

Appendices

Appendix A

Table A.1: 20 most frequent *i*Pfam domain pairs in protein interactions of 5 species.

Accession A	Name A	Accession B	Name B	Frequency
<i>E. coli</i>				
PF00005	ABC_tran	PF00005	ABC_tran	21
PF00072	Response_reg	PF00072	Response_reg	19
PF00126	HTH_1	PF00126	HTH_1	17
PF03466	LysR_substrate	PF00126	HTH_1	16
PF03466	LysR_substrate	PF03466	LysR_substrate	16
PF00271	Helicase_C	PF00271	Helicase_C	15
PF00313	CSD	PF00313	CSD	14
PF00106	adh_short	PF00106	adh_short	12
PF00532	Peripla_BP_1	PF00532	Peripla_BP_1	11
PF00293	NUDIX	PF00293	NUDIX	10
PF00271	Helicase_C	PF00270	DEAD	10
PF00216	Bac_DNA_binding	PF00216	Bac_DNA_binding	9
PF00392	GntR	PF00392	GntR	9
PF00575	S1	PF00575	S1	9
PF00009	GTP_EFTU	PF00009	GTP_EFTU	9
PF00158	Sigma54_activat	PF00158	Sigma54_activat	9
PF02518	HATPase_c	PF02518	HATPase_c	9
PF03144	GTP_EFTU_D2	PF03144	GTP_EFTU_D2	9
PF03144	GTP_EFTU_D2	PF00009	GTP_EFTU	9
PF00270	DEAD	PF00270	DEAD	9
<i>S. cerevisiae</i>				
PF00069	Pkinase	PF00069	Pkinase	266
PF00400	WD40	PF00400	WD40	141
PF00227	Proteasome	PF00227	Proteasome	96
PF01423	LSM	PF01423	LSM	84
PF00076	RRM_1	PF00076	RRM_1	79
PF00271	Helicase_C	PF00271	Helicase_C	74
PF00134	Cyclin_N	PF00069	Pkinase	65

Accession A	Name A	Accession B	Name B	Frequency
PF00271	Helicase_C	PF00270	DEAD	51
PF00018	SH3_1	PF00018	SH3_1	49
PF00004	AAA	PF00004	AAA	46
PF00270	DEAD	PF00270	DEAD	41
PF02984	Cyclin_C	PF00069	Pkinase	35
PF00069	Pkinase	PF00023	Ank	32
PF00433	Pkinase_C	PF00069	Pkinase	30
PF00172	Zn_clus	PF00172	Zn_clus	27
PF05739	SNARE	PF00957	Synaptobrevin	26
PF02985	HEAT	PF02985	HEAT	25
PF00125	Histone	PF00125	Histone	24
PF00271	Helicase_C	PF00176	SNF2_N	21
PF00575	S1	PF00069	Pkinase	20
<i>C. elegans</i>				
PF00105	zf-C4	PF00105	zf-C4	33
PF00104	Hormone_recep	PF00104	Hormone_recep	31
PF00105	zf-C4	PF00104	Hormone_recep	31
PF00595	PDZ	PF00595	PDZ	12
PF00076	RRM_1	PF00076	RRM_1	12
PF00227	Proteasome	PF00227	Proteasome	11
PF00069	Pkinase	PF00069	Pkinase	11
PF02932	Neur_chan_memb	PF02932	Neur_chan_memb	9
PF02931	Neur_chan_LBD	PF02931	Neur_chan_LBD	9
PF02932	Neur_chan_memb	PF02931	Neur_chan_LBD	9
PF00004	AAA	PF00004	AAA	6
PF00412	LIM	PF00018	SH3_1	5
PF01423	LSM	PF01423	LSM	5
PF02188	GoLoco	PF00503	G-alpha	5
PF00651	BTB	PF00651	BTB	4
PF01849	NAC	PF01849	NAC	4
PF00412	LIM	PF00412	LIM	4
PF00595	PDZ	PF00071	Ras	4
PF01849	NAC	PF00627	UBA	4
PF01466	Skp1	PF00646	F-box	4
<i>D. melanogaster</i>				
PF00096	zf-C2H2	PF00096	zf-C2H2	117
PF00134	Cyclin_N	PF00069	Pkinase	63
PF00076	RRM_1	PF00076	RRM_1	54
PF00010	HLH	PF00010	HLH	54
PF01466	Skp1	PF00646	F-box	48
PF00069	Pkinase	PF00069	Pkinase	38
PF00595	PDZ	PF00071	Ras	22
PF00788	RA	PF00071	Ras	21
PF02984	Cyclin_C	PF00069	Pkinase	21
PF00612	IQ	PF00036	efhand	21
PF00179	UQ_con	PF00097	zf-C3HC4	21
PF00046	Homeobox	PF00046	Homeobox	20

Accession A	Name A	Accession B	Name B	Frequency
PF00069	Pkinase	PF00023	Ank	20
PF00651	BTB	PF00651	BTB	14
PF01423	LSM	PF01423	LSM	14
PF00063	Myosin_head	PF00036	efhand	13
PF00134	Cyclin_N	PF00134	Cyclin_N	11
PF00018	SH3_1	PF00017	SH2	11
PF03931	Skp1_POZ	PF00560	LRR_1	10
PF02179	BAG	PF00012	HSP70	10
<i>H. sapiens</i>				
PF07714	Pkinase_Tyr	PF00017	SH2	464
PF00069	Pkinase	PF00069	Pkinase	386
PF07714	Pkinase_Tyr	PF00018	SH3_1	318
PF00018	SH3_1	PF00017	SH2	241
PF00017	SH2	PF00017	SH2	200
PF00018	SH3_1	PF00018	SH3_1	179
PF07714	Pkinase_Tyr	PF07714	Pkinase_Tyr	162
PF00076	RRM_1	PF00076	RRM_1	147
PF00433	Pkinase_C	PF00069	Pkinase	112
PF00010	HLH	PF00010	HLH	95
PF00069	Pkinase	PF00023	Ank	74
PF00105	zf-C4	PF00104	Hormone_recep	72
PF00104	Hormone_recep	PF00104	Hormone_recep	71
PF00096	zf-C2H2	PF00096	zf-C2H2	71
PF00089	Trypsin	PF00079	Serpin	66
PF00105	zf-C4	PF00105	zf-C4	60
PF07714	Pkinase_Tyr	PF00102	Y_phosphatase	58
PF00169	PH	PF00071	Ras	56
PF00046	Homeobox	PF00046	Homeobox	54
PF00619	CARD	PF00619	CARD	54

Table A.2: 20 most frequent iPfam domain pairs in protein interactions of 5 species, excluding intrachain structures.

Accession A	Name A	Accession B	Name B	Frequency
<i>E. coli</i>				
PF00005	ABC_tran	PF00005	ABC_tran	21
PF00072	Response_reg	PF00072	Response_reg	19
PF00126	HTH_1	PF00126	HTH_1	17
PF03466	LysR_substrate	PF00126	HTH_1	16
PF03466	LysR_substrate	PF03466	LysR_substrate	16
PF00271	Helicase_C	PF00271	Helicase_C	15
PF00313	CSD	PF00313	CSD	14
PF00106	adh_short	PF00106	adh_short	12
PF00532	Peripla_BP_1	PF00532	Peripla_BP_1	11

Accession A	Name A	Accession B	Name B	Frequency
PF00293	NUDIX	PF00293	NUDIX	10
PF00392	GntR	PF00392	GntR	9
PF02518	HATPase_c	PF02518	HATPase_c	9
PF00575	S1	PF00575	S1	9
PF00009	GTP_EFTU	PF00009	GTP_EFTU	9
PF00158	Sigma54_activat	PF00158	Sigma54_activat	9
PF00270	DEAD	PF00270	DEAD	9
PF03144	GTP_EFTU_D2	PF03144	GTP_EFTU_D2	9
PF00216	Bac_DNA_binding	PF00216	Bac_DNA_binding	9
PF03144	GTP_EFTU_D2	PF00009	GTP_EFTU	9
PF00004	AAA	PF00004	AAA	8
<i>S. cerevisiae</i>				
PF00069	Pkinase	PF00069	Pkinase	266
PF00400	WD40	PF00400	WD40	141
PF00227	Proteasome	PF00227	Proteasome	96
PF01423	LSM	PF01423	LSM	84
PF00076	RRM.1	PF00076	RRM.1	79
PF00271	Helicase_C	PF00271	Helicase_C	74
PF00134	Cyclin_N	PF00069	Pkinase	65
PF00018	SH3.1	PF00018	SH3.1	49
PF00004	AAA	PF00004	AAA	46
PF00270	DEAD	PF00270	DEAD	41
PF02984	Cyclin_C	PF00069	Pkinase	35
PF00069	Pkinase	PF00023	Ank	32
PF00433	Pkinase_C	PF00069	Pkinase	30
PF00172	Zn_clus	PF00172	Zn_clus	27
PF05739	SNARE	PF00957	Synaptobrevin	26
PF02985	HEAT	PF02985	HEAT	25
PF00125	Histone	PF00125	Histone	24
PF00575	S1	PF00069	Pkinase	20
PF01138	RNase_PH	PF01138	RNase_PH	19
PF03725	RNase_PH_C	PF01138	RNase_PH	19
<i>C. elegans</i>				
PF00105	zf-C4	PF00105	zf-C4	33
PF00105	zf-C4	PF00104	Hormone_recep	31
PF00104	Hormone_recep	PF00104	Hormone_recep	31
PF00076	RRM.1	PF00076	RRM.1	12
PF00595	PDZ	PF00595	PDZ	12
PF00227	Proteasome	PF00227	Proteasome	11
PF00069	Pkinase	PF00069	Pkinase	11
PF02932	Neur_chan_memb	PF02932	Neur_chan_memb	9
PF02931	Neur_chan_LBD	PF02931	Neur_chan_LBD	9
PF02932	Neur_chan_memb	PF02931	Neur_chan_LBD	9
PF00004	AAA	PF00004	AAA	6
PF01423	LSM	PF01423	LSM	5
PF00412	LIM	PF00018	SH3.1	5
PF02188	GoLoco	PF00503	G-alpha	5

Accession A	Name A	Accession B	Name B	Frequency
PF00595	PDZ	PF00071	Ras	4
PF01849	NAC	PF01849	NAC	4
PF01466	Skp1	PF00646	F-box	4
PF00651	BTB	PF00651	BTB	4
PF00210	Ferritin	PF00210	Ferritin	3
PF00017	SH2	PF00017	SH2	3
<i>D. melanogaster</i>				
PF00096	zf-C2H2	PF00096	zf-C2H2	117
PF00134	Cyclin_N	PF00069	Pkinase	63
PF00076	RRM_1	PF00076	RRM_1	54
PF00010	HLH	PF00010	HLH	54
PF01466	Skp1	PF00646	F-box	48
PF00069	Pkinase	PF00069	Pkinase	38
PF00595	PDZ	PF00071	Ras	22
PF02984	Cyclin_C	PF00069	Pkinase	21
PF00179	UQ_con	PF00097	zf-C3HC4	21
PF00788	RA	PF00071	Ras	21
PF00612	IQ	PF00036	efhand	21
PF00046	Homeobox	PF00046	Homeobox	20
PF00069	Pkinase	PF00023	Ank	20
PF00651	BTB	PF00651	BTB	14
PF01423	LSM	PF01423	LSM	14
PF00063	Myosin_head	PF00036	efhand	13
PF00134	Cyclin_N	PF00134	Cyclin_N	11
PF00018	SH3_1	PF00017	SH2	11
PF02179	BAG	PF00012	HSP70	10
PF02196	RBD	PF00071	Ras	10
<i>H. sapiens</i>				
PF07714	Pkinase_Tyr	PF00017	SH2	464
PF00069	Pkinase	PF00069	Pkinase	386
PF00018	SH3_1	PF00017	SH2	241
PF00017	SH2	PF00017	SH2	200
PF00018	SH3_1	PF00018	SH3_1	179
PF07714	Pkinase_Tyr	PF07714	Pkinase_Tyr	162
PF00076	RRM_1	PF00076	RRM_1	147
PF00433	Pkinase_C	PF00069	Pkinase	112
PF00010	HLH	PF00010	HLH	95
PF00069	Pkinase	PF00023	Ank	74
PF00105	zf-C4	PF00104	Hormone_recep	72
PF00104	Hormone_recep	PF00104	Hormone_recep	71
PF00096	zf-C2H2	PF00096	zf-C2H2	71
PF00089	Trypsin	PF00079	Serpin	66
PF00105	zf-C4	PF00105	zf-C4	60
PF07714	Pkinase_Tyr	PF00102	Y_phosphatase	58
PF00169	PH	PF00071	Ras	56
PF00046	Homeobox	PF00046	Homeobox	54
PF00619	CARD	PF00619	CARD	54

Accession A	Name A	Accession B	Name B	Frequency
PF00531	Death	PF00531	Death	53

Appendix B

Table B.1: All structures

E coli			Yeast			Worm			Fly			Human		
Accession	Name	Freq.	Accession	Name	Freq.	Accession	Name	Freq.	Accession	Name	Freq.	Accession	Name	Freq.
PF000005	ABC-tran	49	PF00069	Pkinase	113	PF00069	Pkinase	82	PF00096	zf-C2H2	234	PF00069	Pkinase	346
PF000072	Response_reg	36	PF00400	WD40	87	PF00105	zf-C4	63	PF00069	Pkinase	171	PF00096	zf-C2H2	205
PF025118	HATPase_c	31	PF00271	Helicase_C	70	PF00104	Hormone_recep	57	PF00076	RRM_1	124	PF00169	PH	174
PF001266	HTH_1	30	PF00172	Zn_c1us	49	PF01391	Collagen	46	PF00400	WD40	97	PF00076	RRM_1	172
PF03466	LysR_substrate	28	PF00270	DEAD	48	PF00076	RRM_1	39	PF00046	Homeobox	79	PF00018	SH3_1	170
PF00672	HAMP	20	PF00076	RRM_1	47	PF00400	WD40	32	PF00089	Trypsin	72	PF00400	WD40	151
PF00512	HisKA	19	PF00096	zf-C2H2	36	PF00595	PDZ	28	PF00036	efhand	67	PF00001	7tm_1	145
PF00486	Trans_reg_C	16	PF00004	AAA	33	PF00097	zf-C3HC4	27	PF00097	zf-C3HC4	65	PF00595	PDZ	132
PF00532	Peripla_BP_1	16	PF00005	ABC_tran	30	PF00096	zf-C2H2	25	PF00595	PDZ	65	PF00097	zf-C3HC4	132
PF00271	Helicase_C	15	PF00153	Mito_carr	30	PF02798	GST_N	25	PF00023	Ank	64	PF00047	ig	131
PF00392	GntR	14	PF02985	HEAT	26	PF00043	GST_C	23	PF00018	SH3_1	60	PF00023	Ank	117
PF00106	adh_short	13	PF00071	Ras	24	PF00646	F-box	23	PF00560	LRR_1	60	PF00017	SH2	109
PF00158	Sigma54_activat	13	PF00169	PH	24	PF00651	BTB	22	PF00271	Helicase_C	57	PF00041	fn3	109
PF00196	GerE	13	PF00018	SH3_1	22	PF00018	SH3_1	20	PF00047	ig	56	PF00036	efhand	107
PF00037	Fer4	12	PF00702	Hydrolase	22	PF00169	PH	20	PF00169	PH	54	PF07686	V-set	104
PF00165	HTH_AraC	12	PF00665	rve	21	PF00023	Ank	19	PF07679	L-set	52	PF07714	Pkinase_Tyr	99
PF04055	Radical_SAM	11	PF00097	zf-C3HC4	21	PF00004	AAA	19	PF00651	BTB	51	PF00008	EGF	92
PF00293	NUDIX	11	PF00515	TPR_1	21	PF00271	Helicase_C	19	PF00010	HLH	45	PF00046	Homeobox	92
PF00455	DeoR	10	PF00149	Metallophos	21	PF00046	Homeobox	19	PF00041	fn3	43	PF00560	LRR_1	91
PF02854	HTH_8	10	PF00226	DnaJ	21	PF00149	Metallophos	19	PF00515	TPR_1	43	PF00168	C2	87

Table B.2: Interchain only

E coli			Yeast			Worm			Fly			Human		
Accession	Name	Freq.	Accession	Name	Freq.	Accession	Name	Freq.	Accession	Name	Freq.	Accession	Name	Freq.
PF000005	ABC-tran	49	PF00069	Pkinase	113	PF00069	Pkinase	82	PF00096	zf-C2H2	234	PF00069	Pkinase	346
PF000072	Response_reg	36	PF00400	WD40	87	PF00105	zf-C4	63	PF00069	Pkinase	171	PF00096	zf-C2H2	205
PF025118	HATPase_c	31	PF00271	Helicase_C	70	PF00104	Hormone_recep	57	PF00076	RRM_1	124	PF00169	PH	174
PF001266	HTH_1	30	PF00172	Zn_c1us	49	PF01391	Collagen	46	PF00400	WD40	97	PF00076	RRM_1	172
PF03466	LysR_substrate	28	PF00270	DEAD	48	PF00076	RRM_1	39	PF00046	Homeobox	79	PF00018	SH3_1	170
PF00672	HAMP	20	PF00076	RRM_1	47	PF00400	WD40	32	PF00089	Trypsin	72	PF00400	WD40	151
PF00486	Trans_reg_C	16	PF00096	zf-C2H2	36	PF00595	PDZ	28	PF00036	efhand	67	PF00001	7tm_1	145
PF00532	Peripla_BP_1	16	PF00004	AAA	33	PF00097	zf-C3HC4	27	PF00097	zf-C3HC4	65	PF00595	PDZ	132
PF00271	Helicase_C	15	PF00005	ABC_tran	30	PF02798	GST_N	25	PF00595	PDZ	65	PF00097	zf-C3HC4	132
PF00392	GntR	14	PF02985	HEAT	26	PF00096	zf-C2H2	25	PF00023	Ank	64	PF00047	ig	131
PF00106	adh_short	13	PF00071	Ras	24	PF00043	GST_C	23	PF00018	SH3_1	60	PF00023	Ank	117
PF00158	Sigma54_activat	13	PF00169	PH	24	PF00646	F-box	23	PF00560	LRR_1	60	PF00017	SH2	109
PF00196	GerE	13	PF00018	SH3_1	22	PF00651	BTB	22	PF00271	Helicase_C	57	PF00041	fn3	109
PF00165	HTH_AraC	12	PF00702	Hydrolase	22	PF00018	SH3_1	20	PF00047	ig	56	PF00036	efhand	107
PF00037	Fer4	12	PF00226	DnaJ	21	PF00169	PH	20	PF00169	PH	54	PF07686	V-set	104
PF00293	NUDIX	11	PF00097	zf-C3HC4	21	PF00023	Ank	19	PF07679	L-set	52	PF07714	Pkinase_Tyr	99
PF00270	DEAD	10	PF00665	rve	21	PF00271	Helicase_C	19	PF00651	BTB	51	PF00008	EGF	92
PF00455	DeoR	10	PF00149	Metallophos	21	PF00004	AAA	19	PF00010	HLH	45	PF00046	Homeobox	92
PF02954	HTH_8	10	PF00515	TPR_1	21	PF00149	Metallophos	19	PF00041	fn3	43	PF00360	LRR_1	91
PF00155	Aminotran_1_2	10	PF08240	ADH_N	20	PF00046	Homeobox	19	PF00515	TPR_1	43	PF00168	C2	87

Table B.3: No crystal contacts

E coli				Yeast				Worm				Fly				Human				
Accession	Name	Freq.	Accession	Name	Freq.	Accession	Name	Accession	Name	Freq.	Accession	Name	Accession	Name	Freq.	Accession	Name	Accession	Name	Freq.
PF000005	ABC-tran	49	PF00069	Pkinase	113	PF00069	Pkinase	PF00069	Pkinase	82	PF00096	zf-C2H2	PF00096	Pkinase	234	PF00069	Pkinase	PF00069	Pkinase	346
PF000072	Response_reg	36	PF00400	WD40	87	PF00105	zf-C4	PF00105	zf-C4	63	PF00069	Pkinase	PF00069	zf-C2H2	171	PF00096	zf-C2H2	PF00096	zf-C2H2	205
PF02518	HATPase_c	31	PF00271	Helicase_C	70	PF00104	Hormone_recep	PF00104	Hormone_recep	57	PF00076	RRM_1	PF00076	RRM_1	124	PF00169	PH	PF00169	PH	174
PF00126	HTH_1	30	PF00172	Zn-clus	49	PF01391	Collagen	PF01391	Collagen	46	PF00400	WD40	PF00400	RRM_1	97	PF00076	RRM_1	PF00076	RRM_1	172
PF03466	LysR_substrate	28	PF00270	DEAD	48	PF00076	RRM_1	PF00076	RRM_1	39	PF00046	Homeobox	PF00046	SH3_1	79	PF00018	SH3_1	PF00018	SH3_1	170
PF00672	HAMP	20	PF00076	RRM_1	47	PF00400	WD40	PF00400	WD40	32	PF00089	Trypsin	PF00089	WD40	72	PF00400	WD40	PF00400	WD40	151
PF00532	Peripla_BP_1	16	PF00096	zf-C2H2	36	PF00595	PDZ	PF00036	efhand	28	PF00036	efhand	PF00036	7tm_1	67	PF00001	7tm_1	PF00001	7tm_1	145
PF00271	Helicase_C	15	PF00004	AAA	33	PF00097	zf-C3HC4	PF00097	zf-C3HC4	27	PF00595	PDZ	PF00097	zf-C3HC4	65	PF00097	zf-C3HC4	PF00097	zf-C3HC4	132
PF00392	GntR	14	PF00005	ABC_tran	30	PF02798	GST_N	PF02798	GST_N	25	PF00097	zf-C3HC4	PF00097	PDZ	65	PF00595	PDZ	PF00595	PDZ	132
PF00106	adh_short	13	PF02985	HEAT	26	PF00096	zf-C2H2	PF00096	zf-C2H2	25	PF00023	Ank	PF00023	ig	64	PF00047	ig	PF00047	ig	131
PF00158	Sigma54_activat	13	PF00071	Ras	24	PF00043	GST_C	PF00043	GST_C	23	PF00018	SH3_1	PF00018	Ank	60	PF00023	Ank	PF00023	Ank	117
PF00196	GerE	13	PF00169	PH	24	PF00646	F-box	PF00646	F-box	23	PF00560	LRR_1	PF00560	SH2	60	PF00017	SH2	PF00017	SH2	109
PF00165	HTH_AraC	12	PF00018	SH3_1	22	PF00651	BTB	PF00651	BTB	22	PF00271	Helicase_C	PF00271	fn3	57	PF00041	fn3	PF00041	fn3	109
PF00037	Fer4	12	PF00702	Hydrolase	22	PF00018	SH3_1	PF00018	SH3_1	20	PF00047	ig	PF00047	efhand	56	PF00036	efhand	PF00036	efhand	107
PF00293	NUDIX	11	PF00665	rve	21	PF00169	PH	PF00169	PH	20	PF00169	PH	PF00169	V-set	54	PF07686	V-set	PF07686	V-set	104
PF00155	Aminotran_1-2	10	PF00097	zf-C3HC4	21	PF00023	Ank	PF00023	Ank	19	PF07679	L-set	PF07679	Pkinase_Tyr	52	PF07714	Pkinase_Tyr	PF07714	Pkinase_Tyr	99
PF00171	Alcedh	10	PF00149	Metallophos	21	PF00004	AAA	PF00004	AAA	19	PF00651	BTB	PF00651	Homeobox	51	PF00046	Homeobox	PF00046	Homeobox	92
PF00270	DEAD	10	PF00515	TPR_1	21	PF00046	Homeobox	PF00046	Homeobox	19	PF00010	HLH	PF00010	EGF	45	PF00008	EGF	PF00008	EGF	92
PF02954	HTH_8	10	PF08240	ADH_N	20	PF00149	Metallophos	PF00149	Metallophos	19	PF00041	fn3	PF00041	LRR_1	43	PF00560	LRR_1	PF00560	LRR_1	91
PF07892	Pyr_redox_2	10	PF00107	ADH_zinc_N	17	PF00271	Helicase_C	PF00271	Helicase_C	19	PF00515	TPR_1	PF00515	C2	43	PF00168	C2	PF00168	C2	87

Appendix C

Table C.1: The 30 most frequent *i*Pfam domain architectures per species. The left column lists the sequence of *i*Pfam domains that comprises a distinct domain architecture, separated by a “—”. Non-*i*Pfam domains are omitted to underline the effect of domain architecture on *i*Pfam domain pair frequency. The right column contains the frequency of the architecture, defined as the number of sequences which share the same architecture.

