELISA cytokine release IL-4 Positive							

Figure 2.14 Screen generated by drilling down on the positive T cell assay hyperlink in the Search Result Summary

Source Organism	
126 item(s) found, displaying 1 to 25 (Click the column headers to adjust the sorting)	
« previous	1 2 3 4 5 6 next »
Export all results: compact full	
Source Organism ↑	
Acanthamoeba castellanii	
Adeno-associated virus	
Adeno-associated virus - 2	
Aspergillus fumigatus	
Bacillus amyloliquefaciens	
Bacillus licheniformis	
Bacillus germanica	
Bordetella pertussis	
Bordetella pertussis Tohama I	
Borrelia burgdorferi	
Borrelia burgdorferi B31	
Borrelia garinii IP90	
Bos taurus	
Bovine herpesvirus type 1.1 (strain Cooper)	
Brucella melitensis	
Chamaecyparis obtusa	
Chlamydia muridarum Nigg	
Chlamydia trachomatis	
Chlamydophila abortus	
Chlamydophila pneumoniae	
Corylus avellana	
Crotalus durissus terrificus	
Cryptomeria japonica	
Cucumis melo	

Figure 2.15 Screen generated by drilling down on the Epitope Source Organism link in the Search Result Summary

Host Organism	
95 item(s) found, displaying 1 to 25 (Click the column headers to adjust the sorting)	
« previous 1 2 3 4 next » Go To » <input type="text" value="1"/>	
Export all results: <input checked="" type="checkbox"/> (compact full)	
Host Organism ↑	
Homo sapiens	
Homo sapiens Australian Aborigines	
Homo sapiens Brazilian	
Homo sapiens Caucasian	
Homo sapiens Caucasian, Aborigine and Thai	
Homo sapiens Caucasians	
Homo sapiens Colombian	
Homo sapiens Endemic region	
Homo sapiens European	
Homo sapiens Finnish	
Homo sapiens Gabonese	
Homo sapiens Gambian	
Homo sapiens Indian	
Homo sapiens Israeli	
Homo sapiens Italian	
Homo sapiens Japanese	
Homo sapiens Nepalese	
Homo sapiens North American	
Homo sapiens South African	
Homo sapiens Sudanese	
Homo sapiens white	
Mus musculus	
Mus musculus (C57BL/6 X DBA/2) Tg(HBV 1.3 genome)chi32	
Mus musculus 129	
Mus musculus A/J	

Figure 2.16 Screen generated by drilling down on the Host Organism link in the Search Result Summary

Restricting MHC Allele								
72 item(s) found, displaying 1 to 25 (Click the column headers to adjust the sorting)								
	< previous	1	2	3	next »	Go To »	1	
Export all results: <input checked="" type="checkbox"/> (compact full)								
MHC Allele ↑	Class	Organism	Synonyms	Restriction Level	Haplotype	Locus	Serotype	Molecule
H-2-a class I	I	mouse (Mus musculus)		HAPLOTYPE	H-2-a			
H-2-b class I	I	mouse (Mus musculus)		HAPLOTYPE	H-2-b			
H-2-b class II	II	mouse (Mus musculus)		HAPLOTYPE	H-2-b			
H-2-d class I	I	mouse (Mus musculus)		HAPLOTYPE	H-2-d			
H-2-d class II	II	mouse (Mus musculus)		HAPLOTYPE	H-2-d			
H-2-D _b	I	mouse (Mus musculus)		COMPLETE MOLECULE	H-2-b	D		Db
H-2-D _d	I	mouse (Mus musculus)		COMPLETE MOLECULE	H-2-d	D		Dd
H-2-g7 class I	I	mouse (Mus musculus)		HAPLOTYPE	H-2-g7			
H-2-I _A b	II	mouse (Mus musculus)		COMPLETE MOLECULE	H-2-b	IA		IAb
H-2-I _A d	II	mouse (Mus musculus)		COMPLETE MOLECULE	H-2-d	IA		IAd
H-2-I _A g7	II	mouse (Mus musculus)		COMPLETE MOLECULE		IA		IAg7
H-2-I _A g7 H56P, S57D mutant	II	mouse (Mus musculus)		COMPLETE MOLECULE	H-2-g7	IA		IAg7
H-2-I _A k	II	mouse (Mus musculus)		COMPLETE MOLECULE	H-2-k	IA		IAk
H-2-I _A s	II	mouse (Mus musculus)		COMPLETE MOLECULE	H-2-s	IA		IAs
H-2-I _E k	II	mouse (Mus musculus)		COMPLETE MOLECULE	H-2-k	IE		IEk
H-2-k class I	I	mouse (Mus musculus)		HAPLOTYPE	H-2-k			
H-2-k class II	II	mouse (Mus musculus)		HAPLOTYPE	H-2-k			

Figure 2.17 Screen generated by drilling down on the Restricting MHC Allele link in the Search Result Summary

Reference List

Sort By: Date Order By: Descending [Submit](#)

391 item(s) found, displaying 1 to 25
 Export all results: ([compact](#) | [full](#))

[« previous](#) [1](#) [2](#) [3](#) [4](#) [5](#) [6](#) [7](#) [8](#) [9](#) ... [15](#) [16](#) [next »](#) [Go To »](#) [1](#)

[Daisuke Suzuki; Toshi Nagata; Ghada Eweda; Sohichi Matsumoto; Makoto Matsumoto; Kunio Tsujimura; Yukio Koide.](#)
 Characterization of murine T-cell epitopes on mycobacterial DNA-binding protein 1 (MDP1) using DNA vaccination.
Vaccine, 2010; 28(1873-2518):2020-5
 PMID: [20188258](#)

[Latisha D Heinlen; Lauren L Ritterhouse; Micah T McClain; Michael P Keith; Barbara R Neas; John B Harley; Judith A James.](#)
 Ribosomal P autoantibodies are present before SLE onset and are directed against non-C-terminal peptides.
J Mol Med, 2010; 88(1432-1440):719-27
 PMID: [20396862](#)

[Wen-Quan Zou; Jan Langeveld; Xiangzhu Xiao; Shuqi Chen; Patrick L McGeer; Jue Yuan; Michael C Payne; Hae-Fun Kang; John McGeehan; Man-Sun Sy; Neil S Greenspan; David Kaplan; Gong-Xian Wang; Piero Parchi; Edward Hoover; Geoff Kneale; Glenn Telling; Witold K Surewicz; Qingzhong Kong; Jian-Ping Guo.](#)
 PrP conformational transitions alter species preference of a PrP-specific antibody.
J Biol Chem. 2010; 285(1083-351X):13874-84
 PMID: [20194495](#)

[Amanda R Burton; Zachary Baquet; George S Eisenbarth; Roland Tisch; Richard Smeyne; Creg J Workman; Dario A A Vignali.](#)
 Central nervous system destruction mediated by glutamic acid decarboxylase-specific CD4+ T cells.
J Immunol. 2010; 184(1550-6606):4863-70
 PMID: [20348424](#)

[E A Pastorello; M Monza; V Pravettoni; R Longhi; P Bonara; J Scibilia; L Primavesi; R Scorza.](#)
 Characterization of the T-cell epitopes of the major peach allergen Pru p 3.
Int Arch Allergy Immunol. 2010; 153(1423-0097):1-12
 PMID: [20357479](#)

[Marina A Durward; Jerome Harms; Diogo M Magnani; Linda Eskra; Gary A Splitter.](#)
 Discordant Brucella melitensis antigens yield cognate CD8+ T cells in vivo.
Infect Immun. 2010; 78(1098-5522):168-76
 PMID: [19884330](#)

[Madhan Kumar; N Meenakshi; Jagadish C Sundaramurthi; Gurvinder Kaur; Narinder K Mehra; Alamelu Raja.](#)
 Immune response to Mycobacterium tuberculosis specific antigen ESAT-6 among south Indians.
Tuberculosis (Edinb). 2010; 90(1873-281X):60-9
 PMID: [19944647](#)

[Véronique Bachy; Clara Ballerini; Pauline Gourdin; Aurélie Prignon; Saci Iken; Nadine Antoine; Martine Rosset; Claude Carnaud.](#)
 Mouse vaccination with dendritic cells loaded with prion protein peptides overcomes tolerance and delays scrapie.
J Gen Virol. 2010; 91(1465-2099):809-20
 PMID: [19864503](#)

Figure 2.18 Screen generated by drilling down on the reference hyperlink in the Search Result Summary

When users click on the number of references on the Search Results Summary screen, the system will display the Reference List, as seen in Figure 2.18. This list will display all the references that matched the user's search criteria. Enough information is listed to identify each reference uniquely, including authors, article title, year, PubMed ID, and journal name. To view all the information related to a given reference, click the highlighted author line. The list can be sorted by date, author, PubMed ID, and title in ascending or descending order. Clicking on the PubMed identifier will open the PubMed citation in a new window.

2.3.4.1 Peptide MHC Binding Motif Displays

A display of peptide MHC binding motif diagrams and amino acid binding charts for a subset of MHC allele records appears on the MHC Allele Information page. This page can be reached from the Restricting MHC Allele page shown in Figure 2.17. Clicking on H-2-Db in the left MHC Allele column in the sixth row, the user is brought to its MHC Allele Information page (Figure 2.19).

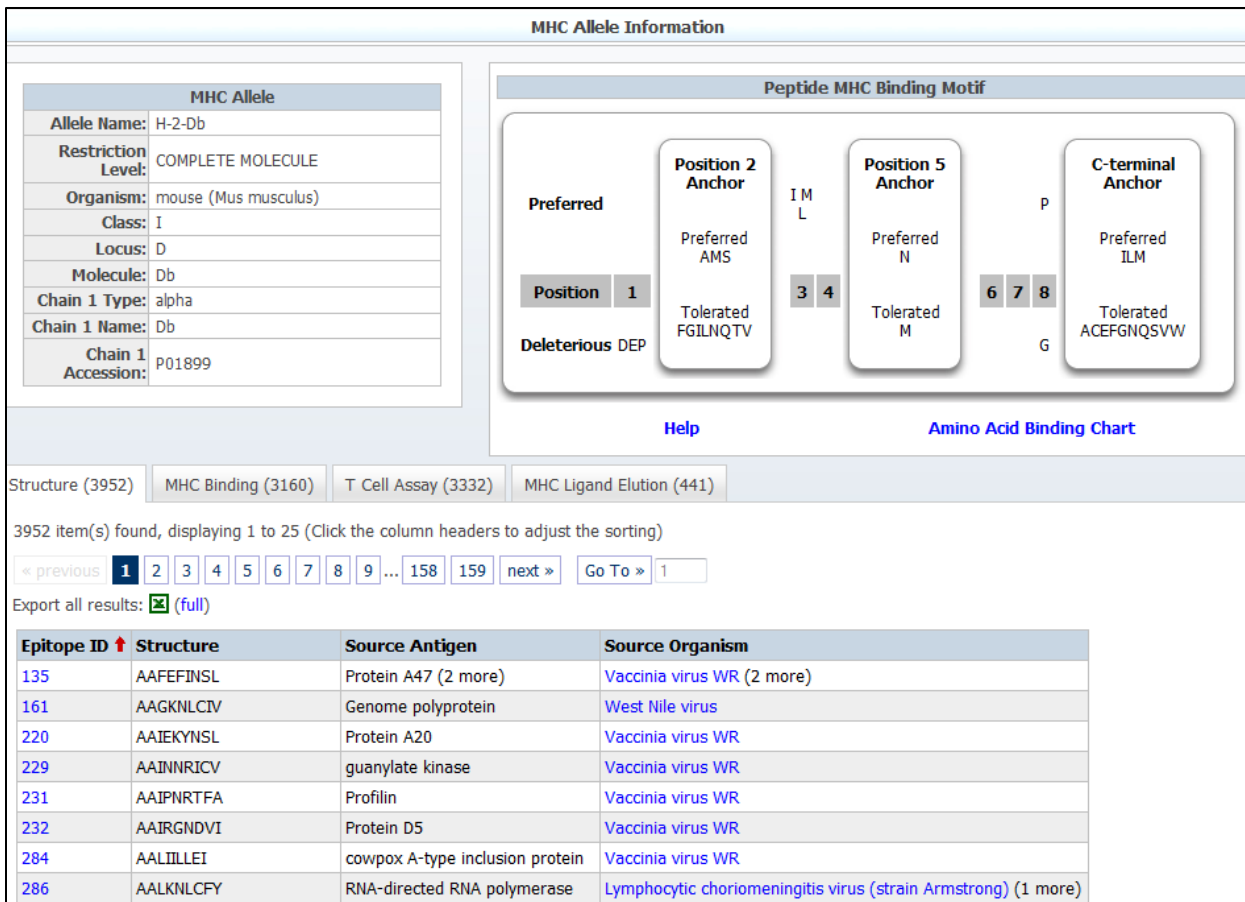


Figure 2.19 Sample MHC Allele Information page with Peptide MHC Binding Motif diagram for H-2_Db. Note the link to the corresponding Amino Acid Binding Chart on the bottom right of the diagram

The Peptide MHC Binding Motif diagram summarizes the results of the amino acid binding for each position, which is shown in Figure 2.20. The amino acids are color-coded for each position, where green indicates the residue is preferred for binding, purple indicates it is deleterious for binding, and white indicates the residue is tolerated at that position. The motif diagram shows which residues are preferred and tolerated at the anchor positions, and which residues are preferred and deleterious at the non-anchor positions.

The residue preference is calculated from the SMM prediction matrix, which is described in Section 2.4.1.1. The following steps are taken in determining the residue preference at each position:

For anchor position:

1. determine the best value of this position from corresponding column of the SMM matrix
2. if a residue's value in this column of the SMM matrix is within 3 fold of the best value, designate this residue as preferred
3. if a residue's value in this column of the SMM matrix is within 10 fold of the best value, designate this residue as tolerated
4. The remaining residues are designated as deleterious

H-2-Db Motif Amino Acid Binding Chart

AMINO ACID	PEPTIDE BINDING MOTIF POSITION								
	1	2	3	4	5	6	7	8	9
A	-0.22	-0.92	0.245	-0.13	0.239	0.013	-0.11	-0.19	0.101
C	0.036	0.191	-0.00	-0.11	0.051	0.080	0.051	0.260	0.132
D	0.456	0.568	0.477	0.387	-0.10	0.059	-0.40	0.175	0.239
E	0.473	0.344	0.283	-0.03	0.200	0.071	-0.16	-0.18	0.088
F	-0.45	0.024	-0.25	0.044	-0.07	-0.08	-0.12	-0.21	-0.25
G	0.156	-0.13	0.277	0.289	0.062	0.082	0.376	0.556	0.120
H	-0.00	0.212	-0.07	0.069	-0.06	0.179	-0.08	0.084	0.325
I	-0.08	0.038	-0.58	-0.13	0.282	0.051	0.115	0.025	-0.84
K	-0.09	0.315	0.426	0.073	0.247	0.088	0.415	0.377	0.246
L	0.028	0.016	-0.52	-0.18	-0.27	-0.09	0.051	0.005	-0.65
M	-0.08	-0.74	-0.50	-0.10	-0.52	-0.20	0.184	0.134	-0.83
N	0.069	0.044	-0.15	0.111	-1.38	0.156	0.077	0.118	0.030
P	0.587	0.541	0.109	0.282	0.130	-0.03	-0.10	-0.66	0.339
Q	0.221	-0.35	0.154	-0.06	0.009	-0.05	0.076	-0.02	0.107
R	-0.11	0.360	0.492	-0.08	0.440	-0.05	0.255	0.449	0.594
S	-0.32	-0.78	0.314	-0.06	0.001	0.106	0.000	-0.03	0.152
T	-0.03	-0.17	0.187	-0.14	0.275	0.003	-0.28	-0.35	0.216
V	-0.07	-0.17	-0.26	-0.35	0.212	0.032	-0.01	-0.14	-0.35
W	-0.08	0.527	-0.31	0.022	-0.05	-0.22	-0.10	0.025	0.023
Y	-0.45	0.112	-0.29	0.135	0.338	-0.17	-0.19	-0.39	0.221

(ROW HEADER) Indicates anchor positions.

(MATRIX BODY) Indicates that the residue is a prefered residues at that position.

Indicates that the residue is a deleterious residue at that position.

Indicates that the residue is a tolerated residue at that position.

Figure 2.20 Amino acid binding matrix for MHC allele H-2-Db

the leftmost ID column, as illustrated by the “View 3D Structure” in Figure 2.21. The 23 B cell response assays shown are a result of using the B Cell Search for Epitope Source Organism = Influenza A and Assay Type = X-Ray Crystallography.

For non-anchor position:

1. determine the median value of this position from corresponding column of the SMM matrix

2. if a residue's value in this column of the SMM matrix is within 3 fold of the median value, designate this residue as tolerated

3. if a residue's value in this column of the SMM matrix is above 3 fold of the median value, designate this residue as prefered

4. if a residue's value in this column of the SMM matrix is below 3 fold of the median value, designate this residue as deleterious

2.3.4.2 Accessing the EpitopeViewer

The EpitopeViewer is an application for three dimensional viewing of receptor-antigen interactions that can be accessed from the Assay List screens. The EpitopeViewer can be used with all assays that have receptor-antigen interaction data available. It is not available for epitopes and assays, and when it is available, a link is provided in

B Cell Response Assays							
23 item(s) found, displaying 1 to 23 (Click the column headers to adjust the sorting) Export all results: compact full							
B Cell ID ↑	Reference	Epitope	Host	Immunization	Assay Antigen	Antigen Epitope Relation	Assay Description
3298 View 3D Structure	R L Malby; Structure 1994	P330, N331, D332, P333, T334, Y342, G344, N345, I3... Neuraminidase Influenza A virus (A/whale/Maine /1/84(H1N9))	Mus musculus	Administration in vivo with Influenza A/tern/Australia /G70C/75 (H1N9) neuraminidase gene (Taxonomic Sibling)	Neuraminidase Neuraminidase Influenza A virus (A/whale/Maine /1/84(H1N9))	Source Antigen	X-Ray Crystallography Characterization of Ab binding Positive
14182 View 3D Structure	M E Churchill; J Mol Biol 1994	DVPDYASL Hemagglutinin precursor (118-125) Influenza A virus (A/Victoria /3/1975(H3N2))	Mus musculus	The immunization procedure is unknown	PYDVPDYASLRS Hemagglutinin precursor (116-127) Influenza A virus (A/Victoria /3/1975(H3N2))	Fragment of Source Antigen	X-Ray Crystallography Characterization of Ab binding Positive
14243 View 3D Structure	U Schulze-Gahmen; J Mol Biol 1993	DVPDYASL Hemagglutinin precursor (118-124) Influenza A virus (A/Victoria /3/1975(H3N2))	Mus musculus	The immunization procedure is unknown	DVPDYASL Hemagglutinin precursor (118-125) Influenza A virus (A/Victoria /3/1975(H3N2))	Fragment of Source Antigen	X-Ray Crystallography Characterization of Ab binding Positive
14757 View 3D Structure	D Fleury; Proteins 2000	K66, L75, D76, I78, D79, P90, H91, V94, F95, R106.... Hemagglutinin Influenza A virus (A/X-31(H3N2))	Mus musculus	Administration in vivo with Influenza A virus (A/X-31(H3N2)) (Source Organism)	QDLPGNDNSTATLCL GHHAVPNGLVKTIT DDQIEVTNATELVQS SSTGK... Hemagglutinin (17-521) Influenza A virus (A/X-31(H3N2))	Fragment of Source Antigen	X-Ray Crystallography Characterization of Ab binding Positive
14758 View 3D Structure	D Fleury; Proteins 2000	G65, K66, L75, D76, I78, D79, P90, H91, V94, F95, ... Hemagglutinin Influenza A virus (A/X-31(H3N2))	Mus musculus	Administration in vivo with Influenza A virus (A/X-31(H3N2)) (Source Organism)	QDLPGNDNSTATLCL GHHAVPNGLVKTIT DDQIEVTNATELVQS SSTGK... Hemagglutinin (17-521) Influenza A virus (A/X-31(H3N2))	Fragment of Source Antigen	X-Ray Crystallography Characterization of Ab binding Positive
14759 View 3D Structure	T Bizebard; Nature 1995	T147, G150, S152, S161, W169, T171, K172, S173, G1... Hemagglutinin precursor Influenza A virus (A/Aichi /2/1968(H3N2))	Mus musculus	The immunization procedure is unknown	Hemagglutinin precursor (-99--99) Influenza A virus (A/Aichi /2/1968(H3N2))	Fragment of Source Antigen	X-Ray Crystallography Characterization of Ab binding Positive

Figure 2.21 B Cell Response screen with Epitope Viewer links in the leftmost column

2.3.4.3 Result Tabs

As users drill down on hyperlinks to further explore the results of a query, they will find the information organized by tabs. As seen in Figure 2.22 for Epitope ID 67436, there are 16 source organisms listed from 47 references. There is also assay information for MHC binding, B cells, T cells, and MHC ligand elution. The Links tab contains links to relevant tools in the Analysis Resource. Links can also appear to the Bioinformatics Resource Centers (BRC).

Epitope Information

Distinct Epitope	
Epitope ID:	67436
Linear Sequence:	TYQRTRALV
Source Antigen:	Nucleoprotein (10 more)
Source Organism:	Influenza A virus (A/Puerto Rico/8/34(H1N1)) (8 more)

Source (16)	Reference (47)	MHC Binding (20)	B Cell Assay (2)	T Cell Assay (92)	MHC Ligand Elution (6)	Links (3)
-------------	----------------	------------------	------------------	-------------------	------------------------	-----------

16 item(s) found, displaying 1 to 16 (Click the column headers to adjust the sorting)
 Export all results: [compact](#) | [full](#)

Source Accession ↑	Source Antigen	Source Organism	3D Structure Homologs
118864386	nucleocapsid protein	Influenza A virus	2 PDB hits
13274623	nucleocapsid protein	Influenza A virus	2 PDB hits
139029	Nucleoprotein	Influenza A virus (A/Anas acuta/Primorje/695/1976(H2N3))	2 PDB hits
139029	Nucleoprotein	Influenza A virus	2 PDB hits
139085	Nucleoprotein	Influenza A virus	2 PDB hits
324710	nucleoprotein	Influenza A virus	2 PDB hits
61250058	Nucleoprotein	Influenza A virus (A/Port Chalmers/1/1973(H3N2))	2 PDB hits
73666582	nucleocapsid protein	Influenza A virus (A/Memphis/1/71H-A/Bellamy/42N)	2 PDB hits
82010653	Nucleocapsid protein	Influenza A virus (A/England/939/69 x A/PR/8/34)	2 PDB hits
8486130	nucleocapsid protein	Influenza A virus (A/Puerto Rico/8/34(H1N1))	2 PDB hits
P03466	Nucleoprotein	Influenza A virus	2 PDB hits
P03466	Nucleoprotein	Influenza A virus (A/Puerto Rico/8/34(H1N1))	2 PDB hits
P03466	Nucleoprotein	Influenza A virus (A/Japan/305/1957(H2N2))	2 PDB hits
P03466	Nucleoprotein	unidentified influenza virus	2 PDB hits
P03466	Nucleoprotein	Influenza A virus (A/Memphis/1/71H-A/Bellamy/42N)	2 PDB hits
P03467	Nucleoprotein	Influenza A virus (A/nt/60/1968(H3N2))	2 PDB hits

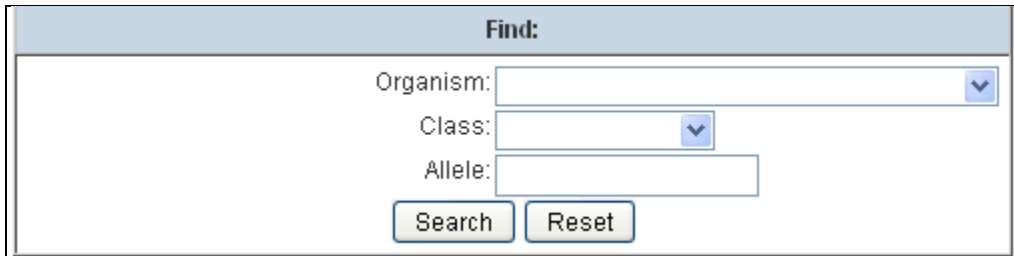
Figure 2.22 Many results pages have information organized into tabs

2.3.5 Finders Overview

Several finders (Allele, Assay, Disease, Molecule, and Organism) are available to help facilitate selections and control vocabulary usage (improves result outputs). At times the potential list of selections can be quite extensive, and the finders help users make selections from large lists. The finders can be utilized when performing all queries except the keyword query. Multiple selections can be made when utilizing finders during a query.