Architecture	Frequency
<i>E. coli</i>	
ABC_tran	32
HTH_1 — LysR_substrate	28
Peripla_BP_1	15
adh_short	13
Response_reg — Trans_reg_C	13
ABC_tran — ABC_tran	11
DeoR	10
NUDIX	10
HAMP — HisKA — HATPase_c	9
Aldedh	9
Aminotran_1_2	8
DEAD — Helicase_C	8
Response_reg — GerE	7
CSD	7
GntR	7
S1	6
Response_reg	6
TPP_enzyme_N — TPP_enzyme_M — TPP_enzyme_C	6
ADH_N — ADH_zinc_N	6
Aminotran_3	6
Fe-ADH	6
GerE	6
Acetyltransf_1	6
Glycos_transf_2	6
Hydrolase	6
Radical_SAM	6

Architecture	Frequency
4HBT	5
NTP_transferase	5
Hydrolase_3	5
Pribosyltran	5
<i>S. cerevisiae</i>	
Pkinase	93
Zn_clus	47
DEAD — Helicase_C	42
RRM_1	29
Mito_carr — Mito_carr — Mito_carr	26
Ras	24
zf-C2H2 — zf-C2H2	22
rve	20
Metallophos	20
WD40 — WD40	18
ADH_N — ADH_zinc_N	16
LSM	16
WD40	14
Aldo_ket_red	14
PH	14
Proteasome	14
HSP70	14
DnaJ	13
AAA	13
UQ_con	13
zf-C3HC4	13
Abhydrolase_1	12
WD40 — WD40 — WD40 — WD40 — WD40	11
ABC_tran — ABC_tran	11
SH3_1	11
adh_short	11
WD40 — WD40 — WD40 — WD40	11
Aminotran_1_2	11
Acetyltransf_1	11
Hydrolase	11
<i>C. elegans</i>	
Pkinase	60
zf-C4 — Hormone_recep	56
Collagen — Collagen — Collagen	30
zf-C3HC4	23
RRM_1	22
GST_N — GST_C	22
F-box	20
Metallophos	18
Collagen — Collagen	15
Homeobox	15
Kinesin	14

Architecture	Frequency
AAA	14
p450	13
K_tetra	12
Proteasome	12
Neur_chan.LBD — Neur_chan_memb	11
BTB	11
Motile.Sperm	10
Filament	10
Ras	10
Arrestin_N — Arrestin_C	10
zf-CCCH — zf-CCCH	10
zf-C2H2	9
ubiquitin	9
MATH — BTB	9
COesterase	9
7tm_1	9
Aminotran_1.2	8
RRM_1 — RRM_1 — RRM_1	8
DEAD — Helicase_C	8
<i>D. melanogaster</i>	
Pkinase	109
Trypsin	66
RRM_1	55
Homeobox	55
zf-C3HC4	45
RRM_1 — RRM_1	44
HLH	42
Ras	38
zf-C2H2 — zf-C2H2 — zf-C2H2 — zf-C2H2	37
zf-C2H2 — zf-C2H2 — zf-C2H2 — zf-C2H2 — zf-C2H2	36
p450	34
UQ_con	29
DEAD — Helicase_C	28
GST_N — GST_C	27
BTB	26
adh_short	25
zf-C2H2 — zf-C2H2 — zf-C2H2	24
Proteasome	23
7tm_1	22
Metallophos	22
zf-C2H2 — zf-C2H2 — zf-C2H2 — zf-C2H2 — zf-C2H2 — zf-C2H2	21
Kinesin	20
zf-C2H2	20
zf-C2H2 — zf-C2H2 — zf-C2H2 — zf-C2H2 — zf-C2H2 — zf-C2H2	19
zf-C4 — Hormone_recep	19

Architecture	Frequency
COesterase	17
AMP-binding	17
zf-C2H2 — zf-C2H2	16
Tetraspannin	16
efhand — efhand — efhand	16
<i>H. sapiens</i>	
Pkinase	200
7tm_1	141
Ras	84
zf-C3HC4	76
RRM_1	68
Homeobox	66
HLH	57
zf-C4 — Hormone_recep	52
RRM_1 — RRM_1	51
IL8	41
Filament	41
DEAD — Helicase_C	38
SH3_1	37
zf-C2H2 — zf-C2H2 — zf-C2H2	32
UQ_con	31
K_tetra	30
PH	28
PDZ	28
MHC_I — C1-set	27
SH2	26
V-set	26
Trypsin	25
UCH	24
Lectin_C	23
C2 — C2	23
TGF_beta	23
WD40 — WD40 — WD40 — WD40 — WD40 — WD40	22
bZIP_1	22
Kinesin	22
RRM_1 — RRM_1 — RRM_1	22

Appendix D

Table D.1: All iPfam domain pairs that are shared between *E. coli*, *S. cerevisiae* and *H. sapiens*.

Accession A	Name A	Accession B	Name B
PF00004	AAA	PF00004	AAA
PF00005	ABC_tran	PF00005	ABC_tran
PF00009	GTP_EFTU	PF00009	GTP_EFTU
PF00011	HSP20	PF00011	HSP20
PF00013	KH1	PF00013	KH1
PF00027	cNMP_binding	PF00027	cNMP_binding
PF00035	dsrm	PF00035	dsrm
PF00043	GST_C	PF00043	GST_C
PF00044	Gp_dh_N	PF00044	Gp_dh_N
PF00056	Ldh_1_N	PF00056	Ldh_1_N
PF00085	Thioredoxin	PF00085	Thioredoxin
PF00091	Tubulin	PF00091	Tubulin
PF00106	adh_short	PF00106	adh_short
PF00107	ADH_zinc_N	PF00107	ADH_zinc_N
PF00117	GATase	PF00117	GATase
PF00118	Cpn60_TCP1	PF00118	Cpn60_TCP1
PF00132	Hexapep	PF00132	Hexapep
PF00149	Metallophos	PF00149	Metallophos
PF00155	Aminotran_1.2	PF00155	Aminotran_1.2
PF00156	Pribosyltran	PF00156	Pribosyltran
PF00160	Pro_isomerase	PF00160	Pro_isomerase
PF00166	Cpn10	PF00118	Cpn60_TCP1
PF00171	Aldedh	PF00171	Aldedh
PF00180	Iso_dh	PF00180	Iso_dh
PF00183	HSP90	PF00183	HSP90
PF00185	OTCace	PF00185	OTCace
PF00199	Catalase	PF00199	Catalase
PF00202	Aminotran_3	PF00202	Aminotran_3
PF00204	DNA_gyraseB	PF00204	DNA_gyraseB
PF00205	TPP_enzyme_M	PF00205	TPP_enzyme_M
PF00206	Lyase_1	PF00206	Lyase_1

Accession A	Name A	Accession B	Name B
PF00208	ELFV_dehydrog	PF00208	ELFV_dehydrog
PF00224	PK	PF00224	PK
PF00227	Proteasome	PF00227	Proteasome
PF00254	FKBP_C	PF00254	FKBP_C
PF00258	Flavodoxin_1	PF00258	Flavodoxin_1
PF00270	DEAD	PF00270	DEAD
PF00271	Helicase_C	PF00176	SNF2_N
PF00271	Helicase_C	PF00270	DEAD
PF00271	Helicase_C	PF00271	Helicase_C
PF00288	GHMP_kinases_N	PF00288	GHMP_kinases_N
PF00289	CPSase_L_chain	PF00289	CPSase_L_chain
PF00291	PALP	PF00291	PALP
PF00293	NUDIX	PF00293	NUDIX
PF00300	PGAM	PF00300	PGAM
PF00317	Ribonuc_red_lgN	PF00317	Ribonuc_red_lgN
PF00328	Acid_phosphat_A	PF00328	Acid_phosphat_A
PF00334	NDK	PF00334	NDK
PF00365	PFK	PF00365	PFK
PF00378	ECH	PF00378	ECH
PF00383	dCMP_cyt_deam_1	PF00383	dCMP_cyt_deam_1
PF00389	2-Hacid_dh	PF00389	2-Hacid_dh
PF00438	S-AdoMet_synt_N	PF00438	S-AdoMet_synt_N
PF00448	SRP54	PF00448	SRP54
PF00456	Transketolase_N	PF00456	Transketolase_N
PF00462	Glutaredoxin	PF00462	Glutaredoxin
PF00479	G6PD_N	PF00479	G6PD_N
PF00483	NTP_transferase	PF00483	NTP_transferase
PF00488	MutS_V	PF00488	MutS_V
PF00491	Arginase	PF00491	Arginase
PF00515	TPR_1	PF00515	TPR_1
PF00533	BRCT	PF00533	BRCT
PF00534	Glycos_transf_1	PF00534	Glycos_transf_1
PF00542	Ribosomal_L12	PF00542	Ribosomal_L12
PF00570	HRDC	PF00570	HRDC
PF00571	CBS	PF00571	CBS
PF00578	AhpC-TSA	PF00578	AhpC-TSA
PF00583	Acetyltransf_1	PF00583	Acetyltransf_1
PF00586	AIRS	PF00586	AIRS
PF00587	tRNA-synt_2b	PF00587	tRNA-synt_2b
PF00596	Aldolase_II	PF00596	Aldolase_II
PF00625	Guanylate_kin	PF00625	Guanylate_kin
PF00627	UBA	PF00009	GTP_EFTU
PF00627	UBA	PF00627	UBA
PF00636	Ribonuclease_3	PF00035	dsrm
PF00664	ABC_membrane	PF00005	ABC_tran
PF00664	ABC_membrane	PF00664	ABC_membrane
PF00676	E1_dh	PF00676	E1_dh
PF00679	EFG_C	PF00009	GTP_EFTU

Accession A	Name A	Accession B	Name B
PF00702	Hydrolase	PF00702	Hydrolase
PF00731	AIRC	PF00731	AIRC
PF00899	ThiF	PF00899	ThiF
PF00923	Transaldolase	PF00923	Transaldolase
PF00929	Exonuc_X-T	PF00929	Exonuc_X-T
PF01000	RNA_pol_A_bac	PF00562	RNA_pol_Rpb2.6
PF01039	Carboxyl_trans	PF01039	Carboxyl_trans
PF01053	Cys_Met_Meta_PP	PF01053	Cys_Met_Meta_PP
PF01063	Aminotran_4	PF01063	Aminotran_4
PF01138	RNase_PH	PF01138	RNase_PH
PF01182	Glucosamine_iso	PF01182	Glucosamine_iso
PF01192	RNA_pol_Rpb6	PF00623	RNA_pol_Rpb1.2
PF01193	RNA_pol_L	PF00562	RNA_pol_Rpb2.6
PF01193	RNA_pol_L	PF01000	RNA_pol_A_bac
PF01193	RNA_pol_L	PF01193	RNA_pol_L
PF01227	GTP_cyclohydroI	PF01227	GTP_cyclohydroI
PF01230	HIT	PF01230	HIT
PF01259	SAICAR_synt	PF01259	SAICAR_synt
PF01423	LSM	PF01423	LSM
PF01467	CTP_transf_2	PF01467	CTP_transf_2
PF01546	Peptidase_M20	PF01546	Peptidase_M20
PF01612	3.5_exonuc	PF00570	HRDC
PF01624	MutS_I	PF01624	MutS_I
PF01751	Toprim	PF00270	DEAD
PF01842	ACT	PF01842	ACT
PF01926	MMR_HSR1	PF01926	MMR_HSR1
PF01965	DJ-1_PfpI	PF01965	DJ-1_PfpI
PF02142	MGS	PF02142	MGS
PF02463	SMC_N	PF02463	SMC_N
PF02518	HATPase_c	PF00183	HSP90
PF02518	HATPase_c	PF00204	DNA_gyraseB
PF02518	HATPase_c	PF01119	DNA_mis_repair
PF02518	HATPase_c	PF02518	HATPase_c
PF02729	OTCace_N	PF00185	OTCace
PF02729	OTCace_N	PF02729	OTCace_N
PF02769	AIRS_C	PF00586	AIRS
PF02769	AIRS_C	PF02769	AIRS_C
PF02772	S-AdoMet_synt_M	PF00438	S-AdoMet_synt_N
PF02772	S-AdoMet_synt_M	PF02772	S-AdoMet_synt_M
PF02773	S-AdoMet_synt_C	PF00438	S-AdoMet_synt_N
PF02773	S-AdoMet_synt_C	PF02772	S-AdoMet_synt_M
PF02773	S-AdoMet_synt_C	PF02773	S-AdoMet_synt_C
PF02775	TPP_enzyme_C	PF00205	TPP_enzyme_M
PF02775	TPP_enzyme_C	PF02775	TPP_enzyme_C
PF02776	TPP_enzyme_N	PF00205	TPP_enzyme_M
PF02776	TPP_enzyme_N	PF02775	TPP_enzyme_C
PF02776	TPP_enzyme_N	PF02776	TPP_enzyme_N
PF02779	Transket_pyr	PF00456	Transketolase_N

Accession A	Name A	Accession B	Name B
PF02779	Transket_pyr	PF00676	E1_dh
PF02779	Transket_pyr	PF02779	Transket_pyr
PF02780	Transketolase_C	PF00456	Transketolase_N
PF02780	Transketolase_C	PF02779	Transket_pyr
PF02780	Transketolase_C	PF02780	Transketolase_C
PF02781	G6PD_C	PF00479	G6PD_N
PF02781	G6PD_C	PF02781	G6PD_C
PF02786	CPSase_L_D2	PF00117	GATase
PF02786	CPSase_L_D2	PF00289	CPSase_L_chain
PF02786	CPSase_L_D2	PF00988	CPSase_sm_chain
PF02786	CPSase_L_D2	PF02142	MGS
PF02786	CPSase_L_D2	PF02786	CPSase_L_D2
PF02787	CPSase_L_D3	PF00117	GATase
PF02787	CPSase_L_D3	PF00289	CPSase_L_chain
PF02787	CPSase_L_D3	PF00988	CPSase_sm_chain
PF02787	CPSase_L_D3	PF02786	CPSase_L_D2
PF02787	CPSase_L_D3	PF02787	CPSase_L_D3
PF02798	GST_N	PF00043	GST_C
PF02798	GST_N	PF02798	GST_N
PF02800	Gp_dh_C	PF00044	Gp_dh_N
PF02800	Gp_dh_C	PF02800	Gp_dh_C
PF02812	ELFV_dehydrog_N	PF00208	ELFV_dehydrog
PF02812	ELFV_dehydrog_N	PF02812	ELFV_dehydrog_N
PF02826	2-Hacid_dh_C	PF00389	2-Hacid_dh
PF02826	2-Hacid_dh_C	PF02826	2-Hacid_dh_C
PF02852	Pyr_redox_dim	PF02817	E3_binding
PF02852	Pyr_redox_dim	PF02852	Pyr_redox_dim
PF02866	Ldh_1_C	PF00056	Ldh_1_N
PF02866	Ldh_1_C	PF02866	Ldh_1_C
PF02867	Ribonuc_red_lgC	PF00317	Ribonuc_red_lgN
PF02867	Ribonuc_red_lgC	PF02867	Ribonuc_red_lgC
PF02881	SRP54_N	PF00448	SRP54
PF02881	SRP54_N	PF02881	SRP54_N
PF02887	PK_C	PF00224	PK
PF02887	PK_C	PF02887	PK_C
PF02978	SRP_SPB	PF00448	SRP54
PF02978	SRP_SPB	PF02881	SRP54_N
PF02978	SRP_SPB	PF02978	SRP_SPB
PF03129	HGTP_anticodon	PF00587	tRNA-synt_2b
PF03129	HGTP_anticodon	PF03129	HGTP_anticodon
PF03144	GTP_EFTU_D2	PF00009	GTP_EFTU
PF03144	GTP_EFTU_D2	PF03144	GTP_EFTU_D2
PF03372	Exo_endo_phos	PF03372	Exo_endo_phos
PF03477	ATP-cone	PF00317	Ribonuc_red_lgN
PF03477	ATP-cone	PF02867	Ribonuc_red_lgC
PF03725	RNase_PH_C	PF01138	RNase_PH
PF03725	RNase_PH_C	PF03725	RNase_PH_C
PF03764	EFG_IV	PF00009	GTP_EFTU

Accession A	Name A	Accession B	Name B
PF03764	EFG_IV	PF00679	EFG_C
PF03764	EFG_IV	PF03144	GTP_EFTU_D2
PF03807	F420_oxidored	PF03807	F420_oxidored
PF03953	Tubulin_C	PF00091	Tubulin
PF03953	Tubulin_C	PF03953	Tubulin_C
PF04983	RNA_pol_Rpb1.3	PF01192	RNA_pol_Rpb6
PF04997	RNA_pol_Rpb1.1	PF01192	RNA_pol_Rpb6
PF04998	RNA_pol_Rpb1.5	PF01192	RNA_pol_Rpb6
PF05188	MutS_II	PF00488	MutS_V
PF05188	MutS_II	PF01624	MutS_I
PF05190	MutS_IV	PF05190	MutS_IV
PF05192	MutS_III	PF00488	MutS_V
PF05192	MutS_III	PF01624	MutS_I
PF05192	MutS_III	PF05188	MutS_II
PF05192	MutS_III	PF05190	MutS_IV
PF06026	Rib_5-P_isom_A	PF06026	Rib_5-P_isom_A
PF06418	CTP_synth_N	PF00117	GATase
PF06418	CTP_synth_N	PF06418	CTP_synth_N
PF07687	M20_dimer	PF01546	Peptidase_M20
PF07687	M20_dimer	PF07687	M20_dimer
PF07973	tRNA_SAD	PF00587	tRNA-synt.2b
PF07992	Pyr_redox.2	PF02852	Pyr_redox_dim
PF07992	Pyr_redox.2	PF07992	Pyr_redox.2
PF08240	ADH_N	PF00107	ADH_zinc_N
PF08240	ADH_N	PF08240	ADH_N
PF08544	GHMP_kinases_C	PF00288	GHMP_kinases_N
PF08544	GHMP_kinases_C	PF08544	GHMP_kinases_C

Appendix E

Table E.1: Most frequent Gene Ontology annotations on all *i*Pfam families shared between *E. coli*, *S. cerevisiae* and *H. sapiens*.