2.3.5.1 Allele Finder

The MHC Allele Finder facilitates the selection of one or more MHC alleles. Initially the Allele Finder lists all alleles ordered by allele name. The Allele Finder allows the user to find alleles by organism, class {I, II, non-classical}, and allele in the Find box (Figure 2.23). After the user supplies their search criteria and clicks the Search button, the system will filter the list of MHC alleles using the organism, class, and allele provided. The allele finder uses wild card characters by default on both ends of criteria entered into the allele field. The system then returns any alleles that contain the value in the name field and match the class selected.

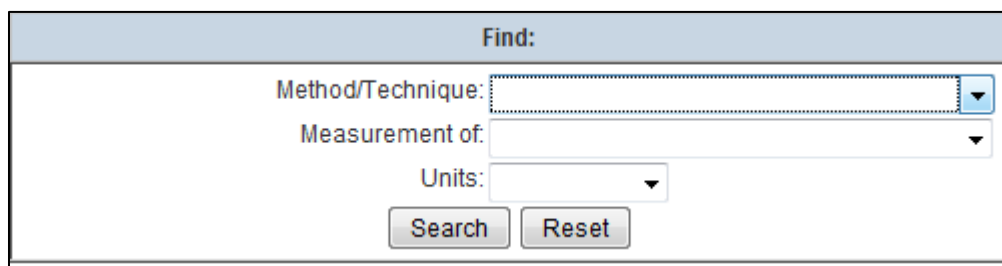


The screenshot shows a search interface titled "Find:" at the top. It has three input fields: "Organism:" with a dropdown arrow, "Class:" with a dropdown arrow, and "Allele:" with a dropdown arrow. Below these fields are two buttons: "Search" and "Reset".

Figure 2.23 Find form on the Allele Finder

2.3.5.2 Assay Finder

The Assay Finder is used to facilitate the selection of one or more assay types and lists all assay types in the selected assay category. The Assay Finder allows the user to find assay by method or technique, by what it measures, and/or units in the Find box (Figure 2.24). After the user supplies their search criteria and clicks the Search button, the system filters the list of assay types using the selections provided in the Find box.



The screenshot shows a search interface titled "Find:" at the top. It has three input fields: "Method/Technique:" with a dropdown arrow, "Measurement of:" with a dropdown arrow, and "Units:" with a dropdown arrow. Below these fields are two buttons: "Search" and "Reset".

Figure 2.24 Search form on the Assay Type Finder

2.3.5.3 Disease Finder

The disease finder is used to facilitate the selection of a disease state and input the selection into a Disease State field. It includes all diseases from The International Statistical Classification of Diseases and Related Health Problems, tenth revision (ICD-10) and displays diseases with their corresponding ICD-10 codes in a hierarchical tree. The first level of the tree displays similar groups of diseases, and each additional level of the tree further breaks down the groups of diseases. Variations of each disease are not included. Searching capabilities are provided so users can quickly select a disease state.

The Disease Finder will allow the user to find diseases using the disease name or ICD-10 code (Figure 2.25). When the user performs a search, the system will display the first match and then allow the user to move forward and backward through the matching records using Next and Previous buttons. When the user provides a name, any disease name that contains the character string provided will be considered a match.

The selections in the disease finder are displayed in a tree. To expand a node of the tree, the user clicks the plus sign next to the name. To collapse a node, a user clicks the corresponding minus sign. The ICD-10 code is displayed next to each selection in square brackets. For example the ICD-10 code for Bartonellosis is A44 and the code for Systemic bartonellosis is A44.0.

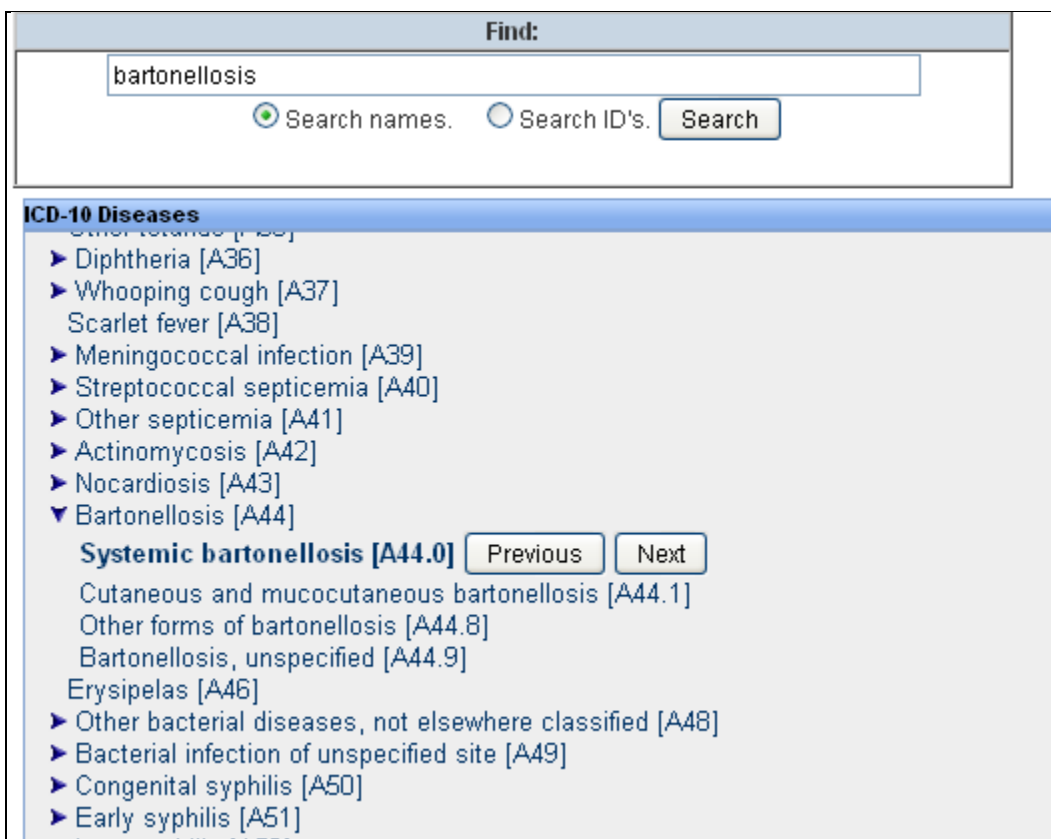


Figure 2.25 Search form and disease list on the Disease Finder

2.3.5.4 Molecule Finder

As mentioned in the description of the Home Page Search (Section 2.3.2) has been updated to the Home Page search. It remains the same for Advanced searches, although that will change in 2011. The Molecule Finder is used to facilitate the selection of source antigens, immunogens, and epitopes. Records in the Source Finder come from GenPept, GenBank, ChEBI, Swiss-Prot, and IEDB curators.

Over a year ago it became apparent to the IEDB team that the source molecule finder did not fully meet the needs of the IEDB user base. In order to select a specific source molecule, the user needed to type its name in a free text field. Although this was a workable method, it proved to be error-prone and assumed that the user was familiar with the contents of the database. Two of the most requested features of our users were: 1) the ability to browse the available source molecules in a meaningful way and 2) the ability to select a source protein and have all related proteins added to the query as well. To this end, the source molecule finder was redesigned to include two parallel trees, one for non-peptides and the other for protein molecules.. The first contains the structures curated by the Chemical Entities of Biological Interest (ChEBI) database. An example is shown in Figure 2.26.

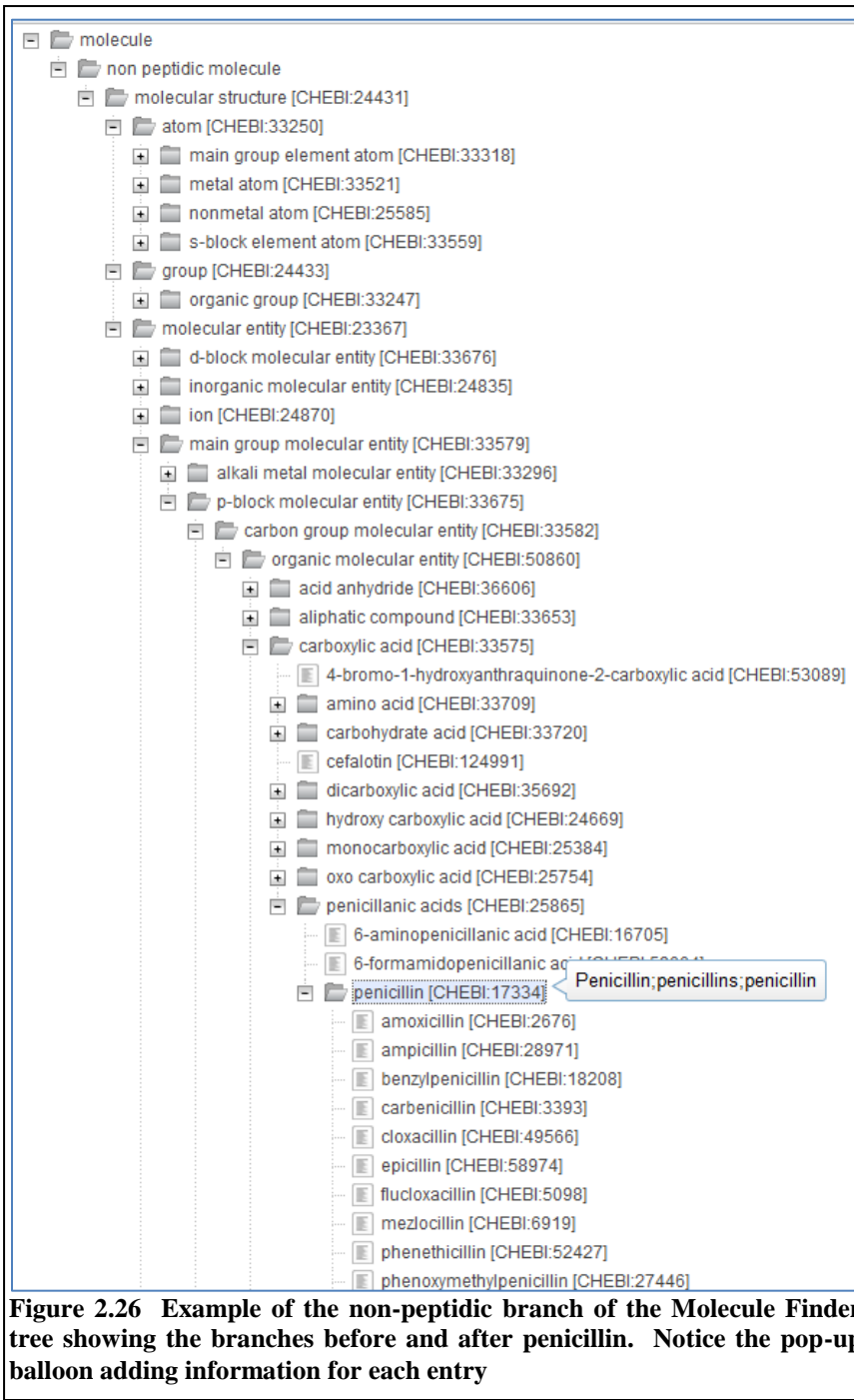


Figure 2.26 Example of the non-peptidic branch of the Molecule Finder tree showing the branches before and after penicillin. Notice the pop-up balloon adding information for each entry

The development team determined that the most logical way to group the proteins was by organism. In order to accomplish this, the NCBI species was determined for each of the proteins in the database. For viruses and bacteria, this involved traversing the NCBI taxonomy from the sub-species (strain) level up to the species level. For each species, a set of reference proteins was selected from the NCBI protein database based upon the availability of a complete genome for the species. All proteins for each species were BLASTed against the reference protein set to determine their homologs. These data were used to build the protein tree in a way that mirrors a

pruned version of the NCBI taxonomy. The result is a coherent tree that is divided along major taxonomic categories and is quickly traversed with proteins grouped logically below each species. The ability to perform a free text search was maintained and is now enhanced by the ability to see where the protein is placed in the tree. It is now possible, for example, to select all Influenza A haemagglutinin (HA) proteins by selecting one node of the tree rather than individually clicking on the 100+ different HA proteins in the database. This can be seen in Figure 2.27.

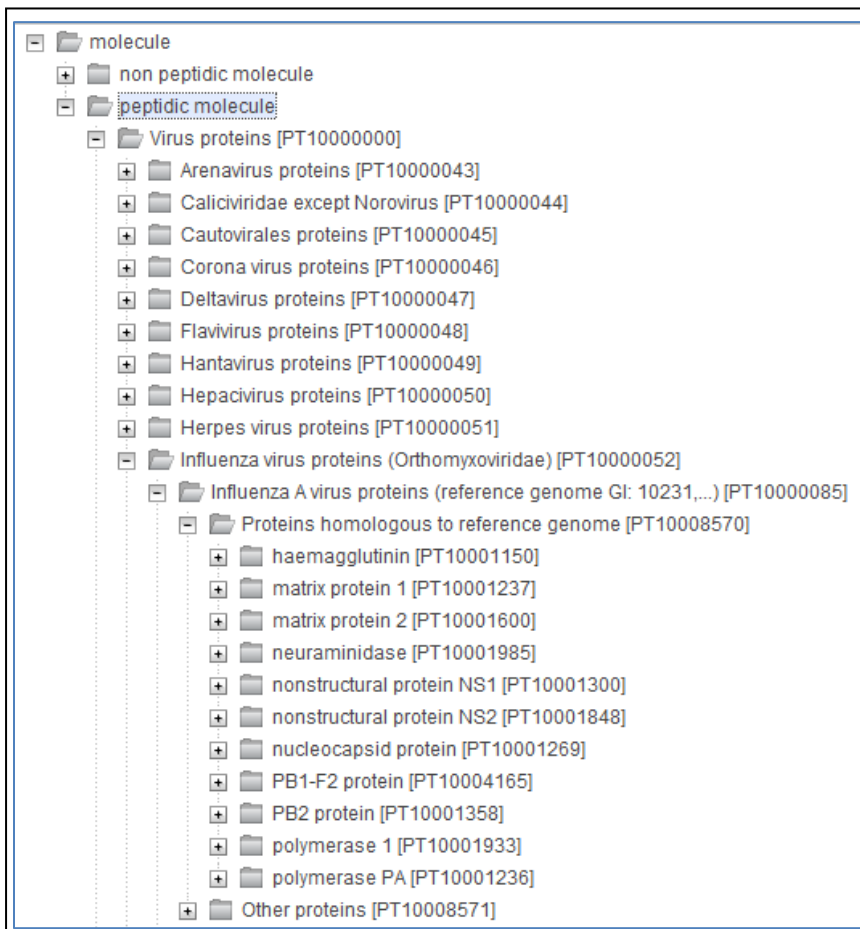


Figure 2.27 Example of the protein tree as found in the Source Antigen Molecule Finder on the IEDB home page

The Molecule Finder on the Advance Search pages remains the same as before. It has a comprehensive and flexible search form, and even includes the Organism Finder. Due to the large volume of possible selections, the Molecule Finder initially won't display any selections. Users need to perform a search to narrow the list down. In the example shown in Figure 2.28, the user has entered search criteria for Influenza A Nucleoproteins that were curated from the Swiss-Prot database. The search on these criteria resulted in a list of five matching sources, and the user can select their desired sources from the list by click on *Select* in the far left column.

Find:

Molecule Accession:	<input type="text"/>	Database:	<input type="button" value="Swiss-Prot ▾"/>
Chemical Type:	Protein	<input type="button" value="▼"/>	
Sequence:	<input type="text"/>		
Molecule Name:	<input type="text" value="np"/>		
Source Organism:	<input type="text" value="Influenza A virus"/>		<input type="button" value="Organism Finder"/>
<input type="button" value="Search"/>			

5 items found, displaying all items.

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Options	Molecule Accession	Database	Molecule Names	Organism Name	Chemical Type	Structure	Position
Select	A0ZT74	Swiss-Prot	Nucleoprotein; NP	Influenza A virus (A/Tern /Australia /I363/2004(H2N5))	Protein		
Select	P03466	Swiss-Prot	Nucleoprotein; NP	Influenza A virus (A/Puerto Rico/8/34(H1N1))	Protein		
Select	P03467	Swiss-Prot	Nucleoprotein; NP	Influenza A virus (A/nt/60 /1968(H3N2))	Protein		
Select	P22435	Swiss-Prot	Nucleoprotein; NP	Influenza A virus (A/Hong Kong/1/1968(H3N2))	Protein		
Select	Q08041	Swiss-Prot	Nucleoprotein; NP	Influenza A virus (A/Memphis /101/1972(H3N2))	Protein		

Figure 2.28 Molecule Finder Search form and resulting source list

2.3.5.5 Organism Finder

The organism finder is used to facilitate the selection of a species or virus from the NCBI Taxonomy Database. Common selections are displayed initially to speed the selection of the usual suspects, as seen in Figure 2.29. To view the entire NCBI taxonomy data set click *NCBI Taxonomy Tree* in the accordion slider. The Common Selection bar will move down to reveal the taxonomy tree. An Allergen tree has also been provided. The Allergen tree consists of species from the NCBI taxonomy, but is organized to help allergists locate common allergen selections more easily. To use the Allergen tree click the Allergen Tree heading in the accordion slider. The Allergen tree was updated in 2010 to display only nodes reference by curated data. In addition, all nodes in the Allergen tree are now selectable.

NCBI Taxonomy Tree
Allergen Tree
Common Selections
Bacillus anthracis
Clostridium tetani
Dermatophagoides pteronyssinus(European house dust mite)
Hepatitis B virus
Hepatitis B virus subtype ADW
Hepatitis B virus subtype ADW2
Hepatitis B virus subtype AYW
Hepatitis C virus
Hepatitis C virus (isolate 1)
Hepatitis C virus subtype 1a
Hepatitis C virus subtype 1b
Homo sapiens(human)
Human herpesvirus 1(Herpes simplex virus type 1)

Figure 2.29 Common selections in Organism finder

The Organism finder will allow the user to find species using their name or taxonomy identifier (assigned by NCBI). When the user performs a search, the system will display the first match then allow the user to move forward and backward through the matching records using Next and Previous buttons as in Figure 2.30. When the user provides a name, any species name or synonym that contains the name provided will be considered a match. Search results will always appear in the NCBI Taxonomy tree, not the Allergen Tree or Common Selections.

The selections in the organism finder are displayed in a tree (Figure 2.30). The taxonomy identifier is displayed next to each node of the tree in square brackets. For example, the taxonomy identifier for the selection of Dengue Virus 2 in the example below is 11060. To see the synonyms for a selection, users can place their computer mouse over the scientific name.

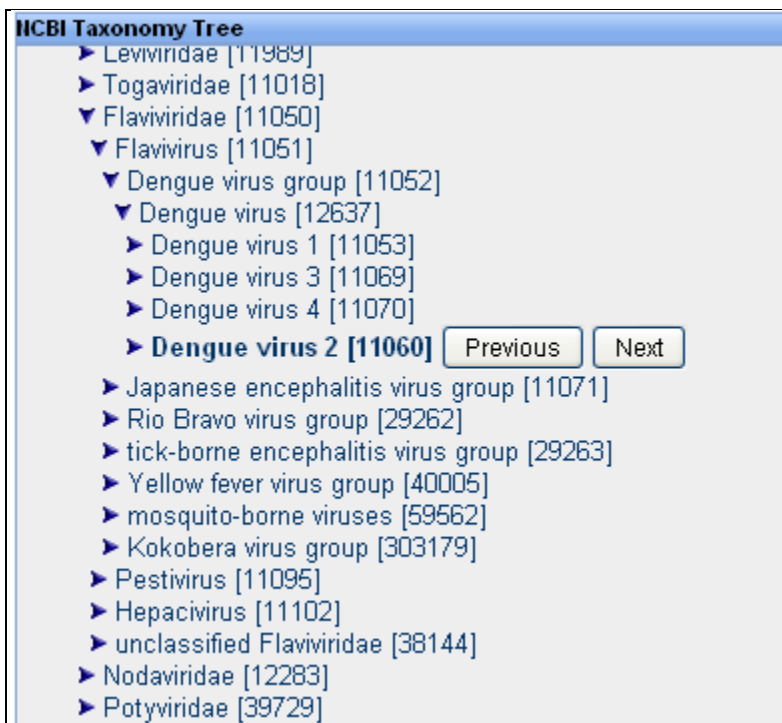


Figure 2.30 NCBI Taxonomy Tree

2.4 Tools

The Tools pull-down menu access the IEDB Analysis Resource and has three items – T Cell Epitope Prediction, B Cell Epitope Prediction, and Analysis Tools. The purpose of the Analysis Resource of the IEDB is to provide computational tools that enhance the value of the IEDB database to the user. Providing access to tools in one centralized location helps make users aware of available solutions to their problems. All of the information contained within the Analysis Resource, including analysis tools and algorithms developed by the IEDB staff, are freely available to the scientific community.

The tools provided in the analysis resource fall into two categories – prediction tools and analysis tools. Prediction tools extrapolate beyond data held in the database. They can be used to predict epitopes in protein sequences or predict properties of known epitopes, such as their MHC binding affinity. Analysis tools help extract and interpret data contained in the database.

For prediction tools, it is important to differentiate between the **tool** making predictions, and the **method** used to generate that tool, given a set of training data. For example, the artificial neural network method or approach, when trained on a particular data set, will yield a prediction tool. As the ANN method is trained on different data sets, different corresponding prediction tools will result. In this way, the ANN method can be used to develop an MHC class I prediction tool and a separate MHC class II prediction tool. These tools can be refined as more data are available for training. One benefit of the IEDB is that it allows implementing methods to automatically generate new prediction tools as the database grows.

Prediction tools can be subdivided into categories by what they predict. The current tools fall into the subcategories listed below:

- T cell epitopes – MHC class I and II binding prediction
- T cell epitopes – Processing prediction
- B-cell epitope prediction

The next subsections will describe each prediction and analytical tool in more detail.

2.4.1 T Cell Epitope Prediction

2.4.1.1 T Cell Epitopes - MHC binding prediction

The Analysis Resource provides tools for predicting peptide binding to MHC class I and II molecules. For class I binding predictions, one new method, NetMHCpan, was introduced in 2010. Users can select predictions performed with tools derived from seven different methods – artificial neural network (ANN), average relative binding (ARB), stabilized matrix method (SMM), SMM with a peptide:MHC binding energy covariance matrix (SMMPMBEC), Scoring matrices derived from combinatorial peptide libraries (Comblib_Sidney2008), Consensus, and NetMHCpan. The last three methods listed are new for the year. For class II binding predictions, one new method, NetMHCIIpan, was added in 2010. Users can select the Combinatorial Library method, the ARB method, the SMM_align method, a method devised by Sturniolo et al. and used in TEPITOPE, and a consensus method derived from the ARB, SMM_align, Sturniolo methods, and NetMHCIIpan. Tutorials and example data are available for both the class I and II tools.