Accession	Function	Freq	Process	Freq	Compartment	Freq
PF00291	catalytic activity	9	metabolic process	13		
PF00702	catalytic activity	9	metabolic process	13		
PF01063	catalytic activity	9	metabolic process	13		
PF00171	oxidoreductase activity	9	metabolic process	13		
PF00378	catalytic activity	9	metabolic process	13		
PF00106	oxidoreductase activity	9	metabolic process	13		
PF00180	oxidoreductase activity, acting on the CH-OH group of donors, NAD or NADP as acceptor	3	metabolic process	13		
PF00389	oxidoreductase activity, acting on the CH-OH group of donors, NAD or NADP as acceptor	3	metabolic process	13		
PF00289	ligase activity	2	metabolic process	13		
PF02817	acyltransferase activity	2	metabolic process	13		
PF01842	amino acid binding	2	metabolic process	13		
PF00676	oxidoreductase activity, acting on the aldehyde or oxo group of donors, disulfide as acceptor	1	metabolic process	13		
PF00583	N-acetyltransferase activity	1	metabolic process	13		

Accession	Function	Freq	Process	Freq	Compartment	Freq
PF00623	DNA-directed RNA polymerase activity	8	transcription	7	nucleus	1
PF04998	DNA-directed RNA polymerase activity	8	transcription	7		
PF01193	DNA-directed RNA polymerase activity	8	transcription	7		
PF01000	DNA-directed RNA polymerase activity	8	transcription	7		
PF04997	DNA-directed RNA polymerase activity	8	transcription	7		
PF04983	DNA-directed RNA polymerase activity	8	transcription	7		
PF00562	DNA-directed RNA polymerase activity	8	transcription	7		
PF01624	ATP binding	26	mismatch repair	6		
PF05190	ATP binding	26	mismatch repair	6		
PF00488	ATP binding	26	mismatch repair	6		
PF05188	ATP binding	26	mismatch repair	6		
PF01119	ATP binding	26	mismatch repair	6		
PF05192	ATP binding	26	mismatch repair	6		
PF00208	oxidoreductase ac- tivity	9	amino acid metabolic pro- cess	5		
PF02812	oxidoreductase ac- tivity	9	amino acid metabolic pro- cess	5		
PF01053	pyridoxal phos- phate binding	4	amino acid metabolic pro- cess	5		
PF02887	magnesium ion binding	3	glycolysis	5		
PF00044	NAD binding	3	glycolysis	5		
PF02800	NAD binding	3	glycolysis	5		
PF00224	magnesium ion binding	3	glycolysis	5		
PF00185	carboxyl- and car- bamoyltransferase activity	2	amino acid metabolic pro- cess	5		
PF02729	carboxyl- and car- bamoyltransferase activity	2	amino acid metabolic pro- cess	5		

Accession	Function	Freq	Process	Freq	Compartment	Freq
PF00365	6-phosphofructokinase activity	1	glycolysis	5	6-phosphofructokinase complex	1
PF00166	ATP binding	26	protein folding	4		
PF00155	pyridoxal phosphate binding	4	biosynthetic process	4		
PF01467	nucleotidyltransferase activity	2	biosynthetic process	4		
PF00483	nucleotidyltransferase activity	2	biosynthetic process	4		
PF00183	unfolded protein binding	1	protein folding	4		
PF00534			biosynthetic process	4		
PF00254			protein folding	4		
PF00160			protein folding	4		
PF00438	ATP binding	26	one-carbon compound metabolic process	3		
PF02772	ATP binding	26	one-carbon compound metabolic process	3		
PF02773	ATP binding	26	one-carbon compound metabolic process	3		
PF03725	RNA binding	7	RNA processing	3		
PF00636	RNA binding	7	RNA processing	3		
PF01138	RNA binding	7	RNA processing	3		
PF03129	ATP binding	26	translation	2		
PF02852	oxidoreductase activity	9	cell redox homeostasis	2	cytoplasm	3
PF00056	oxidoreductase activity	9	tricarboxylic acid cycle intermediate metabolic process	2		
PF02866	oxidoreductase activity	9	tricarboxylic acid cycle intermediate metabolic process	2		
PF02881	GTP binding	8	SRP-dependent cotranslational protein targeting to membrane	2	signal recognition particle, endoplasmic reticulum targeting	2
PF00448	GTP binding	8	SRP-dependent cotranslational protein targeting to membrane	2	membrane	1

Accession	Function	Freq	Process	Freq	Compartment	Freq
PF02781	glucose-6-phosphate dehydrogenase activity	2	glucose metabolic process	2		
PF00479	glucose-6-phosphate dehydrogenase activity	2	glucose metabolic process	2		
PF00542	structural constituent of ribosome	1	translation	2	intracellular	6
PF00199	catalase activity	1	electron transport	2		
PF00462	protein disulfide oxidoreductase activity	1	cell redox homeostasis	2		
PF03807			electron transport	2		
PF01182			carbohydrate metabolic process	2		
PF00923			carbohydrate metabolic process	2		
PF02463	ATP binding	26	DNA metabolic process	1	chromosome	2
PF00204	ATP binding	26	DNA topological change	1	chromosome	2
PF00664	ATP binding	26	transport	1	integral to membrane	1
PF00334	ATP binding	26	UTP biosynthetic process	1		
PF00118	ATP binding	26	cellular protein metabolic process	1		
PF00587	ATP binding	26	tRNA aminoacylation for protein translation	1		
PF00288	ATP binding	26	phosphorylation	1		
PF00988	ATP binding	26	nitrogen compound metabolic process	1		
PF03953	GTP binding	8	protein polymerization	1	protein complex	1
PF01192	DNA-directed RNA polymerase activity	8	transcription, DNA-dependent	1		
PF02978	RNA binding	7	protein targeting	1	signal recognition particle, endoplasmic reticulum targeting	2
PF01751	nucleic acid binding	5	DNA modification	1		

Accession	Function	Freq	Process	Freq	Compartment	Freq
PF02787	carbamoyl-phosphate synthase activity	2	arginine biosynthetic process	1	cytoplasm	3
PF01227	GTP cyclohydrolase I activity	1	aromatic compound biosynthetic process	1	cytoplasm	3
PF00731	phosphoribosylaminoimidazole carboxylase activity	1	de novo' IMP biosynthetic process	1	phosphoribosylaminoimidazole carboxylase complex	1
PF02867	ribonucleoside-diphosphate reductase activity	1	DNA replication	1	ribonucleoside-diphosphate reductase complex	1
PF00227	threonine endopeptidase activity	1	ubiquitin-dependent protein catabolic process	1	proteasome core complex (sensu Eukaryota)	1
PF01259	phosphoribosylaminoimidazolesuccinocarboxamide synthase activity	1	purine nucleotide biosynthetic process	1		
PF01546	metallopeptidase activity	1	proteolysis	1		
PF06026	ribose-5-phosphate isomerase activity	1	pentose-phosphate shunt, non-oxidative branch	1		
PF06418	CTP synthase activity	1	pyrimidine nucleotide biosynthetic process	1		
PF01423			mRNA metabolic process	1	ribonucleoprotein complex	1
PF00156			nucleoside metabolic process	1		
PF00004	ATP binding	26				
PF02786	ATP binding	26				
PF02518	ATP binding	26				
PF00270	ATP binding	26				
PF00271	ATP binding	26				
PF00176	ATP binding	26				
PF00005	ATP binding	26				
PF02775	catalytic activity	9				
PF00586	catalytic activity	9				
PF00206	catalytic activity	9				
PF00258	oxidoreductase activity	9				
PF00899	catalytic activity	9				
PF00117	catalytic activity	9				

Accession	Function	Freq	Process	Freq	Compartment	Freq
PF00578	oxidoreductase activity	9				
PF01926	GTP binding	8			intracellular	6
PF00679	GTP binding	8				
PF00009	GTP binding	8				
PF03144	GTP binding	8				
PF03764	GTP binding	8				
PF00013	RNA binding	7				
PF00570	nucleic acid binding	5			intracellular	6
PF01612	nucleic acid binding	5			intracellular	6
PF00383	hydrolase activity	4				
PF00202	pyridoxal phosphate binding	4				
PF00149	hydrolase activity	4				
PF07687	hydrolase activity	4				
PF00293	hydrolase activity	4				
PF02776	thiamin pyrophosphate binding	3				
PF02826	oxidoreductase activity, acting on the CH-OH group of donors, NAD or NADP as acceptor	3				
PF00205	magnesium ion binding	3				
PF00596	metal ion binding	2				
PF01039	ligase activity	2				
PF00132	acyltransferase activity	2				
PF00491	metal ion binding	2				
PF00035	double-stranded RNA binding	1			intracellular	6
PF00328	acid phosphatase activity	1				
PF00533					intracellular	6

Appendix F

Table F.1: List of disease mutations linked to protein interaction defects, derived from the scientific literature.

MutationVariant	Seq ID	Description	Inh.	Mech.
604312 .0001	P01034	In patients with Icelandic-type cerebroarterial amyloidosis (105150), Abrahamson et al. (1987) identified a 358T-A transversion in the CST3 gene, resulting in a leu68-to-gln (L68Q) substitution. The dimerization was highly temperature-dependent, with a rise in incubation temperature from 37 to 40 degrees centigrade resulting in a 150% increase in dimerization rate.		GF
107300 .0021	P01008	Antithrombin III deficiency	AD	LF
121011 .0020	P29033	gap-junction protein (no direct functional link)	AD	LF
123580 .0001	P02489	Crystallin change of preference in polymers	AD	CF
123590 .0001	P02511	Crystallin change of preference in polymers	AD	CF
123680 .0001	Q53R50	Crystallin change of preference in polymers in Cop-pock cataract	AD	LF
125647 .0002	Q4LE79	Desmoplakin; This region of the desmoplakin protein interacts with intermediate filaments to anchor them to the desmosome	AR	LF
134850 .0010	P02679	Fibrinogen G, impaired polymerisation	AR	LF
134850 .0017	P02679	Fibrinogen G, impaired polymerisation	AR	LF
138040 .0009	P04150	GLUCOCORTICOID receptor, reduced cofactor binding	AD	LF
139250 .0020	P01241	Growth Hormone, in a prepubertal Spanish child with familial short stature (604271), Lewis et al. (2004) found an ile179-to-met (I179M) amino acid substitution. Molecular modeling studies suggested that the I179M substitution might perturb interactions between GH and the GH receptor loop containing residue trp169, thereby affecting signal transduction.	AD	LF
139320 .0032	Q5JWD2	GNAS	IM	LF

MutationVariant	Seq ID	Description	Inh.	Mech.
139350 .0004	P04264	Keratin	AD	LF
139350 .0015	P04264	Keratin	AD	LF
141800 .0179	Q5R9M5	HBA1; Hb Yuda has a very low oxygen affinity and slightly decreased cooperative subunit interaction.		LF
141900 .0038	P68871	HBB; HEMOGLOBIN C [HBB, GLU6LYS]	IM	GF
147545 .0002	P35568	IRS1	AR	LF
147557 .0014	P16144	Koster et al. (2001) reported that this mutation renders integrin beta-4 unable to interact with plectin (601282) and prevents the localization of plectin in hemidesmosomes.	AR	LF
147557 .0015	P16144	Koster et al. (2001) reported that this mutation renders integrin beta-4 unable to interact with plectin (601282) and prevents the localization of plectin in hemidesmosomes.	AR	LF
600576 .0001	P43694	Garg et al. (2003) demonstrated that GATA4 (600576) interacts with TBX5 and showed that a missense mutation in GATA4, G296S (600576.0001), abrogated this interaction.	AD	LF
235200 .0011	NP_620575	By performing immunoprecipitation studies in HeLa cells, Ka et al. (2005) found that the Q283P mutation prevented the normal interaction between HFE protein and beta-2-microglobulin (B2M; 109700) and between HFE protein and transferrin receptor (TFRC; 190010).	CH	LF
300300 .0025	Q32ML5	de Weers et al. (1994) identified a C-to-T transition at position 993, resulting in a substitution of tryptophan for arginine-288. This mutation was found in the SH2-like domain where arg288 is highly conserved and crucial for the interaction with the aromatic ring of phosphotyrosine. Therefore, the replacement of arg288 by a nonpolar tryptophan would entirely abrogate the formation of the high-affinity comosine binding pocket. The change to a neutral glycine residue is highly likely to disrupt the binding potential of this region. This patient has less than 1% B cells and undetectable immunoglobulin levels, indicating that the replacement of this highly conserved arginine residue completely abolishes the functioning of Btk.	XL	LF
300490 .0013	O60880	SH2 Domain Protein 1A; Based on the molecular structure of the SH2D1A-SLAM (603492) interaction, this mutation was predicted to disrupt binding between the SH2 domain of SH2D1A and the cytoplasmic domain of SLAM. The mutation was also predicted to interfere with SH2D1A-2B4 (605554) binding because of the strong amino acid homology shared by SLAM and 2B4.	XL	LF

MutationVariant	Seq ID	Description	Inh.	Mech.
305371 .0002	P15976	Freson et al. (2001) described a family with isolated X-linked macrothrombocytopenia without anemia but with some dyserythropoietic features (see 300367) in 13 males in 9 sibships of 3 generations connected through carrier females. A novel mutation in the GATA1 gene, asp218 to gly (D218G), resulted in a weaker interaction with FOG1	XL	LF
305371 .0005	P15976	Freson et al. (2002) described a 2-generation family with deep macrothrombocytopenia (see 300367), marked anemia, and early mortality. The mutation is predicted to result in substitution of tyrosine for aspartate-218 (D218Y). Zinc finger interaction studies revealed a stronger loss of affinity of D218Y-GATA1 than of D218G-GATA1 (305371.0002) for the essential transcription factor FOG1 (601950) and a disturbed GATA1 self-association.	XL	LF
600160 .0016	P42771	CDK inhib 2a; A val59-to-gly mutation in the CDKN2A gene was found in 4 families segregating cutaneous malignant melanoma; The mutation, which occurs in a hydrophobic region with the second ankyrin repeat, impairs p16-INK4a function, as shown by studies of protein-protein interactions and cell proliferation assays.	AD	LF
601130 .0002	P11712	Cytochrome P450; the CYP2C9*3 variant is less than 5% as efficient as the wildtype enzyme, while CYP2C9*2 shows about 12% of wildtype activity, apparently as a result of the amino acid substitution altering the interaction of the enzyme with cytochrome P450 oxidoreductase. Aithal et al. (1999) studied the frequency of the 2 variant alleles in individuals with a low warfarin dose requirement; see 122700. Patients in the low-dose group were more likely to have difficulties at the time of induction of warfarin therapy and had an increased risk of major bleeding complications.	PM	LF
601769 .0010	P11473	Whitfield et al. (1996) identified a mutation in the VDR gene, resulting in an ile314-to-ser (I314S) substitution in the hormone-binding domain of the protein. The mutation caused decreased 1,25-(OH)2D3-dependent transactivation of the VDR and impaired heterodimeric interaction with the retinoid X receptor	AR	LF

MutationVariant	Seq ID	Description	Inh.	Mech.
601769 .0011	P11473	In a patient with vitamin D-dependent rickets type II (277440), Whitfield et al. (1996) identified a mutation in the VDR gene, resulting in an arg391-to-cys (R391C) substitution in the hormone-binding domain of the protein. The mutation caused decreased 1,25-(OH)2D3-dependent transactivation of the VDR and impaired heterodimeric interaction with the retinoid X receptor (RXR; 180245).	AR	LF
603273 .0009	Q9H3D4	In a 6-year-old patient with Hay-Wells syndrome (106260) who lacked any limb defects, McGrath et al. (2001) identified an A-to-T transversion at nucleotide 1542 of the TP63 gene, resulting in a leu518-to-phe substitution in the sterile alpha motif (SAM) domain. Molecular modeling suggested that the substitution would alter protein-protein interactions.	AD	LF
603273 .0010	Q9H3D4	In a 10-month-old infant with typical features of Hay-Wells syndrome (106260), McGrath et al. (2001) identified a T-to-G transversion at nucleotide 1564 of the TP63 gene, resulting in a cys526-to-gly substitution in the sterile alpha motif (SAM) domain. Molecular modeling suggested that the substitution would alter protein-protein interactions.	AD	LF
603714 .0002	O95343	Laflamme et al. (2004) demonstrated that the SIX3 protein carrying this mutation did not interact with NOR1 (600542) in vivo.	AD	LF
606860 .0002	P05155	Complement Component 1 Inhib; Davis et al. (1992) showed that the dysfunction demonstrated by this mutation results from a block in the interaction with target protease.		LF
608014 .0001	Q9UJY1	HS 22kd Prot, Increased binding!	AD	GF
608014 .0002	Q9UJY1	HS 22kd Prot, Increased binding!	AD	GF
608537 .0019	P40337	Ang et al. (2002) concluded that the R200W substitution impairs the interaction of VHL with HIF1-alpha	AR	LF
103850 .0002	P04075	Mutation of Glu to Arg in subunit interface. However, this is not proved to disrupt protein-protein interaction but it seems likely and the authors argue this is the case	AR	LF
256540 .0014	P10619	A structural model of the mutant PPCA was constructed by amino acid substitution of 453glutamic acid for lysine in the crystal structure of the wild type PPCA precursor reported. The results show that the K453E mutation is located at the dimer interface of the PPCA and reduces the hydrogen bond formation in the dimer. This structural change may cause instability of the PPCA dimer.		

MutationVariant	Seq ID	Description	Inh.	Mech.
305900	.0051	NP_000393 In a study of the causative mutation in 12 cases of G6PD deficiency associated with chronic nonspherocytic hemolytic anemia, Vulliamy et al. (1998) found 1 patient to have a novel mutation, which they called G6PD Serres: a 1082C-T change, causing an ala361-to-val substitution in the dimer interface where most other severe G6PD mutations are found. <i>Blood</i> . 2000 Feb 15;95(4):1499-501.	XL	LF
193400	.0013	NP_000543 <i>J Biol Chem</i> . 1991 Jul 25;266(21):13499-502. In previous studies, we have mapped the epitope for an anti-vWF monoclonal antibody which inhibits the interaction between FVIII and vWF to a region spanning Thr78 to Thr96 of the mature protein. We now report the identification of a mutation within this region of vWF that results in decreased FVIII binding.	CH	LF
606869	.0009	P06865 Paw et al. (1990) identified a G-to-A transition at nucleotide 1511 resulting in substitution of histidine for arginine at position 504 in the HEXA molecule. Cultured fibroblasts from the patient synthesized an alpha subunit that could acquire mannose 6-phosphate and be secreted, but which failed to associate with the beta-subunit to form the enzymatically active heterodimer.	AR	LF
193400	.0024	NP_000543 Schneppenheim et al. (1996) demonstrated a heterozygous cys2010-to-arg mutation in the mature vWF subunit causing the type IID von Willebrand disease phenotype in 2 unrelated patients. Recombinant expression of mutant vWF fragments demonstrated that the mutation was responsible for defective disulfide bonding of the C-terminal domains, thus impairing dimer formation.	AD	LF
141850	.0005	P69905 Goossens et al. (1982) described another nondeletion mechanism: mutation in the 125th codon of the alpha-2 gene resulted in substitution of proline for leucine in a region of the H helix of the alpha-globin chain, which is critical for alpha-beta contact, resulting in impediment to alpha-beta dimer formation, the initial step in hemoglobin tetramer assembly.	AR	LF

MutationVariant	Seq ID	Description	Inh.	Mech.
125660 .0006	Q53SB5	The leu345-to-pro mutation (L345P) in this kindred was located in an evolutionarily highly conserved position of the desmin coiled-coil rod domain important for dimer formation. L345P desmin was incapable of forming filamentous networks in transfected HeLa and SW13 cells. Sjoberg et al. (1999) concluded that the L345P mutation causes myopathy by interfering in a dominant-negative manner with the dimerization-polymerization process of intermediate filament assembly.	AD	LF
190160 NA	P10828	Proc Natl Acad Sci U S A. 1997 Jan 7;94(1):248-53 Here we describe a novel leucine to valine mutation in codon 454 (L454V) of the thyroid hormone beta receptor. [...] indicating that the interaction of this residue with accessory proteins is critical for transcriptional activation. (Not in OMIM nor UniProt)	AD	LF
235200 .0001	NP_620575J	Biol Chem. 1997 May 30;272(22):14025-8: Co-immunoprecipitation studies demonstrate that wild-type HLA-H binds beta2-microglobulin and that the C282Y mutation completely abrogates this interaction.	AR	LF
607008 .0001	P11310	MCAD DEFICIENCY; the amount of K329E tetramer formed was distinctly less than wildtype at any point up to 60 minutes after import, indicating that the assembly of K329E was defective. After further incubation, K304E decayed more rapidly than did wildtype, indicating a reduced stability. In similar studies K329R behaved like the wildtype, while K329D closely resembled K329E, indicating that a basic residue at 304 is essential for tetramer formation and intramitochondrial stability of mature MCAD.	AR	LF
176300 .0039	P02766	Jenne et al. (1996) identified a 'new' amyloidogenic val20ile mutation of the TTR gene. tetramer stability was significantly reduced in agreement with the expected change in the interactions between 2 opposing dimers via the side chain of ile20.		CF
174763 .0002	P54098	J Biol Chem. 2005 Sep 9;280(36):31341-6: the A467T mutant enzyme failed to interact with and was not stimulated by the accessory subunit.	AR	LF
157140 .0003	P10636	J Neurochem. 2000 Jun;74(6):2583-9: Mutated tau is less phosphorylated than its normal counterpart at serines 396 and 404. Furthermore, the phosphorylated mutant protein is unable to bind to microtubules.	AD	LF

MutationVariant	Seq ID	Description	Inh.	Mech.	
191044	.0001	P19429	Biochemistry. 2002 Jun 11;41(23):7267-74: the affinity is reduced by approximately 14-fold by the T142 phosphorylation and approximately 4-fold by the mutation R145G.	AD	LF
P51587	VAR_020705P51587	Oncogene. 2003 Jan 9;22(1):28-33: the cancer-predisposing mutation Y42C in BRCA2 significantly compromised the interaction between RPA and BRCA2		LF	
276000	.0006	Q5NV57	Hum Mutat. 2004 Jan;23(1):22-31: E79K markedly inhibited autoactivation of cationic trypsinogen. Remarkably, however, E79K trypsin activated anionic trypsinogen PRSS2 (601564) 2-fold.	AD	CF
P00156	VAR_013653P00156	European Journal of Biochemistry, Volume 271, Issue 7, April 2004, Pages 1292-1298: The mitochondrial cytochrome b missense mutation, G167E, has been reported in a patient with cardiomyopathy. The residue G167 is located in an extramembranous helix close to the hinge region of the iron-sulfur protein. Analysis of the enzyme activity indicated that the mutation affected its stability, which could be the result of an altered binding of the iron-sulfur protein on the complex. [...]This suggested that the mutation G167E could hinder the movement of the iron-sulfur protein, probably by distorting the structure of the hinge region.		LF	
238331	NA	P09622	Biochem Biophys Res Commun. 1999 Aug 19;262(1):163-6: Asp for Val at position 479 of the precursor form - the mutation resides within the interface domain and likely perturbs stable dimerization		LF
P04275	VAR_005800P04275	PubMed=1409710: von Willebrand disease type B: a missense mutation selectively abolishes ristocetin-induced von Willebrand factor binding to platelet glycoprotein Ib; J Thromb Haemost. 2006 Feb;4(2):417-25: The interaction of von Willebrand factor-A1 domain with collagen: mutation G1324S (type 2M von Willebrand disease) impairs the conformational change in A1 domain induced by collagen	CH	LF	
606672	.0003	P07359	J Thromb Haemost. 2003 Oct;1(10):2198-205: The 125I-labeled VWF binding to mutant compared with the wild type displayed three patterns, gain-of-function (G233S, G233V, and M239V), equivalent function (G233A), and loss-of-function (G233K and G233D)	AD	GF