2.4.1.1.1 Peptide Binding to MHC Class I Molecules

Users can select from seven different methods for predicting class I epitopes – ANN, ARB, SMM, SMMPMBEC, Comblib_Sidney2008, Consensus, and NetMHCpan, which are described further below.

Artificial Neural Network

Artificial neural networks (ANN) are computer algorithms modeled after the brain. They consist of many simple processing units which are wired together in a communication network. Each unit is a simplified model of a neuron which sends off a new signal if it receives a sufficiently strong input signal from the other units to which it is connected. The strength of these connections can be varied in order for the network to perform a desired pattern of node signal activity, which is learned from a set of input training data. The training data in this case are peptide sequences with quantitative affinities for a specific MHC molecule.

Many different implementations of artificial neural networks exist. The one utilized here is described for HLA-A2 binding predictions by Nielsen et al. (Protein Science, 2003) and has been applied to a number of different alleles (<http://www.cbs.dtu.dk/services/NetMHC/>).

Average Relative Binding (ARB)

Average relative binding (ARB) matrix binding prediction method is based on the assumption that each residue along the peptide molecule independently contributes to binding affinity. When a residue R occurs at position i in the peptide, it is assumed to contribute a constant amount of R_i to the free energy of binding of the peptide. The effect of each of the 20 possible

amino acids at each possible position along the peptide sequence, therefore, can be estimated by a matrix of coefficients. The overall binding propensity of each peptide sequence, an algorithm “score”, is calculated by multiplying the R_i coefficients. Predicted IC₅₀ values, which provide quantitative K_D (IC₅₀) predictions, are then calculated by mathematical transformations of the algorithm scores (Bui et al., Immunogenetics 2005).

Stabilized Matrix Method (SMM)

The Stabilized Matrix Method (SMM) described by Peters and Sette (BMC Bioinformatics, 2005) can be applied to calculate matrices from quantitative affinity data of peptides binding to MHC molecules. The advantage of this method is that it suppresses the noise present in the training data, caused by the inevitable experimental error as well as the limited number of data points.

Stabilized matrix Method with a Peptide:MHC Binding Energy Covariance matrix (SMMPMBEC)

SMMPMBEC is an improved version of SMM. It is different from SMM in that it addresses sparseness of peptide sequence coverage that is often found in binding data sets by using Peptide:MHC Binding Energy Covariance (PMBEC). The PMBEC matrix was derived from experimentally determined binding affinity measurements using combinatorial peptide libraries. SMMPMBEC is described in Kim et al. BMC Bioinformatics 2009.

Scoring matrices derived from combinatorial peptide libraries (Comblib_Sidney2008)

Comblib_Sidney2008 refers to a set of predictors (i.e. scoring matrices) that were derived from binding affinity measurements of combinatorial peptide libraries against a panel of MHC alleles. This work is described in Sidney et al. Immunome Res. 2008. This class of predictors is unique in that average binding energy contribution of a given residue at a position is directly measured, without worrying about limited peptide sequence coverage.

Consensus

The Consensus predictor was motivated by an idea that predictions made by consulting “consensus” of individual predictions from multiple predictors may result in improved performance over that of any individual ones. For MHC-I, a work describing an early implementation can be found in Moutaftsi M et al. Nat Biotech 2006.

NetMHCpan

NetMHCpan predicts binding of peptides to a MHC class I molecule using artificial neural networks (ANN). It predicts binding for over 1,650 alleles, including HLA-A, B, C, E, G; non-human primates; mouse; pig; and user-supplied MHC sequence. Predictions can be made for peptide sequences of 8 to 11 residues in length. The method has been trained on over 110,000 peptide/MHC interactions.

2.4.1.1.1 MHC Class I Binding Prediction Resource

In addition to prediction tools, the Analysis Resource makes data sets and method evaluations available to users. The IEDB MHC Class I Binding Prediction Resource (<http://mhcbindingpredictions.immuneepitope.org>) contains training data, test data, and other resources for tool developers interested in predictions of peptide binding to MHC class I

molecules. The user can follow links to a manuscript describing the resource in detail (Peters et al., PLoS Comput. Biol. 2006), a dataset of experimental affinities of peptide to MHC molecules, and a description of the framework used for the evaluation of prediction methods. A link to this site appears at the bottom of the T Cell Tools tab in the Analysis Resource.

As described in the manuscript, predictions were obtained from public web-servers for all relevant peptide-MHC affinities in the dataset. The correlation between predicted and measured affinities was evaluated using scatter plots, linear regression, and ROC analyses. The evaluation of these external tools can be accessed on the site by name of the method or the MHC allele. As carefully noted in the manuscript, this is not a fair evaluation of the value of each method, primarily because the data available to each method are highly divergent.

A similar evaluation of the prediction performance of three prediction methods available in the IEDB Analysis Resource (ANN, ARB, and SMM) was carried out using cross-validation on the dataset. In contrast to the comparison of external predictions, this is a fair evaluation of prediction performance of the three methods, since these three internal methods all used the same training data and testing data, while the external methods used a variety of training and testing data sets. Again, the evaluations of these three internal methods can be accessed by name of the method or the MHC allele.

2.4.1.1.2 Peptide Binding to MHC Class II Molecules

Users can select from six different methods for predicting class II epitopes – ARB, SMM-align, Sturniolo, Combinatorial Library, Consensus, NN-align, and NetMHCIIpan. NetMHCIIpan was added in 2010. The Consensus method has been selected as the default method. An evaluation conducted by the IEDB team has indicated that this method generally performs better than the others. The six methods are described further below. The allele selection option was also modified for all the prediction methods, which now allows users to submit multiple alleles.

SMM-align

The MHC class II binding groove is open at both ends making the correct alignment of a peptide in the binding groove a crucial part of identifying the core of an MHC class II binding motif. The stabilization matrix alignment method, SMM-align, allows for direct prediction of peptide:MHC binding affinities. The method uses amino terminal peptide flanking residues (PFR) to get a consistent gain in predictive performance by favoring binding registers with a minimum PFR length of two amino acids. The method predicts quantitative peptide:MHC binding affinity values. The method has been trained and evaluated on a data set that covers the nine HLA-DR supertypes suggested and three mouse H2-IA allele. The method is described by Nielsen et al. (BMC Bioinformatics, 2007).

Sturniolo

This matrix-based approach is used in the TEPIPOPE class II epitope prediction program. It is described in Sturniolo et al. (Nat. Biotechnol., 1999).

Combinatorial Library

The positional scanning combinatorial libraries approach utilized a pool of random peptide libraries to systematically measure the contribution to MHC binding from each amino acid at

each of the nine positions at the binding peptide. Each pool in the library contains 9-mer peptides with one fixed residue at a single position. With each of the 20 naturally occurring residues represented at each position along the 9-mer backbone, the entire library consisted of 180 peptide mixtures. Competitive binding assays were then carried out to determine the IC₅₀ values for each pool. IC₅₀ values for each mixture were standardized as a ratio to the geometric mean IC₅₀ value of the entire set of 180 mixtures, and then normalized at each position so that the value associated with the optimal value at each position corresponds to 1. For each position, an average (geometric) relative binding affinity (ARB) was calculated, and then the ratio of the ARB for the entire library to the ARB for each position was derived. The final results are a set of 9 by 20 scoring matrices which could predict the binding of novel peptides to MHC molecules.

The methodology is the same as that used for the MHC class I combinatorial library tool as described in Sidney et al. Immunome Res. 2008. A paper specifically describing the class II tool will be published in 2010.

Consensus

The consensus method was developed by the IEDB team by exploiting features of the other three aforementioned methods. The method was updated with the introduction of NN-align, so the revised Consensus method uses NN-align, SMM-align, and the combinatorial peptide scanning library. When the scanning library is not available for an allele, the Sturniolo method is used instead. A paper describing the original method was published by Wang et al. (PLoS Comput Biol, 2008).

NN-align

NN-align is an artificial neural network-based alignment algorithm for MHC class II peptide binding prediction. It simultaneously identifies the MHC class II binding core and binding affinity. The method is trained using an algorithm that corrects bias in the training data caused by redundancy in binding core representation. Prediction accuracy has been shown to improve significantly when information about the residues flanking the peptide-binding core is taken into account. A 2009 paper in BMC Bioinformatics by Nielsen and Lund describe the method in detail (PMID: 19765293).

NetMHCIIpan

NetMHCIIpan predicts binding of peptides for over 500 HLA-DR alleles using artificial neural networks. A paper describing the method was published by Nielsen et al. in Immunome Research, November 2010.

The datasets used in assessing the performance of the ARB, SMM-align, and Sturniolo methods and in developing the Consensus method, as described in Wang et al., can be found at <http://mhcbindingpredictions.immunopeptope.org/MHCII/>. The three datasets can be used for developing algorithms that predict peptides binding to MHC class II molecules and/or activating CD4+ T cells. The first is a comprehensive dataset consisting of more than 10,000 previously unpublished MHC-peptide binding affinities for 16 alleles (peptide_affinity_dataset.zip). The second dataset is a text file of 29 peptide/MHC crystal structures found in the PDB that can be used for binding core predictions (non_redundant_pdb_core_pep_allele.txt). The third dataset

contains 664 peptide sequences experimentally tested for CD4+ T-cell responses (LCMV_T_cell_activation.txt).

2.4.1.2 T Cell Epitopes – MHC I Processing Prediction

2.4.1.2.1 *Proteasomal cleavage/TAP transport/MHC class I combined predictor*

For the prediction of antigen processing through the MHC class I antigen presentation pathway, we incorporated predictions of proteasomal cleavage and TAP transport similar to the MHCPATHWAY website described in (Tenzer et al., CMLS, 2005). The predictions are based on in vitro experiments characterizing the sequence specificity of proteasomal cleavage and TAP transport. The goal of the prediction is to identify MHC-I ligands (peptides that are naturally processed from their source proteins and presented by MHC class I molecules).

The proteasomal cleavage predictions evaluate how efficiently a peptide or its N-terminally prolonged precursors can be liberated from its source protein. The TAP transport predictions evaluate how efficiently a peptide or its N-terminal prolonged precursors are transported into the ER by TAP (Peters et al., Immunol, 2003). When this information is taken together and combined with MHC class I binding predictions, the tool yields a prediction of the efficiency with which a peptide is presented on the cell surface. The same seven methods described in Section 2.4.1.1.1 are available.

2.4.1.2.2 *Neural network based prediction of proteasomal cleavage sites (NetChop) and T cell epitopes (NetCTL)*

NetChop produces neural network predictions for cleavage sites of the human proteasome (Kesmir et al., 2002). NetChop takes into account the characteristics of the structurally modified proteasomes found in cells stimulated by gamma-interferon under physiological conditions. The NetChop algorithm was trained on in vitro data and MHC Class I ligand data. The use of this training set, combined with the artificial neural network methodology, makes the prediction of cleavage sites more accurate. NetChop has been trained only on human data, but since the proteasome structure is quite conserved, the algorithm developers believe that the tool is capable of making reliable predictions for at least the other mammalian proteasomes.

NetCTL predicts CTL epitopes in protein sequences integrating prediction of peptide MHC binding, proteasomal C terminal cleavage and TAP transport efficiency. The method is described in detail in Larsen et al. (Eur J Immunol., 2005).

2.4.2 B Cell Epitope Prediction

2.4.2.1 Prediction of linear epitopes from protein sequence

Six different tools are provided that predict antibody epitope candidates from amino acid sequences. Five are based on amino acid property scales and a sixth method uses a Hidden Markov Model. Parameters such as hydrophilicity, flexibility, accessibility, and antigenic propensity of polypeptides chains have been correlated with the location of continuous epitopes in a few well-characterized proteins. Based on these observations, amino acid property scales have been developed to predict antigenic determinants. Each scale consists of 20 values

assigned to each of the amino acid residues on the basis of their relative propensity to possess the property described by the scale. The following amino acid property scales have been selected and implemented based on their popularity and coverage of different categories.

- Secondary structure - Chou and Fasman beta turn prediction
- Surface exposure - Emini surface accessibility prediction
- Flexibility - Karplus and Schulz flexibility prediction
- Antigenicity - Kolaskar and Tongaonkar antigenicity prediction
- Hydrophobicity/hydrophilicity - Parker hydrophilicity prediction

BepiPred combines the predictions of a hidden Markov model and the propensity scale of Parker et al. It is described in Larsen et al. (Immunome Research, 2006).

2.4.2.2 DiscoTope - Prediction of epitopes from protein structure

DiscoTope is designed specifically to predict discontinuous epitopes. It uses protein three-dimensional structural data in addition sequence data. The method is based on amino acid statistics, spatial information, and surface accessibility in a compiled data set of discontinuous epitopes determined by X-ray crystallography of antibody/antigen protein complexes. The method is described in Haste Andersen et al. (Protein Sci., 2006).

2.4.2.3 ElliPro - Epitope prediction based upon structural protrusion

ElliPro predicts linear and discontinuous antibody epitopes based on a protein antigen's 3D structure. ElliPro accepts either a protein structure (preferred) or a protein sequence as an input. If a protein sequence is used, ElliPro will predict its 3D structure by homology modeling. Its use is described in the Tutorial tab of the ElliPro section of the Analysis Resource. The method is described in Julia Ponomarenko et al. (BMC Bioinformatics, 2008).

2.4.3 Epitope Analysis Tools

2.4.3.1 Population coverage

T cells recognize a complex between a specific MHC type and a particular pathogen-derived epitope and thus a given epitope will elicit a response only in individuals that express an MHC molecule capable of binding that particular epitope. MHC molecules are extremely polymorphic (over a thousand different variants are known in humans). Therefore, selecting multiple peptides with different MHC binding specificities will afford increased coverage of the patient population targeted as vaccine recipients. The issue of population coverage in relation to MHC polymorphism is further complicated by the fact that different MHC types are expressed at dramatically different frequencies in different ethnicities. Thus, without careful consideration, a vaccine with ethnically biased population coverage could result. To address this issue, the actual/predicted binding capacity of potential epitopes to as many different MHC molecules possible (and when available, also restriction data of T cell responses recognizing the epitope) can be used to project the population coverage in different ethnicities of different vaccine candidates or epitope sets. Accordingly, epitope-based vaccines or diagnostics can be designed to maximize population coverage, while minimizing complexity (that is, the number of different epitopes included in the diagnostic or vaccine), and also minimizing the variability of coverage obtained or projected in different ethnic groups.

An important consideration in the process of epitope selection is that the patient population coverage afforded by a given set is not simply corresponding to the sum of the coverage of its individual components. Thus, to calculate the coverage afforded by a given mixture of epitopes, a more comprehensive approach and a suitable algorithm has been developed for this specific purpose (Bui et al. BMC Bioinformatics 2006). This method calculates the fraction of individuals predicted to respond to a given epitope set on the basis of HLA genotypic frequencies, assuming non-linkage disequilibrium between HLA loci, and on the basis of MHC binding and/or T cell restriction data. The algorithm is briefly explained here. First, genotypic frequencies of various MHC are tabulated. Each time a peptide binds to a given MHC, a “hit” is recorded for that MHC. The process is repeated for all peptides. Then the hits for MHC are tallied. Next, the frequency of each possible diploid MHC combination (phenotype) is calculated. For n MHC types, this corresponds to an $n \times n$ tabulation of the frequency at which each specific pair of MHCs will be found in the population from which the MHC frequencies are derived. A similar table is generated to contain the number of hits per each of the MHC combinations by adding the number of hits associated with each of the two alleles of MHC in the combination (a simple exception is the case of homozygous combinations, where the number of hits is simply the number of hits of the given MHC). From these two tables, a frequency distribution is assembled, tabulating the genotypic frequency of all MHC combinations associated with a certain number of hits. The result of the analysis is displayed as a frequency distribution histogram and a cumulative frequency plot.

We have derived HLA allele genotypic frequencies from the dbMHC database (<http://www.ncbi.nlm.nih.gov/mhc/>) and stored them in a database on the IEDB tool server. At present, dbMHC provides allele frequencies for 78 populations and 11 different geographical areas. It is envisioned that the compiled data will be updated regularly as further HLA frequency data become available. Furthermore, customized frequency data can be utilized in the calculation, should studies of specific and particular patient populations be of interest to a given user. Multiple population coverages can be simultaneously calculated and an average population coverage is generated. Since MHC class I and II restricted epitopes elicit immune responses from two different T cell populations (CTL and Th cells, respectively), the program provides three different coverage calculation modes – (1) class I separate, (2) class II separate, and (3) class I and class II combined.

2.4.3.2 Epitope conservancy

In a diagnostic or epitope-based vaccine setting, focusing on conserved epitopes allows for targeting responses around pathogen variability, whether it exists prior to infection, or develops in the natural course of disease. The use of conserved epitopes would be expected to focus the immune response on sequences crucial for retaining biological function of the pathogen proteins, and thus with intrinsically lower variability, even under immune pressure. The epitope conservancy analysis tools implemented here aims to address the issue of variability (or conservation) of epitopes, and to assist in the selection of epitopes with the desired pattern of conservation. The algorithm has been implemented to calculate the degree of conservancy of an epitope within a given protein sequence set at different degree of identities. The degree of conservation is defined as the number of protein sequences that contain the epitope at a given

identity level, divided by the total number of protein sequences found in the dataset analyzed (Bui et al. BMC Bioinformatics 2007).

2.4.3.3 Epitope Cluster Analysis

This tool groups epitopes into clusters based on sequence identity. A cluster is defined as a group of sequences that has a sequence similarity greater than the minimum sequence identity threshold specified. Epitope sequences can be either directly entered in the text area or uploaded from a file. Two acceptable sequence formats are PLAIN and FASTA. The user can select the sequence identity threshold at which they want to calculate epitope clusters. Clusters are displayed in a table format where clusters are indicated by table rows which have the same color. All calculated cluster results can be saved to a file by clicking on the "Download data to file" button.

2.4.3.4 Homology Mapping Tool and EpitopeViewer

This Homology Mapping Tool maps linear epitopes to 3D structures of proteins (Beaver, et al., Immunome Res 2007). This is done by comparing the epitope source protein sequence with that of proteins with known 3D structures in the PDB. The tool generates an alignment between the query sequence of the epitope source sequence and a homologous sequence from the PDB, and visualizes the result in an EpitopeViewer. For input, the tool uses the SwissProt ID of the antigen protein, the epitope sequence, and the position of the epitope in the antigen sequence as curated within the IEDB or input by the user. The tool applies the NCBI BLAST algorithm for performing sequence homology search, and provides options for the sophisticated user to choose cutoff values on parameters used in the search programs (such as e-value and penalty on gap initiation and gap extension). The tool output page displays the alignment between the query sequence of the antigen containing the epitope and the sequence from the PDB representing significant hits (matches). The region within the epitope is highlighted in the alignment, and the sequence identity for the epitope and homologous region is provided. The EpitopeViewer application for visualization of homologous epitope/antigen and its further structural analysis is launched from the output page.

Functionality was added to the homology mapping tool in 2010 to allow users to input their own conformational epitopes and map them to protein structures. This feature was documented on the "Example 5" tab of the homology mapping web page. The "Help" tab was also updated.

The convenient and easy to use EpitopeViewer, a Java application running JOGL, has been developed for three-dimensional visualization of immune epitopes and analyses of their interactions with antigen-specific receptors of the immune system (antibodies, T cell receptors, MHC molecules) for structures available in the Protein Data Bank (PDB). The EpitopeViewer is based on the Molecular Biology Toolkit (MBT; <http://mbt.sdsc.edu/>) developed at the San Diego Supercomputer Center (SDSC). It uses data both from the PDB and the IEDB, and visualizes one epitope at a time from a particular PDB structure (Beaver, et al., Immunome Res 2007).

The EpitopeViewer provides the following functionality:

- Link to the PDB web-page displaying a particular structure.

- Visualization of the 3D structure of epitope/antigen in complex with immune receptor(s) as curated within the IEDB and available in the PDB.
- Visualization of the 3D structure of epitope and antigen mapped to a PDB structure using the Homology Mapping tool.
- Visualization of sequences of epitope/antigen and immune receptor(s).
- 3D-visualization of intermolecular (epitope-paratope, epitope-antibody CDR, epitope-MHC, pMHC-TCR, pMHC-TCR CDR), inter-atom and inter-residue interactions curated within the IEDB and/or calculated on the fly from the PDB file with essential details (contact type, atoms, distance) provided;
- 2D-plot of inter-residue interactions between epitope and immune receptor.
- Generation of publication-quality pictures of structures, sequences, and plots of contacting residues.

2.5 Support Overview

The Support pull-down menu contains items that aid the user to understand and utilize the features of the website. Users can access relevant documentation, request help, and provide feedback.

2.5.1 Solutions Center

The IEDB Solutions Center is the primary resource for information on using the website's features. The user can submit help requests, check on the status of requests, browse and search the knowledge base and forums, and link to help documentation, such as the Curation Manual. In order to submit and subsequently track a help request via the Solutions Center, users must follow a simple registration procedure in order to provide an email address, name, and password. The Solutions Center can also be accessed by a link at the bottom of each web page.

2.5.2 Provide Feedback

All users are able to submit feedback, which will transmit an email message to the IEDB team. Feedback is intended for questions, input, and suggestions, such as new features they would like to see added in the future. Feedback helps the IEDB team update the system to provide users with the best possible experience. If users need help using the system or handling an unexpected result, a Help Request is more appropriate. The feedback feature can be accessed with the Support pull-down menu and the “Provide Feedback” at the bottom of each web page. Both initiate an email to feedback@iedb.org.

2.5.3 Help Request

Users can submit help requests in three ways. At the bottom of most pages is a “Help request” link. Selecting this will initiate an email to help@iedb.org. The email help request can also be initiated by selecting “Help Request” from the Support pull-down menu. The user can describe their request and send the email. The third method utilizes the IEDB Solutions Center. The top menu bar on the IEDB Solutions Center web page has a “SUBMIT A REQUEST TO THE

HELP DESK" link. Selecting this link brings the user to a "Submit a request" web page. The user must fill in their email address, the subject line, and a description of the help request. All help requests are forwarded to the same help desk.

Help requests should consist of problems that users have with the application, such as a certain function of the system not working. Questions on how to use features of the application should be covered in the online help available in the Solutions Center. After submitting a help request, a confirmation e-mail will be sent to the user's e-mail address, which will include the help request number. When the help request issue has been resolved, an e-mail will be sent to notify the user that the help request has been satisfied.

Help requests are generally responded to within one business day. Requests that are specific to analytical tools or the method used to curate data are answered initially to inform the requestor that their question/comment is being forwarded to team specialists, and an approximate date of full response is provided. Based on the complexity of the request or if that request prompts the team to make changes to the system or curate additional data/source organisms, requests are resolved immediately in some cases, while others are resolved in future IEDB system builds, or later curation.

2.5.4 Data Field Descriptions

The Data Field Descriptions item in the Support pull-down menu gives the user access to the Curation Manual. Using the Find function of their web browser, users can search for the data field name of interest in order to gain a fuller understanding of the field definition and the possible values.

2.5.5 Help with Common Queries

A new help feature was introduced in 2010. A series of 16 queries has been designed to parallel the IEDB meta-analysis efforts. Meta-analyses are performed at intervals to query the database for all data relating to subjects of interest (e.g. [Influenza](#), [Malaria](#), [Tuberculosis](#)). The queries represent a simplified version of such analysis. Two examples from the series are "How do I generate a list of all the different species and strains reported for an organism of interest?" and "How do I search for human data and specify geographic region?" The user can click on the button (+) next to each query for step-by-step instructions on how to find the answer. Data generated through use of the tutorial can be downloaded for further analysis using Excel. When available, the option to download data appears in the upper left-hand side of the page (Excel icon). Tools for performing specific analyses are also available through the Analysis Resource on the IEDB website.

2.6 More IEDB

The "More IEDB" pull-down menu contains links to other relevant information that may be of interest to users. Its component links are described below.

2.6.1 Acknowledgements

A host of talented individuals have worked hard to make the Immune Epitope Database a reality. A roster of the current team members can be viewed on this page.

2.6.2 Citing the IEDB

Data and tools within the IEDB are presented as a public resource. Users are requested to consider citing the IEDB when they present information obtained from the IEDB or use tools contained in the Analysis Resource. It is expected that the authors of an entry as well as the IEDB are properly cited whenever their work is referred to:

1. The IEDB website should be cited using the URL: www.iedb.org
2. The journal reference for the IEDB was updated after the release of IEDB 2.0. The IEDB should be cited as:

Vita R, Zarebski L, Greenbaum JA, Emami H, Hoof I, Salimi N, Damle R, Sette A, Peters B. The immune epitope database 2.0. *Nucleic Acids Res.* 2010 Jan; 38(Database issue):D854-62. Epub 2009 Nov 11.

2.6.3 Database Export

The contents of the Immune Epitope Database are exported weekly to files in XML format. The database export page (Figure 2.31) contains a complete database export in XML format along with exports of the various records that are referenced in the complete database export. The complete dataset export is provided as a ZIP archive (`iedb_export.zip`) containing one XML file for each reference contained in the database. The export also contains ZIP archives of XML files that provide users with the full record for the various identifiers that are referenced in the full database export, such as the IEDB source organism accession identifier list, the MHC allele name list, and the organism list. The supporting XML files only contain full records when the record is not otherwise available from the original source.

The database export page also provides the relevant XML Schema Definition (XSD) files for each of the XML files provided. The XSD file(s) for a particular XML file will be located in the second column of the export table. In the case of the complete database export, multiple schemas are provided, with the primary schema being listed first, followed by any supporting schemas.

One of the new features introduced in 2009 was the addition of two new database export formats. The contents of the IEDB are exported weekly as a MySQL Structure Query Language (SQL) database. The database export page contains the complete database export in two formats, a single SQL script (`iedb_public.sql.gz`), and a TAR archive of MyISAM binary tables (`iedb_public.tar.gz`). Both export formats are generated from a MySQL v5.0 database server. A physical entity relationship diagram (ERD) for the MySQL database is also available as a PDF file.

Archives of the data as they appeared in IEDB 1.0, 2.0, 2.1, and 2.3 are available by clicking on the *Archived Versions*. For each version, there is a compress file that contains an XML for each reference. The corresponding XSD files are also available for download.

Database Export		
Information about our database export features can be found in our developer's forum .		
Version 2.4		
XML Database Export		IEDB Schema
Complete Database Export	61MB	Curation.xsd (Primary IEDB schema) 46kB
		CurationSimpleTypes.xsd 418kB
		IedbPDBViewerSchema.xsd 7kB
IedbAccessionList.zip	21kB	IedbAccessionList.xsd 909B
MhcAlleleNameList.zip	18kB	MhcAlleleNameList.xsd 2kB
OrganismList.zip	11MB	OrganismList.xsd 751B
AssayTypeList.zip	4kB	AssayTypeList.xsd 771B
MySQL Database Export		Physical Entity Relationship Diagram
SQL Statement Export	67MB	iedb_public_erd.pdf 25kB
MyISAM Binary Export	130MB	
Archived Versions		

Figure 2.31 IEDB Database Export web page

2.6.4 Documents

A variety of IEDB reference materials is available for download by all users. The Documents page lists the files available for download by category. The files available for download include an Introduction to IEDB and Analysis Resource, IEDB Annual Workshop Executive Summaries, the Annual Compendia, and quarterly newsletters. Additional reference materials will be added for download over time.

2.6.5 IEDB Ontology

Selecting IEDB Ontology from the pull-down menu takes the user to <http://ontology.iedb.org>, the home page for the Ontology of Immune Epitopes (ONTIE). The IEDB ontology is available to download in a format that can be viewed with the Protégé OWL editor. The file ONTIE.owl contains the classes and relations used in the IEDB export. IEDB-export.owl is the ONTIE-annotated export of the IEDB T cell epitope data. Smaller versions of the file are available for download.

2.6.6 Links to External Sources

The IEDB system provides a list of links to external resources solely for the convenience of Immune Epitope Database visitors. The Immune Epitope Database has no interest in, responsibility for, or control over the linked-site. The Immune Epitope Database makes no promises or warranties of any kind, express or implied, including those of fitness for any particular purpose, as to the content of the linked-site. To view the links available, select Links under the Resources heading on the main menu. The hyperlinks on the links page are grouped by category:

- Public Databases, Prediction Algorithms, and Other Tools
- Antibody Related Links
- Bioinformatics Resource Centers
- MHC and TCR Related Links

- Protein Related Links
- Laboratory Resources
- Biodefense Resources

2.6.7 Patent List

As part of the IEDB curation effort, the Derwent World Patent Index has been searched for potentially relevant patent items. These are presented in a table on the Patent List web page that lists the publication number, patent title, inventors, assignee name, patent abstract, date filed, and date published. The enhanced abstracts of all of the 774 listed patent items have been reviewed, but the actual patents have not been read or curated. This information is presented for those users who wish to explore these patent items further.

The list mostly includes patents related to Category A-C priority pathogens, emerging and re-emerging infectious diseases, Malaria, Hepatitis B, Clostridium tetani, Leishmania, and Candida albicans, as well as other diseases. Users can search the table by using the "find" feature of their browser.

2.6.8 Publications

A list of publications relevant to the IEDB can be found on the Publications page. They have been grouped into four categories – General IEDB, Epitope Meta-analyses, Analysis Resource, and Curation.

2.6.9 Terms of Use

The Terms of Use page is a collection of statements that outline the conditions related to using the IEDB system. The Terms of use includes our privacy notice, copyright information, and various disclaimers. The IEDB privacy policy describes what user information is collected, the circumstances for collecting it, and how it is used. Personal information about users is not collected unless the users choose to provide it.

3 Scientific Publications

This section lists the scientific publications for which the IEDB played a contributory role. The first section lists publications authored by the IEDB contractor team over the past seven years. The second section lists references that cited the IEDB. This list was compiled by using Google Scholar and the ISI Web of Knowledge to find citations for each one of the IEDB team authored papers.

3.1 Publications of the IEDB team by Year

Scientific articles written by the IEDB team members that are relevant to the IEDB are listed below by year of publication. In 2010, nine articles were published, while seven were published in 2009, nine were published in 2008, twelve in 2007, five in 2006, six in 2005, and one in 2004. In all, the team has written 49 scientific articles about the IEDB in the last seven years.

3.1.1 2010

1. Brinkman RR, Courtot M, Derom D, Fostel JM, He Y, Lord P, Malone J, Parkinson H, Peters B, Rocca-Serra P, Ruttenberg A, Sansone SA, Soldatova LN, Stoeckert CJ Jr, Turner JA, Zheng J; OBI consortium. Modeling biomedical experimental processes with OBI. *J Biomed Semantics*. 2010 Jun 22;1 Suppl 1:S7. PubMed PMID: 20626927
2. Diehl AD, Augustine AD, Blake JA, Cowell LG, Gold ES, Gondré-Lewis TA, Masci AM, Meehan TF, Morel PA, Nijnik A, Peters B, Pulendran B, Scheuermann RH, Yao QA, Zand MS, Mungall CJ. "Hematopoietic cell types: Prototype for a revised cell ontology. *J Biomed Inform*. 2010 Feb 1. [Epub ahead of print] PMID: 20123131
3. Kim Y, Sette A, Peters B. Applications for T-cell epitope queries and tools in the Immune Epitope Database and Analysis Resource. *J Immunol Methods*. 2010 Oct 31. [Epub ahead of print] PubMed PMID: 21047510.
4. Nielsen M, Lund O, Buus S, Lundsgaard C. MHC class II epitope predictive algorithms. *Immunology*. 2010 Jul;130(3):319-28. Epub 2010 Apr 12. Review. PubMed PMID: 20408898;
5. Ponomarenko J, Papangelopoulos N, Zajonc DM, Peters B, Sette A, Bourne PE. IEDB-3D: structural data within the immune epitope database. *Nucleic Acids Res*. 2010 Oct 28. [Epub ahead of print] PubMed PMID: 21030437.
6. Salimi N, Fleri W, Peters B, Sette A. Design and utilization of epitope-based databases and predictive tools. *Immunogenetics*. 2010 Apr;62(4):185-96. Epub 2010 Mar 6. Review. PubMed PMID: 20213141; PubMed Central PMCID: PMC2843836.
7. Vaughan K, Greenbaum J, Blythe M, Peters B, Sette A. Meta-analysis of all immune epitope data in the Flavivirus genus: inventory of current immune epitope data status in the context of virus immunity and immunopathology. *Viral Immunol*. 2010 Jun;23(3):259-84. PubMed PMID: 20565291; PubMed Central PMCID: PMC2942863.
8. Wang P, Sidney J, Kim Y, Sette A, Lund O, Nielsen M, Peters B. Peptide binding predictions for HLA DR, DP and DQ molecules. *BMC Bioinformatics*. 2010 Nov 22;11:568. PubMed PMID: 21092157; PubMed Central PMCID: PMC2998531.

9. Zhang H, Wang P, Papangelopoulos N, Xu Y, Sette A, Bourne PE, Lund O, Ponomarenko J, Nielsen M, Peters B. Limitations of Ab initio predictions of peptide binding to MHC class II molecules. *PLoS One*. 2010 Feb 17;5(2):e9272. PubMed PMID: 20174654; PubMed Central PMCID: PMC2822856.

3.1.2 2009

10. Davies V, Vaughan K, Damle R, Peters B, Sette A. Classification of the universe of immune epitope literature: representation and knowledge gaps. *PLoS One*. 2009 Sep 14;4(9):e6948. PubMed PMID: 19774228;
11. Greenbaum JA, Kotturi MF, Kim Y, Oseroff C, Vaughan K, Salimi N, Vita R, Ponomarenko J, Scheuermann RH, Sette A, Peters B. Pre-existing immunity against swine-origin H1N1 influenza viruses in the general human population. *Proc Natl Acad Sci U S A*. 2009 Dec 1;106(48):20365-70. Epub 2009 Nov 16. PubMed PMID: 19918065; PubMed Central PMCID: PMC2777968.
12. Hoof I, Peters B, Sidney J, Pedersen LE, Sette A, Lund O, Buus S, Nielsen M, NetMHCpan, a method for MHC class I binding prediction beyond humans. *Immunogenetics*. 2009 Jan;61(1):1-13. PMID: 19002680
13. Kim Y, Sidney J, Pinilla C, Sette A, Peters B. Derivation of an amino acid similarity matrix for peptide: MHC binding and its application as a Bayesian prior. *BMC Bioinformatics*. 2009 Nov 30;10:394. PubMed PMID: 19948066.
14. Vaughan K, Blythe M, Greenbaum J, Zhang Q, Peters B, Doolan DL, Sette A, Meta-analysis of immune epitope data for all Plasmodia: overview and applications for malarial immunobiology and vaccine-related issues. *Parasite Immunol*. 2009 Feb;31(2):78-97. [PMID: 19149776]
15. Vita R, Zarebski L, Greenbaum JA, Emami H, Hoof I, Salimi N, Damle R, Sette A, Peters B. The Immune Epitope Database 2.0. *Nucleic Acids Res*. 2009 Nov 11. [Epub ahead of print] PubMed PMID: 19906713.
16. Zhang H, Lund O, Nielsen M. The PickPocket method for predicting binding specificities for receptors based on receptor pocket similarities: Application to MHC-peptide binding. *Bioinformatics*. 2009 Mar 17. PMID: 19297351

3.1.3 2008

17. Lundegaard C., K. Lamberth, M. Harndahl, S. Buus, O. Lund, M. Nielsen. (2008). "NetMHC-3.0: accurate web accessible predictions of human, mouse and monkey MHC class I affinities for peptides of length 8-11." *Nucleic Acids Res*. 2008 36:W509-12. PMID: 18996943
18. Nielsen, M., C. Lundegaard, T. Blicher, B. Peters, A. Sette, S. Justesen, S. Buus, O. Lund. (2008). "Quantitative Predictions of Peptide Binding to Any HLA-DR Molecule of Known Sequence: NetMHCIIpan." *PLoS Computational Biology* 4(7). PMID: 18604266
19. Ponomarenko J, H. H. Bui, W. Li, N. Fusseder, P. E. Bourne, A. Sette, B. Peters. (2008). "ElliPro: a new structure-based tool for the prediction of antibody epitopes." *BMC Bioinformatics*, 9(514). PMID: 19055730

20. Ponomarenko J. V., M. H. C. van Regenmortel (2008) "B-cell epitope prediction." Structural Bioinformatics, Ed: Bourne PE, Gu J. Wiley-Liss; 2 edition.
21. Rapin N., I. Hoof, O. Lund, M. Nielsen. (2008). "MHC motif viewer." Immunogenetics. 60(12): 759-65. PMID: 18766337
22. Vita R., B. Peters, A. Sette. (2008). "The Curation Guidelines of the Immune Epitope Database and Analysis Resource." Cytometry A 73(11): 1066-1070. PMID: 18688821
23. Wang, P., J. Sidney, C. Dow, B. Mothe, A. Sette, B. Peters. (2008). "A Systematic Assessment of MHC Class II Peptide Binding Predictions and Evaluation of a Consensus Approach." PLoS Computational Biology 4(4). PMID: 18389056
24. Zarebski L. M., K. Vaughan, J. Sidney, B. Peters, H. Grey, K. D. Janda, A. Casadevall, A. Sette. (2008). "Analysis of epitope information related to Bacillus anthracis and Clostridium botulinum." Expert Rev Vaccines 7(1): 55-74. PMID: 18251694
25. Zhang, Q., P. Wang, et al. (2008). "Immune epitope database analysis resource (IEDB-AR)." Nucleic Acids Research 36: W513-W518. PMID: 18515843

3.1.4 2007

26. Beaver JE, Bourne PE, Ponomarenko JV. EpitopeViewer: a Java application for the visualization and analysis of immune epitopes in the Immune Epitope Database and Analysis Resource (IEDB). Immunome Res. 2007 Feb 21;3:3. PMID: 17313688
27. Blythe MJ, Zhang Q, Vaughan K, de Castro R Jr, Salimi N, Bui HH, Lewinsohn DM, Ernst JD, Peters B, Sette A. An analysis of the epitope knowledge related to Mycobacteria. Immunome Res. 2007 Dec 14;3(1):10. PMID: 18081934
28. Bui HH, Peters B, Assarsson E, Mbawuike I, Sette A. Ab and T cell epitopes of influenza A virus, knowledge and opportunities. Proc Natl Acad Sci U S A. 2007 Jan 2;104(1):246-51. PMID: 17200302
29. Bui HH, Sidney J, Li W, Fusseder N, Sette A. Development of an epitope conservancy analysis tool to facilitate the design of epitope-based diagnostics and vaccines. BMC Bioinformatics. 2007 Sep 26;8(1):361. PMID: 17897458
30. Ernst JD, Lewinsohn DM, Behar S, Blythe M, Schlesinger LS, Kornfeld H, Sette A. Meeting Report: NIH Workshop on the Tuberculosis Immune Epitope Database. Tuberculosis (Edinb). 2007 Dec 6. PMID: 18068490
31. Greenbaum JA, Andersen PH, Blythe M, Bui HH, Cachau RE, Crowe J, Davies M, Kolaskar AS, Lund O, Morrison S, Mumey B, Ofran Y, Pellequer JL, Pinilla C, Ponomarenko JV, Raghava GP, van Regenmortel MH, Roggen EL, Sette A, Schlessinger A, Sollner J, Zand M, Peters B. Towards a consensus on datasets and evaluation metrics for developing B-cell epitope prediction tools. J Mol Recognit. 2007 Mar-Apr;20(2):75-82. PMID: 17205610
32. Lundsgaard C, Lund O, Kesmir C, Brunak S, Nielsen M. Modeling the adaptive immune system: predictions and simulations. Bioinformatics. 2007 Dec 15;23(24):3265-75. PMID: 18045832

33. Nielsen M, Lundsgaard C, Blicher T, Lamberth K, Harndahl M, Justesen S, Røder G, Peters B, Sette A, Lund O, Buus S. NetMHCpan, a method for quantitative predictions of peptide binding to any HLA-A and -B locus protein of known sequence. PLoS ONE. 2007 Aug 29;2(8):e796. PMID: 17726526
34. Peters B, Sette A. Integrating epitope data into the emerging web of biomedical knowledge resources. Nat Rev Immunol. 2007 Jun;7(6):485-90. PMID: 17479127
35. Ponomarenko JV, Bourne PE., Antibody-protein interactions: benchmark datasets and prediction tools evaluation. BMC Struct Biol. 2007 Oct 2;7(1):64. PMID: 17910770
36. Sette A, Peters B., Immune epitope mapping in the post-genomic era: lessons for vaccine development. Curr Opin Immunol. 2007 Feb;19(1):106-10. PMID: 17113275
37. Wang P, Morgan AA, Zhang Q, Sette A, Peters B. Automating document classification for the Immune Epitope Database. BMC Bioinformatics. 2007 Jul 26;8:269. PMID: 17655769

3.1.5 2006

38. Bui HH, Sidney J, Dinh K, Southwood S, Newman MJ, Sette A. Predicting population coverage of T-cell epitope-based diagnostics and vaccines. BMC Bioinformatics. 2006 Mar 17;7(1):153. PMID: 16545123
39. Peters B, Bui HH, Frankild S, Nielson M, Lundsgaard C, Kostem E, Basch D, Lamberth K, Harndahl M, Fleri W, Wilson SS, Sidney J, Lund O, Buus S, Sette A., A community resource benchmarking predictions of peptide binding to MHC-I molecules, PLoS Comput Biol. 2006 Jun 9;2(6):e65. Epub 2006 Jun 9. PMID: 16789818
40. Salimi N, R. Vita. (2006). "The biocurator: connecting and enhancing scientific data". PLoS Comput Biol 2(10) :e125. PMID: 17069454
41. Sette, A., H. H. Bui, S. Buus, W. Fleri, R. Kubo, O. Lund, D. Nemazee, J. V. Ponomarenko, M. Sathiamurthy, S. Stewart, S. Way, S. S. Wilson, B. Peters. (2006). "The Immune Epitope Database and Analysis Resource." Pattern Recognition in Bioinformatics: International Workshop, Hong Kong, China, August 20, 2006 : Proceedings, Springer, Jagath C. Rajapakse, Limsoon Wong, Raj Acharya (Eds). ISBN: 3540374469 9783540374466
42. Vita R, K. Vaughan, L. Zarebski, N. Salimi, W. Fleri , H. Grey, M. Sathiamurthy, J. Mokili, H. H. Bui, P. E. Bourne, J. V. Ponomarenko, R. de Castro Jr, R. K. Chan, J. Sidney, S. S. Wilson, S. Stewart, S. Way, B. Peters, A. Sette. (2006) "Curation of complex, context-dependent immunological data." BMC Bioinformatics. 12(7): 341. PMID: 16836764

3.1.6 2005

43. Bui HH, Sidney J, Peters B, Sathiamurthy M, Sinichi A, Purton KA, Mothé BR, Chisari FV, Watkins DI, Sette A. Automated generation and evaluation of specific MHC binding predictive tools: ARB matrix applications. Immunogenetics. 2005 Jun;57(5):304-14. Epub 2005 May 3. PMID: 15868141

44. Peters B, Sette A. Generating quantitative models describing the sequence specificity of biological processes with the stabilized matrix method. *BMC Bioinformatics*. 2005 May 31;6:132. PMID: 15927070
45. Peters, B., J. Sidney, P. Bourne, H. H. Bui, S. Buus, G. Doh, W. Flerl, M. Kronenberg, R.Kubo, O. Lund, D. Nemazee, J. V. Ponomarenko, M. Sathiamurthy, S. Schoenberger, S. Stewart, P. Surko, S. Way, S. Wilson, A. Sette. (2005). "The design and implementation of the immune epitope database and analysis resource." *Immunogenetics* 57(5): 326-336. PMID: 15895191
46. Peters, B., J. Sidney, P. Bourne, H. H. Bui, S. Buus, G. Doh, W. Flerl, M. Kronenberg, R.Kubo, O. Lund, D. Nemazee, J. V. Ponomarenko, M. Sathiamurthy, S. Schoenberger, S. Stewart, P. Surko, S. Way, S. Wilson, A. Sette. (2005). "The Immune Epitope Database and Analysis Resource: From Vision to Blueprint." *PLoS Biology* 3(3). PMID: 15760272
47. Sathiamurthy, M., B. Peters, H. H. Bui, J. Sidney, J. Mokili, S. S. Wilson, W. Flerl, D. McGuinness, P. Bourne, A. Sette. (2005). "An ontology for immune epitopes: application to the design of a broad scope database of immune reactivities." *Immunome Res* 1(2): 1745-7580. PMID: 16305755
48. Sette, A., W. Flerl, B. Peters, M. Sathiamurthy, H. H. Bui, S. Wilson. (2005). "A Roadmap for the Immunomics of Category A-C Pathogens." *Immunity* 22(2): 155-161. PMID: 15773067

3.1.7 2004

49. Sette A. (2004). "The immune epitope database and analysis resource: from vision to blueprint." *Genome Inform Ser Workshop Genome Inform* 15(2):299. PMID: 16312048

3.2 Publications Citing the IEDB in 2010

In 2010, 277 references cited at least one of the 49 papers written by the IEDB team over the past six years. This list was compiled by using Google Scholar and the ISI Web of Knowledge to find citations for each one of the 49 papers. The list excludes self-citations by the 49 IEDB papers. This total may increase over time since these databases may not include references published late in 2010.

The papers have been categorized by the type of IEDB paper they cited – General IEDB (G), Analysis Resource (RA), Curation (C), or Meta-analysis (MA). All papers listed below denote their category or categories with these abbreviations since a paper can cite several IEDB papers that belong in different categories. For example, a paper that cites an analysis resource paper may also cite the primary IEDB PLoS Biology (2005) paper, and thus is listed under General IEDB and has an [AR, G] to indicate the Analysis Resource citation. Any paper that has “General” as one of its categories has been listed in the General category below. All papers having “Analysis Resource” as a category and not having “General” have been listed in the “Analysis Resource” list below. A similar hierarchy has been applied for the Curation papers. That means all papers listed under “Meta-analysis” below only belong to that category. belonging to the General category and one orIf papers belong to more than one category, they have been placed in the first Papers belonging to more than one category are listed a paper cites

an additional IEDB reference that is in a different category, the additional category is denoted with one of the three initials given above.

The “General” category IEDB papers were cited 97 times in 2010. The Analysis Resource papers received 143 citations, more than any other category. The Curation category papers received the fewest citations at eight. The meta-analysis papers received the most growth in papers at 87, as opposed to 23 in 2009, representing a growth of 280%.