MutationVariant	Seq ID	Description	Inh.	Mech.
193400	.0018	NP_000543 Proc Natl Acad Sci U S A. 1992 Oct 15;89(20):9846-9: the type B variant VWF displayed an aberrant interaction with the gpIb platelet receptor that seemed to be independent of multimeric structure.	AD	LF
193400	.0008	NP_000543 J Biol Chem. 1992 Oct 15;267(29):21187-92: the Arg578→Gln mutation increases the affinity of vWF for GPIb but does not directly impair vWF interaction with collagen or heparin. Arg578 may therefore be necessary to prevent normal vWF from interacting with GPIb.	AD	GF
193400	.00012	NP_000543 Blood. 1992 Feb 1;79(3):563-7: These results illustrate the importance of Arg 53 of the mature vWF subunit for the binding of FVIII to vWF	AR	LF
O75695	VAR_008499	O75695 Structure 2006 Feb;14:367-378: The abilities of RP2 to bind Arl3 and cause retinitis pigmentosa seem to be correlated, since both the R118H and E138G mutants show drastically reduced affinity to Arl3		LF
O75695	VAR_018074	O75695 Structure 2006 Feb;14:367-378: The abilities of RP2 to bind Arl3 and cause retinitis pigmentosa seem to be correlated, since both the R118H and E138G mutants show drastically reduced affinity to Arl3		LF
603693	.0001	Q8WW38 Although the mutant protein retained the ability to bind the partner protein GATA4 (600576) and repress GATA4-mediated gene activation, it was subtly impaired in this function.		LF
607759	NA	P08514 J Thromb Haemost. 2004 Jul;2(7):1167-75: A novel Phe171Cys mutation in the alpha(IIb) gene of patients with GT is associated with abrogation of alpha(IIb)beta(3) complex formation	AR	LF
300384	.0008	P50402 Hum Genet. 1999 Mar;104(3):262-8: Biochemical analysis has demonstrated that the mobility and expression levels of the mutant forms of emerin are indistinguishable from that of wild-type emerin, but that they have weakened interactions with nuclear lamina components		LF
300384	.0009	P50402 Hum Genet. 1999 Mar;104(3):262-8: Biochemical analysis has demonstrated that the mobility and expression levels of the mutant forms of emerin are indistinguishable from that of wild-type emerin, but that they have weakened interactions with nuclear lamina components		LF
605906	.0009	O75112 J Biol Chem. 2004 Feb 20;279(8):6746-52: o reveal the biochemical changes due to the mutation, we performed a yeast two-hybrid assay and a pull-down assay. It was demonstrated by both assays that the D626N mutation of Cypher/ZASP increased the affinity of the LIM domain for protein kinase C	AD	GF

MutationVariant	Seq ID	Description	Inh.	Mech.	
603959	.0015	Q9Y5I7	Am J Hum Genet. 2003 Dec;73(6):1293-301: The T233R mutation was found to abolish binding of CLDN16 to ZO1	AR	LF
193400	NA	NP_000543	J Thromb Haemost. 2004 Jul;2(7):1135-42: The mutation L1503Q does not significantly disrupt the conformation of the protein; thus the subtle loss of multimers in this patient may be due to altered interactions with the ADAMTS13 protease.		LF
P04275	VAR_010242	P04275	Blood. 2000 Jul 15;96(2):560-8: Multimer analysis showed that rVWF R273W failed to form high-molecular-weight multimers present in wild-type rVWF	AR	LF
P98172	VAR_023135	P98172	Twigg S.R.F., Kan R., Babbs C., Bochukova E.G., Robertson S.P., Wall S.A., Morriss-Kay G.M., Wilkie A.O.M. "Mutations of ephrin-B1 (EFNB1), a marker of tissue boundary formation, cause craniofrontonasal syndrome." Proc. Natl. Acad. Sci. U.S.A. 101:8652-8657(2004)	AR	LF
P98172	VAR_023130	P98172	Twigg S.R.F., Kan R., Babbs C., Bochukova E.G., Robertson S.P., Wall S.A., Morriss-Kay G.M., Wilkie A.O.M. "Mutations of ephrin-B1 (EFNB1), a marker of tissue boundary formation, cause craniofrontonasal syndrome." Proc. Natl. Acad. Sci. U.S.A. 101:8652-8657(2004)	AR	LF
P98172	VAR_023131	P98172	Am J Hum Genet. June 2004; 74(6): 12091215	AR	LF
P98172	VAR_023128	P98172	Am J Hum Genet. June 2004; 74(6): 12091215	AR	LF
Q99574	VAR_008520	Q99574	Nature. 1999 Sep 23;401(6751):376-9; polymerization disease: Familial dementia caused by polymerization of mutant neuroserpin.	AD	GF
P98172	VAR_023132	P98172	Am J Hum Genet. June 2004; 74(6): 12091215	AR	LF
P98172	VAR_023133	P98172	Am J Hum Genet. June 2004; 74(6): 12091215	AR	LF
P98172	VAR_023134	P98172	Am J Hum Genet. June 2004; 74(6): 12091215	AR	LF
610550	.0007	Q00266	J. Biol. Chem., Vol. 276, Issue 17, 13803-13809, April 27, 2001; Chamberlin et al. (1997) identified a heterozygous 791G-A transition in the MAT1A gene, resulting in an arg264-to-his (R264H) substitution. In vitro studies suggested that residue 264 is involved in salt bridge formation essential for subunit dimerization and that the dominant effect of the R264H mutation is exerted by the formation of enzymatically inactive R264/R264H dimers.	AD	LF
610550	.0009	Q00266	Unlike the R264H (610550.0007) mutation, which behaves as an autosomal dominant, the authors found that the R264C mutation behaves as an autosomal recessive.	AR	LF

MutationVariant	Seq ID	Description	Inh.	Mech.
134850 .0004	P02679	Ebert and Bell (1988) identified Baltimore-3 as a congenital abnormal fibrinogen with defective fibrin monomer polymerization. Bantia et al. (1990) demonstrated an asn308-to-ile mutation. Polymerization is also affected by asn308-to-lys (Kyoto-1).	AD	LF
134850 .0005	P02679	Ebert and Bell (1988) identified Baltimore-3 as a congenital abnormal fibrinogen with defective fibrin monomer polymerization. Bantia et al. (1990) demonstrated an asn308-to-ile mutation. Polymerization is also affected by asn308-to-lys (Kyoto-1).	AD	LF
P05166 VAR_000280P05166		Molecular Genetics and Metabolism, Volume 74, Number 4, December 2001, pp. 476-483(8): To clarify the molecular effect associated with gene alterations causing propionic acidemia, 12 different mutations affecting the PCCB gene were analyzed for their involvement in alpha-beta heteromeric and beta-beta homomeric assembly.	AR	LF
P05166 VAR_000281P05166		Molecular Genetics and Metabolism, Volume 74, Number 4, December 2001, pp. 476-483(8): To clarify the molecular effect associated with gene alterations causing propionic acidemia, 12 different mutations affecting the PCCB gene were analyzed for their involvement in alpha-beta heteromeric and beta-beta homomeric assembly.	AR	LF
P05166 VAR_009086P05166		Molecular Genetics and Metabolism, Volume 74, Number 4, December 2001, pp. 476-483(8): To clarify the molecular effect associated with gene alterations causing propionic acidemia, 12 different mutations affecting the PCCB gene were analyzed for their involvement in alpha-beta heteromeric and beta-beta homomeric assembly.	AR	LF
600160 .0007	P42771	Oncogene. 1999 Sep 23;18(39):5423-34; Harland et al. (1997) identified a met53-to-ile mutation in the CDKN2A gene in affected members of a family with melanoma. They showed that the protein expressed from this previously described mutation did not bind to CDK4/CDK6, confirming its role as a causal mutation in melanoma.	AD	LF
P42771 VAR_001409P42771		Oncogene. 1999 Sep 23;18(39):5423-34		LF
P42771 VAR_001410P42771		Oncogene. 1999 Sep 23;18(39):5423-34		LF
P42771 VAR_001411P42771		Oncogene. 1999 Sep 23;18(39):5423-34		LF
P42771 VAR_001420P42771		Oncogene. 1999 Sep 23;18(39):5423-34		LF
P42771 VAR_001424P42771		Oncogene. 1999 Sep 23;18(39):5423-34		LF
P42771 VAR_001449P42771		Oncogene. 1999 Sep 23;18(39):5423-34		LF
P42771 VAR_001447P42771		Oncogene. 1999 Sep 23;18(39):5423-34		LF
P42771 VAR_001448P42771		Oncogene. 1999 Sep 23;18(39):5423-34		LF

MutationVariant	Seq ID	Description	Inh.	Mech.
603868 .0010	P51159	In a study of the spectrum of mutations in children with primary emophagocytic lymphohistiocytosis (267700), Zur Stadt et al. 2006) identified 2 mutations in RAB27A in 3 patients with Griscelli syndrome type 2 (607624), which can present with hemophagocytic lymphohistiocytosis. One of these was the missense mutation ala87 to pro (A87P). In functional studies using a mammalian 2-hybrid system, they found that the A87P mutation in RAB27A and leu403 to pro in UNC13D (608897.0007) each prevented the formation of a stable UNC13D/RAB27A complex in vitro.	AR	LF
608897 0007	Q70J99	In a study of the spectrum of mutations in children with primary emophagocytic lymphohistiocytosis (267700), Zur Stadt et al. 2006) identified 2 mutations in RAB27A in 3 patients with Griscelli syndrome type 2 (607624), which can present with hemophagocytic lymphohistiocytosis. One of these was the missense mutation ala87 to pro (A87P). In functional studies using a mammalian 2-hybrid system, they found that the A87P mutation in RAB27A and leu403 to pro in UNC13D (608897.0007) each prevented the formation of a stable UNC13D/RAB27A complex in vitro.	AR	LF
Q99574 VAR_008521Q99574		Nature. 1999 Sep 23;401(6751):376-9; polymerization disease: Familial dementia caused by polymerization of mutant neuroserpin.	AD	GF
100710 .0003	P11230	Quiram et al. (1999) demonstrated that the mutation impairs AChR assembly by disrupting a specific interaction between the beta and delta (100720) subunits.	CH	LF
O15273 VAR_029446		J Am Coll Cardiol. 2004 Dec 7;44(11):2192-201: Two TCAP mutations, T137I and R153H, were found in patients with HCM, and another TCAP mutation, E132Q, was identified in a patient with DCM. It was demonstrated by the qualitative assays that the HCM-associated mutations augment the ability of Tcap to interact with titin and calsarcin-1, whereas the DCM-associated mutations impair the interaction of Tcap with MLP, titin, and calsarcin-1		
104760 .0001	P05067	In Levy et al. (1990) identified a 1852G-C transversion in the APP gene, resulting in a glu693-to-gln (E693Q) substitution. Miravalle et al. (2000) demonstrated in vitro that the E693Q mutation resulted in a high content of beta-sheet amyloid conformation and fast aggregation/fibrillization properties.	AD	GF

MutationVariant	Seq ID	Description	Inh.	Mech.
104760 .0013	P05067	In a patient with early-onset familial Alzheimer disease (104300), Kamino et al. (1992) identified an A-to-G transition in the APP gene, resulting in a glu693-to-gly (E693G) substitution. <i>n vitro</i> , the Arctic mutant form of A-beta forms protofibrils and fibrils at higher rates and in larger quantities than wildtype A-beta. In transgenic mice that expressed the Arctic mutant in neurons, Cheng et al. (2004) found that amyloid plaques formed faster and were more extensive compared to control mice. Cheng et al. (2004) concluded that the Arctic mutation is highly amyloidogenic <i>in vivo</i> .	AD	GF
176640 .0001	Q53YK7	The PRNP gene has an unstable region of 5 variant tandem octapeptide coding repeats between codons 51 and 91. Extension of this repeat causes rapid formation of amyloid plaques and neurodegeneration	AD	GF
141900 .0243	P68871	HEMOGLOBIN S [HBB, GLU6VAL] The classic sickle cell anaemia	IM	GF
P00439 VAR_000900P00439		Molecular Genetics and Metabolism 73, 230-238 (2001) : The R157N mutation, associated here with the most marked decrease in two-hybrid interaction, also showed in all other expression systems the most severe effects, including rapid and very extensive aggregation.	AD	GF
P00441 VAR_007131P00441		Proc Natl Acad Sci U S A. 2004 April 20; 101(16): 59765981: The crystal structures of the A4V and I113T mutants of SOD1 reveal a significant reorientation of the two subunits at the monomer-monomer interface. This destabilization of the dimeric interface may result in an increased tendency to unfold or lose metals <i>in vivo</i> .	AD	LF
P00441 VAR_007155P00441		Proc Natl Acad Sci U S A. 2004 April 20; 101(16): 59765981: The crystal structures of the A4V and I113T mutants of SOD1 reveal a significant reorientation of the two subunits at the monomer-monomer interface. This destabilization of the dimeric interface may result in an increased tendency to unfold or lose metals <i>in vivo</i> .	AD	LF
602533 .0002	Q99497	Mutations in DJ-1, a human gene with homologues in organisms from all kingdoms of life, have been shown to be associated with autosomal recessive, early onset Parkinson's disease. The structure suggests that the loss of function caused by the Parkinson's-associated mutation L166P in DJ-1 is due to destabilization of the dimer interface.	AR	LF
P00492 VAR_006756P00492		Human Mutation 23 (6), pp. 599-611: Destroys the helix thus the dimerization		LF

MutationVariant	Seq ID	Description	Inh.	Mech.	
P00492	VAR_006802P00492	Human Mutation 23 (6), pp. 599-611: Interrupts hydrogen bond in A-B interface		LF	
P00492	VAR_006803P00492	Human Mutation 23 (6), pp. 599-611: Interrupts hydrogen bond in A-B interface		LF	
P00492	VAR_006765P00492	Human Mutation 23 (6), pp. 599-611: Removes one hydrogen bond, but not severe effect		LF	
P01241	VAR_015805P01241	Hum Mutat. 2003 Apr;21(4):424-40: two of the amino acids involved (K41 and T175) are among eight key residues identified as being necessary for tight binding affinity between site 1 of GH and the GHR.	AR	LF	
P01241	VAR_015814P01241	Hum Mutat. 2003 Apr;21(4):424-40: two of the amino acids involved (K41 and T175) are among eight key residues identified as being necessary for tight binding affinity between site 1 of GH and the GHR.	AR	LF	
107680	.0016	P02647	In an English family with autosomal dominant nonneuropathic systemic amyloidosis, Soutar et al. (1992) identified a CTG (leu)-to-CGG (arg) transversion at codon 60. The affected individuals were heterozygotes.	AD	GF
107680	.0010	P02647	n a family of English-Scottish-Irish extraction, Van Allen et al. (1968) studied a form of amyloidosis in which neuropathy dominated the clinical picture early in the course and nephropathy late in the course.	AD	GF
107680	.0024	P02647	Hamidi Asl et al. (1999) found that autosomal dominant hereditary amyloidosis with a unique cutaneous and cardiac presentation and death from heart failure by the sixth or seventh decade was associated with a 1389T-C transition in exon 4 of the APOA1 gene. The predicted substitution of leu90-to-pro (L90P) substitution was confirmed by structural analysis of amyloid protein isolated from cardiac deposits of amyloid. The subunit protein was composed exclusively of NH2-terminal fragments of the variant APOA1 with the longest ending at residue 94 in the wildtype sequence. Amyloid fibrils derived from 4 previously described APOA1 variants were composed of similar fragments with carboxy-terminal heterogeneity, but contrary to those variants, which all carry one extra positive charge, the leu90-to-pro substitution did not result in any charge modification.	AD	GF

MutationVariant	Seq ID	Description	Inh.	Mech.
601145 .0004	P04080	Lalioti et al. (1997) identified a homozygous G-to-C transversion at nucleotide 426 in exon 1 of the cystatin B gene in non-Finnish EPM1 (254800) families from northern Africa and Europe. The mutation resulted in a gly4-to-arg substitution and was the first missense mutation described in association with EPM1. Molecular modeling predicted that this substitution severely affected the contact of cystatin B with papain. Alakurtti et al. (2005) transiently expressed the G4R mutation in BHK-21 cells. The mutant protein failed to associate with lysosomes.	AR	LF
152780 .0004	P01229	In a 30-year-old man who presented with delayed puberty and infertility and was found to have hypogonadism associated with absence of circulating luteinizing hormone, Valdes-Socin et al. (2004) identified a homozygous gly36-to-asp (G36D) substitution in the LHB gene; the mutation disrupted a vital cysteine knot motif and abrogated the heterodimerization and secretion of luteinizing hormone.	AR	LF
O15273 VAR_029447		J Am Coll Cardiol. 2004 Dec 7;44(11):2192-201: Two TCAP mutations, T137I and R153H, were found in patients with HCM, and another TCAP mutation, E132Q, was identified in a patient with DCM. It was demonstrated by the qualitative assays that the HCM-associated mutations augment the ability of Tcap to interact with titin and calsarcin-1, whereas the DCM-associated mutations impair the interaction of Tcap with MLP, titin, and calsarcin-1		

Appendix G

Table G.1: All predicted interacting mutations with the respective structural template sequences, percentage identity between query and target sequence as well as the predicted crystal contact status of the template interaction.