3.2.1 General IEDB

1. Anastas, P., Bejatolah, M. K., and Gajendra, P. S. (2010). "Bridging Innate and Adaptive Antitumor Immunity Targeting Glycans." *Journal of Biomedicine and Biotechnology* 2010: 354068. **[AR, G]**
2. Ansari, H. R., Flower, D. R., and Raghava, G. P. S. (2010). "AntigenDB: an immunoinformatics database of pathogen antigens." *Nucleic Acids Research* 38(Database issue): D847-D853. **[G]**
3. Antonets, D. V. and Maksyutov, A. Z. (2010). "TEpredict: Software for T-Cell epitope prediction." *Molecular Biology* 44(1): 119-127. **[AR, G]**
4. Arens, R. and Schoenberger, S. P. (2010). "Plasticity in programming of effector and memory CD8+ T cell formation." *Immunological Reviews* 235(1): 190-205. **[G]**
5. Augustine, A. D., Cassetti, M. C., Ennis, F. A., Harris, E., Hildebrand, W. H., and Repik, P. M. (2010). "NIAID Workshop on Flavivirus Immunity." *Viral Immunology* 23(3): 235-240. **[MA, G]**
6. Baillie, L. W., Huwar, T. B., Moore, S., Mellado-Sanchez, G., Rodriguez, L., Neeson, B. N., Flick-Smith, H. C., Jenner, D. C., Atkins, H. S., and Ingram, R. J. (2010). "An anthrax subunit vaccine candidate based on protective regions of *Bacillus anthracis* protective antigen and lethal factor." *Vaccine* 28(41): 6740-6748. **[G]**
7. Blanchard, N. and Shastri, N. (2010). "Cross presentation of peptides from intracellular pathogens by MHC class I molecules." *Annals of the New York Academy of Sciences* 1183(1): 237-250. **[G]**
8. Bodewes, R., Osterhaus, A. D. M. E., and Rimmelzwaan, G. F. (2010). "Targets for the Induction of Protective Immunity Against Influenza A Viruses." *Viruses-Basel* 2(1): 166-188. **[G]**
9. Bordner, A. J. and Mittelmann, H. D. (2010). "MultiRTA: A simple yet reliable method for predicting peptide binding affinities for multiple class II MHC allotypes." *BMC Bioinformatics* 11(1): 482. **[AR, G]**
10. Bordner, A. J. and Mittelmann, H. D. (2010). "Prediction of the binding affinities of peptides to class II MHC using a regularized thermodynamic model." *BMC Bioinformatics* 11(1): 41. **[AR, G]**
11. Bordner, A. J. and Morris, R. J. (2010). "Towards Universal Structure-Based Prediction of Class II MHC Epitopes for Diverse Allotypes." *PloS ONE* 5(12): 599-604. **[AR, G]**
12. Botten, J., Sidney, J., Mothe, B. R., Peters, B., Sette, A., and Kotturi, M. F. (2010). "Coverage of related pathogenic species by multivalent and cross-protective vaccine design: arenaviruses as a model system." *Microbiology and Molecular Biology Reviews* 74(2): 157-170. **[AR, G]**
13. Bremel, R. D. and Homan, E. J. (2010). "An integrated approach to epitope analysis I: Dimensional reduction, visualization and prediction of MHC binding using amino acid principal components and regression approaches." *Immunome Research* 6:7. **[AR, G]**
14. Brinkac, L. M., Davidsen, T., Beck, E., Ganapathy, A., Caler, E., Dodson, R. J., Durkin, A. S., Harkins, D. M., Lorenzi, H., and Madupu, R. (2010). "Pathema: a clade-specific bioinformatics resource center for pathogen research." *Nucleic Acids Research* 38(Database issue): D408-D414. **[G]**

15. Brinkman, R. R., Courtot, M., Derom, D., Fostel, J. M., He, Y., Lord, P., Malone, J., Parkinson, H., Peters, B., and Rocca-Serra, P. (2010). "Modeling biomedical experimental processes with OBI." *Journal of Biomedical Semantics* 1(Suppl 1): S7. **[G]**
16. Bryson, C. J., Jones, T. D., and Baker, M. P. (2010). "Prediction of Immunogenicity of Therapeutic Proteins: Validity of Computational Tools." *BioDrugs* 24(1): 1-8. **[AR, G]**
17. Burek, P., Herre, H., Roeder, I., Glauche, I., Scherf, N., and Loffler, M. (2010). "Towards a Cellular Genealogy Ontology." Proceedings of the 2nd Workshop on 'Ontologien in Biomedizin und Lebenswissenschaften' (OBML), Mannheim, Germany; IMISE-Reports 2: 59-63; www.glauche.org/download/obml2010report_burek.pdf. **[G]**
18. Calis, J. J. A., Sanchez-Perez, G. F., and Kesmir, C. (2010). "MHC class I molecules exploit the low G+C content of pathogen genomes for enhanced presentation." *European Journal of Immunology* 40(10): 2699-2709. **[AR, G]**
19. Caoili, S.E.C. (2010). "Benchmarking B-Cell Epitope Prediction for the Design of Peptide-Based Vaccines: Problems and Prospects." *Journal of Biomedicine and Biotechnology* 2010: 910524; 10.1155/2010/910524. **[AR, C]**
20. Caoili, S.E.C. (2010). "Immunization with peptide-protein conjugates: impact on benchmarking B-cell epitope prediction for vaccine design." *Protein and Peptide Letters* 17(3): 386-398. **[AR, C]**
21. Cardenas, C., Bidon-Chanal, A., Conejeros, P., Arenas, G., Marshall, S., and Luque, F. J. (2010). "Molecular modeling of class I and II alleles of the major histocompatibility complex in *Salmo salar*." *Journal of Computer-Aided Molecular Design* 24(12): 1035-1051. **[AR, G]**
22. Chung, J. L., Sun, J., Sidney, J., Sette, A., and Peters, B. (2010). "IMMUNOCAT-a data management system for epitope mapping studies." *Journal of Biomedicine and Biotechnology* 2010: 856842. **[MA, G]**
23. Cochrane, G. R. and Galperin, M. Y. (2010). "The 2010 Nucleic Acids Research database issue and online database collection: a community of data resources." *Nucleic Acids Research* 38(Database issue): D1-D4. **[G]**
24. de Verteuil, D., Muratore-Schroeder, T. L., Granados, D. P., Fortier, M. H., Hardy, M. P., Bramouille, A., Caron E., Vincent, K., Mader, S., Lemieux, S., Thibault, P., and Perreault, C. (2010). "Deletion of immunoproteasome subunits imprints on the transcriptome and has a broad impact on peptides presented by major histocompatibility complex I molecules." *Molecular & Cellular Proteomics* 9(9): 2034-2047. **[G]**
25. Diez-Rivero, C. M., Chenlo, B., Zuluaga, P., and Reche, P. A. (2010). "Quantitative modeling of peptide binding to TAP using support vector machine." *Proteins: Structure, Function, and Bioinformatics* 78(1): 63-72. **[G]**
26. Diez-Rivero, C. M., Lafuente, E. M., and Reche, P. A. (2010). "Computational analysis and modeling of cleavage by the immunoproteasome and the constitutive proteasome." *BMC Bioinformatics* 11: 479. **[AR, G]**
27. Dimitrov, I., Garnev, P., Flower, D. R., and Doytchinova, I. (2010). "EpiTOP-a proteochemometric tool for MHC class II binding prediction." *Bioinformatics* 26(16): 2066-2068. **[AR, G]**
28. Dinakarpandian, D. and Dinakar, C. (2010). "Bioinformatics in Allergy: A Powerful Tool Joining Science and Clinical Applications." *Allergy Frontiers: Future Perspectives*, Springer Japan: 19-30; doi: 10.1007/978-4-431-99365-0. **[G]**
29. Durward, M. A., Harms, J., Magnani, D. M., Eskra, L., and Splitter, G. A. (2010). "Discordant *Brucella melitensis* Antigens Yield Cognate CD8+ T Cells In Vivo." *Infection and Immunity* 78(1): 168-176. **[AR, G]**
30. Duvvuri, V. R. S. K., Moghadas, S. M., Guo, H., Duvvuri, B., Heffernan, J. M., Fisman, D. N., Wu, G. E., and Wu, J. (2010). "Original Article: Highly conserved cross reactive CD4+ T cell HA

- epitopes of seasonal and the 2009 pandemic influenza viruses." *Influenza and Other Respiratory Viruses* 4(5): 249-258. [AR, MA, G]
31. Flower, D., Macdonald, I., Ramakrishnan, K., Davies, M., and Doytchinova, I. (2010). "Computer aided selection of candidate vaccine antigens." *Immunome Research* 6(Suppl 2): S1. [AR, C, G]
 32. Garcia, A., O'Neill, K., Garcia, L. J., Lord, P., Stevens, R., Corcho, O., and Gibson, F. (2010). "Developing Ontologies within Decentralised Settings." *Semantic E-Science, Annals of Information Systems* 11: 99-139; doi: 10.1007/978-1-4419-5908-9-4. [G]
 33. Garcia-Boronat, M., Diez-Rivero, C.M., and Reche, P. A. (2010). "TEPIDAS: A DAS server for integrating T-cell epitope annotations." *Immunomics Review: Bioinformatics for Immunomics*, Springer New York 3: 57-65; doi: 10.1007/978-1-4419-0540-6. [G]
 34. Gowthaman, U., Chodisetti, S. B., Parihar, P., and Agrewala, J. N. "Evaluation of different generic in silico methods for predicting HLA class I binding peptide vaccine candidates using a reverse approach." *Amino Acids* 39(5): 1333-1342. [AR, G]
 35. Halling-Brown, M., Pappalardo, F., Rapin, N., Zhang, P., Alemani, D., Emerson, A., Castiglione, F., Duroux, P., Pennisi, M., and Miotto, O. (2010). "ImmunoGrid: towards agent-based simulations of the human immune system at a natural scale." *Philosophical Transactions of the Royal Society A* 368(1920): 2799-2815; doi: 10.1098/rsta.2010.0067. [AR, G]
 36. He, Y. and Xiang, Z. (2010). "Bioinformatics analysis of Brucella vaccines and vaccine targets using VIOLIN." *Immunome Research* 6(Suppl 1): S5. [G]
 37. He, Y., Xiang, Z., and Mobley, H.L. (2010). "Vaxign: the first web-based vaccine design program for reverse vaccinology and applications for vaccine development." *Journal of Biomedicine and Biotechnology*: 297505; doi: 10.1155/2010/297505. [AR, G]
 38. Hemavathi, K., Seenivasagam, R., Praba, V. C., and Lavanyashree, K. (2010). "Identification and analysis of epitope-based malarial vaccine targets using computational biology." *International Journal of Immunological Studies* 1(2): 195-213. [AR, G]
 39. Infantes, S., Lorente, E., Barnea, E., Beer, I., Cragnolini, J. J., Garcia, R., Lasala, F., Jimenez, M., Admon, A., and Lopez, D. (2010). "Multiple, Non-conserved, Internal Viral Ligands Naturally Presented by HLA-B27 in Human Respiratory Syncytial Virus-infected Cells." *Molecular & Cellular Proteomics* 9(7): 1533-1539. [G]
 40. Ingram, R. J., Metan, G., Maillere, B., Doganay, M., Ozkul, Y., Kim, L. U., Baillie, L., Dyson, H., Williamson, E. D., and Chu, K. K. (2010). "Natural Exposure to Cutaneous Anthrax Gives Long-Lasting T Cell Immunity Encompassing Infection-Specific Epitopes." *The Journal of Immunology* 184(7): 3814-3821. [G]
 41. Jaaskelainen, S., Riikonen, P., Salakoski, T., and Vihinen, M. (2010). "Accuracy of protein hydropathy predictions." *International Journal of Data Mining and Bioinformatics* 4(6): 735-754. [G]
 42. Jager, N., Wisniewska, J. M., Hiss, J. A., Freier, A., Losch, F. O., Walden, P., Wrede, P., and Schneider, G. (2010). "Attractors in Sequence Space: Agent Based Exploration of MHC I Binding Peptides." *Molecular Informatics* 29(1-2): 65-74. [G]
 43. James, E. A., Moustakas, A. K., Bui, J., Papadopoulos, G. K., Bondinas, G., Buckner, J. H., and Kwok, W. W. "DR1001 presents "altered-self" peptides derived from joint associated proteins by accepting citrulline in three of its binding pockets." *Arthritis & Rheumatism* 62(10): 2909-2918. [G]
 44. Karunakaran, K. P., Yu, H., Foster, L. J., and Brunham, R. C. (2010). "Development of a Chlamydia trachomatis T cell vaccine." *Human Vaccines* 6(8): 676-680. [G]
 45. Kosmrlj, A., Read, E. L., Qi, Y., Allen, T. M., Altfeld, M., Deeks, S. G., Pereyra, F., Carrington, M., Walker, B. D., and Chakraborty, A. K. (2010). "Effects of thymic selection of the T-cell repertoire on HLA class I-associated control of HIV infection." *Nature* 465(7296): 350-354. [AR, G]

46. Kumar, N. and Mohanty, D. (2010). "Structure-based identification of MHC binding peptides: Benchmarking of prediction accuracy." *Molecular Biosystems* 6(12): 2508-2520. [AR, G]
47. Lata, S. and Raghava, G. P. S. (2010). "Databases and Web-Based Tools for Innate Immunity." *Immunomics Reviews: Bioinformatics for Immunomics*, Springer 3: 67-76; doi: 10.1007/978-1-4419-0540-6-6. [G]
48. Liepe, J., Mishto, M., Textoris-Taube, K., Janek, K., Keller, C., Henklein, P., Kloetzel, P. M., and Zaikin, A. (2010). "The 20S proteasome splicing activity discovered by SpliceMet." *PLoS Comput Biol* 6(6): e1000830. [G]
49. Li Pira, G., Federico, I., Paolo, M., and Fabrizio, M. (2010). "High Throughput T Epitope Mapping and Vaccine Development." *Journal of Biomedicine and Biotechnology* 2010: 325720. [AR, C, G]
50. Liu, J., Wang, L., and Zhu, S. (2010). "SVR-PAIRWISE method to predict MHC-II binding peptides." *International Journal of Bioinformatics Research and Applications* 6(1): 58-68. [AR, G]
51. Lombardi, R., Harakava, R., and Colariccio, A. (2010). "Cloning and purification of Banana streak OL virus coat protein fragment." *Pesquisa Agropecuaria Brasileira* 45(8): 811-817. [G]
52. Lucchese, G., Stufano, A., and Kanduc, D. (2010). "Proposing low-similarity peptide vaccines against Mycobacterium tuberculosis." *Journal of Biomedicine and Biotechnology* 2010: 832341. [MA, G]
53. Lundsgaard, C., Lund, O., Buus, S., and Nielsen, M. (2010). "Major histocompatibility complex class I binding predictions as a tool in epitope discovery." *Immunology* 130(3): 309-318. [AR, G]
54. Lundsgaard, C., Hoof, I., Lund, O., and Nielsen, M. (2010). "State of the art and challenges in sequence based T-cell epitope prediction." *Immunome Research* 6(Suppl 2): S3. [AR, G]
55. Madhumathi, J., Pradiba, D., Prince, P. R., Jeyaprita, P. J., Rao, D. N., and Kaliraj, P. "Crucial epitopes of Wuchereria bancrofti abundant larval transcript recognized in natural infection." *European Journal of Clinical Microbiology & Infectious Diseases* 29(12): 1481-1486. [G]
56. Madhumathi, J., Prince, P. R., Anugraha, G., Kiran, P., Rao, D. N., Reddy, M. V. R., and Kaliraj, P. (2010). "Identification and characterization of nematode specific protective epitopes of Brugia malayi TRX towards development of synthetic vaccine construct for lymphatic filariasis." *Vaccine* 28(31): 5038-5048. [G]
57. Madhurantakam, C., Nilsson, O. B., Uchtenhagen, H., Konradsen, J., Saarne, T., Hogbom, E., Sandalova, T., Gronlund, H., and Achour, A. (2010). "Crystal Structure of the Dog Lipocalin Allergen Can f 2: Implications for Cross-reactivity to the Cat Allergen Fel d 4." *Journal of Molecular Biology* 401(1): 68-83. [G]
58. Magnan, C. N., Zeller, M., Kayala, M. A., Vigil, A., Randall, A., Felgner, P. L., and Baldi, P. (2010). "High-throughput prediction of protein antigenicity using protein microarray data." *Bioinformatics* 26(23): 2936-2943. [AR, G]
59. Moutaftsi, M., Tscharke, D. C., Vaughan, K., Koelle, D. M., Stern, L., Calvo-Calle, M., Ennis, F., Terajima, M., Sutter, G., and Crotty, S. (2010). "Uncovering the interplay between CD8, CD4 and antibody responses to complex pathogens." *Future Microbiology* 5(2): 221-239. [AR, MA, G]
60. Musson, J. A., Ingram, R., Durand, G., Ascough, S., Waters, E. L., Hartley, M. G., Robson, T., Maillere, B., Williamson, E. D., Sriskandan, S., Altmann, D., and Robinson, J.H. (2010). "Repertoire of HLA-DR1-Restricted CD4 T-Cell Responses to Capsular Caf1 Antigen of Yersinia pestis in Human Leukocyte Antigen Transgenic Mice." *Infection and immunity* 78(10): 4356-4362. [G]
61. Nakamura, Y., Komiyama, T., Furue, M., Gojobori, T., and Akiyama, Y. (2010). "CIG-DB: the database for human or mouse immunoglobulin and T cell receptor genes available for cancer studies." *BMC Bioinformatics* 11: 398; doi: 10.1186/1471-2105-11-398. [G]

62. Ng, X. R., Tan, T. W., and Tong, J. C. "Pathogen-Specific Motifs of HLA Class I T-Cell Epitopes: their possibilities and values." http://www.nus.edu.sg/nurop/2010/Proceedings/FoS/Biochemistry/Ng%20Xin%20Ru%20Marie_U062300W.pdf. [G]
63. Nielsen, M., Lund, O., Buus, S., and Lundsgaard, C. (2010). "MHC Class II epitope predictive algorithms." *Immunology* 130(3): 319-328. [AR, G]
64. Nijveen, H., Kester, M. G. D., Hassan, C., Viars, A., de Ru, A. H., de Jager, M., Falkenburg, J. H. F., Leunissen, J. A. M., and van Veelen, P. A. (2010). "HSPVdb - the Human Short Peptide Variation Database for improved mass spectrometry-based detection of polymorphic HLA-ligands." *Immunogenetics: e*Pub ahead of print; doi: 10.1007/s00251-010-0497-1. [G]
65. Panov, P., Dzeroski, S., and Soldatova, L. N. (2010). "Representing Entities in the OntoDM Data Mining Ontology." *Inductive Databases and Constraint-Based Data Mining*, Springer New York: 27-58, doi: 10.1007/978-1-4419-7738-0_2. http://kt.ijs.si/pance_panov/OntoDM/. [G]
66. Parker, A. S., Zheng, W., Griswold, K. E., and Bailey-Kellogg, C. (2010). "Optimization algorithms for functional deimmunization of therapeutic proteins." *BMC Bioinformatics* 11: 180; doi: 10.1186/1471-2105-11-180. [AR, G]
67. Paul, S. and Piontkivska, H. (2010). "Frequent associations between CTL and T-Helper epitopes in HIV-1 genomes and implications for multi-epitope vaccine designs." *BMC Microbiology* 10: 212; doi: 10.1186/1471-2180-10-212. [AR, G]
68. Prechl, J., Papp, K., and Erdei, A. (2010). "Antigen microarrays: descriptive chemistry or functional immunomics?" *Trends in Immunology* 31(4): 133-137. [G]
69. Qian, Y., Campbell, J., Kong, M., Liu, Y., and Scheuermann, R. H. "FLOCK: a density-based clustering method for automated identification and comparison of cell populations in high-dimensional flow cytometry data." FlowCAP Summit 2010, U.S. NIH Campus: flowcap.flowsite.org/summit2010/slides/10_Qian.pdf. [G]
70. Rapin, N., Hoof, I., Lund, O., and Nielsen, M. (2010). "The MHC motif viewer: a visualization tool for MHC binding motifs." *Current Protocols in Immunology* 88: 18.17.1-18.17.13; doi: 10.1002/0471142735.im1817s88. [AR, G]
71. Ridge, P. G. and Crockett, D. K. (2010). "Feature selection for characterizing HLA class I peptide motif anchors." *Proceedings of the First ACM International Conference on Bioinformatics and Computational Biology*: 644-645; doi: 10.1145/1854776.1854896. [G]
72. Roomp, K., Antes, I., and Lengauer, T. (2010). "Predicting MHC class I epitopes in large datasets." *BMC Bioinformatics* 11: 90; 10.1186/1471-2105-11-90. [AR, G]
73. Sassetti, C. M. and Rubin, E. J. (2010). "Relics of selection in the mycobacterial genome." *Nature Genetics* 42(6): 476-478. [G]
74. Schotsaert, M., Ibanez, L.I., Fiers, W., and Saelens, X. (2010). "Controlling Influenza by Cytotoxic T-Cells: Calling for Help from Destroyers." *Journal of Biomedicine and Biotechnology* 2010: 863985; doi: 10.1155/2010/863985. [AR, C, MA, G]
75. Siddiqui, S., Tarrab, E., Lamarre, A., and Basta, S. (2010). "Altered immunodominance hierarchies of CD8+ T cells in the spleen after infection at different sites is contingent on high virus inoculum." *Microbes and Infection* 12(4): 324-330. [G]
76. Sidney, J., Steen, A., Moore, C., Ngo, S., Chung, J., Peters, B., and Sette, A. (2010). "Divergent Motifs but Overlapping Binding Repertoires of Six HLA-DQ Molecules Frequently Expressed in the Worldwide Human Population." *The Journal of Immunology* 185(7): 4189-4198. [AR, G]
77. Sidney, J., Steen, A., Moore, C., Ngo, S., Chung, J., Peters, B., and Sette, A. (2010). "Five HLA-DP molecules frequently expressed in the worldwide human population share a common HLA supertypic binding specificity." *The Journal of Immunology* 184(5): 2492-2503. [G]