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
100650	.0001	P05091	504	2.27	P05091	504	100.00	No
100690	.0003	NP_000070	274	3.75	P02711	278	81.77	No
100690	.0006	NP_000070	269	2.58	P02711	273	81.77	No
100690	.0009	NP_000070	276	3.79	P02711	280	81.77	No
100710	.0001	P11230	289	2.93	Q6S3I0	285	60.53	No
100720	.0002	Q07001	271	3.54	P02711	260	34.31	No
102540	.0002	P68032	363	3.71	P68135	363	98.93	Yes
102560	.0003	NP_001605	332	2.59	P07830	332	94.91	No
102600	.0004	NP_000476	65	2.02	P49435	67	47.41	No
102610	.0002	P68133	117	2.62	P68135	117	100.00	No
102610	.0006	P68133	359	2.63	P68135	359	100.00	Yes
102610	.0010	P68133	336	3.26	P68135	336	100.00	No
102610	.0010	P68133	336	3.26	P68139	336	100.00	No
102610	.0013	P68133	334	2.59	P68135	334	100.00	No
102610	.0013	P68133	334	2.59	P68139	334	100.00	No
103600	.0007	P02768	143	2.26	P02768	143	100.00	Yes
103600	.0011	P02768	345	2.44	P02768	345	100.00	No
103850	.0001	P04075	128	3.95	P00883	128	99.14	No
103850	.0002	P04075	206	3.20	P00883	206	99.14	No
107280	.0001	NP_001076	414	3.77	P01011	414	100.00	No
107300	.0007	P01008	416	2.84	P05619	335	40.27	No
107300	.0010	P01008	425	2.09	P01008	425	100.00	No
107300	.0011	P01008	426	2.65	P01008	426	100.00	No
107300	.0019	P01008	439	4.17	P05619	357	40.27	No
107300	.0020	P01008	425	2.09	P01008	425	100.00	No
107300	.0021	P01008	425	2.09	P01008	425	100.00	No
107300	.0022	P01008	414	2.94	P05619	333	40.27	No
107300	.0027	P01008	416	2.84	P05619	335	40.27	No
107300	.0041	P01008	402	3.09	P05120	357	35.39	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
107400	.0004	P01009	400	2.37	P01009	400	100.00	No
107400	.0012	P01009	76	3.59	P01009	76	100.00	No
107400	.0013	P01009	288	3.24	P01009	288	100.00	No
107400	.0014	P01009	393	4.17	P01009	393	100.00	No
107400	.0017	P01009	76	3.59	P01009	76	100.00	No
107400	.0019	P01009	280	2.40	P01009	280	100.00	No
107400	.0026	P01009	382	2.73	P01009	382	100.00	No
107400	.0029	P01009	360	2.98	P01011	361	45.55	No
107400	.0037	P01009	280	2.40	P01009	280	100.00	No
107400	.0039	P01009	77	3.58	P01009	77	100.00	No
107680	.0005	P02647	131	2.04	P02647	131	100.00	No
107680	.0016	P02647	84	2.73	P02647	84	100.00	No
107680	.0021	P02647	74	3.82	P02647	74	100.00	No
107680	.0022	P02647	180	3.02	P02647	180	100.00	No
107680	.0024	P02647	114	2.44	P02647	114	100.00	No
107680	.0026	P02647	198	2.75	P02647	198	100.00	No
107930	.0003	P20711	309	2.41	P80041	309	91.82	No
109270	.0003	P02730	327	2.02	P02730	327	100.00	No
114240	.0010	P20807	490	3.92	Q07009	416	56.86	No
114800	.0002	P00915	246	4.22	O43570	275	36.80	No
118504	.0004	P43681	280	2.61	P02711	272	50.00	No
120130	.0001	P02462	1408	3.86	P02452	148	38.60	Yes
120130	.0002	P02462	921	3.86	P02452	151	38.60	No
120140	.0044	NP_001835	717	3.86	P02452	145	38.60	No
120150	.0021	P02452	1178	3.86	P02452	145	43.86	Yes
120160	.0008	NP_000080	907	3.86	P02452	142	40.35	No
120160	.0010	NP_000080	547	3.86	P02452	145	42.11	No
120160	.0015	NP_000080	976	3.86	P02452	151	38.60	No
120160	.0030	NP_000080	661	3.86	P02452	139	42.11	No
120190	.0003	P05997	960	3.86	P02452	136	42.11	No
120290	.0004	NP_542412	977	3.86	P02452	148	43.86	No
120550	.0001	P02745	208	2.06	P02746	213	37.82	No
120580	.0002	NP_958850	534	2.12	P00734	467	36.20	No
121050	.0008	NP_001990	1169	5.87	Q9JJS8	152	30.77	Yes
122500	.0002	P08185	389	2.51	P01011	405	47.98	No
123101	.0003	P35548	172	3.65	P06601	243	46.43	No
123610	.0002	P05813	91	3.67	P53674	118	52.44	No
123620	.0001	NP_000487	155	2.02	P02522	154	95.12	No
123690	.0001	P07320	14	2.66	P08209	14	87.34	No
123690	.0004	P07320	23	3.23	P62697	129	39.24	No
123690	.0006	P07320	23	3.23	P62697	129	39.24	No
123940	.0003	P19013	449	3.97	P08670	395	38.76	No
124020	.0003	P33261	212	3.14	P10632	212	77.97	No
125240	.0001	P08174	87	6.30	P20023	75	30.19	No
125270	.0004	P13716	240	2.28	P13716	240	100.00	No
125660	.0003	NP_001918	393	3.13	P08670	387	73.38	No
125660	.0006	NP_001918	345	2.79	P08670	339	73.38	No
125660	.0007	NP_001918	406	4.16	P08670	400	73.38	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
125660	.0010	NP_001918	385	2.43	P08670	379	73.38	No
125660	.0011	NP_001918	389	3.89	P08670	383	73.38	Yes
130130	.0002	P08246	206	2.49	P00761	189	36.27	No
130130	.0006	P08246	139	3.20	P00761	114	36.27	No
130130	.0007	P08246	101	2.42	P08246	101	100.00	No
130130	.0009	P08246	71	5.41	P08246	71	100.00	No
130410	.0001	NP_001976	164	3.92	P38117	164	100.00	No
130410	.0003	NP_001976	128	3.97	P38117	127	100.00	No
130410	.0003	NP_001976	128	3.97	P38117	128	100.00	No
131399	.0001	NP_000493	286	4.08	P05164	314	72.21	No
131550	.0004	NP_005219	719	3.81	Q06187	408	36.48	Yes
131550	.0005	NP_005219	719	3.81	Q06187	408	36.48	Yes
134370	.0007	P08603	1207	2.30	P68638	240	31.37	Yes
134797	.0005	NP_000129	1249	5.87	Q9JJS8	152	37.50	Yes
134797	.0011	NP_000129	723	4.22	P07204	441	42.42	No
134850	.0001	P02679	301	2.72	P02679	301	100.00	No
134850	.0002	P02679	301	2.72	P02679	301	100.00	No
134850	.0004	P02679	334	3.47	P02679	334	100.00	No
134850	.0005	P02679	334	3.47	P02679	334	100.00	No
134850	.0006	P02679	336	4.27	P02679	336	100.00	No
134850	.0018	P02679	191	2.95	Q02020	227	43.46	No
134850	.0019	P02679	335	3.07	P02679	335	100.00	No
136351	.0003	NP_004110	835	2.30	P06213	1183	38.35	Yes
136351	.0004	NP_004110	835	2.30	P06213	1183	38.35	Yes
136351	.0005	NP_004110	835	2.30	P06213	1183	38.35	Yes
136351	.0006	NP_004110	835	2.30	P06213	1183	38.35	Yes
136351	.0007	NP_004110	835	2.30	P06213	1183	38.35	Yes
136352	.0003	NP_891555	1041	4.05	Q07912	256	35.04	Yes
136352	.0005	NP_891555	1114	4.23	Q06187	596	36.69	Yes
136530	.0002	P01225	69	5.86	P01225	69	100.00	No
136850	.0006	NP_000134	343	3.15	P05042	296	60.79	No
136850	.0007	NP_000134	233	3.61	P05042	186	60.79	No
136850	.0008	NP_000134	233	3.61	P05042	186	60.79	No
137780	.0010	P14136	362	3.97	P08670	395	63.96	No
137780	.0012	P14136	352	3.39	P08670	385	63.96	No
138079	.0001	P35557	279	2.31	P05708	283	51.88	No
139250	.0020	P01241	205	2.01	P01241	205	100.00	No
139320	.0003	P63092	272	3.01	P04896	272	99.74	No
139320	.0008	P63092	201	4.27	P63096	177	41.62	No
139320	.0009	P63092	201	4.27	P63096	177	41.62	No
139320	.0010	P63092	227	4.62	P10824	203	41.91	No
139320	.0012	P63092	227	4.62	P10824	203	41.91	No
139320	.0013	P63092	201	4.27	P63096	177	41.62	No
139320	.0018	P63092	170	3.46	P63096	146	41.62	No
139320	.0020	P63092	231	4.27	P04896	231	99.74	No
139320	.0021	P63092	201	4.27	P63096	177	41.62	No
139330	.0001	NP_653082	38	3.58	P63096	41	67.44	No
139340	.0001	NP_005263	79	2.81	P63096	78	69.74	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
139350	.0003	P04264	481	4.94	P08670	399	38.76	No
139360	.0001	NP_002061	179	4.27	P63096	177	87.90	No
139360	.0002	NP_002061	179	4.27	P63096	177	87.90	No
139360	.0003	NP_002061	179	4.27	P63096	177	87.90	No
139360	.0004	NP_002061	200	3.56	P10824	198	87.90	No
140100	.0005	P00738	265	2.36	P00742	341	32.38	No
141800	.0028	NP_000549	75	2.25	P02089	79	42.19	No
141800	.0095	NP_000549	75	2.25	P02089	79	42.19	No
141800	.0100	NP_000549	75	2.25	P02089	79	42.19	No
141800	.0122	NP_000549	75	2.25	P02089	79	42.19	No
141800	.0157	NP_000549	75	2.25	P02089	79	42.19	No
141850	.0006	P69905	62	3.12	P02089	67	42.19	No
141850	.0007	P69905	109	2.56	P02089	114	42.19	No
141850	.0008	P69905	61	2.40	P02089	66	42.19	No
141850	.0009	P69905	27	2.10	P01958	27	87.69	No
141850	.0011	P69905	16	2.30	P01990	16	67.69	Yes
141850	.0012	P69905	47	2.60	P02208	57	30.77	Yes
141850	.0025	P69905	47	2.60	P02208	57	30.77	Yes
141850	.0031	P69905	104	3.59	P69905	104	100.00	No
141850	.0034	P69905	74	2.25	P02089	79	42.19	No
141850	.0035	P69905	80	2.20	P02089	85	42.19	No
141850	.0037	P69905	126	2.03	P69905	126	100.00	No
141850	.0042	P69905	20	2.28	P02118	19	41.41	Yes
141850	.0045	P69905	66	2.37	P02089	71	42.19	No
141850	.0049	P69905	72	3.17	P02089	77	42.19	No
141850	.0052	P69905	95	3.08	P69905	95	100.00	No
141850	.0053	P69905	37	3.98	P01965	37	83.85	No
141850	.0055	P69905	31	2.86	P69905	31	100.00	No
141850	.0060	P69905	65	2.18	P02089	70	42.19	No
141850	.0065	P69905	59	2.52	P02089	64	42.19	No
141900	.0005	P68871	19	2.34	P02118	19	69.40	Yes
141900	.0019	P68871	15	5.78	P02118	15	69.40	Yes
141900	.0021	P68871	102	2.60	P68871	102	100.00	No
141900	.0025	P68871	88	2.86	P68871	88	100.00	No
141900	.0026	P68871	119	2.23	P68871	119	100.00	No
141900	.0027	P68871	127	2.16	P68871	127	100.00	No
141900	.0028	P68871	100	3.08	P68871	100	100.00	No
141900	.0030	P68871	67	3.12	P02089	67	79.85	No
141900	.0046	P68871	99	2.45	P68871	99	100.00	No
141900	.0048	P68871	66	2.40	P02089	66	79.85	No
141900	.0064	P68871	19	2.34	P02118	19	69.40	Yes
141900	.0068	P68871	98	2.88	P68871	98	100.00	No
141900	.0071	P68871	26	2.10	P68871	26	100.00	No
141900	.0078	P68871	77	3.17	P02089	77	79.85	No
141900	.0079	P68871	79	2.25	P02089	79	79.85	No
141900	.0084	P68871	79	2.25	P02089	79	79.85	No
141900	.0096	P68871	127	2.16	P68871	127	100.00	No
141900	.0104	P68871	26	2.10	P68871	26	100.00	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
141900	.0109	P68871	37	3.09	P68871	37	100.00	No
141900	.0114	P68871	99	2.45	P68871	99	100.00	No
141900	.0116	P68871	66	2.40	P02089	66	79.85	No
141900	.0118	P68871	92	5.44	P02089	92	79.85	No
141900	.0119	P68871	92	5.44	P02089	92	79.85	No
141900	.0120	P68871	17	2.30	P68871	17	100.00	No
141900	.0126	P68871	64	2.52	P02089	64	79.85	No
141900	.0130	P68871	128	2.12	P68871	128	100.00	No
141900	.0131	P68871	77	3.17	P02089	77	79.85	No
141900	.0143	P68871	132	2.60	P68871	132	100.00	No
141900	.0144	P68871	30	2.86	P68871	30	100.00	No
141900	.0145	P68871	102	2.60	P68871	102	100.00	No
141900	.0146	P68871	99	2.45	P68871	99	100.00	No
141900	.0148	P68871	124	2.47	P68871	124	100.00	No
141900	.0151	P68871	98	2.88	P68871	98	100.00	No
141900	.0158	P68871	36	3.98	P68871	36	100.00	No
141900	.0162	P68871	17	2.30	P68871	17	100.00	No
141900	.0163	P68871	67	3.12	P02089	67	79.85	No
141900	.0164	P68871	92	5.44	P02089	92	79.85	No
141900	.0168	P68871	19	2.34	P02118	19	69.40	Yes
141900	.0169	P68871	97	3.47	P68871	97	100.00	No
141900	.0172	P68871	114	2.56	P02089	114	79.85	No
141900	.0184	P68871	97	3.47	P68871	97	100.00	No
141900	.0186	P68871	92	5.44	P02089	92	79.85	No
141900	.0192	P68871	17	2.30	P68871	17	100.00	No
141900	.0193	P68871	97	3.47	P68871	97	100.00	No
141900	.0195	P68871	100	3.08	P68871	100	100.00	No
141900	.0197	P68871	92	5.44	P02089	92	79.85	No
141900	.0199	P68871	36	3.98	P68871	36	100.00	No
141900	.0201	P68871	98	2.88	P68871	98	100.00	No
141900	.0203	P68871	52	2.02	P02118	52	69.40	No
141900	.0212	P68871	52	2.02	P02118	52	69.40	No
141900	.0213	P68871	117	3.30	P68871	117	100.00	No
141900	.0220	P68871	35	2.99	P68871	35	100.00	No
141900	.0229	P68871	78	2.52	P02089	78	79.85	No
141900	.0230	P68871	99	2.45	P68871	99	100.00	No
141900	.0234	P68871	15	5.78	P02118	15	69.40	Yes
141900	.0236	P68871	102	2.60	P68871	102	100.00	No
141900	.0241	P68871	37	3.09	P68871	37	100.00	No
141900	.0250	P68871	117	3.30	P68871	117	100.00	No
141900	.0253	P68871	88	2.86	P68871	88	100.00	No
141900	.0256	P68871	70	2.18	P02089	70	79.85	No
141900	.0269	P68871	102	2.60	P68871	102	100.00	No
141900	.0272	P68871	52	2.02	P02118	52	69.40	No
141900	.0273	P68871	36	3.98	P68871	36	100.00	No
141900	.0274	P68871	67	3.12	P02089	67	79.85	No
141900	.0276	P68871	26	2.10	P68871	26	100.00	No
141900	.0278	P68871	30	2.86	P68871	30	100.00	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
141900	.0281	P68871	79	2.25	P02089	79	79.85	No
141900	.0288	P68871	124	2.47	P68871	124	100.00	No
141900	.0289	P68871	124	2.47	P68871	124	100.00	No
141900	.0294	P68871	123	2.16	P68871	123	100.00	No
141900	.0300	P68871	97	3.47	P68871	97	100.00	No
141900	.0301	P68871	99	2.45	P68871	99	100.00	No
141900	.0302	P68871	132	2.60	P68871	132	100.00	No
141900	.0307	P68871	99	2.45	P68871	99	100.00	No
141900	.0311	P68871	17	2.30	P68871	17	100.00	No
141900	.0313	P68871	15	5.78	P02118	15	69.40	Yes
141900	.0315	P68871	37	3.09	P68871	37	100.00	No
141900	.0318	P68871	35	2.99	P68871	35	100.00	No
141900	.0319	P68871	127	2.16	P68871	127	100.00	No
141900	.0320	P68871	127	2.16	P68871	127	100.00	No
141900	.0394	P68871	119	2.23	P68871	119	100.00	No
141900	.0397	P68871	114	2.56	P02089	114	79.85	No
141900	.0404	P68871	92	5.44	P02089	92	79.85	No
141900	.0405	P68871	99	2.45	P68871	99	100.00	No
141900	.0411	P68871	17	2.30	P68871	17	100.00	No
141900	.0424	P68871	114	2.56	P02089	114	79.85	No
141900	.0427	P68871	92	5.44	P02089	92	79.85	No
141900	.0428	P68871	18	2.32	P02118	18	69.40	Yes
141900	.0433	P68871	79	2.25	P02089	79	79.85	No
141900	.0438	P68871	67	3.12	P02089	67	79.85	No
141900	.0440	P68871	37	3.09	P68871	37	100.00	No
141900	.0447	P68871	67	3.12	P02089	67	79.85	No
141900	.0448	P68871	127	2.16	P68871	127	100.00	No
141900	.0452	P68871	98	2.88	P68871	98	100.00	No
141900	.0453	P68871	79	2.25	P02089	79	79.85	No
141900	.0466	P68871	26	2.10	P68871	26	100.00	No
141900	.0469	P68871	77	3.17	P02089	77	79.85	No
141900	.0481	P68871	124	2.47	P68871	124	100.00	No
141900	.0487	P68871	36	3.98	P68871	36	100.00	No
141900	.0490	P68871	36	3.98	P68871	36	100.00	No
141900	.0492	P68871	122	3.09	P68871	122	100.00	No
141900	.0494	P68871	117	3.30	P68871	117	100.00	No
141900	.0495	P68871	123	2.16	P68871	123	100.00	No
141900	.0499	P68871	128	2.12	P68871	128	100.00	No
141900	.0500	P68871	128	2.12	P68871	128	100.00	No
141900	.0512	P68871	64	2.52	P02089	64	79.85	No
141900	.0518	P68871	97	3.47	P68871	97	100.00	No
141900	.0525	P68871	26	2.10	P68871	26	100.00	No
141900	.0531	P68871	52	2.02	P02118	52	69.40	No
142000	.0004	P02042	99	2.45	P68871	99	92.54	No
142000	.0016	P02042	98	2.88	P68871	98	92.54	No
142000	.0029	P02042	30	2.86	P68871	30	92.54	No
142000	.0034	P02042	26	2.10	P68871	26	92.54	No
142000	.0035	P02042	37	3.09	P68871	37	92.54	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
142000	.0039	NP_000510	37	3.98	P68871	36	92.54	No
142000	.0041	NP_000510	89	2.86	P68871	88	92.54	No
142200	.0001	P69891	75	2.38	P02089	75	70.15	No
142200	.0002	P69891	128	2.12	P69891	128	100.00	No
142200	.0006	P69891	37	3.09	P02070	36	74.63	No
142200	.0006	P69891	37	3.09	P68871	37	74.63	No
142200	.0007	P69891	79	2.25	P02089	79	70.15	No
142200	.0008	P69891	97	3.47	P02070	96	74.63	No
142200	.0008	P69891	97	3.47	P68871	97	74.63	No
142200	.0016	P69891	36	3.98	P02070	35	74.63	No
142200	.0016	P69891	36	3.98	P68871	36	74.63	No
142200	.0018	P69891	75	2.38	P02089	75	70.15	No
142200	.0032	P69891	75	2.38	P02089	75	70.15	No
142250	.0009	P69892	77	3.17	P02089	77	70.90	No
142250	.0014	P69892	117	3.30	P68871	117	75.37	No
142250	.0019	P69892	26	2.10	P68871	26	75.37	No
142250	.0021	P69892	125	2.06	P69891	125	99.25	No
142250	.0022	P69892	66	2.40	P02089	66	70.90	No
142250	.0031	P69892	66	2.40	P02089	66	70.90	No
142250	.0034	P69892	92	5.44	P02089	92	70.90	No
142250	.0036	P69892	15	5.78	P02118	15	73.13	Yes
142250	.0039	P69892	75	2.38	P02089	75	70.90	No
142250	.0045	P69892	75	2.38	P02089	75	70.90	No
142250	.0048	P69892	17	2.30	P68871	17	75.37	No
142250	.0049	P69892	19	2.34	P02118	19	73.13	Yes
142360	.0004	P05546	462	2.49	P05546	462	100.00	No
142410	.0005	P20823	272	4.16	P40424	288	34.62	No
142984	.0001	P28358	319	2.82	Q6B2C0	185	37.04	Yes
142989	.0004	P35453	314	2.38	P02836	500	33.93	No
142989	.0007	P35453	298	3.65	P06601	243	30.36	No
142993	.0001	P58304	200	4.16	P40424	288	32.14	No
142993	.0002	P58304	200	4.16	P40424	288	32.14	No
142994	.0008	P50219	248	3.39	P02836	459	50.00	Yes
147450	.0001	P00441	37	3.51	P00441	37	100.00	No
147450	.0002	P00441	38	2.87	P00441	38	100.00	No
147450	.0003	P00441	41	3.40	P00441	41	100.00	Yes
147450	.0004	P00441	41	3.40	P00441	41	100.00	Yes
147450	.0006	P00441	85	3.55	P53636	117	30.71	No
147450	.0007	P00441	93	3.69	P00441	93	100.00	Yes
147450	.0008	P00441	93	3.69	P00441	93	100.00	Yes
147450	.0011	P00441	113	2.73	P00441	113	100.00	No
147450	.0016	P00441	104	3.07	P00441	104	100.00	Yes
147450	.0017	P00441	144	2.22	P00446	167	31.16	Yes
147450	.0020	P00441	6	3.81	P00442	6	83.33	No
147450	.0026	P00441	126	3.05	P00441	126	100.00	No
147450	.0033	P00441	93	3.69	P00441	93	100.00	Yes
148040	.0016	P13647	472	3.36	P08670	401	37.13	No
148041	.0004	NP_005545	469	3.39	P08670	403	37.13	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
148066	.0001	NP_000517	384	2.70	P08670	368	35.83	No
148066	.0011	NP_000517	415	4.94	P08670	399	35.83	No
148066	.0012	NP_000517	419	3.39	P08670	403	35.83	No
148067	.0008	NP_005548	354	2.41	P08670	336	34.53	No
148080	.0012	P13645	439	3.93	P08670	389	36.16	No
150330	.0017	P02545	377	4.16	P08670	400	30.39	No
151385	.0006	Q01196	107	2.92	Q01196	107	100.00	No
151385	.0008	Q01196	58	2.90	Q01196	58	100.00	No
152780	.0001	P01229	74	4.14	P01233	74	86.54	No
152780	.0004	P01229	56	3.85	P01233	56	86.54	No
153450	.0002	P61626	85	2.78	P61626	85	100.00	No
153450	.0003	P61626	82	6.08	P61626	82	100.00	No
153450	.0005	P61626	82	6.08	P61626	82	100.00	No
156845	.0003	NP_000239	217	3.51	Q12772	343	40.00	No
160710	.0002	P13533	795	4.13	P13538	795	60.00	No
160760	.0001	P12883	403	2.33	P10587	405	51.93	No
160760	.0014	P12883	403	2.33	P10587	405	51.93	No
160760	.0015	P12883	403	2.33	P10587	405	51.93	No
160760	.0022	P12883	532	2.27	P13538	534	81.80	No
160760	.0024	P12883	743	2.52	P10587	753	51.93	Yes
162280	.0001	NP_006149	333	2.92	P08670	342	53.25	No
164761	.0013	NP_066124	918	4.05	Q06187	563	35.74	Yes
164790	.0001	P01111	13	3.01	P01112	13	91.88	No
164790	.0002	P01111	61	4.56	P01112	61	91.88	No
164840	.0003	P04198	394	3.51	Q12772	343	34.00	No
164860	.0007	P08581	1136	5.25	P00520	295	40.40	Yes
171060	.0007	P21439	1161	3.24	Q9CHL8	473	46.15	Yes
171760	.0002	P05186	71	4.02	Q9BHT8	45	47.20	No
171760	.0004	P05186	71	4.02	Q9BHT8	45	47.20	No
172400	.0001	NP_000166	158	3.79	P06744	157	100.00	No
172400	.0002	P06744	346	2.58	P06744	346	100.00	No
172400	.0003	P06744	524	2.99	P06744	524	100.00	No
172400	.0004	NP_000166	539	3.10	P06744	538	100.00	No
172471	.0002	P15735	189	3.47	P05132	200	33.06	No
172471	.0002	P15735	189	3.47	P00517	200	33.06	No
173110	.0001	P28069	172	4.06	P10037	172	98.65	No
173110	.0005	P28069	143	4.06	P14859	299	58.11	Yes
173350	.0005	P00747	616	5.08	P00761	42	45.59	No
173350	.0007	P00747	751	2.42	P00747	751	100.00	Yes
173515	.0005	P14770	24	5.79	P07359	24	34.62	No
175100	.0008	P25054	713	2.29	P35222	646	32.43	Yes
176300	.0007	P02766	131	2.94	O93330	133	57.27	No
176300	.0008	P02766	136	4.70	P02766	136	100.00	No
176300	.0011	P02766	134	3.84	P02766	134	100.00	No
176300	.0033	P02766	134	3.84	P02766	134	100.00	No
176300	.0034	P02766	127	2.12	P02766	127	100.00	No
176300	.0046	P02766	73	3.86	P02766	73	100.00	No
176300	.0047	P02766	38	4.03	P02766	38	100.00	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
176300	.0050	P02766	89	4.94	O93330	91	57.27	No
176730	.0001	NP_000198	49	3.54	P01308	49	100.00	No
176730	.0002	NP_000198	48	3.24	P01308	48	100.00	No
176730	.0003	NP_000198	34	3.49	P01308	34	100.00	No
176860	.0002	P04070	444	5.62	P00735	615	40.61	No
176860	.0005	P04070	301	3.00	P00735	467	40.61	No
176860	.0008	P04070	343	3.37	P00735	508	40.61	No
176860	.0011	P04070	334	2.89	P00735	499	40.61	No
176860	.0012	P04070	289	2.88	P00735	454	40.61	No
176860	.0019	P04070	226	3.30	P00735	381	40.61	No
176860	.0022	P04070	339	2.75	P00735	504	40.61	No
176860	.0024	P04070	149	3.20	P00743	138	54.84	No
176930	.0004	P00734	425	2.37	P00734	425	100.00	No
176930	.0005	P00734	601	2.93	P00734	601	100.00	No
176947	.0005	P43403	465	4.05	Q07912	256	38.98	Yes
180200	.0003	P06400	445	3.83	P06400	445	100.00	Yes
180200	.0004	P06400	567	3.66	P06400	567	100.00	No
180200	.0019	P06400	661	4.22	P06400	661	100.00	No
188540	.0001	P01222	49	3.85	P01225	48	42.31	No
188540	.0002	P01222	32	3.57	P01225	31	42.31	No
188540	.0004	P01222	69	4.14	P01225	66	42.31	No
188826	.0002	P35625	191	2.22	P16035	200	45.88	No
189980	.0006	P00519	351	5.05	P54763	742	42.80	Yes
190020	.0001	P01112	12	2.54	P01112	12	100.00	No
190020	.0002	P01112	61	4.56	P01112	61	100.00	No
190020	.0003	P01112	12	2.54	P01112	12	100.00	No
190020	.0004	P01112	12	2.54	P01112	12	100.00	No
190020	.0005	P01112	13	3.01	P01112	13	100.00	No
190070	.0001	NP_004976	12	2.54	P01112	12	94.38	No
190070	.0002	NP_004976	12	2.54	P01112	12	94.38	No
190070	.0003	NP_004976	13	3.01	P01112	13	94.38	No
190070	.0004	NP_004976	59	3.71	P01112	59	94.38	No
190070	.0005	NP_004976	12	2.54	P01112	12	94.38	No
190070	.0006	NP_004976	12	2.54	P01112	12	94.38	No
190070	.0007	NP_004976	12	2.54	P01112	12	94.38	No
190070	.0009	NP_004976	60	3.86	P01112	60	94.38	No
190070	.0011	NP_004976	58	4.11	P01112	58	94.38	No
190070	.0013	NP_004976	34	3.06	P01112	34	94.38	No
190450	.0005	P60174	170	3.99	P04789	172	53.59	No
190990	.0001	P07951	91	3.43	P42639	91	85.59	No
190990	.0002	P07951	147	4.37	P42639	147	85.59	No
190990	.0003	P07951	117	2.08	P42639	117	85.59	Yes
191010	.0002	P09493	175	2.93	P42639	175	98.73	No
191010	.0003	P09493	95	2.48	P42639	95	98.73	No
191030	.0001	NP_689476	9	3.30	P04268	50	78.39	No
191044	.0003	NP_000354	82	2.49	P19429	81	99.24	No
191170	.0001	P04637	248	3.89	P04637	248	100.00	No
191170	.0003	P04637	245	3.65	P02340	242	88.66	Yes

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
191170	.0006	P04637	249	3.89	P04637	249	100.00	Yes
191170	.0008	P04637	242	5.61	P04637	242	100.00	No
191170	.0009	P04637	245	3.65	P02340	242	88.66	Yes
191170	.0010	P04637	248	3.89	P04637	248	100.00	No
191170	.0013	P04637	241	3.27	P04637	241	100.00	No
191170	.0019	P04637	245	3.65	P02340	242	88.66	Yes
191170	.0024	P04637	280	3.89	P04637	280	100.00	No
191170	.0030	P04637	175	3.89	P04637	175	100.00	Yes
191170	.0032	P04637	138	3.26	P04637	138	100.00	Yes
191170	.0038	P04637	189	2.04	P04637	189	100.00	Yes
191306	.0001	NP_002244	1147	3.80	P08631	497	42.57	Yes
191315	.0008	NP_001007793	604	4.59	P32577	304	37.19	Yes
217030	.0001	P05156	418	4.66	P03951	469	37.73	No
218030	.0007	P80365	227	2.70	P19992	147	30.52	No
227500	.0003	P08709	238	5.53	P00763	48	42.99	No
227500	.0004	P08709	307	2.12	P00760	109	42.06	Yes
227500	.0006	P08709	304	3.00	P00761	94	41.12	No
227500	.0007	P08709	117	2.99	P00740	104	61.29	No
227500	.0018	P08709	121	5.88	P00740	108	61.29	No
227500	.0023	P08709	414	2.45	Q9Y5Y6	816	37.67	Yes
229700	.0004	NP_000498	30	2.48	P09467	29	99.69	No
232050	.0005	P05166	168	2.88	Q8GBW6	146	52.71	No
232050	.0008	P05166	435	4.05	Q9X4K7	417	57.77	No
232800	.0003	NP_000280	39	3.77	P00512	25	48.00	No
232800	.0004	NP_000280	543	3.07	P00512	140	35.96	No
232800	.0006	NP_000280	39	3.77	P00512	25	48.00	No
234000	.0001	P00748	590	5.02	P00747	784	43.93	Yes
238331	.0002	P09622	488	4.27	P09624	479	77.98	No
250850	.0001	Q00266	322	3.04	P13444	323	97.08	No
250850	.0002	Q00266	55	2.81	P13444	56	97.98	No
250850	.0007	Q00266	264	3.43	P13444	265	97.08	No
250850	.0009	Q00266	264	3.43	P13444	265	97.08	No
256540	.0009	P10619	132	3.11	Q8W4X3	166	30.87	No
259730	.0007	NP_000058	40	3.94	P23589	70	52.36	No
264900	.0010	P03951	430	2.81	P00761	47	40.38	No
264900	.0011	P03951	594	3.22	P03951	594	100.00	No
264900	.0014	P03951	418	3.53	P00761	35	40.38	No
264900	.0015	P03951	587	4.76	P00766	207	40.18	No
300039	.0003	P49335	202	3.78	P14859	296	78.38	Yes
300075	.0017	P51812	268	3.79	P49137	263	30.96	No
300104	.0002	P31150	70	3.80	P39958	78	55.76	No
300206	.0002	NP_055086	487	5.60	Q9NZN1	487	100.00	No
300300	.0001	NP_000052	525	4.05	Q07912	256	41.30	Yes
300300	.0005	NP_000052	28	3.31	Q06187	27	100.00	No
300300	.0021	NP_000052	252	5.44	P08631	114	54.72	Yes
300300	.0022	NP_000052	255	2.28	O89100	304	33.33	No
300300	.0025	NP_000052	288	3.80	O60880	13	30.14	No
300300	.0026	NP_000052	307	4.27	P35235	32	32.43	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
300300	.0027	NP_000052	334	3.75	P27986	670	33.80	No
300300	.0032	NP_000052	408	2.92	Q06187	407	100.00	Yes
300300	.0036	NP_000052	520	4.10	P08069	1134	34.94	Yes
300300	.0037	NP_000052	520	4.10	P08069	1134	34.94	Yes
300300	.0047	NP_000052	613	3.26	P00520	455	48.19	No
300300	.0047	NP_000052	613	3.26	P00519	455	48.19	Yes
300382	.0008	Q96QS3	373	2.62	P06601	258	69.64	No
300382	.0015	Q96QS3	333	3.39	P02836	459	42.86	Yes
300382	.0016	Q96QS3	369	2.89	P06601	254	69.64	No
300386	.0003	P29965	227	2.84	P29965	227	100.00	No
300461	.0004	NP_000522	111	2.51	P04391	76	43.26	No
300461	.0025	NP_000522	129	3.02	P04391	94	43.26	No
300490	.0001	O60880	55	2.35	O60880	55	100.00	No
300490	.0004	O60880	32	4.27	P35235	32	30.14	No
300490	.0013	O60880	55	2.35	O60880	55	100.00	No
303900	.0001	P04000	247	2.14	P02699	231	45.97	No
305900	.0011	NP_000393	216	4.53	P11413	215	100.00	No
305900	.0015	NP_000393	410	3.54	P11413	409	100.00	No
305900	.0024	NP_000393	213	2.72	P11413	212	100.00	No
305900	.0027	NP_000393	227	3.21	P11413	226	100.00	Yes
305900	.0029	NP_000393	463	2.34	P11413	462	100.00	Yes
305900	.0035	NP_000393	227	3.21	P11413	226	100.00	Yes
305900	.0039	NP_000393	410	3.54	P11413	409	100.00	No
305900	.0040	NP_000393	439	3.55	P11413	438	100.00	No
305900	.0050	NP_000393	467	3.56	P11413	466	100.00	Yes
306900	.0015	P00740	75	2.42	P00741	29	92.68	No
306900	.0016	P00740	75	2.42	P00741	29	92.68	No
306900	.0022	P00740	106	3.30	P00740	106	100.00	No
306900	.0024	P00740	160	3.29	P09871	161	31.43	No
306900	.0062	NP_000124	363	2.78	P00743	370	47.00	No
308000	.0016	NP_000185	70	2.67	P00492	69	100.00	No
308000	.0017	NP_000185	71	3.25	Q26997	81	38.10	No
312865	.0007	O15266	173	2.86	P02836	510	48.21	No
313700	.0024	P10275	608	4.26	P03372	234	53.33	No
314200	.0003	P05543	303	2.59	P01011	311	44.39	No
516020	.0007	P00156	166	3.28	P00157	166	81.54	No
516030	.0008	P00395	196	2.01	P00396	196	93.64	No
516050	.0006	P00414	58	5.16	P06030	66	50.00	No
600046	.0014	O95477	935	3.62	Q9YGA6	38	31.25	No
600046	.0015	O95477	935	3.62	Q9YGA6	38	31.25	No
600194	.0004	P35908	485	3.45	P08670	398	35.18	No
600194	.0006	P35908	482	3.97	P08670	395	35.18	No
600211	.0010	Q13950	200	3.21	Q01196	149	91.04	No
600211	.0012	Q13950	169	3.35	Q01196	118	91.04	No
600225	.0002	P30793	134	2.33	P30793	134	100.00	No
600225	.0008	P30793	144	3.94	P30793	144	100.00	No
600225	.0015	P30793	135	2.72	P30793	135	100.00	No
600225	.0017	P30793	211	4.32	P22288	202	97.12	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
600509	.0003	NP_000343	716	3.84	P68187	39	33.72	No
600509	.0011	NP_000343	1506	3.85	Q9KQW9	506	33.33	Yes
600529	.0001	NP_001689	197	2.66	Q13825	197	100.00	No
600584	.0001	P52952	178	2.30	P02836	494	48.21	Yes
600584	.0013	P52952	190	4.16	P40424	288	30.36	No
600644	.0001	NP_976030	185	6.16	P15151	179	31.33	No
600871	.0002	Q99684	403	2.74	P03001	166	31.82	No
600983	.0010	P08235	645	5.86	P06536	482	89.19	No
600993	.0001	Q13485	358	2.11	Q13485	358	100.00	No
600993	.0003	Q13485	493	2.48	Q13485	493	100.00	No
600993	.0011	Q13485	352	3.51	Q13485	352	100.00	No
601107	.0001	Q92887	768	3.65	Q58206	153	31.65	Yes
601107	.0005	Q92887	1382	4.20	Q9CHL8	430	37.99	Yes
601145	.0004	P04080	4	3.43	P04080	4	100.00	No
601538	.0006	O75360	88	4.46	P40424	252	33.93	No
601538	.0011	O75360	99	3.65	P06601	243	67.86	No
601538	.0012	O75360	99	3.65	P06601	243	67.86	No
601542	.0005	NP_700476	91	4.16	P40424	288	33.93	No
601545	.0001	NP_000421	149	4.99	P62871	53	33.33	No
601545	.0006	NP_000421	31	3.95	P63005	30	100.00	No
601615	.0005	Q99758	568	3.62	P68187	38	33.33	No
601622	.0010	Q15672	156	3.54	P01106	403	44.90	No
601687	.0005	Q99456	429	4.94	P08670	399	33.11	No
601769	.0002	P11473	73	4.26	P03372	234	46.67	No
601769	.0011	P11473	391	2.95	Q13133	415	39.44	No
601789	.0002	Q92968	326	2.44	P08631	127	32.08	Yes
601802	.0001	Q9UBX0	160	4.16	P40424	288	33.93	No
601928	.0003	O43790	402	3.97	P08670	395	37.66	No
601928	.0005	O43790	402	3.97	P08670	395	37.66	No
602018	.0001	Q99748	191	2.04	Q07731	205	45.26	No
602049	.0001	P15153	57	4.23	P15153	57	100.00	No
602153	.0002	Q14533	402	3.97	P08670	395	37.66	No
602225	.0001	O43186	80	2.89	P06601	254	64.29	No
602225	.0005	O43186	41	3.12	P06601	215	64.29	No
602225	.0006	O43186	41	3.12	P06601	215	64.29	No
602298	.0001	P51149	129	2.78	P62825	126	32.91	No
602298	.0002	P51149	162	3.47	P62826	156	32.91	No
602298	.0002	P51149	162	3.47	P62826	157	32.91	No
602298	.0003	P51149	161	3.86	P11233	163	36.25	No
602421	.0010	P13569	549	3.70	P13569	549	100.00	No
602421	.0011	P13569	549	3.70	P13569	549	100.00	No
602421	.0012	P13569	549	3.70	P13569	549	100.00	No
602421	.0022	P13569	1282	2.52	Q9CHL8	421	31.67	Yes
602421	.0032	P13569	1303	3.39	Q9CHL8	442	31.67	Yes
602421	.0048	P13569	1291	4.20	Q9CHL8	430	31.67	Yes
602421	.0063	P13569	1283	2.74	Q9CHL8	422	31.67	Yes
602421	.0114	P13569	1303	3.39	Q9CHL8	442	31.67	Yes
602438	.0003	Q9ULV5	20	2.65	P22121	196	36.98	Yes

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
602445	.0001	NP_005016	49	3.58	O35684	49	86.93	No
602445	.0002	NP_005016	52	3.19	O35684	52	86.93	No
602533	.0004	Q99497	149	3.28	Q99497	149	100.00	Yes
602575	.0001	O60663	246	4.42	P02836	504	37.50	No
602575	.0002	O60663	198	3.12	P06601	215	35.71	No
602575	.0011	O60663	226	3.65	P06601	243	35.71	No
602765	.0001	P78385	407	3.97	P08670	395	37.66	No
602821	.0002	Q12840	280	4.20	P33173	307	44.04	Yes
603234	.0017	O95255	1339	2.74	Q9CHL8	422	36.11	Yes
603470	.0008	P00966	363	3.87	Q9X2A1	361	60.00	No
603470	.0009	P00966	390	3.83	Q9X2A1	388	60.00	No
603470	.0010	P00966	304	3.22	Q9X2A1	302	60.00	No
603470	.0012	P00966	86	2.52	Q9X2A1	84	60.00	No
603470	.0013	P00966	279	4.22	Q9X2A1	277	60.00	No
603470	.0016	P00966	362	3.21	Q9X2A1	360	60.00	No
603470	.0019	P00966	310	3.26	Q9X2A1	308	60.00	No
603851	.0005	Q99453	100	3.12	P06601	215	69.64	No
603868	.0001	P51159	73	5.75	P63012	76	46.88	No
603868	.0006	P51159	152	3.49	P01112	134	33.96	Yes
604277	.0004	Q9UBP0	499	3.83	Q01853	637	40.66	No
604720	.0005	Q9UP52	690	2.50	P02786	658	53.38	No
605020	.0001	Q9NZR4	166	3.12	P06601	215	62.50	No
605271	.0001	Q9UK55	324	5.69	P01011	299	32.34	No
605481	.0005	Q8IZT6	3060	4.58	Q02440	775	36.84	Yes
605481	.0006	Q8IZT6	1326	4.17	Q02440	778	35.00	Yes
605481	.0008	Q8IZT6	2063	3.41	Q02440	787	35.00	Yes
605511	.0003	P57727	251	5.08	P00761	42	40.38	No
605511	.0004	P57727	404	3.88	P07338	216	41.28	No
606765	.0005	NP_783651	453	3.61	P05164	462	47.52	No
606873	.0012	P07686	183	2.03	P07686	183	100.00	Yes
606885	.0005	P16219	383	3.62	P15651	383	94.63	No
606989	.0002	NP_000241	173	4.41	P05164	173	100.00	No
606989	.0003	NP_000241	251	3.39	P05164	251	100.00	No
606999	.0008	P07902	171	3.78	P09148	151	52.30	No
606999	.0011	P07902	183	2.41	P09148	163	52.30	No
606999	.0016	P07902	194	3.47	P09148	174	52.30	No
607379	.0005	P35240	535	2.05	P26038	517	38.76	No
607379	.0006	P35240	538	3.26	P26038	520	38.76	No
607809	.0002	P24752	183	3.07	P07097	146	43.14	No
608053	.0002	P13804	266	3.56	P13804	266	100.00	No
608310	.0002	P04424	286	2.79	P04424	286	100.00	No
608348	.0003	P12694	290	3.12	P84129	227	37.04	No
608537	.0025	P40337	155	2.93	P40337	155	100.00	No
608801	.0007	Q92947	337	3.08	Q06319	290	31.03	No
608845	.0003	Q9H0F7	31	3.88	P84080	30	42.20	No
608845	.0003	Q9H0F7	31	3.88	P84079	30	42.20	No
608845	.0003	Q9H0F7	31	3.88	P84077	30	42.20	No
608845	.0005	Q9H0F7	31	3.88	P84080	30	42.20	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
608845	.0005	Q9H0F7	31	3.88	P84079	30	42.20	No
608845	.0005	Q9H0F7	31	3.88	P84077	30	42.20	No
609014	.0002	Q9HCC0	99	3.09	Q8GBW6	61	35.11	No
609014	.0003	Q9HCC0	155	2.11	Q9X4K7	123	32.40	No
609712	.0003	NP_870986	353	4.05	P30613	384	100.00	No
609712	.0004	P30613	384	4.05	P30613	384	100.00	No
609712	.0006	P30613	479	2.41	P11974	435	59.17	Yes
O15266	VAR_012346	O15266	173	2.86	P02836	510	48.21	No
O43186	VAR_003750	O43186	41	3.12	P06601	215	64.29	No
O43186	VAR_003751	O43186	80	2.89	P06601	254	64.29	No
O43186	VAR_007946	O43186	41	3.12	P06601	215	64.29	No
O43790	VAR_018126	O43790	402	3.97	P08670	395	37.66	No
O43790	VAR_018127	O43790	402	3.97	P08670	395	37.66	No
O60663	VAR_004203	O60663	226	3.65	P06601	243	35.71	No
O60663	VAR_004205	O60663	246	4.42	P02836	504	37.50	No
O60806	VAR_018387	O60806	128	3.32	P24781	127	81.36	No
O60880	VAR_005612	O60880	32	4.27	P35235	32	30.14	No
O60880	VAR_018307	O60880	55	2.35	O60880	55	100.00	No
O95255	VAR_013390	O95255	1339	2.74	Q9CHL8	422	36.11	Yes
O95255	VAR_013391	O95255	1347	4.20	Q9CHL8	430	36.11	Yes
O95342	VAR_013334	O95342	461	4.04	Q9YGA6	42	30.82	No
O95477	VAR_009150	O95477	935	3.62	Q9YGA6	38	31.25	No
P00156	VAR_013653	P00156	166	3.28	P00157	166	81.54	No
P00414	VAR_002167	P00414	78	3.54	P00415	78	87.84	No
P00441	VAR_007132	P00441	7	3.01	P00441	7	100.00	No
P00441	VAR_007136	P00441	37	3.51	P00441	37	100.00	No
P00441	VAR_007137	P00441	38	2.87	P00441	38	100.00	No
P00441	VAR_007138	P00441	41	3.40	P00441	41	100.00	Yes
P00441	VAR_007139	P00441	41	3.40	P00441	41	100.00	Yes
P00441	VAR_007144	P00441	85	3.55	P53636	117	30.71	No
P00441	VAR_007146	P00441	93	3.69	P00441	93	100.00	Yes
P00441	VAR_007147	P00441	93	3.69	P00441	93	100.00	Yes
P00441	VAR_007148	P00441	93	3.69	P00441	93	100.00	Yes
P00441	VAR_007149	P00441	93	3.69	P00441	93	100.00	Yes
P00441	VAR_007155	P00441	113	2.73	P00441	113	100.00	No
P00441	VAR_007156	P00441	115	3.75	P00441	115	100.00	No
P00441	VAR_007157	P00441	125	3.92	P00441	125	100.00	Yes
P00441	VAR_007159	P00441	139	4.10	P00441	139	100.00	Yes
P00441	VAR_007160	P00441	144	2.22	P00446	167	31.16	Yes
P00441	VAR_007161	P00441	148	3.38	P00441	148	100.00	No
P00441	VAR_007162	P00441	148	3.38	P00441	148	100.00	No
P00441	VAR_007163	P00441	149	3.53	P00441	149	100.00	No
P00441	VAR_007164	P00441	151	2.66	P00441	151	100.00	No
P00441	VAR_008717	P00441	6	3.81	P00442	6	83.33	No
P00441	VAR_008719	P00441	93	3.69	P00441	93	100.00	Yes
P00441	VAR_008720	P00441	104	3.07	P00441	104	100.00	Yes
P00441	VAR_008722	P00441	124	4.20	P00441	124	100.00	Yes
P00441	VAR_008724	P00441	144	2.22	P00446	167	31.16	Yes