78. Smith, B. and Ceusters, W. (2010). "Ontological realism: A methodology for coordinated evolution of scientific ontologies." *Applied Ontology* 5(3): 139-188. **[G]**
79. Soldatova, L. N., Lord, P., Sansone, S. A., Stephens, S. M., and Shah, N. H. (2010). "Selected papers from the 12th annual Bio-Ontologies meeting." *Journal of Biomedical Semantics* 1(Suppl 1): I1. **[G]**
80. Stocki, P., Morris, N. J., Preisinger, C., Wang, X. N., Kolch, W., Multhoff, G., and Dickinson, A. M. (2010). "Identification of potential HLA class I and class II epitope precursors associated with heat shock protein 70 (HSPA)." *Cell Stress and Chaperones* 15(5): 729-741. **[G]**
81. Swertz, M. A., Velde, K. J., Tesson, B., Scheltema, R., Arends, D., Vera, G., Alberts, R., Dijkstra, M., Schofield, P., Schughart, K., Hancock, J.M., Smedley, D., Wolstencroft, K., Goble, C., de Brock, E.O., Jones, A.R., Parkinson, H.E., members of the CASIMIR, GEN2PHEN consortiums, and Jansen, R.C. (2010). "XGAP: a uniform and extensible data model and software platform for genotype and phenotype experiments." *Genome Biology* 11(3): R27; doi: 10.1186/gb-2010-11-3-r27. **[G]**
82. Tan, P. T. J., Heiny, A. T., Miotto, O., Salmon, J., Marques, E. T., Lemonnier, F., and August, J. T. (2010). "Conservation and diversity of influenza A H1N1 HLA-restricted T cell epitope candidates for epitope-based vaccines." *PloS ONE* 5(1): e8754; doi: 10.1371/journal.pone.0008754. **[G]**
83. Tomar, N. and De, R. K. "Immunoinformatics: an integrated scenario." *Immunology* 131(2): 153-168; doi: 10.1111/j.1365-2567.2010.03330.x. **[AR, G]**
84. Tong, J. C., Simarmata, D., Lin, R. T. P., Renia, L., and Ng, L. F. P. (2010). "HLA Class I Restriction as a Possible Driving Force for Chikungunya Evolution." *PloS ONE* 5(2): e9291; doi: 10.1371/journal.pone.0009291. **[AR, G]**
85. Totrov, M., Jiang, X. Q., Kong, X. P., Cohen, S., Krachmarov, C., Salomon, A., Williams, C., Seaman, M. S., Cardozo, T., Gorny, M. K., Wang, S. X., Lu, S., Pinter, A., and Zolla-Pazner, S. (2010). "Structure-guided design and immunological characterization of immunogens presenting the HIV-1 gp120 V3 loop on a CTB scaffold." *Virology* 405(2): 513-523. **[AR, G]**
86. Vielemeyer, O., Nizak, C., Jimenez, A. J., Echard, A., Goud, B., Camonis, J., Rain, J. C., and Perez, F. (2010). "Characterization of single chain antibody targets through yeast two hybrid." *BMC Biotechnology* 10: 59; doi: 10.1186/1472-6750-10-59. **[G]**
87. Wahl, A., Schafer, F., Bardet, W., and Hildebrand, W. H. (2010). "HLA class I molecules reflect an altered host proteome after influenza virus infection." *Human Immunology* 71(1): 14-22. **[G]**
88. Warren, R. L. and Holt, R. A. (2010). "A census of predicted mutational epitopes suitable for immunologic cancer control." *Human Immunology* 71(3): 245-254. **[AR, G]**
89. Wisniewska, J. M., Jager, N., Freier, A., Losch, F. O., Wiesmller, K. H., Walden, P., Wrede, P., Schneider, G., and Hiss, J. A. (2010). "MHC I Stabilizing Potential of Computer-Designed Octapeptides." *Journal of Biomedicine and Biotechnology* 2010: 396847; doi: 10.1155/2010/396847. **[AR, G]**
90. Wullner, D., Zhou, L., Bramhall, E., Kuck, A., Goletz, T. J., Swanson, S., Chirmule, N., and Jawa, V. (2010). "Considerations for optimization and validation of an in vitro PBMC derived T cell assay for immunogenicity prediction of biotherapeutics." *Clinical Immunology* 137(1): 5-14. **[AR, G]**
91. Yan, Q. (2010). "Immunoinformatics and systems biology methods for personalized medicine." *Systems Biology in Drug Discovery and Development, Methods in Molecular Biology*, Springer 662(Pt 2): 203-220; doi: 10.1007/978-1-60761-800-3_10. **[G]**
92. Yang, B., Sayers, S., Xiang, Z., and He, Y. (2010). "Protegen: a web-based protective antigen database and analysis system." *Nucleic Acids Research* 39(Database issue): D1073-D1078; 10.1093/nar/gkq944. **[G]**

93. Yasser, E. L. M., Dobbs, D., and Honavar, V. "Predicting MHC-II Binding Affinity Using Multiple Instance Regression." IEEE/ACM Transactions on Computational Biology and Bioinformatics, IEEE Computer Society: <http://doi.ieeecomputersociety.org/10.1109/TCBB.2010.94>. [AR, G]
94. Yue, C., Oelke, M., Paulaitis, M. E., and Schneck, J. P. (2010). "Novel Cellular Microarray Assay for Profiling T-Cell Peptide Antigen Specificities." *Journal of Proteome Research* 9(11): 5629-5637. [G]
95. Zaiss, D. M. W., Boog, C. J. P., van Eden, W., and Sijts, A. (2010). "Considerations in the design of vaccines that induce CD8 T cell mediated immunity." *Vaccine* 28(49): 7716-7720. [G]
96. Zhao, L. and Li, J. (2010). "Mining for the antibody-antigen interacting associations that predict the B cell epitopes." *BMC Structural Biology* 10(Suppl 1): S6. [AR, G]
97. Zvi, A., Ariel, N., and Shafferman, A. (2010). "Data Mining, Bioinformatic and Immunoinformatic Analyses of *Francisella tularensis* Schu S4 Genome in Search for Novel Vaccine Candidates." *The Challenge of Highly Pathogenic Microorganisms*, Springer: 229-243; doi: 10.1007/978-90-481-9054-6-25. [AR, G]

3.2.2 Analysis Resource

1. Abbott, W. G. H., Tsai, P., Leung, E., Trevarton, A., Ofanoa, M., Hornell, J., Gane, E. J., Munn, S. R., and Rodrigo, A. G. (2010). "Associations between HLA Class I Alleles and Escape Mutations in the Hepatitis B Virus Core Gene in New Zealand-Resident Tongans." *Journal of Virology* 84(1): 621-629. [AR]
2. Alexander, J., Bilsel, P., del Guercio, M. F., Marinkovic-Petrovic, A., Southwood, S., Stewart, S., Ishioka, G., Kotturi, M. F., Botten, J., and Sidney, J. (2010). "Identification of broad binding class I HLA supertype epitopes to provide universal coverage of influenza A virus." *Human Immunology* 71(5): 468-474. [AR]
3. Alexander, J., Bilsel, P., del Guercio, M. F., Stewart, S., Marinkovic-Petrovic, A., Southwood, S., Crimi, C., Vang, L., Walker, L., and Ishioka, G. (2010). "Universal influenza DNA vaccine encoding conserved CD4+ T cell epitopes protects against lethal viral challenge in HLA-DR transgenic mice." *Vaccine* 28(3): 664-672. [AR]
4. Anderson, R., Huang, Y., and Langley, J. M. (2010). "Prospects for defined epitope vaccines for respiratory syncytial virus." *Future Microbiology* 5(4): 585-602. [AR]
5. Ansari, H. R. and Raghava, G. P. S. (2010). "Identification of conformational B-cell Epitopes in an antigen from its primary sequence." *Immunome Research* 6: 6. [AR]
6. Antunes, D. A., Vieira, G. F., Rigo, M. M., Cibulski, S. P., Sinigaglia, M., Chies, J. A. B., and Goldberg, A. C. (2010). "Structural Allele-Specific Patterns Adopted by Epitopes in the MHC-I Cleft and Reconstruction of MHC: peptide Complexes to Cross-Reactivity Assessment." *PloS ONE* 5(4): e10353. [AR]
7. Berry, J. D., Hay, K., Rini, J. M., Yu, M., Wang, L. F., Plummer, F. A., Corbett, C. R., and Andonov, A. (2010). "Neutralizing epitopes of the SARS-CoV S-protein cluster independent of repertoire, antigen structure or mAb technology." *Mabs* 2(1): 53-66. [AR]
8. Black, M., Trent, A., Tirrell, M., and Olive, C. (2010). "Advances in the design and delivery of peptide subunit vaccines with a focus on Toll-like receptor agonists." *Expert Review of Vaccines* 9(2): 157-173. [AR]
9. Brennan, F. R., Morton, L. D., Spindeldreher, S., Kiessling, A., Allenspach, R., Hey, A., Muller, P. Y., Frings, W., and Sims, J. (2010). "Safety and immunotoxicity assessment of immunomodulatory monoclonal antibodies." *Mabs* 2(3): 233-255. [AR]
10. Bruner, B. F., Vista, E. S., Wynn, D. M., Harley, J. B., and James, J. A. (2010). "Anti-neutrophil cytoplasmic antibodies target sequential functional proteinase 3 epitopes in the sera of patients with Wegener's granulomatosis." *Clinical & Experimental Immunology* 162(2): 262-270. [AR]

11. Bui, M. R., Hodson, V., King, T., Leopold, D., Dai, S., Fiolkoski, V., Oakes, S., Duke, R., Apelian, D., and Franzusoff, A. (2010). "Mutation-specific control of BCR-ABL T315I positive leukemia with a recombinant yeast-based therapeutic vaccine in a murine model." *Vaccine* 28(37): 6028-6035. **[AR]**
12. Cohen, T., Moise, L., Martin, W., and Groot, A. S. (2010). "Immunoinformatics: The Next Step in Vaccine Design." *Infectious Disease Informatics*, Springer New York: 223-244; doi: 10.1007/978-1-4419-1327-2. **[AR]**
13. Cong, H., Mui, E. J., Witola, W. H., Sidney, J., Alexander, J., Sette, A., Maewal, A., and McLeod, R. (2010). "Towards an immunosense vaccine to prevent toxoplasmosis: Protective Toxoplasma gondii epitopes restricted by HLA-A* 0201." *Vaccine* 29(4): 754-762. **[AR]**
14. Davies, M. N. and Flower, D. R. (2010). "Computational Epitope Mapping." *Infectious Disease Informatics*, Springer New York: 187-202; doi: 10.1007/978-1-4419-1327-2_9. **[AR]**
15. Davies, M. N. and Flower, D. R. (2010). "Computational Vaccinology." *Immunomics Reviews: Bioinformatics for Immunomics*, Springer New York: 1-20; doi: 10.1007/978-1-4419-0540-6_1. **[AR]**
16. de Groot, N. G., Heijmans, C., Zoet, Y. M., de Ru, A. H., Verreck, F. A., van Veelen, P. A., Drijfhout, J. W., Doxiadis, G. G. M., Remarque, E. J., and Doxiadis, I. I. N. (2010). "AIDS-protective HLA-B* 27/B* 57 and chimpanzee MHC class I molecules target analogous conserved areas of HIV-1/SIVcpz." *Proceedings of the National Academy of Sciences* 107(34): 15175-15180. **[AR]**
17. Deenadayalan, A., Sundaramurthi, J. C., and Raja, A. (2010). "Immunological and proteomic analysis of preparative isoelectric focusing separated culture filtrate antigens of Mycobacterium tuberculosis." *Experimental and Molecular Pathology* 88(1): 156-162. **[AR]**
18. Denisova, G., Denisov, D., and Bramson, J. (2010). "Applying bioinformatics for antibody epitope prediction using affinity-selected mimotopes-relevance for vaccine design." *Immunome Research* 6(Suppl 2): S6. **[AR]**
19. Dimitrov, I., Garnev, P., Flower, D. R., and Doytchinova, I. (2010). "MHC Class II Binding Prediction-A Little Help from a Friend." *Journal of Biomedicine and Biotechnology* 2010: 705821. **[AR]**
20. Dimitrov, I., Garnev, P., Flower, D. R., and Doytchinova, I. (2010). "Peptide binding to the HLA-DRB1 supertype: a proteochemometrics analysis." *European Journal of Medicinal Chemistry* 45(1): 236-243. **[AR]**
21. Ding, J., Wang, Y., Cheng, T., Chen, X., and Gao, B. (2010). "Identification of HLA-A24-Binding Peptides of Mycobacterium tuberculosis Derived Proteins with Beta 2m Linked HLA-A24 Single Chain Expressing Cells." *Immunological Investigations* 39(2): 103-113. **[AR]**
22. Dudek, N. L., Perlmutter, P., Aguilar, M. I., Croft, N. P., and Purcell, A. W. (2010). "Epitope Discovery and Their Use in Peptide Based Vaccines." *Current Pharmaceutical Design* 16(28): 3149-3157. **[AR]**
23. Eberhard, H. P., Feldmann, U., Bochtler, W., Baier, D., Rutt, C., Schmidt, A. H., and Muller, C. R. "Estimating unbiased haplotype frequencies from stem cell donor samples typed at heterogeneous resolutions: a practical study based on over 1 million German donors." *Tissue Antigens* 76(5): 352-361; doi: 10.1111/j.1399-0039.2010.01518.x. **[AR]**
24. Erup Larsen, M., Kloverpris, H., Stryhn, A., Koophethile, C. K., Sims, S., Ndung'u, T., Goulder, P., Buus, S., and Nielsen, M. (2010). "HLArestrictor-a tool for patient-specific predictions of HLA restriction elements and optimal epitopes within peptides." *Immunogenetics* 63(1): 43-55. **[AR]**
25. Felder, K. M., Hoelzle, K., Heinritzi, K., Ritzmann, M., and Hoelzle, L. E. (2010). "Antibodies to actin in autoimmune haemolytic anaemia." *BMC Veterinary Research* 6: 18. **[AR]**

26. Fiorucci, S. and Zacharias, M. (2010). "Prediction of Protein-Protein Interaction Sites Using Electrostatic Desolvation Profiles." *Biophysical Journal* 98(9): 1921-1930. **[AR]**
27. Flower, D., Phadwal, K., Macdonald, I., Coveney, P., Davies, M., and Wan, S. (2010). "T-cell epitope prediction and immune complex simulation using molecular dynamics: state of the art and persisting challenges." *Immunome Research* 6(Suppl 2): S4. **[AR]**
28. Fontaine Costa, A. I., Rao, X., LeChenadec, E., van Baarle, D., and Kesmir, C. (2010). "HLA-B molecules target more conserved regions of the HIV-1 proteome." *AIDS* 24(2): 211-215. **[AR]**
29. Gaikwad Vishnu, J., Bhosale, S. S., Patel, S. S., Kadam, D. S., and Dodamani, G. O. (2010). "Prediction of Interspecies Relationship and Antigenicity of Segment 1 Protein of Influenza A Virus from India." *E-International Scientific International Research Journal* 2(3): 207-209; http://www.eisrjc.com/journals/journal_1/eisrj-vol-2-issue-3-5.pdf. **[AR]**
30. Gunther, S., Schlundt, A., Sticht, J., Roske, Y., Heinemann, U., Wiesmller, K. H., Jung, G., Falk, K., Rotzschke, O., and Freund, C. (2010). "Bidirectional binding of invariant chain peptides to an MHC class II molecule." *Proceedings of the National Academy of Sciences* 107(51): 22219-22224. **[AR]**
31. Gupta, S. K., Smita, S., Sarangi, A. N., Srivastava, M., Akhoon, B. A., Rahman, Q., and Gupta, S. K. (2010). "In silico CD4+ T-cell epitope prediction and HLA distribution analysis for the potential proteins of *Neisseria meningitidis* Serogroup B--A clue for vaccine development." *Vaccine* 28(43): 7092-7097. **[AR]**
32. Hadrup, S. R. and Schumacher, T. N. (2010). "MHC-based detection of antigen-specific CD8+ T cell responses." *Cancer Immunology, Immunotherapy* 59(9): 1425-1433. **[AR]**
33. Haslam, N. J. and Gibson, T. J. (2010). "EpiC: An Open Resource for Exploring Epitopes To Aid Antibody-Based Experiments." *Journal of Proteome Research* 9(7): 3759-3763. **[AR]**
34. Hellqvist, E., Kvarnstrom, M., Soderberg, A., Vrethem, M., Ernerudh, J., and Rosen, A. (2010). "Myelin protein zero is naturally processed in the B cells of monoclonal gammopathy of undetermined significance of immunoglobulin M isotype: aberrant triggering of a patient's T cells." *Haematologica* 95(4): 627-636. **[AR]**
35. Hoof, I., Perez, C. L., Buggert, M., Gustafsson, R. K. L., Nielsen, M., Lund, O., and Karlsson, A. C. (2010). "Interdisciplinary analysis of HIV-specific CD8+ T cell responses against variant epitopes reveals restricted TCR promiscuity." *The Journal of Immunology* 184(9): 5383-5391. **[AR]**
36. Hu, X., Zhou, W., Udaka, K., Mamitsuka, H., and Zhu, S. (2010). "MetaMHC: a meta approach to predict peptides binding to MHC molecules." *Nucleic Acids Research* 38(Web Server issue): W474-W479. **[AR]**
37. Ivanenkov, V. V., Crawford, P. A., Toyama, A., Sevigny, J., and Kirley, T. L. (2010). "Epitope mapping in cell surface proteins by site-directed masking: defining the structural elements of NTPDase3 inhibition by a monoclonal antibody." *Protein Engineering Design and Selection* 23(7): 579-588. **[AR]**
38. Jan, M., Meng, S., Chen, N. C., Mai, J., Wang, H., and Yang, X. F. (2010). "Inflammatory and Autoimmune Reactions in Atherosclerosis and Vaccine Design Informatics." *Journal of Biomedicine and Biotechnology* 2010: 459798. **[AR]**
39. Kobayashi, H., Azumi, M., Hayashi, S., Sato, K., Aoki, N., Kimura, S., Kakizaki, H., Nagato, T., Harabuchi, Y., Tateno, M., and Celis, E. (2010). "Characterization of human CD4 helper T cell responses against Aurora kinase A." *Cancer Immunology, Immunotherapy* 59(7): 1029-1039. **[AR]**
40. Kotturi, M. F., Botten, J., Maybeno, M., Sidney, J., Glenn, J., Bui, H. H., Oseroff, C., Crotty, S., Peters, B., and Grey, H. (2010). "Polyfunctional CD4+ T cell responses to a set of pathogenic arenaviruses provide broad population coverage." *Immunome Research* 6: 4; doi: 10.1186/1745-7580-6-4. **[AR]**

41. Kumar, M., Sundaramurthi, J. C., Mehra, N. K., Kaur, G., and Raja, A. (2010). "Cellular immune response to Mycobacterium tuberculosis-specific antigen culture filtrate protein-10 in south India." *Medical Microbiology and Immunology* 199(1): 11-25. **[AR]**
42. Laing, K. J., Magaret, A. S., Mueller, D. E., Zhao, L., Johnston, C., De Rosa, S. C., Koelle, D. M., Wald, A., and Corey, L. (2010). "Diversity in CD8+ T Cell Function and Epitope Breadth Among Persons with Genital Herpes." *Journal of Clinical Immunology* 30(5): 703-722. **[AR]**
43. Lambeck, A. J. A., Nijman, H. W., Hoogeboom, B. N., Regts, J., de Mare, A., Wilschut, J., and Daemen, T. (2010). "Role of T cell competition in the induction of cytotoxic T lymphocyte activity during viral vector-based immunization regimens." *Vaccine* 28(26): 4275-4282. **[AR]**
44. Lampen, M. H., Verweij, M. C., Querido, B., van der Burg, S. H., Wiertz, E. J. H. J., and van Hall, T. (2010). "CD8+ T Cell Responses against TAP-Inhibited Cells Are Readily Detected in the Human Population." *The Journal of Immunology* 185(11): 6508-6517. **[AR]**
45. Larsen, M. E., Kornblit, B., Larsen, M. V., Masmas, T. N., Nielsen, M., Thiim, M., Garred, P., Stryhn, A., Lund, O., and Buus, S. (2010). "Degree of Predicted Minor Histocompatibility Antigen Mismatch Correlates with Poorer Clinical Outcomes in Nonmyeloablative Allogeneic Hematopoietic Cell Transplantation." *Biology of Blood and Marrow Transplantation* 16(10): 1370-1381. **[AR]**
46. Larsen, M. V., Lelic, A., Parsons, R., Nielsen, M., Hoof, I., Lamberth, K., Loeb, M.B., Buus, S., Bramson, J., and Lund, O. (2010). "Identification of CD8+ T cell epitopes in the West Nile virus polyprotein by reverse-immunology using NetCTL." *PLoS ONE* 5(9): e12697. **[AR]**
47. Liang, P., Bach, F., Bouchard, G., and Jordan, M. I. (2010). "Asymptotically optimal regularization in smooth parametric models." *Advances in Neural Information Processing Systems (NIPS)*, Cambridge, MA; www.di.ens.fr/~fbach/regularization-nips2010.pdf. **[AR]**
48. Liang, S., Zheng, D., Standley, D. M., Yao, B., Zacharias, M., and Zhang, C. (2010). "EPSVR and EPMeta: prediction of antigenic epitopes using support vector regression and multiple server results." *BMC Bioinformatics* 11: 381. **[AR]**
49. Liu, J., Li, Q. J., and Zhang, W. (2010). "A novel locally linear embedding and wavelet transform based encoding method for prediction of MHC-II binding affinity." *Interdisciplinary Sciences: Computational Life Sciences* 2(2): 145-150. **[AR]**
50. Lundegaard, C. (2010). "Mice, men and MHC supertypes." *Expert Review of Vaccines* 9(7): 713-718. **[AR]**
51. MacNamara, A., Rowan, A., Hilburn, S., Kadolsky, U., Fujiwara, H., Suemori, K., Yasukawa, M., Taylor, G., Bangham, C. R. M., and Asquith, B. (2010). "HLA Class I Binding of HBZ Determines Outcome in HTLV-1 Infection." *PLoS Pathogens* 6(9): 5121-5129. **[AR]**
52. Maier, R. H., Maier, C. J., Rid, R., Hintner, H., Bauer, J. W., and Onder, K. (2010). "Epitope Mapping of Antibodies Using a Cell Array-Based Polypeptide Library." *Journal of Biomolecular Screening* 15(4): 418-426. **[AR]**
53. Malnick, S. D. H., Bar-Ilan, A., Goland, S., Somin, M., Doniger, T., Basevitz, A., and Unger, R. (2010). "Perimyocarditis following streptococcal group A infection: From clinical cases to bioinformatics analysis." *European Journal of Internal Medicine* 21(4): 354-356. **[AR]**
54. Matera, L. (2010). "The choice of the antigen in the dendritic cell-based vaccine therapy for prostate cancer." *Cancer Treatment Reviews* 36(2): 131-141. **[AR]**
55. McNamara, L. A., He, Y., and Yang, Z. (2010). "Using epitope predictions to evaluate efficacy and population coverage of the Mtb 72 f vaccine for tuberculosis." *BMC Immunology* 11(1): 18. **[AR, MA]**
56. Mejia, J. S., Arthun, E. N., and Titus, R. G. (2010). "Cysteine-Free Proteins in the Immunobiology of Arthropod-Borne Diseases." *Journal of Biomedicine and Biotechnology* 2010: 171537. **[AR]**