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P00441	VAR_013524	P00441	38	2.87	P00441	38	100.00	No
P00441	VAR_013526	P00441	49	2.28	P00441	49	100.00	No
P00441	VAR_013529	P00441	76	2.97	P00441	76	100.00	Yes
P00441	VAR_013531	P00441	86	4.15	P00441	86	100.00	No
P00441	VAR_013532	P00441	89	2.76	P00441	89	100.00	Yes
P00441	VAR_013535	P00441	105	2.16	P00441	105	100.00	Yes
P00441	VAR_013536	P00441	108	3.52	P00441	108	100.00	No
P00441	VAR_013538	P00441	114	3.85	P00441	114	100.00	No
P00441	VAR_013539	P00441	126	3.05	P00441	126	100.00	No
P00451	VAR_015134	P00451	2307	2.35	P12259	2183	42.54	Yes
P00480	VAR_004864	P00480	90	3.76	P04391	55	43.26	No
P00480	VAR_004875	P00480	126	4.22	P04391	91	43.26	No
P00480	VAR_004876	P00480	129	3.02	P04391	94	43.26	No
P00480	VAR_004922	P00480	264	2.20	P04391	232	38.06	No
P00480	VAR_004923	P00480	264	2.20	P04391	232	38.06	No
P00480	VAR_004924	P00480	267	2.66	P00480	267	100.00	Yes
P00480	VAR_004925	P00480	268	4.55	P00480	268	100.00	Yes
P00480	VAR_004926	P00480	269	3.04	P00480	269	100.00	Yes
P00492	VAR_006773	P00492	69	2.67	P00492	69	100.00	No
P00492	VAR_006774	P00492	70	3.25	Q26997	81	38.10	No
P00533	VAR_019297	P00533	719	3.81	Q06187	408	36.48	Yes
P00734	VAR_006715	P00734	425	2.37	P00734	425	100.00	No
P00734	VAR_006719	P00734	601	2.93	P00734	601	100.00	No
P00740	VAR_006543	P00740	91	4.57	P00741	45	92.68	No
P00740	VAR_006548	P00740	102	5.88	P09871	143	37.93	No
P00740	VAR_006549	P00740	106	3.30	P00740	106	100.00	No
P00740	VAR_006550	P00740	108	5.88	P00740	108	100.00	No
P00740	VAR_006564	P00740	160	3.29	P09871	161	31.43	No
P00740	VAR_006575	P00740	241	3.30	P00761	23	45.71	No
P00740	VAR_006576	P00740	253	3.36	P00761	34	45.71	No
P00740	VAR_006577	P00740	253	3.36	P00761	34	45.71	No
P00740	VAR_006578	P00740	265	3.24	P00761	46	45.71	No
P00740	VAR_006580	P00740	283	2.03	P00761	62	45.71	No
P00740	VAR_006584	P00740	302	4.66	P00761	81	45.71	No
P00740	VAR_006585	P00740	316	2.96	P00761	93	45.71	No
P00740	VAR_006586	P00740	321	2.66	P00761	98	45.71	No
P00740	VAR_006587	P00740	333	3.26	P00761	110	45.71	No
P00740	VAR_006591	P00740	356	5.33	P00761	129	45.71	No
P00740	VAR_006592	P00740	357	3.37	P00761	130	45.71	No
P00740	VAR_006594	P00740	363	2.78	P00743	370	47.00	No
P00740	VAR_006600	P00740	390	2.38	P08709	383	42.79	No
P00740	VAR_006601	P00740	394	3.78	P00761	168	45.71	No
P00740	VAR_006604	P00740	407	5.21	P00761	181	45.71	No
P00740	VAR_006605	P00740	413	3.64	P00761	187	45.71	No
P00740	VAR_006609	P00740	430	3.22	P00761	200	45.71	No
P00740	VAR_006610	P00740	431	4.69	P00761	201	45.71	No
P00740	VAR_006611	P00740	431	4.69	P00761	201	45.71	No
P00740	VAR_006612	P00740	432	3.14	P00761	202	45.71	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P00740	VAR_006613	P00740	432	3.14	P00761	202	45.71	No
P00740	VAR_006620	P00740	450	3.32	P00761	220	45.71	No
P00740	VAR_017308	P00740	75	2.42	P00741	29	92.68	No
P00740	VAR_017312	P00740	252	5.53	P00761	33	45.71	No
P00740	VAR_017315	P00740	306	2.61	P00761	85	45.71	No
P00740	VAR_017316	P00740	357	3.37	P00761	130	45.71	No
P00740	VAR_017317	P00740	397	2.61	P00761	171	45.71	No
P00740	VAR_017318	P00740	410	3.74	P00761	184	45.71	No
P00740	VAR_017319	P00740	411	3.65	P00761	185	45.71	No
P00740	VAR_017320	P00740	411	3.65	P00761	185	45.71	No
P00740	VAR_017321	P00740	414	3.88	P00761	188	45.71	No
P00740	VAR_017322	P00740	442	2.93	P00761	212	45.71	No
P00740	VAR_017324	P00740	453	5.62	P00761	223	45.71	No
P00740	VAR_017344	P00740	52	2.02	P00741	6	92.68	Yes
P00740	VAR_017346	P00740	106	3.30	P00740	106	100.00	No
P00740	VAR_017352	P00740	241	3.30	P00761	23	45.71	No
P00740	VAR_017353	P00740	252	5.53	P00761	33	45.71	No
P00740	VAR_017354	P00740	318	2.64	P00761	95	45.71	No
P00740	VAR_017355	P00740	333	3.26	P00761	110	45.71	No
P00740	VAR_017362	P00740	407	5.21	P00761	181	45.71	No
P00740	VAR_017363	P00740	412	3.87	P00761	186	45.71	No
P00740	VAR_017364	P00740	435	5.02	P00761	205	45.71	No
P00740	VAR_017365	P00740	442	2.93	P00761	212	45.71	No
P00747	VAR_006629	P00747	620	3.24	P00761	46	45.59	No
P00747	VAR_006630	P00747	751	2.42	P00747	751	100.00	Yes
P00748	VAR_006624	P00748	590	5.02	P00747	784	43.93	Yes
P00790	VAR_006483	P00790	92	2.09	P07339	95	50.00	No
P00813	VAR_002222	P00813	141	2.39	P56658	141	89.05	No
P00966	VAR_000683	P00966	86	2.52	Q9X2A1	84	60.00	No
P00966	VAR_000688	P00966	272	4.22	Q9X2A1	270	60.00	No
P00966	VAR_000690	P00966	304	3.22	Q9X2A1	302	60.00	No
P00966	VAR_000692	P00966	363	3.87	Q9X2A1	361	60.00	No
P00966	VAR_000693	P00966	363	3.87	Q9X2A1	361	60.00	No
P00966	VAR_000694	P00966	390	3.83	Q9X2A1	388	60.00	No
P00966	VAR_015892	P00966	86	2.52	Q9X2A1	84	60.00	No
P00966	VAR_015900	P00966	265	3.86	Q9X2A1	263	60.00	No
P00966	VAR_015901	P00966	269	2.91	Q9X2A1	267	60.00	No
P00966	VAR_015903	P00966	310	3.26	Q9X2A1	308	60.00	No
P00966	VAR_015904	P00966	362	3.21	Q9X2A1	360	60.00	No
P00966	VAR_016008	P00966	279	4.22	Q9X2A1	277	60.00	No
P00966	VAR_016009	P00966	310	3.26	Q9X2A1	308	60.00	No
P00966	VAR_016010	P00966	363	3.87	Q9X2A1	361	60.00	No
P00966	VAR_016011	P00966	363	3.87	Q9X2A1	361	60.00	No
P00966	VAR_016015	P00966	119	4.03	P59846	116	54.26	No
P01008	VAR_007042	P01008	90	4.16	O35684	29	32.34	No
P01008	VAR_007044	P01008	112	4.27	P05619	32	40.27	No
P01008	VAR_007047	P01008	133	3.75	P07385	41	30.91	No
P01008	VAR_007053	P01008	158	2.95	P05619	73	40.27	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P01008	VAR_007056	P01008	198	4.46	O35684	132	32.34	No
P01008	VAR_007062	P01008	283	5.14	P01008	283	100.00	No
P01008	VAR_007063	P01008	302	2.12	P01008	302	100.00	No
P01008	VAR_007065	P01008	334	2.34	P01012	262	31.81	No
P01008	VAR_007069	P01008	414	2.94	P05619	333	40.27	No
P01008	VAR_007070	P01008	416	2.84	P05619	335	40.27	No
P01008	VAR_007071	P01008	416	2.84	P05619	335	40.27	No
P01008	VAR_007074	P01008	425	2.09	P01008	425	100.00	No
P01008	VAR_007075	P01008	425	2.09	P01008	425	100.00	No
P01008	VAR_007076	P01008	425	2.09	P01008	425	100.00	No
P01008	VAR_007077	P01008	426	2.65	P01008	426	100.00	No
P01008	VAR_007078	P01008	434	3.53	P05619	352	40.27	No
P01008	VAR_007079	P01008	434	3.53	P05619	352	40.27	No
P01008	VAR_007080	P01008	434	3.53	P05619	352	40.27	No
P01008	VAR_007082	P01008	437	3.08	P01008	437	100.00	No
P01008	VAR_007083	P01008	438	3.31	P01008	438	100.00	No
P01008	VAR_007084	P01008	439	4.17	P05619	357	40.27	No
P01008	VAR_007085	P01008	439	4.17	P05619	357	40.27	No
P01008	VAR_007087	P01008	456	3.57	P05619	374	40.27	No
P01008	VAR_007088	P01008	457	2.58	P01008	457	100.00	Yes
P01008	VAR_009258	P01008	438	3.31	P01008	438	100.00	No
P01008	VAR_012316	P01008	95	4.09	P05619	16	40.27	No
P01009	VAR_006980	P01009	58	2.83	P01009	58	100.00	No
P01009	VAR_006985	P01009	77	3.58	P01009	77	100.00	No
P01009	VAR_006986	P01009	84	3.01	P05120	38	31.15	No
P01009	VAR_006999	P01009	280	2.40	P01009	280	100.00	No
P01009	VAR_007001	P01009	354	3.32	P05120	352	31.15	No
P01009	VAR_007005	P01009	382	2.73	P01009	382	100.00	No
P01009	VAR_007006	P01009	386	2.33	P01009	386	100.00	No
P01009	VAR_007007	P01009	386	2.33	P01009	386	100.00	No
P01009	VAR_007009	P01009	393	4.17	P01009	393	100.00	No
P01111	VAR_006845	P01111	13	3.01	P01112	13	91.88	No
P01111	VAR_006846	P01111	61	4.56	P01112	61	91.88	No
P01111	VAR_021194	P01111	12	2.54	P01112	12	91.88	No
P01112	VAR_006836	P01112	12	2.54	P01112	12	100.00	No
P01112	VAR_006837	P01112	12	2.54	P01112	12	100.00	No
P01112	VAR_006838	P01112	61	4.56	P01112	61	100.00	No
P01116	VAR_006839	P01116	12	2.54	P01112	12	94.38	No
P01116	VAR_006840	P01116	12	2.54	P01112	12	94.38	No
P01116	VAR_006841	P01116	61	4.56	P01112	61	94.38	No
P01116	VAR_016026	P01116	12	2.54	P01112	12	94.38	No
P01116	VAR_016027	P01116	12	2.54	P01112	12	94.38	No
P01116	VAR_016028	P01116	12	2.54	P01112	12	94.38	No
P01116	VAR_016029	P01116	13	3.01	P01112	13	94.38	No
P01116	VAR_016030	P01116	59	3.71	P01112	59	94.38	No
P01130	VAR_005361	P01130	327	3.20	P09871	145	38.24	No
P01130	VAR_005362	P01130	329	5.88	P09871	147	38.24	No
P01130	VAR_005367	P01130	343	3.29	P09871	161	38.24	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P01130	VAR_005373	P01130	364	5.87	Q9JJS8	152	40.63	No
P01241	VAR_015814	P01241	201	2.34	P01241	201	100.00	No
P01241	VAR_015815	P01241	209	3.80	P01241	209	100.00	No
P01308	VAR_003971	P01308	34	3.49	P01308	34	100.00	No
P01308	VAR_003972	P01308	48	3.24	P01308	48	100.00	No
P01308	VAR_003973	P01308	49	3.54	P01308	49	100.00	No
P01308	VAR_003976	P01308	92	3.28	P01308	92	100.00	No
P01857	VAR_003888	P01857	241	2.07	P01865	240	61.63	No
P02042	VAR_003104	P02042	26	2.10	P68871	26	92.54	No
P02042	VAR_003113	P02042	98	2.88	P68871	98	92.54	No
P02042	VAR_003114	P02042	99	2.45	P68871	99	92.54	No
P02452	VAR_001644	P02452	221	3.86	P02452	151	47.37	No
P02452	VAR_001646	P02452	263	3.76	P02452	133	35.09	No
P02452	VAR_001647	P02452	263	3.76	P02452	133	35.09	No
P02452	VAR_001648	P02452	272	3.86	P02452	142	35.09	No
P02452	VAR_001649	P02452	275	3.86	P02452	145	35.09	No
P02452	VAR_001650	P02452	332	3.86	P02452	142	45.61	No
P02452	VAR_001654	P02452	383	3.76	P02452	133	40.35	No
P02452	VAR_001655	P02452	389	3.86	P02452	139	40.35	No
P02452	VAR_001656	P02452	389	3.86	P02452	139	40.35	No
P02452	VAR_001657	P02452	398	3.86	P02452	148	40.35	No
P02452	VAR_001658	P02452	398	3.86	P02452	148	40.35	No
P02452	VAR_001659	P02452	401	3.86	P02452	151	40.35	No
P02452	VAR_001672	P02452	569	3.86	P02452	139	43.86	No
P02452	VAR_001675	P02452	638	3.86	P02452	148	43.86	No
P02452	VAR_001677	P02452	701	3.86	P02452	151	40.35	No
P02452	VAR_001683	P02452	743	3.76	P02452	133	40.35	No
P02452	VAR_001684	P02452	743	3.76	P02452	133	40.35	No
P02452	VAR_001688	P02452	809	3.86	P02452	136	42.11	No
P02452	VAR_001689	P02452	815	3.86	P02452	142	42.11	No
P02452	VAR_001690	P02452	821	3.86	P02452	148	42.11	No
P02452	VAR_001696	P02452	869	3.86	P02452	136	43.86	No
P02452	VAR_001697	P02452	884	3.86	P02452	151	43.86	No
P02452	VAR_001699	P02452	926	3.76	P02452	133	42.11	No
P02452	VAR_001708	P02452	1049	3.86	P02452	136	43.86	Yes
P02452	VAR_001709	P02452	1058	3.86	P02452	145	43.86	Yes
P02452	VAR_001710	P02452	1061	3.86	P02452	148	43.86	Yes
P02452	VAR_001711	P02452	1061	3.86	P02452	148	43.86	Yes
P02452	VAR_001719	P02452	1106	3.76	P02452	133	45.61	Yes
P02452	VAR_001720	P02452	1124	3.86	P02452	151	45.61	Yes
P02452	VAR_001725	P02452	1166	3.76	P02452	133	43.86	Yes
P02452	VAR_001726	P02452	1172	3.86	P02452	139	43.86	Yes
P02452	VAR_001727	P02452	1181	3.86	P02452	148	43.86	Yes
P02452	VAR_001728	P02452	1184	3.86	P02452	151	43.86	Yes
P02452	VAR_008118	P02452	866	3.76	P02452	133	43.86	No
P02458	VAR_001742	P02458	285	3.86	P02452	142	49.12	No
P02458	VAR_001749	P02458	705	3.86	P02452	142	42.11	No
P02458	VAR_001752	P02458	822	3.86	P02452	136	47.37	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P02458	VAR_001757	P02458	948	3.86	P02452	142	45.61	No
P02458	VAR_001761	P02458	1074	3.86	P02452	148	45.61	Yes
P02458	VAR_001764	P02458	1119	3.76	P02452	133	43.86	Yes
P02458	VAR_001765	P02458	1128	3.86	P02452	142	43.86	Yes
P02458	VAR_017641	P02458	702	3.86	P02452	139	42.11	No
P02458	VAR_017642	P02458	711	3.86	P02452	148	42.11	No
P02458	VAR_017644	P02458	825	3.86	P02452	139	47.37	No
P02458	VAR_017646	P02458	879	3.76	P02452	133	43.86	No
P02458	VAR_023929	P02458	648	3.86	P02452	145	38.60	No
P02458	VAR_023931	P02458	828	3.86	P02452	142	47.37	No
P02458	VAR_024820	P02458	648	3.86	P02452	145	38.60	No
P02458	VAR_024821	P02458	702	3.86	P02452	139	42.11	No
P02461	VAR_001769	P02461	201	3.86	P02452	139	49.12	No
P02461	VAR_001773	P02461	567	3.86	P02452	139	42.11	No
P02461	VAR_001780	P02461	756	3.86	P02452	148	42.11	No
P02461	VAR_001783	P02461	804	3.76	P02452	133	40.35	No
P02461	VAR_001786	P02461	936	3.86	P02452	145	42.11	No
P02461	VAR_001787	P02461	936	3.86	P02452	145	42.11	No
P02461	VAR_001788	P02461	939	3.86	P02452	148	42.11	No
P02461	VAR_001791	P02461	996	3.86	P02452	145	38.60	No
P02461	VAR_001793	P02461	1050	3.86	P02452	139	43.86	Yes
P02461	VAR_001797	P02461	1104	3.76	P02452	133	43.86	Yes
P02461	VAR_001798	P02461	1164	3.76	P02452	133	38.60	Yes
P02461	VAR_001799	P02461	1167	3.86	P02452	136	38.60	Yes
P02461	VAR_001800	P02461	1170	3.86	P02452	139	38.60	Yes
P02461	VAR_001801	P02461	1173	3.86	P02452	142	38.60	Yes
P02461	VAR_001802	P02461	1176	3.86	P02452	145	38.60	Yes
P02461	VAR_001803	P02461	1182	3.86	P02452	151	38.60	Yes
P02461	VAR_011098	P02461	204	3.86	P02452	142	49.12	No
P02461	VAR_011099	P02461	204	3.86	P02452	142	49.12	No
P02461	VAR_011100	P02461	210	3.86	P02452	148	49.12	No
P02461	VAR_011111	P02461	264	3.86	P02452	136	40.35	No
P02461	VAR_011112	P02461	267	3.86	P02452	139	40.35	No
P02461	VAR_011113	P02461	321	3.76	P02452	133	42.11	No
P02461	VAR_011114	P02461	327	3.86	P02452	139	42.11	No
P02461	VAR_011117	P02461	444	3.86	P02452	136	36.84	No
P02461	VAR_011119	P02461	501	3.76	P02452	133	42.11	No
P02461	VAR_011120	P02461	519	3.86	P02452	151	42.11	No
P02461	VAR_011124	P02461	636	3.86	P02452	148	42.11	No
P02461	VAR_011128	P02461	699	3.86	P02452	151	43.86	No
P02461	VAR_011131	P02461	744	3.86	P02452	136	42.11	No
P02461	VAR_011134	P02461	879	3.86	P02452	148	52.63	No
P02461	VAR_011135	P02461	882	3.86	P02452	151	52.63	No
P02461	VAR_011140	P02461	924	3.76	P02452	133	42.11	No
P02461	VAR_011141	P02461	942	3.86	P02452	151	42.11	No
P02461	VAR_011144	P02461	984	3.76	P02452	133	38.60	No
P02461	VAR_011145	P02461	999	3.86	P02452	148	38.60	No
P02461	VAR_011149	P02461	1044	3.76	P02452	133	43.86	Yes