57. Mitra, I. and Cui, Y. (2010). "Understanding molecular recognition and epitope prediction from Information Theoretic approach." *BMC Bioinformatics* 11(Suppl 4): P22; doi: 10.1186/1471-2105-11-S4-P22. **[AR]**
58. Moosavi, F., Mohabatkar, H., and Mohsenzadeh, S. (2010). "Computer-aided analysis of structural properties and epitopes of Iranian HPV-16 E7 oncoprotein." *Interdisciplinary Sciences: Computational Life Sciences* 2(4): 367-372. **[AR]**
59. Nelson, P. A., Khodadoust, M., Prodhomme, T., Spencer, C., Patarroyo, J. C., Varrin-Doyer, M., Ho, J. D., Stroud, R. M., Zamvil, S. S., and Kleinschmitz, C. (2010). "Immunodominant T Cell Determinants of Aquaporin-4, the Autoantigen Associated with Neuromyelitis Optica." *PloS ONE* 5(11): 383-392. **[AR]**
60. Neumann-Haefelin, C., Timm, J., Schmidt, J., Kersting, N., Fitzmaurice, K., Oniangue-Ndza, C., Kemper, M. N., Humphreys, I., McKiernan, S., Kelleher, D., Lohmann, V., Bowness, P., Huzly, D., Rosen H.R., Kim, A.Y., Lauer, G.M., Allen, T.M., Barnes, E., Roggendorf, M., Blum, H.E., and Thimme, R. (2010). "Protective effect of human leukocyte antigen B27 in hepatitis C virus infection requires the presence of a genotype specific immunodominant CD8+ T cell epitope." *Hepatology* 51(1): 54-62. **[AR]**
61. Nielsen, M., Justesen, S., Lund, O., Lundsgaard, C., and Buus, S. (2010). "NetMHCIIpan-2.0 - Improved pan-specific HLA-DR predictions using a novel concurrent alignment and weight optimization training procedure." *Immunome Research* 6: 9; doi: 10.1186/1745-7580-6-9. **[AR]**
62. Oseroff, C., Sidney, J., Kotturi, M. F., Kolla, R., Alam, R., Broide, D. H., Wasserman, S. I., Weiskopf, D., McKinney, D. M., and Chung, J. L. (2010). "Molecular Determinants of T Cell Epitope Recognition to the Common Timothy Grass Allergen." *The Journal of Immunology* 185(2): 943-955. **[AR]**
63. Pappalardo, F., Halling-Brown, M., Pennisi, M., Chiacchio, F., Sansom, C., Shepherd, A., Moss, D., Motta, S., and Brusic, V. (2010). "The ImmunoGrid Simulator: How to Use It." *Lecture Notes in Computer Science: Computational Intelligence Methods for Bioinformatics and Biostatistics*, Springer 6160: 1-19; doi: 10.1007/978-3-642-14571-1-1. **[AR]**
64. Patch, J. R., Pedersen, L. E., Toka, F. N., Moreas, M., Grubman, M. J., Nielsen, M., Jungersten, G., Buus, S., and Golde, W. T. (2010). "Induction of foot-and-mouth disease virus (FMDV) specific cytotoxic T cell killing by vaccination." *Clinical and Vaccine Immunology*: ePub ahead of print; doi: 10.1128/CVI.00417-10. **[AR]**
65. Pomes, A. (2010). "Relevant B Cell Epitopes in Allergic Disease." *International Archives of Allergy and Immunology* 152(1): 1-11. **[AR]**
66. Rapin, N., Lund, O., Bernaschi, M., and Castiglione, F. (2010). "Computational Immunology Meets Bioinformatics: The Use of Prediction Tools for Molecular Binding in the Simulation of the Immune System." *PloS ONE* 5(4): e9862; doi: 10.1371/journal.pone.0009862. **[AR]**
67. Ravindranath, M. H., Kaneku, H., El-Awar, N., Morales-Buenrostro, L. E., and Terasaki, P. I. (2010). "Antibodies to HLA-E in nonalloimmunized males: pattern of HLA-Ia reactivity of anti-HLA-E-positive sera." *The Journal of Immunology* 185(3): 1935-1948. **[AR]**
68. Richards, K. A., Topham, D., Chaves, F. A., and Sant, A. J. (2010). "Cutting Edge: CD4 T Cells Generated from Encounter with Seasonal Influenza Viruses and Vaccines Have Broad Protein Specificity and Can Directly Recognize Naturally Generated Epitopes Derived from the Live Pandemic H1N1 Virus." *The Journal of Immunology* 185(9): 4998-5002. **[AR, MA]**
69. Riemer, A. B., Keskin, D. B., Zhang, G., Handley, M., Anderson, K. S., Brusic, V., Reinhold, B., and Reinherz, E. L. (2010). "A Conserved E7-derived Cytotoxic T Lymphocyte Epitope Expressed on Human Papillomavirus 16-transformed HLA-A2+ Epithelial Cancers." *Journal of Biological Chemistry* 285(38): 29608-29622. **[AR]**

70. Roblek, M., Schnchner, S., Huber, V., Ollram, K., and Vlcek-Vesely, S. (2010). "Monoclonal Antibodies Specific for Disease-Associated Point-Mutants: Lamin A/C R453W and R482W." 5(5): e10604. **[AR]**
71. Rojas, O. L., Rojas-Villarraga, A., Cruz-Tapias, P., Sanchez, J. L., Suarez-Escudero, J. C., and Patarroyo and Anaya, J. M. (2010). "HLA class II polymorphism in Latin American patients with multiple sclerosis." Autoimmunity Reviews 9(6): 407-413. **[AR]**
72. Rosa, D. S., Ribeiro, S. P., and Cunha-Neto, E. (2010). "CD4+ T cell epitope discovery and rational vaccine design." Archivum Immunologiae et Therapiae Experimentalis 58(2): 121-130; doi: 10.1007/s00005-010-0067-0. **[AR]**
73. Scarabelli, G., Morra, G., and Colombo, G. (2010). "Predicting Interaction Sites from the Energetics of Isolated Proteins: A New Approach to Epitope Mapping." Biophysical Journal 98(9): 1966-1975. **[AR]**
74. Scott, D. W. and De Groot, A. S. (2010). "Can we prevent immunogenicity of human protein drugs?" British Medical Journal 69(Suppl 1): i72-i76. **[AR]**
75. Sedegah, M., Kim, Y., Peters, B., McGrath, S., Ganeshan, H., Lejano, J., Abot, E., Banania, G., Belmonte, M., and Sayo, R. (2010). "Identification and localization of minimal MHC-restricted CD8+ T cell epitopes within the Plasmodium falciparum AMA1 protein." Malaria Journal 9: 241; doi: 10.1186/1475-2875-9-241. **[AR]**
76. Sette, A. and Rappuoli, R. (2010). "Reverse Vaccinology: Developing Vaccines in the Era of Genomics." Immunity 33(4): 530-541. **[AR, MA]**
77. Seward, R. J., Drouin, E. E., Steere, A. C., and Costello, C. E. (2010). "Peptides presented by HLA-DR molecules in synovia of patients with rheumatoid arthritis or antibiotic-refractory Lyme arthritis." Molecular & Cellular Proteomics, in press; doi: 10.1074/mcp.M110.002477 mcp.M110.002477. **[AR]**
78. Singh, S. P., Khan, F., and Mishra, B. N. (2010). "Computational characterization of Plasmodium falciparum proteomic data for screening of potential vaccine candidates." Human Immunology 71(2): 136-143. **[AR]**
79. Sirskyj, D., Diaz-Mitoma, F., Golshani, A., Kumar, A., and Azizi, A. (2010). "Innovative bioinformatic approaches for developing peptide-based vaccines against hypervariable viruses." Immunology and Cell Biology 89(1): 81-89. **[AR, MA]**
80. Sollner, J., Heinzel, A., Summer, G., Fechete, R., Stipkovits, L., Szathmary, S., and Mayer, B. (2010). "Concept and application of a computational vaccinology workflow." Immunome Research 6(Suppl 2): S7. **[AR]**
81. Solomon, C., Southwood, S., Hoof, I., Rudersdorf, R., Peters, B., Sidney, J., Pinilla, C., Marcondes, M. C. G., Ling, B., Marx, P., Sette, A., and Mothe, B. (2010). "The most common Chinese rhesus macaque MHC class I molecule shares peptide binding repertoire with the HLA-B7 supertype." Immunogenetics 62(7): 451-464. **[AR]**
82. Stas, P., Gansemans, Y., and Lasters, I. (2010). "Immunogenicity assessment of antibody therapeutics." Current Trends in Monoclonal Antibody Development and Manufacturing, Biotechnology: Pharmaceutical Aspects, Springer 11(6): 271-291; doi: 10.1007/978-0-387-76643-0-16. **[AR]**
83. Stranzl, T., Larsen, M. V., Lundsgaard, C., and Nielsen, M. (2010). "NetCTLpan: pan-specific MHC class I pathway epitope predictions." Immunogenetics 62(6): 357-368. **[AR]**
84. Tarosso, L. F., Sauer, M. M., Sanabani, S., Giret, M. T., Tomiyama, H. I., Sidney, J., Piaskowski, S. M., Diaz, R. S., Sabino, E. C., Sette, A., Kalil-Filho, J., Watkins, D.I., and Kallas, E.G. (2010). "Unexpected Diversity of Cellular Immune Responses against Nef and Vif in HIV-1-Infected Patients Who Spontaneously Control Viral Replication." PloS ONE 5(7): e11436; doi: 10.1371/journal.pone.0011436. **[AR]**

85. Voigt, A., Jakel, S., Textoris-Taube, K., Keller, C., Drung, I., Szalay, G., Klingel, K., Henklein, P., Stangl, K., Kloetzel, P. M., and Kuckelkorn, U. (2010). "Generation of *in silico* predicted coxsackievirus B3-derived MHC class I epitopes by proteasomes." *Amino Acids* 39(1): 243-255. **[AR]**
86. Wang, S. T., Chang, C. C., Yen, M. C., Tu, C. F., Chu, C. L., Peng, Y. T., Chen, D. Y., Lan, J. L., and Lin, C. C. (2010). "RNA interference-mediated silencing of Foxo3 in antigen-presenting cells as a strategy for the enhancement of DNA vaccine potency." *Gene Therapy*, advanced online publication; doi: 10.1038/gt.2010.146. **[AR]**
87. Widmer, C., Toussaint, N., Altun, Y., Kohlbacher, O., and Ratsch, G. (2010). "Novel Machine Learning Methods for MHC Class I Binding Prediction." *Lecture Notes in Computer Science, Pattern Recognition in Bioinformatics*, Springer (6282): 98-109; doi: 10.1007/978-3-642-16001-1-9. **[AR]**
88. Wu, C. Y., Chen, Y. C., and Lim, C. (2010). "A structural-alphabet-based strategy for finding structural motifs across protein families." *Nucleic Acids Research* 38(14): e150. **[AR]**
89. Xu, W., Li, H. Z., Liu, J. J., Guo, Z., Zhang, B. F., Chen, F. F., Pei, D. S., and Zheng, J. N. (2010). "Identification of HLA-A* 0201-restricted cytotoxic T lymphocyte epitope from proliferating cell nuclear antigen." *Tumor Biology* 32(1): 63-69. **[AR]**
90. Xu, X. L., Sun, J., Liu, Q., Wang, X. J., Xu, T. L., Zhu, R. X., Wu, D., and Cao, Z. W. (2010). "Evaluation of spatial epitope computational tools based on experimentally-confirmed dataset for protein antigens." *Chinese Science Bulletin* 55(20): 2169-2174. **[AR]**
91. Yang, M. and Mine, Y. (2010). "Peptide-Based Immunotherapy for Food Allergy." *Bioactive Proteins and Peptides as Functional Foods and Nutraceuticals*, Wiley-Blackwell: Chapter 8; doi: 10.1002/9780813811048.ch8. **[AR]**
92. Yanover, C., Malkki, M., Gooley, T., Petersdorf, E. W., and Bradley, P. (2010). "How do amino acid mismatches affect the outcome of hematopoietic cell transplants?: a structural perspective." *Proceedings of the First ACM International Conference on Bioinformatics and Computational Biology*: 627-633, <http://delivery.acm.org/10.1145/1860000/1854893/p627-yanover.pdf>; doi: 10.1145/1854776.1854893. **[AR]**
93. Yauch, L. E., Prestwood, T. R., May, M. M., Morar, M. M., Zellweger, R. M., Peters, B., Sette, A., and Shresta, S. (2010). "CD4+ T Cells Are Not Required for the Induction of Dengue Virus-Specific CD8+ T Cell or Antibody Responses but Contribute to Protection after Vaccination." *The Journal of Immunology* 185(9): 5405-5416. **[AR]**
94. You, L. W., Brusic, V., Gallagher, M., and Boden, M. (2010). "Using Gaussian Process with Test Rejection to Detect T-Cell Epitopes in Pathogen Genomes." *IEEE/ACM Transactions on Computational Biology and Bioinformatics* 7(4): 741-751. **[AR]**
95. Zhang, H., Lund, O., Nielsen, M., Wang, P., Papangelopoulos, N., Xu, Y., Sette, A., Bourne, P. E., Ponomarenko, J., and Peters, B. (2010). "Limitations of *ab initio* predictions of peptide binding to MHC class II molecules." *PloS ONE* 5(2): 2927; doi: 10.1371/journal.pone.0009272. **[AR]**
96. Zhang, W., Choi, J., Zeng, W., Rogers, S. A., Alyea, E. P., Rheinwald, J. G., Canning, C. M., Brusic, V., Sasada, T., and Reinerherz, E. L. (2010). "Graft-versus-Leukemia Antigen CML66 Elicits Coordinated B-Cell and T-Cell Immunity after Donor Lymphocyte Infusion." *Clinical Cancer Research* 16(10): 2729-2739. **[AR]**
97. Zhang, Z. W., Zhang, Y. G., Wang, Y. L., Pan, L., Fang, Y. Z., Jiang, S. T., Lu, J. L., and Zhou, P. (2010). "Screening and identification of B cell epitopes of structural proteins of foot-and-mouth disease virus serotype Asia1." *Veterinary Microbiology* 140(1-2): 25-33. **[AR]**
98. Zheng, Y., Wen, Y., George, A. M., Steinbach, A. M., Phillips, B. E., Giannoukakis, N., Gawalt, E. S., and Meng, W. S. (2010). "A peptide-based material platform for displaying antibodies to engage T cells." *Biomaterials* 32(1): 249-257. **[AR]**

3.2.3 Curation

1. Kastrin, A., Peterlin, B., and Hristovski, D. (2010). "Chi-square-based Scoring Function for Categorization of MEDLINE Citations." *Methods of Information in Medicine* 49(4): 371-378. **[C]**
2. Lourenco, A., Carreira, R., Glez-Pena, D., Mendez, J. R., Carneiro, S., and Rocha, L. M. (2010). "BioDR: Semantic indexing networks for biomedical document retrieval." *Expert Systems with Applications* 37(4): 3444-3453; doi: 10.1016/j.eswa.2009.10.044. **[C]**
3. Mazumder, R., Natale, D. A., Julio, J. A. E., Yeh, L. S., and Wu, C. H. (2010). "Community annotation in biology." *Biology Direct* 5: 12, <http://www.biology-direct.com/content/5/1/12>; doi: 10.1186/1745-6150-5-12. **[C]**
4. Sorani, M. D., Ortmann, W. A., Bierwagen, E. P., and Behrens, T. W. (2010). "Clinical and biological data integration for biomarker discovery." *Drug Discovery Today* 15(17-18): 741-748. **[C]**

3.2.4 Epitope Meta-Analyses

1. Agrati, C., Gioia, C., Lalle, E., Cimini, E., Castilletti, C., Armignacco, O., Lauria, F. N., Ferraro, F., Antonini, M., and Ippolito, G. (2010). "Association of Profoundly Impaired Immune Competence in H1N1v-Infected Patients with a Severe or Fatal Clinical Course." *The Journal of Infectious Diseases* 202(5): 681-689. **[MA]**
2. Allwinn, R. and Doerr, H. W. (2010). "Die Schweinegrippe-Betrachtungen zur Epidemiologie und Pathogenität Swine Influenza-Comments on Epidemiology and Pathogenicity." *Klinikarzt* 39(4): 192-196. **[MA]**
3. Bodewes, R., Kreijtz, J. H. C. M., and Hillaire, M. (2010). "Vaccination with whole inactivated virus vaccine affects the induction of heterosubtypic immunity against influenza virus A/H5N1 and immunodominance of virus-specific CD8+ T-cell responses in mice." *Journal of General Virology* 91(7): 1743-1753. **[MA]**
4. Boesteanu, A. C., Babu, N. S., Wheatley, M., Papazoglou, E. S., and Katsikis, P. D. (2010). "Biopolymer encapsulated live influenza virus as a universal CD8+ T cell vaccine against influenza virus." *Vaccine* 29(2): 314-322. **[MA]**
5. Campanini, G., Piralla, A., Paolucci, S., Rovida, F., Percivalle, E., Maga, G., and Baldanti, F. (2010). "Genetic divergence of influenza A NS1 gene in pandemic 2009 H1N1 isolates with respect to H1N1 and H3N2 isolates from previous seasonal epidemics." *Virology Journal* 7: 209. **[MA]**
6. Cattamanchi, A., Ssewenyana, I., Davis, J. L., Huang, L., Worordria, W., Den Boon, S., Yoo, S., Andama, A., Hopewell, P., and Cao, H. (2010). "Role of interferon-gamma release assays in the diagnosis of pulmonary tuberculosis in patients with advanced HIV infection." *BMC Infectious Diseases* 10: 75. **[MA]**
7. Chan, J., Holmes, A., and Rabadian, R. (2010). "Network Analysis of Global Influenza Spread." *PloS Computational Biology* 6(11): e1001005. **[MA]**
8. Chan, Y. J., Lee, C. L., Hwang, S. J., Fung, C. P., Wang, F. D., Yen, D. H. T., Tsai, C. H., Chen, Y. M. A., and Lee, S. D. (2010). "Seroprevalence of antibodies to pandemic (H1N1) 2009 influenza virus among hospital staff in a medical center in Taiwan." *Journal of the Chinese Medical Association* 73(2): 62-66. **[MA]**
9. Chen, M., Xiao, W., Ai, B. A., He, H., Liu, X. Z., Dong, H., Xiong, J., and Xu, L. (2010). "Sensitivity Assessment of Rapid Influenza Diagnostic Tests for the Detection of the 2009 Pandemic Influenza A (H1N1) Virus in Clinical Specimens." *Lab Medicine* 41: 731-734. **[MA]**
10. Chi, C. Y., Liu, C. C., Lin, C. C., Wang, H. C., Cheng, Y. T., Chang, C. M., and Wang, J. R. (2010). "Preexisting Antibody Response against 2009 Pandemic Influenza H1N1 Viruses in the Taiwanese Population." *Clinical and Vaccine Immunology* 17(12): 1958-1962. **[MA]**

11. Christakis, N. A. and Fowler, J. H. (2010). "Social Network Sensors for Early Detection of Contagious Outbreaks." *PLoS ONE* 5(9): e12948. **[MA]**
12. Comas, I., Chakravarti, J., Small, P. M., Galagan, J., Niemann, S., Kremer, K., Ernst, J. D., and Gagneux, S. (2010). "Human T cell epitopes of *Mycobacterium tuberculosis* are evolutionarily hyperconserved." *Nature Genetics* 42(6): 498-503. **[MA]**
13. Combadiere, B., Siberil, S., and Duffy, D. (2010). "Keeping the memory of influenza viruses." *Pathologie Biologie* 58(2): e79-e86. **[MA]**
14. Combadiere, B., Vogt, A., Mahe, B., Costagliola, D., Hadam, S., Bonduelle, O., Sterry, W., Staszewski, S., Schaefer, H., van der Werf, S., Katlama, C., Autran, B., and Blume-Peytavi, U. (2010). "Preferential amplification of CD8 effector-T cells after transcutaneous application of an inactivated influenza vaccine: a randomized phase I trial." *PLoS ONE* 5(5): e10818. **[MA]**
15. De Groot, A. S., McClaine, E., Moise, L., and Martin, W. (2010). "Time for T? Thoughts about the 2009 novel H1N1 influenza outbreak and the role of T cell epitopes in the next generation of influenza vaccines." *Human Vaccines* 6(2): 161-163. **[MA]**
16. Ding, X., Jiang, L., Ke, C., Yang, Z., Lei, C., Cao, K., Xu, J., Xu, L., Yang, X., and Zhang, Y. "Amino acid sequence analysis and identification of mutations under positive selection in hemagglutinin of 2009 influenza A (H1N1) isolates." *Virus Genes* 41(3): 329-340. **[MA]**
17. Ebrahim, G. J. (2010). "Swine Flu (S-OIV) Pandemic." *Journal of Tropical Pediatrics* 56(1): 1-3; doi: 10.1093/tropej/fmp130. **[MA]**
18. Echevarria-Zuno, S., Mejia-Arangure, J. M., Grajales-Muniz, C., Gonzalez-Bonilla, C., and Borja-Aburto, V. H. (2010). "Seasonal vaccine effectiveness against pandemic A/H1N1-Authors' reply." *The Lancet* 375(9717): 802-803; doi: 10.1016/S0140-6736(10)60339-X. **[MA]**
19. Ellebedy, A. H., Duceppe, M. F., Duan, S., Stigler-Rosser, E., Rubrum, A. M., Govorkova, E. A., Webster, R. G., and Webby, R. J. (2010). "Impact of prior seasonal influenza vaccination and infection on pandemic A (H1N1) influenza virus replication in ferrets." *Vaccine*, article in press; doi: 10.1016/j.vaccine.2010.08.067. **[MA]**
20. Ellebedy, A. H., Fabrizio, T. P., Kayali, G., Oguin III, T. H., Brown, S. A., Rehg, J., Thomas, P. G., and Webby, R. J. (2010). "Contemporary Seasonal Influenza A (H1N1) Virus Infection Primes for a More Robust Response To Split Inactivated Pandemic Influenza A (H1N1) Virus Vaccination in Ferrets." *Clinical and Vaccine Immunology* 17(12): 1998-2006. **[MA]**
21. Epstein, S. L. and Price, G. E. (2010). "Cross-protective immunity to influenza A viruses." *Expert Review of Vaccines* 9(11): 1325-1341. **[MA]**
22. Frobert, E., Bouscambert-Duchamp, M., Escuret, V., Mundweiler, S., Barthelemy, M., Morfin, F., Valette, M., Gerdil, C., Lina, B., and Ferraris, O. (2010). "Anti N1 cross-protecting antibodies against H5N1 detected in H1N1 infected people." *Current Microbiology* 61(1): 25-28. **[MA]**
23. Gaseitsiwe, S., Valentini, D., Mahdavifar, S., Reilly, M., Ehrnst, A., and Maeurer, M. (2010). "Peptide Microarray-Based Identification of *Mycobacterium tuberculosis* Epitope Binding to HLA-DRB1* 0101, DRB1* 1501, and DRB1* 0401." *Clinical and Vaccine Immunology* 17(1): 168-175. **[MA]**
24. Ge, X., Tan, V., Bollyky, P. L., Standifer, N. E., James, E. A., and Kwok, W. W. (2010). "Assessment of seasonal influenza A virus-specific CD4 T-cell responses to 2009 pandemic H1N1 swine-origin influenza A virus." *Journal of Virology* 84(7): 3312-3319. **[MA]**
25. Gras, S., Kedzierski, L., Valkenburg, S. A., Laurie, K., Liu, Y. C., Denholm, J. T., Richards, M. J., Rimmelzwaan, G. F., Kelso, A., and Doherty, P. C. (2010). "Cross-reactive CD8+ T-cell immunity between the pandemic H1N1-2009 and H1N1-1918 influenza A viruses." *Proceedings of the National Academy of Sciences* 107(28): 12599-12604. **[MA]**

26. Jain, S., Patrick, A. J., and Rosenthal, K. L. (2010). "Multiple tandem copies of conserved gp41 epitopes incorporated in gag virus-like particles elicit systemic and mucosal antibodies in an optimized heterologous vector delivery regimen." *Vaccine* 28(43): 7070-7080. **[MA]**
27. Johns, M. C., Eick, A. A., Blazes, D. L., Lee, S., Perdue, C. L., Lipnick, R., Vest, K. G., Russell, K. L., DeFraites, R. F., and Sanchez, J. L. (2010). "Seasonal influenza vaccine and protection against pandemic (H1N1) 2009-associated illness among US military personnel." *PloS ONE* 5(5): e10722. **[MA]**
28. Kelly, H. and Barr, I. (2010). "Large trials confirm immunogenicity of H1N1 vaccines." *Lancet* 375(9708): 6-9. **[MA]**
29. Koyama, S., Aoshi, T., Tanimoto, T., Kumagai, Y., Kobiyama, K., Tougan, T., Sakurai, K., Coban, C., Horii, T., Akira, S., and Ishii, K. J. (2010). "Plasmacytoid Dendritic Cells Delineate Immunogenicity of Influenza Vaccine Subtypes." *Science Translational Medicine* 2(25): 25ra24. **[MA]**
30. Kuroda, M., Katano, H., Nakajima, N., Tobiume, M., Ainai, A., Sekizuka, T., Hasegawa, H., Tashiro, M., Sasaki, Y., Arakawa, Y., Hata, S., Watanabe, M., and Sata, T. (2010). "Characterization of Quasispecies of Pandemic 2009 Influenza A Virus (A/H1N1/2009) by De Novo Sequencing Using a Next-Generation DNA Sequencer." *PloS ONE* 5(4): 2605-2615. **[MA]**
31. Labrosse, B., Tourdjman, M., Porcher, R., LeGoff, J., De Lamballerie, X., Simon, F., Molina, J. M., and Clavel, F. (2010). "Detection of Extensive Cross-Neutralization between Pandemic and Seasonal A/H1N1 Influenza Viruses Using a Pseudotype Neutralization Assay." *PloS ONE* 5(6): e11036. **[MA]**
32. Langley, W. A., Li, Z. N., Bradley, K. C., Talekar, G. R., Galloway, S. E., and Steinhauer, D. A. (2010). "The effects of preexisting immunity to influenza on responses to influenza vectors in mice." *Vaccine* 28(38): 6305-6313. **[MA]**
33. Laurie, K. L., Carolan, L. A., Middleton, D., Lowther, S., Kelso, A., and Barr, I. G. (2010). "Multiple Infections with Seasonal Influenza A Virus Induce Cross-Protective Immunity against A (H1N1) Pandemic Influenza Virus in a Ferret Model." *The Journal of Infectious Diseases* 202(7): 1011-1020. **[MA]**
34. Liu, S. L., Zhang, Z. R., Wang, C. M., Dong, Y., Cui, L. B., Yang, X. H., Sun, Z., Wang, J., Chen, J., Huang, R. J., Miao, F., Ruan, B., Xie, L., He, H. X., and Deng, J. (2010). "2009 Pandemic Characteristics and Controlling Experiences of Influenza H1N1 Virus 1 Year After the Inception in Hangzhou, China." *Journal of Medical Virology* 82(12): 1985-1995. **[MA]**
35. Liu, Y., Liu, X., Fang, J., Shen, X., Chen, W., Lin, X., Li, H., Tan, W., Wang, Y., Zhao, P., and Qi, Z. (2010). "Characterization of antibodies specific for hemagglutinin and neuraminidase proteins of the 1918 and 2009 pandemic H1N1 viruses." *Vaccine* 29(2): 183-190. **[MA]**
36. Londt, B. Z., Nunez, A., Banks, J., Alexander, D. J., Russell, C., Richard-Londt, A. C., and Brown, I. H. (2010). "The effect of age on the pathogenesis of a highly pathogenic avian influenza (HPAI) H5N1 virus in Pekin ducks (*Anas platyrhynchos*) infected experimentally." *Influenza and Other Respiratory Viruses* 4(1): 17-25. **[MA]**
37. Lv, H., Gao, Y., Wu, Y., Zhai, M., Li, L., Zhu, Y., Liu, W., Wu, Z., Chen, F., and Qi, Y. (2010). "Identification of a novel cytotoxic T lymphocyte epitope from CFP21, a secreted protein of *Mycobacterium tuberculosis*." *Immunology Letters* 133(2): 94-98. **[MA]**
38. MacLeod, M. K. L., Kappler, J. W., and Marrack, P. (2010). "Memory CD4 T cells: generation, reactivation and re-assignment." *Immunology* 130(1): 10-15. **[MA]**
39. Mahajan, B., Berzofsky, J. A., Boykins, R. A., Majam, V., Zheng, H., Chattopadhyay, R., de la Vega, P., Moch, J. K., Haynes, J. D., Belyakov, I. M., Nakhси, H. L., and Kumar, S. (2010). "Multiple Antigen Peptide Vaccines against *Plasmodium falciparum* Malaria." *Infection and immunity* 78(11): 4613-4624. **[MA]**

40. Mathews, J. D., McBryde, E. S., McVernon, J., Pallaghy, P. K., and McCaw, J. M. (2010). "Prior immunity helps to explain wave-like behaviour of pandemic influenza in 1918-9." *BMC Infectious Diseases* 10(1): 128. **[MA]**
41. McCullers, J. A., Van De Velde, L. A., Allison, K. J., Branum, K. C., Webby, R. J., and Flynn, P. M. (2010). "Recipients of Vaccine against the 1976 Swine Flu Have Enhanced Neutralization Responses to the 2009 Novel H1N1 Influenza Virus." *Clinical Infectious Diseases* 50(11): 1487-1492. **[MA]**
42. Mei, S., van de Vijver, D., Xuan, L., Zhu, Y., and Sloot, P. M. A. (2010). "Quantitatively evaluating interventions in the influenza A (H1N1) epidemic on China campus grounded on individual-based simulations." *Procedia Computer Science* 1(1): 1669-1676. **[MA]**
43. Micozzi, A. (2010). "Pubblichiamo gli interventi di tre ricercatori indipendenti sull' H1N1-Influenza A." *Bollettino Notiziario* 2: 14-19; www.odmbologna.it/bollettini/files/BOLLETTINO_N_2_FEB_10.pdf. **[MA]**
44. Moghadas, S. M., Pizzi, N. J., Wu, J., Tamblyn, S. E., and Fisman, D. N. "Canada in the face of the 2009 H1N1 pandemic." *Influenza and Other Respiratory Viruses*; doi: 10.1111/j.1750-2659.2010.00184.x. **[MA]**
45. Morens, D., Taubenberger, J. K., Harvey, H. A., and Memoli, M. J. (2010). "The 1918 influenza pandemic: Lessons for 2009 and the future." *Crit Care Med* 38(4 Suppl): e10-e20. **[MA]**
46. Morens, D. M., Taubenberger, J. K., and Fauci, A. S. (2010). "The 2009 H1N1 Pandemic Influenza Virus: What Next?" *mBio* 1(4): e00211-e00210; doi: 10.1128/mBio.00211-10. **[MA]**
47. Olson, M. R., Russ, B. E., Doherty, P. C., Turner, S. J., and Stambas, J. (2010). "Influenza A virus-specific CD8+ T-cell responses: from induction to function." *Future Virol.* 5(2): 175-183. **[MA]**
48. Peacock, C. (2010). "Host and parasite genomics, an Australasian perspective." *Parasite Immunology* 32(8): 599-606. **[MA]**
49. Pearton, M., Kang, S. M., Song, J. M., Kim, Y. C., Quan, F. S., Anstey, A., Ivory, M., Prausnitz, M. R., Compans, R. W., and Birchall, J. C. (2010). "Influenza Virus-Like Particles coated onto microneedles can elicit stimulatory effects on Langerhans cells in human skin." *Vaccine* 28(37): 6104-6113. **[MA]**
50. Pitak-Arnlop, P., Schubert, S., Dhanuthai, K., Sappayatosok, K., Bauer, U., Ngamwannagul, P., Liebert, U. G., and Hemprich, A. (2010). "Swine-origin H1N1 influenza A virus and dental practice: a critical review." *Clinical Oral Investigations* 14(1): 11-17. **[MA]**
51. Price, G. E., Soboleski, M. R., Lo, C. Y., Misplon, J. A., Quirion, M. R., Houser, K. V., Pearce, M. B., Pappas, C., Tumpey, T. M., and Epstein, S. L. (2010). "Single-Dose Mucosal Immunization with a Candidate Universal Influenza Vaccine Provides Rapid Protection from Virulent H5N1, H3N2 and H1N1 Viruses." *PloS ONE* 5(10): 6852-6862. **[MA]**
52. Radovanovi , G. and Radovanovi , D. "Fatalni pandemijski grip A-H1N1 kod bolesnice sa faktorima visokog rizika (Fatal pandemic flu A-H1N1 in a female patient with high risk factors)." *PONS Med J* 7(1): 22-25. **[MA]**
53. Rappuoli, R., Medini, D., Siena, E., Budroni, S., Dormitzer, P. R., and Del Giudice, G. (2010). "Building an insurance against modern pandemics." *Current Opinion in Investigational Drugs* 11(2): 126-130. **[MA]**
54. Reed, C. and Katz, M. (2010). "Serological surveys for 2009 pandemic influenza A H1N1." *Lancet* 375(9720): 1062-1063. **[MA]**
55. Reichert, T., Chowell, G., Nishiura, H., Christensen, R., and McCullers, J. (2010). "Does Glycosylation as a modifier of Original Antigenic Sin explain the case age distribution and unusual toxicity in pandemic novel H1N1 influenza?" *BMC Infectious Diseases* 10: 5; doi: 10.1186/1471-2334-10-5. **[MA]**

56. Romero, S. O. (2010). "Desarrollo y autorizacion de vacunas pandemicas de la gripe A (H1N1) 2009 (Development and licensing of pandemic influenza A vaccines (H1N1) 2009)." *Vacunas.Investigacion y Practica* 11(1): 17-24. **[MA]**
57. Rumenyantsev, A. A., Zhang, Z. X., Gao, Q. S., Moretti, N., Brown, N., Kleanthous, H., Delagrave, S., Guirakhoo, F., Collett, M. S., and Pugachev, K. V. (2010). "Direct random insertion of an influenza virus immunologic determinant into the NS1 glycoprotein of a vaccine flavivirus." *Virology* 396(2): 329-338. **[MA]**
58. Schinazi, R. B. (2010). "Will the swine strain crowd out the seasonal influenza strain?" Arxiv preprint arXiv:1004.5569; <http://arxiv.org/abs/1004.5569>. **[MA]**
59. Shin, J. S., Do Hwang, S., Kim, H. S., Cho, S. W., and Seo, S. H. (2010). "Protection of Ferrets from Infection by Swine-Origin 2009 A (H1N1) Influenza Virus by the Inactivated Vaccine." *Viral Immunology* 23(4): 395-402. **[MA]**
60. Sintchenko, V., Anthony, S., Phan, X. H., Lin, F., and Coiera, E. W. (2010). "A PubMed-Wide Associational Study of Infectious Diseases." *PloS ONE* 5(3): e9535. **[MA]**
61. Skowronski, D. M., De Serres, G., Crowcroft, N. S., Janjua, N. Z., Boulianne, N., Hottes, T. S., Rosella, L. C., Dickinson, J. A., Gilca, R., Sethi, P., Ouhoummane, N., Willison, D.J., Rouleau, I., Petric, M., Fonseca, K., Drews, S.J., Rebabpragada, A., Charest, H., Hamelin, M.E., Boivin, G., Gardy, J.L., Li, Y., Kwindt, T.L., Patrick, D.M., and Brunham, R.C. (2010). "Association between the 2008-09 Seasonal Influenza Vaccine and Pandemic H1N1 Illness during Spring-Summer 2009: Four Observational Studies from Canada." *PLoS Medicine* 7(4): e1000258; doi: 10.1371/journal.pmed.1000258. **[MA]**
62. Song, H., Wittman, V., Byers, A., Tapia, T., Zhou, B., Warren, W., Heaton, P., and Connolly, K. (2010). "In vitro stimulation of human influenza-specific CD8+ T cells by dendritic cells pulsed with an influenza virus-like particle (VLP) vaccine." *Vaccine* 28(34): 5524-5532. **[MA]**
63. Stanekova, Z. and Vareckova, E. (2010). "Conserved epitopes of influenza A virus inducing protective immunity and their prospects for universal vaccine development." *Virology Journal* 7: 351; doi: 10.1186/1743-422X-7-351. **[MA]**
64. Stoyanov, C. T., Boscardin, S. B., Deroubaix, S., Barba-Spaeth, G., Franco, D., Nussenzweig, R. S., Nussenzweig, M., and Rice, C. M. (2010). "Immunogenicity and protective efficacy of a recombinant yellow fever vaccine against the murine malarial parasite Plasmodium yoelii." *Vaccine* 28(29): 4644-4652. **[MA]**
65. Sun, Y., Liu, J., Yang, M., Gao, F., Zhou, J., Kitamura, Y., Gao, B., Tien, P., Shu, Y., Iwamoto, A., Chen, Z., and Gao, G.F. (2010). "Identification and structural definition of H5-specific CTL epitopes restricted by HLA-A* 0201 derived from the H5N1 subtype of influenza A viruses." *Journal of General Virology* 91(Pt 4): 919-930. **[MA]**
66. Sun, Y. P., Shi, Y., Zhang, W., Li, Q., Liu, D., Vavricka, C., Yan, J. H., and Gao, G. F. (2010). "In silico characterization of the functional and structural modules of the hemagglutinin protein from the swine-origin influenza virus A (H1N1)-2009." *Science China-Life Sciences* 53(6): 633-642. **[MA]**
67. Suwannakarn, K., Chieochansin, T., Thongmee, C., Makkoch, J., Praianantathavorn, K., Theamboonlers, A., Sreevatsan, S., and Poovorawan, Y. (2010). "Molecular Evolution of Human H1N1 and H3N2 Influenza A Virus in Thailand, 2006-2009." *PloS ONE* 5(3): e9717; doi: 10.1371/journal.pone.0009717. **[MA]**
68. Talaat, K. X., Greenberg, M. X., Lai, M. X., Hartel, G. X., Wichems, C. X., Rockman, S., Jeanfreau, R. X., Ghosh, M. X., Kabongo, M. X., Gittleson, C., and Karron, R.A. (2010). "A Single Dose of Unadjuvanted Novel 2009 H1N1 Vaccine Is Immunogenic and Well Tolerated in Young and Elderly Adults." *The Journal of Infectious Diseases* 202(9): 1327-1337. **[MA]**

69. Tang, J. W., Lee, C. K., Lee, H. K., Loh, T. P., Chiu, L., Tambyah, P. A., and Koay, E. S. (2010). "Tracking the emergence of pandemic Influenza A/H1N1/2009 and its interaction with seasonal influenza viruses in Singapore." *Ann Acad Med Singapore* 39(4): 291-294. **[MA]**
70. Tang, J. W., Shetty, N., and Lam, T. T. Y. (2010). "Features of the new pandemic influenza A/H1N1/2009 virus: virology, epidemiology, clinical and public health aspects." *Current Opinion in Pulmonary Medicine* 16(3): 235-241. **[MA]**
71. Teijaro, J. R., Verhoeven, D., Page, C. A., Turner, D., and Farber, D. L. (2010). "Memory CD4 T Cells Direct Protective Responses to Influenza Virus in the Lungs through Helper-Independent Mechanisms." *Journal of Virology* 84(18): 9217-9226. **[MA]**
72. To, K. K. W., Li, I. W. S., Hung, I. F. N., Cheng, V. C. C., and Yuen, K. Y. (2010). "Pathogenesis of pandemic H1N1 2009 influenza virus infection and the implication on management." *Frontiers of Medicine in China* 4(2): 147-156. **[MA]**
73. Tu, W., Mao, H., Zheng, J., Liu, Y., Chiu, S. S., Qin, G., Chan, P. L., Lam, K. T., Guan, J., Zhang, L., Guan, Y., Yuen, K.Y., Peiris, J.S., and Lau, Y.L. (2010). "Cytotoxic T Lymphocytes Established by Seasonal Human Influenza Cross-React against 2009 Pandemic H1N1 Influenza Virus." *Journal of Virology* 84(13): 6527-6535. **[MA]**
74. Turner, S. J., Doherty, P. C., and Kelso, A. (2010). "Q & A: H1N1 pandemic influenza- what's new?" *BMC biology* 8: 130, <http://www.biomedcentral.com/1741-7007/8/130>; doi: 10.1186/1741-7007-8-130. **[MA]**
75. Valentini, D., Gaseitsiwe, S., and Maeurer, M. (2010). "Humoral 'reactome' profiles using peptide microarray chips." *Trends in Immunology* 31(11): 399-400; doi: doi:10.1016/j.it.2010.08.007. **[MA]**
76. Valleron, A. J. and Guidet, B. (2010). "Real-time comparative monitoring of the A/H1N1 pandemic in France." *Clinical Microbiology and Infection* 16(4): 393-396. **[MA]**
77. Xu, R., Ekiert, D. C., Krause, J. C., Hai, R., Crowe Jr, J. E., and Wilson, I. A. (2010). "Structural basis of preexisting immunity to the 2009 H1N1 pandemic influenza virus." *Science* 328(5976): 357-360. **[MA]**
78. Zichel, R., Mimran, A., Keren, A., Barnea, A., Steinberger-Levy, I., Marcus, D., Turgeman, A., and Reuveny, S. (2010). "Efficacy of a Potential Trivalent Vaccine Based on Hc Fragments of Botulinum Toxins A, B, and E Produced in a Cell-Free Expression System." *Clinical and Vaccine Immunology* 17(5): 784-792. **[MA]**

4 References

- Beaver JE, Bourne PE, Ponomarenko JV. EpitopeViewer: a Java application for the visualization and analysis of immune epitopes in the Immune Epitope Database and Analysis Resource (IEDB). *Immunome Res.* 2007 Feb 21;3:3. PMID: 17313688
- Bui HH, Sidney J, Peters B, Sathiamurthy M, Sinichi A, Purton KA, Mothe BR, Chisari FV, Watkins DI, Sette A. Automated generation and evaluation of specific MHC binding predictive tools: ARB matrix applications. *Immunogenetics.* 2005 Jun;57(5):304-14. Epub 2005 May 3.
- Bui HH, Sidney J, Dinh K, Southwood S, Newman MJ, Sette A. Predicting population coverage of T-cell epitope-based diagnostics and vaccines. *BMC Bioinformatics.* 2006 Mar 17;7(1):153. PMID: 16545123
- Bui HH, Sidney J, Li W, Fusseeder N, Sette A. Development of an epitope conservancy analysis tool to facilitate the design of epitope-based diagnostics and vaccines. *BMC Bioinformatics.* 2007 Sep 26;8(1):361. PMID: 17897458
- Chou P, Fasman G. Prediction of the secondary structure of proteins from their amino acid sequence. *Adv Enzymol Relat Areas Mol Biol.* 1978:45–148. PMID: 364941
- Emini E, Hughes J, Perlow D, Boger J. Induction of hepatitis A virus-neutralizing antibody by a virus specific synthetic peptide. *J Virol.* 1985;55:836–839. PMID: 2991600
- Haste Andersen P, Nielsen M, Lund O. Prediction of residues in discontinuous B-cell epitopes using protein 3D structures. *Protein Sci.* 2006 Nov;15(11):2558-67. Epub 2006 Sep 25. PMID: 17001032
- Karplus PA, Schulz GE. Prediction of chain flexibility in proteins. *Naturwissenschaften.* 1985. 72:212-213.
- Keşmir C, Nussbaum AK, Schild H, Detours V, Brunak S. Prediction of proteasome cleavage motifs by neural networks. *Protein Eng.* 2002 Apr;15(4):287-96. PMID: 11983929
- Kim Y, Sidney J, Pinilla C, Sette A, Peters B. Derivation of an amino acid similarity matrix for peptide: MHC binding and its application as a Bayesian prior. *BMC Bioinformatics.* 2009 Nov 30;10:394. PMID: 19948066
- Kolaskar AS, Tongaonkar PC. A semi-empirical method for prediction of antigenic determinants on protein antigens. *FEBS Lett.* 1990 Dec 10;276(1-2):172-4. PMID: 1702393
- Larsen JE, Lund O, Nielsen M. Improved method for predicting linear B-cell epitopes. *Immunome Res.* 2006 Apr 24;2:2. PMID: 16635264
- Larsen MV, Lundsgaard C, Lamberth K, Buus S, Brunak S, Lund O, Nielsen M. An integrative approach to CTL epitope prediction: a combined algorithm integrating MHC class I binding, TAP transport efficiency, and proteasomal cleavage predictions. *Eur J Immunol.* 2005 Aug;35(8):2295-303. PMID: 15997466

- Moutafsi M, Peters B, Pasquetto V, Tscharke DC, Sidney J, Bui HH, Grey H, Sette A. A consensus epitope prediction approach identifies the breadth of murine T(CD8+)-cell responses to vaccinia virus. *Nat Biotechnol.* 2006 Jul;24(7):817-9. PMID: 16767078
- Nielsen M, Justesen S, Lund O, Lundsgaard C, Buus S. NetMHCIIpan-2.0 - Improved pan-specific HLA-DR predictions using a novel concurrent alignment and weight optimization training procedure. *Immunome Res.* 2010 Nov 13;6:9. PMID: 21073747
- Nielsen M, Lundsgaard C, Blicher T, Lamberth K, Harndahl M, Justesen S, Roder G, Peters B, Sette A, Lund O, Buus S. 2007. NetMHCpan, a method for quantitative predictions of peptide binding to any HLA-A and -B locus protein of known sequence. *PLoS ONE* 2:e796. PMID: 17726526
- Nielsen M, Lundsgaard C, Lund O. Prediction of MHC class II binding affinity using SMM-align, a novel stabilization matrix alignment method. *BMC Bioinformatics.* 2007 Jul 4;8:238. PMID: 17608956
- Nielsen M, Lundsgaard C, Worning P, Lauemoller SL, Lamberth K, Buus S, Brunak S, Lund O., Reliable prediction of T-cell epitopes using neural networks with novel sequence representations, *Protein Sci.* 2003 May;12(5):1007-17
- Parker J, Guo D, Hodges R. New hydrophilicity scale derived from High-Performance Liquid Chromatography peptide retention data: correlation of predicted surface residues with antigenicity and X-ray-derived accessible sites. *Biochemistry.* 1986;25:5425–5432. PMID: 2430611
- Peters B, Sidney J, Bourne P, Bui HH, Buus S, Doh G, Fleri W, Kronenberg M, Kubo R, Lund O, Nemazee D, Ponomarenko JV, Sathiamurthy M, Schoenberger S, Stewart S, Surko P, Way S, Wilson S, Sette A. The immune epitope database and analysis resource: from vision to blueprint. *PLoS Biol.* 2005 Mar;3(3):e91. PMID: 15760272.
- Peters B, Bui HH, Frankild S, Nielson M, Lundsgaard C, Kostem E, Basch D, Lamberth K, Harndahl M, Fleri W, Wilson SS, Sidney J, Lund O, Buus S, Sette A., A community resource benchmarking predictions of peptide binding to MHC-I molecules, *PLoS Comput. Biol.* 2006 Jun 9;2(6):e65. Epub 2006 Jun 9. PMID: 16789818
- Peters B, Sette A., "Generating quantitative models describing the sequence specificity of biological processes with the stabilized matrix method." *BMC Bioinformatics* 2005 May 31;6(1):132
- Peters B, Bulik S, Tampe R, Van Endert PM, Holzhutter HG. Identifying MHC class I epitopes by predicting the TAP transport efficiency of epitope precursors. *J Immunol.* 2003 Aug 15;171(4):1741-9.
- Ponomarenko J, H. H. Bui, W. Li, N. Fusseeder, P. E. Bourne, A. Sette, B. Peters. (2008). "ElliPro: a new structure-based tool for the prediction of antibody epitopes." *BMC Bioinformatics*, 9(514). PMID: 19055730
- Sidney J, Assarsson E, Moore C, Ngo S, Pinilla C, Sette A, Peters B. Quantitative peptide binding motifs for 19 human and mouse MHC class I molecules derived using positional scanning combinatorial peptide libraries. *Immunome Res.* 2008 Jan 25;4:2. PMID: 18221540

Sturniolo T, Bono E, Ding J, Raddrizzani L, Tuereci O, Sahin U, Braxenthaler M, Gallazzi F, Protti MP, Sinigaglia F, Hammer J. Generation of tissue-specific and promiscuous HLA ligand databases using DNA microarrays and virtual HLA class II matrices. *Nat Biotechnol.* 1999 Jun;17(6):555-61. PMID: 10385319

Tenzer S, Peters B, Bulik S, Schoor O, Lemmel C, Schatz MM, Kloetzel PM, Rammensee HG, Schild H, Holzhutter HG. Modeling the MHC class I pathway by combining predictions of proteasomal cleavage, TAP transport and MHC class I binding. *Cell Mol Life Sci.* 2005 May;62(9):1025-37.

Wang, P., J. Sidney, C. Dow, B. Mothe, A. Sette, B. Peters. (2008). "A Systematic Assessment of MHC Class II Peptide Binding Predictions and Evaluation of a Consensus Approach." *PLoS Computational Biology* 4(4). PMID: 18389056