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P02461	VAR_011150	P02461	1050	3.86	P02452	139	43.86	Yes
P02461	VAR_011155	P02461	1164	3.76	P02452	133	38.60	Yes
P02461	VAR_011156	P02461	1164	3.76	P02452	133	38.60	Yes
P02461	VAR_011157	P02461	1170	3.86	P02452	139	38.60	Yes
P02461	VAR_011158	P02461	1173	3.86	P02452	142	38.60	Yes
P02461	VAR_011159	P02461	1179	3.86	P02452	148	38.60	Yes
P02533	VAR_003843	P02533	383	2.70	P08670	368	35.83	No
P02533	VAR_003844	P02533	414	4.94	P08670	399	35.83	No
P02533	VAR_003845	P02533	418	3.39	P08670	403	35.83	No
P02533	VAR_010450	P02533	376	2.96	P08670	361	35.83	No
P02533	VAR_010451	P02533	387	3.13	P08670	372	35.83	No
P02533	VAR_023724	P02533	407	3.26	P08670	392	35.83	No
P02533	VAR_023725	P02533	412	3.03	P08670	397	35.83	No
P02538	VAR_017076	P02538	468	3.39	P08670	403	37.13	No
P02545	VAR_009985	P02545	358	3.66	P08670	381	30.39	No
P02545	VAR_009986	P02545	371	2.86	P08670	394	30.39	No
P02545	VAR_016205	P02545	377	4.16	P08670	400	30.39	No
P02647	VAR_000610	P02647	84	2.73	P02647	84	100.00	No
P02647	VAR_000616	P02647	132	4.16	P02647	132	100.00	No
P02647	VAR_021362	P02647	180	3.02	P02647	180	100.00	No
P02679	VAR_002409	P02679	301	2.72	P02679	301	100.00	No
P02679	VAR_002410	P02679	301	2.72	P02679	301	100.00	No
P02679	VAR_002412	P02679	334	3.47	P02679	334	100.00	No
P02679	VAR_002413	P02679	334	3.47	P02679	334	100.00	No
P02679	VAR_002414	P02679	336	4.27	P02679	336	100.00	No
P02679	VAR_015853	P02679	335	3.07	P02679	335	100.00	No
P02708	VAR_000285	P02708	299	3.75	P02711	278	81.77	No
P02708	VAR_021207	P02708	294	2.58	P02711	273	81.77	No
P02708	VAR_021208	P02708	301	3.79	P02711	280	81.77	No
P02730	VAR_000800	P02730	327	2.02	P02730	327	100.00	No
P02730	VAR_013786	P02730	147	2.32	P02730	147	100.00	Yes
P02766	VAR_007548	P02766	38	4.03	P02766	38	100.00	No
P02766	VAR_007549	P02766	38	4.03	P02766	38	100.00	No
P02766	VAR_007551	P02766	44	4.22	P02766	44	100.00	No
P02766	VAR_007577	P02766	89	4.94	O93330	91	57.27	No
P02766	VAR_007592	P02766	127	2.12	P02766	127	100.00	No
P02766	VAR_007594	P02766	131	2.94	O93330	133	57.27	No
P02766	VAR_007595	P02766	134	3.84	P02766	134	100.00	No
P02766	VAR_007596	P02766	136	4.70	P02766	136	100.00	No
P02766	VAR_007597	P02766	136	4.70	P02766	136	100.00	No
P02766	VAR_007598	P02766	134	3.84	P02766	134	100.00	No
P02768	VAR_000511	P02768	143	2.26	P02768	143	100.00	Yes
P02768	VAR_000523	P02768	345	2.44	P02768	345	100.00	No
P03951	VAR_012093	P03951	430	2.81	P00761	47	40.38	No
P03951	VAR_012096	P03951	594	3.22	P03951	594	100.00	No
P04070	VAR_006648	P04070	108	2.19	P00740	105	43.33	No
P04070	VAR_006649	P04070	109	3.30	P00740	106	43.33	No
P04070	VAR_006657	P04070	147	5.88	P00742	136	48.39	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P04070	VAR_006658	P04070	149	3.20	P00743	138	54.84	No
P04070	VAR_006670	P04070	226	3.30	P00735	381	40.61	No
P04070	VAR_006671	P04070	243	2.86	P00735	399	40.61	No
P04070	VAR_006673	P04070	253	5.33	P00735	409	40.61	No
P04070	VAR_006679	P04070	289	2.88	P00735	454	40.61	No
P04070	VAR_006681	P04070	298	3.22	P00735	464	40.61	No
P04070	VAR_006682	P04070	301	3.00	P00735	467	40.61	No
P04070	VAR_006683	P04070	301	3.00	P00735	467	40.61	No
P04070	VAR_006687	P04070	321	3.20	P00735	487	40.61	No
P04070	VAR_006691	P04070	334	2.89	P00735	499	40.61	No
P04070	VAR_006693	P04070	343	3.37	P00735	508	40.61	No
P04070	VAR_006695	P04070	367	2.03	P00735	533	40.61	No
P04070	VAR_006697	P04070	385	3.78	P00735	551	40.61	No
P04070	VAR_006698	P04070	388	2.61	P00761	171	37.67	No
P04070	VAR_006699	P04070	388	2.61	P00761	171	37.67	No
P04070	VAR_006702	P04070	401	3.74	P00735	570	40.61	No
P04070	VAR_006704	P04070	423	3.14	P00735	594	40.61	No
P04070	VAR_006705	P04070	426	5.02	P00735	597	40.61	No
P04070	VAR_006706	P04070	433	2.93	P00735	604	40.61	No
P04070	VAR_006707	P04070	436	2.77	P00735	607	40.61	No
P04070	VAR_006708	P04070	441	3.32	P00735	612	40.61	No
P04070	VAR_006709	P04070	444	5.62	P00735	615	40.61	No
P04075	VAR_000550	P04075	128	3.95	P00883	128	99.14	No
P04080	VAR_002206	P04080	4	3.43	P04080	4	100.00	No
P04181	VAR_000568	P04181	93	3.59	P04181	93	100.00	No
P04181	VAR_000569	P04181	154	3.07	P04181	154	100.00	No
P04181	VAR_000570	P04181	180	3.11	P04181	180	100.00	No
P04181	VAR_000579	P04181	319	4.95	P04181	319	100.00	No
P04264	VAR_017825	P04264	478	4.09	P08670	396	38.76	No
P04264	VAR_017826	P04264	478	4.09	P08670	396	38.76	No
P04264	VAR_017827	P04264	481	4.94	P08670	399	38.76	No
P04264	VAR_017828	P04264	485	3.39	P08670	403	38.76	No
P04275	VAR_005802	P04275	1374	3.59	P04275	1374	100.00	Yes
P04275	VAR_005803	P04275	1374	3.59	P04275	1374	100.00	Yes
P04424	VAR_000677	P04424	111	2.39	P11447	107	47.62	No
P04424	VAR_000678	P04424	193	2.22	P24058	195	71.77	No
P04424	VAR_000679	P04424	286	2.79	P04424	286	100.00	No
P04629	VAR_009630	P04629	649	4.10	P08069	1134	43.61	Yes
P04629	VAR_009631	P04629	654	4.05	Q07912	256	39.92	Yes
P04629	VAR_009632	P04629	674	2.30	P06213	1183	44.91	Yes
P04637	VAR_005880	P04637	137	2.41	P04637	137	100.00	Yes
P04637	VAR_005881	P04637	138	3.26	P04637	138	100.00	Yes
P04637	VAR_005923	P04637	172	3.34	P04637	172	100.00	Yes
P04637	VAR_005927	P04637	174	2.74	P04637	174	100.00	Yes
P04637	VAR_005928	P04637	175	3.89	P04637	175	100.00	Yes
P04637	VAR_005929	P04637	175	3.89	P04637	175	100.00	Yes
P04637	VAR_005930	P04637	175	3.89	P04637	175	100.00	Yes
P04637	VAR_005931	P04637	175	3.89	P04637	175	100.00	Yes

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P04637	VAR_005932	P04637	175	3.89	P04637	175	100.00	Yes
P04637	VAR_005933	P04637	176	5.61	P04637	176	100.00	Yes
P04637	VAR_005934	P04637	176	5.61	P04637	176	100.00	Yes
P04637	VAR_005935	P04637	177	4.12	P04637	177	100.00	No
P04637	VAR_005939	P04637	184	2.31	P04637	184	100.00	No
P04637	VAR_005943	P04637	189	2.04	P04637	189	100.00	Yes
P04637	VAR_005944	P04637	190	2.53	P02340	187	88.66	Yes
P04637	VAR_005952	P04637	198	3.54	P04637	198	100.00	Yes
P04637	VAR_005955	P04637	213	3.89	P04637	213	100.00	Yes
P04637	VAR_005965	P04637	237	4.78	P04637	237	100.00	Yes
P04637	VAR_005969	P04637	241	3.27	P04637	241	100.00	No
P04637	VAR_005970	P04637	242	5.61	P04637	242	100.00	No
P04637	VAR_005971	P04637	245	3.65	P02340	242	88.66	Yes
P04637	VAR_005972	P04637	245	3.65	P02340	242	88.66	Yes
P04637	VAR_005973	P04637	245	3.65	P02340	242	88.66	Yes
P04637	VAR_005974	P04637	245	3.65	P02340	242	88.66	Yes
P04637	VAR_005975	P04637	245	3.65	P02340	242	88.66	Yes
P04637	VAR_005980	P04637	247	4.00	P04637	247	100.00	No
P04637	VAR_005981	P04637	248	3.89	P04637	248	100.00	No
P04637	VAR_005982	P04637	248	3.89	P04637	248	100.00	No
P04637	VAR_005983	P04637	248	3.89	P04637	248	100.00	No
P04637	VAR_005984	P04637	248	3.89	P04637	248	100.00	No
P04637	VAR_005985	P04637	249	3.89	P04637	249	100.00	Yes
P04637	VAR_005986	P04637	249	3.89	P04637	249	100.00	Yes
P04637	VAR_006000	P04637	277	5.61	P04637	277	100.00	No
P04637	VAR_006007	P04637	280	3.89	P04637	280	100.00	No
P04637	VAR_006008	P04637	280	3.89	P04637	280	100.00	No
P04637	VAR_006009	P04637	280	3.89	P04637	280	100.00	No
P05164	VAR_015377	P05164	173	4.41	P05164	173	100.00	No
P05164	VAR_015378	P05164	251	3.39	P05164	251	100.00	No
P05165	VAR_009088	P05165	52	2.72	P24182	16	55.36	Yes
P05166	VAR_000274	P05166	165	3.66	Q8GBW6	143	52.71	No
P05166	VAR_000275	P05166	168	2.88	Q8GBW6	146	52.71	No
P05166	VAR_000278	P05166	410	2.90	Q9X4K7	392	57.77	No
P05166	VAR_000279	P05166	497	2.64	Q9X4K7	488	57.77	No
P05166	VAR_000281	P05166	519	2.12	Q8GBW6	503	52.71	No
P05166	VAR_009082	P05166	205	2.47	Q9X4K7	185	57.77	No
P05166	VAR_009086	P05166	536	2.68	Q9X4K7	527	57.77	No
P05166	VAR_023849	P05166	112	3.86	Q9X4K7	92	57.77	No
P05166	VAR_023851	P05166	165	3.66	Q8GBW6	143	52.71	No
P05166	VAR_023852	P05166	188	2.45	Q9X4K7	168	57.77	No
P05166	VAR_023856	P05166	435	4.05	Q9X4K7	417	57.77	No
P05166	VAR_023857	P05166	439	3.66	Q9X4K7	421	57.77	No
P05166	VAR_023858	P05166	468	2.95	Q9X4K7	450	57.77	No
P05186	VAR_006149	P05186	71	4.02	Q9BHT8	45	47.20	No
P05186	VAR_006150	P05186	71	4.02	Q9BHT8	45	47.20	No
P05186	VAR_011087	P05186	381	3.47	P00634	394	32.61	No
P05186	VAR_013975	P05186	71	4.02	Q9BHT8	45	47.20	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P05997	VAR_013588	P05997	960	3.86	P02452	136	42.11	No
P06213	VAR_015927	P06213	1158	4.10	P08069	1134	82.71	Yes
P06213	VAR_015928	P06213	1158	4.10	P08069	1134	82.71	Yes
P06400	VAR_005579	P06400	567	3.66	P06400	567	100.00	No
P06400	VAR_005581	P06400	654	2.94	P06400	654	100.00	No
P06400	VAR_005582	P06400	661	4.22	P06400	661	100.00	No
P06400	VAR_010049	P06400	530	3.97	P06400	530	100.00	No
P06400	VAR_010050	P06400	657	2.96	P06400	657	100.00	No
P06744	VAR_002528	P06744	342	3.27	P06744	342	100.00	No
P06744	VAR_002529	P06744	346	2.58	P06744	346	100.00	No
P06744	VAR_002530	P06744	346	2.58	P06744	346	100.00	No
P06744	VAR_002531	P06744	374	2.60	P06744	374	100.00	No
P06744	VAR_002532	P06744	388	5.37	P06744	388	100.00	No
P06744	VAR_002536	P06744	516	2.65	P06744	516	100.00	No
P06744	VAR_002537	P06744	524	2.99	P06744	524	100.00	No
P06744	VAR_002538	P06744	538	3.10	P06744	538	100.00	No
P06865	VAR_003203	P06865	39	2.01	P07686	72	38.46	Yes
P07195	VAR_004177	P07195	171	2.01	P07195	171	100.00	No
P07195	VAR_011634	P07195	68	2.07	P07195	68	100.00	No
P07195	VAR_011636	P07195	171	2.01	P07195	171	100.00	No
P07196	VAR_009703	P07196	331	2.92	P08670	342	53.25	No
P07202	VAR_006060	P07202	453	3.61	P05164	462	47.52	No
P07202	VAR_021623	P07202	240	3.95	P05164	262	47.52	No
P07202	VAR_021625	P07202	326	2.35	P05164	339	47.52	No
P07202	VAR_021629	P07202	493	3.30	P05164	501	47.52	No
P07202	VAR_021632	P07202	660	4.30	P05164	668	47.52	No
P07320	VAR_010733	P07320	14	2.66	P08209	14	87.34	No
P07320	VAR_021145	P07320	23	3.23	P62697	129	39.24	No
P07477	VAR_011656	P07477	139	4.20	P00761	124	78.97	No
P07741	VAR_006747	P07741	64	2.02	P49435	67	47.41	No
P07902	VAR_002553	P07902	51	3.44	P09148	31	52.30	No
P07902	VAR_002554	P07902	55	2.49	P09148	35	52.30	No
P07902	VAR_002559	P07902	97	3.15	P09148	77	52.30	No
P07902	VAR_002560	P07902	98	3.15	P09148	78	52.30	No
P07902	VAR_002563	P07902	117	3.46	P09148	97	52.30	No
P07902	VAR_002564	P07902	118	2.31	P09148	98	52.30	No
P07902	VAR_002583	P07902	171	3.78	P09148	151	52.30	No
P07902	VAR_002584	P07902	179	3.23	P09148	159	52.30	No
P07902	VAR_002585	P07902	183	2.41	P09148	163	52.30	No
P07902	VAR_002589	P07902	194	3.47	P09148	174	52.30	No
P07902	VAR_002594	P07902	201	2.00	P09148	181	60.36	No
P07902	VAR_002596	P07902	209	4.60	P09148	189	60.36	No
P07902	VAR_002597	P07902	209	4.60	P09148	189	60.36	No
P07902	VAR_002599	P07902	217	2.50	P09148	197	60.36	No
P07902	VAR_002601	P07902	231	3.48	P09148	211	60.36	No
P07902	VAR_002602	P07902	249	6.28	P09148	229	60.36	No
P07902	VAR_002618	P07902	323	3.43	P09148	300	60.36	No
P07902	VAR_002619	P07902	323	3.43	P09148	300	60.36	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P07902	VAR_008042	P07902	45	2.27	P09148	25	52.30	No
P07902	VAR_023328	P07902	51	3.44	P09148	31	52.30	No
P07949	VAR_006338	P07949	873	4.10	P08069	1134	41.57	Yes
P07949	VAR_006342	P07949	918	4.05	Q06187	563	35.74	Yes
P07949	VAR_006345	P07949	946	3.01	Q07912	325	37.01	Yes
P07949	VAR_006347	P07949	973	4.13	P11362	722	55.31	No
P07951	VAR_013468	P07951	117	2.08	P42639	117	85.59	Yes
P07951	VAR_013469	P07951	147	4.37	P42639	147	85.59	No
P07951	VAR_016086	P07951	91	3.43	P42639	91	85.59	No
P07954	VAR_002447	P07954	312	2.47	Q9LCC6	265	46.34	No
P07954	VAR_013501	P07954	233	3.61	P05042	186	60.79	No
P08123	VAR_001862	P08123	433	3.86	P02452	151	42.11	No
P08123	VAR_001866	P08123	547	3.86	P02452	145	40.35	No
P08123	VAR_001874	P08123	670	3.86	P02452	148	42.11	No
P08123	VAR_001878	P08123	730	3.86	P02452	145	40.35	No
P08123	VAR_001879	P08123	736	3.86	P02452	151	40.35	No
P08123	VAR_001884	P08123	778	3.76	P02452	133	47.37	No
P08123	VAR_001885	P08123	784	3.86	P02452	139	47.37	No
P08123	VAR_001886	P08123	787	3.86	P02452	142	47.37	No
P08123	VAR_001887	P08123	790	3.86	P02452	145	47.37	No
P08123	VAR_001888	P08123	796	3.86	P02452	151	47.37	No
P08123	VAR_001900	P08123	1078	3.76	P02452	133	45.61	Yes
P08123	VAR_001901	P08123	1096	3.86	P02452	151	45.61	Yes
P08123	VAR_008120	P08123	973	3.86	P02452	148	38.60	No
P08185	VAR_016223	P08185	389	2.51	P01011	405	47.98	No
P08237	VAR_006063	P08237	38	3.77	P00512	25	48.00	No
P08237	VAR_006064	P08237	38	3.77	P00512	25	48.00	No
P08237	VAR_006067	P08237	542	3.07	P00512	140	35.96	No
P08246	VAR_009538	P08246	32	3.08	P00747	583	38.31	No
P08246	VAR_009539	P08246	177	2.19	P00747	724	38.31	Yes
P08519	VAR_006633	P08519	4193	4.61	P00747	173	58.11	Yes
P08559	VAR_004952	P08559	167	2.36	P08559	167	100.00	No
P08559	VAR_004954	P08559	205	3.02	P08559	205	100.00	Yes
P08559	VAR_004957	P08559	231	3.70	P08559	231	100.00	No
P08581	VAR_006290	P08581	1228	2.30	P06213	1183	41.18	Yes
P08581	VAR_006291	P08581	1228	2.30	P06213	1183	41.18	Yes
P08581	VAR_006292	P08581	1230	3.29	P06213	1185	41.18	Yes
P08581	VAR_006293	P08581	1230	3.29	P06213	1185	41.18	Yes
P08581	VAR_006294	P08581	1250	4.05	Q06187	563	35.34	Yes
P08603	VAR_019406	P08603	959	5.88	P08174	253	37.74	No
P08603	VAR_025865	P08603	630	5.88	P08174	225	37.74	Yes
P08603	VAR_025868	P08603	951	2.61	P08174	245	37.74	No
P08603	VAR_025876	P08603	1142	3.98	P68638	180	38.46	Yes
P08603	VAR_025877	P08603	1157	6.30	P20023	75	36.54	Yes
P08709	VAR_006506	P08709	238	5.53	P00763	48	42.99	No
P08709	VAR_006508	P08709	304	3.00	P00761	94	41.12	No
P08709	VAR_006509	P08709	307	2.12	P00760	109	42.06	Yes
P08709	VAR_006516	P08709	402	3.76	P00763	198	42.99	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P08709	VAR_006517	P08709	402	3.76	P00763	198	42.99	No
P08709	VAR_014407	P08709	121	5.88	P00740	108	61.29	No
P08709	VAR_014416	P08709	304	3.00	P00761	94	41.12	No
P08709	VAR_014417	P08709	307	2.12	P00760	109	42.06	Yes
P08709	VAR_014419	P08709	391	2.73	P00747	747	36.57	Yes
P08709	VAR_014420	P08709	435	2.93	P00763	227	42.99	No
P08709	VAR_015141	P08709	312	2.32	P00761	102	41.12	No
P08709	VAR_015143	P08709	363	2.71	P00761	149	41.12	No
P08709	VAR_015144	P08709	403	3.74	P00763	199	42.99	No
P08779	VAR_017067	P08779	353	2.41	P08670	336	34.53	No
P09417	VAR_006965	P09417	145	2.70	P11348	142	96.84	No
P09493	VAR_007601	P09493	175	2.93	P42639	175	98.73	No
P09622	VAR_006908	P09622	488	4.27	P09624	479	77.98	No
P10153	VAR_013150	P10153	156	5.45	P61823	145	39.47	No
P10275	VAR_004685	P10275	608	4.26	P03372	234	53.33	No
P10275	VAR_009746	P10275	601	5.86	P15207	584	100.00	No
P10275	VAR_009747	P10275	604	3.09	P34021	309	42.67	Yes
P10275	VAR_009749	P10275	611	5.86	P06536	492	76.00	No
P10275	VAR_009783	P10275	720	3.13	P10275	720	100.00	No
P10275	VAR_009788	P10275	725	4.05	P10275	725	100.00	No
P10275	VAR_009792	P10275	733	3.70	P10275	733	100.00	No
P10619	VAR_001386	P10619	65	5.35	Q8W4X3	97	30.87	No
P10619	VAR_001389	P10619	395	4.19	Q8W4X3	409	30.87	No
P10721	VAR_004107	P10721	791	4.10	P08069	1134	35.71	Yes
P10721	VAR_004109	P10721	816	2.30	P06213	1183	39.10	Yes
P10721	VAR_023828	P10721	816	2.30	P06213	1183	39.10	Yes
P11177	VAR_021057	P11177	132	2.52	P11177	132	100.00	No
P11217	VAR_014004	P11217	291	3.32	P00490	267	47.36	No
P11230	VAR_000287	P11230	285	3.28	Q6S3I0	281	60.53	No
P11230	VAR_000288	P11230	289	2.93	Q6S3I0	285	60.53	No
P11362	VAR_017890	P11362	666	6.26	Q06187	562	36.55	Yes
P11362	VAR_017891	P11362	719	3.75	P00519	458	41.83	Yes
P11413	VAR_002470	P11413	175	3.95	P11413	175	100.00	Yes
P11413	VAR_002476	P11413	211	2.66	P11413	211	100.00	No
P11413	VAR_002477	P11413	212	2.72	P11413	212	100.00	No
P11413	VAR_002478	P11413	215	4.53	P11413	215	100.00	No
P11413	VAR_002479	P11413	226	3.21	P11413	226	100.00	Yes
P11413	VAR_002480	P11413	226	3.21	P11413	226	100.00	Yes
P11413	VAR_002482	P11413	256	3.89	P11413	256	100.00	Yes
P11413	VAR_002483	P11413	273	3.59	P11413	273	100.00	Yes
P11413	VAR_002484	P11413	277	2.74	P11413	277	100.00	No
P11413	VAR_002503	P11413	409	3.54	P11413	409	100.00	No
P11413	VAR_002504	P11413	409	3.54	P11413	409	100.00	No
P11413	VAR_002506	P11413	438	3.55	P11413	438	100.00	No
P11413	VAR_002514	P11413	462	2.34	P11413	462	100.00	Yes
P11473	VAR_004662	P11473	73	4.26	P03372	234	46.67	No
P11473	VAR_004667	P11473	391	2.95	Q13133	415	39.44	No
P11488	VAR_009279	P11488	37	3.58	P63096	41	67.44	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P11498	VAR_015200	P11498	451	2.07	P24182	410	31.43	No
P12107	VAR_013583	P12107	625	3.86	P02452	148	42.11	No
P12107	VAR_013584	P12107	676	3.86	P02452	139	52.63	No
P12107	VAR_013587	P12107	1516	3.86	P02452	136	42.11	Yes
P12694	VAR_004969	P12694	190	2.44	P12694	190	100.00	No
P12694	VAR_015101	P12694	290	3.12	P84129	227	37.04	No
P12883	VAR_004573	P12883	403	2.33	P10587	405	51.93	No
P12883	VAR_004574	P12883	403	2.33	P10587	405	51.93	No
P12883	VAR_004586	P12883	731	2.19	P13538	733	81.80	No
P12883	VAR_014199	P12883	743	2.52	P10587	753	51.93	Yes
P12883	VAR_017747	P12883	532	2.27	P13538	534	81.80	No
P12883	VAR_020803	P12883	320	2.88	P10587	322	51.93	Yes
P13569	VAR_000167	P13569	504	2.89	P26361	504	85.96	No
P13569	VAR_000176	P13569	549	3.70	P13569	549	100.00	No
P13569	VAR_000177	P13569	549	3.70	P13569	549	100.00	No
P13569	VAR_000178	P13569	549	3.70	P13569	549	100.00	No
P13569	VAR_000197	P13569	579	3.99	P13569	579	100.00	No
P13569	VAR_000200	P13569	613	2.15	P13569	613	100.00	No
P13569	VAR_000201	P13569	614	3.21	P13569	614	100.00	No
P13569	VAR_000261	P13569	1282	2.52	Q9CHL8	421	31.67	Yes
P13569	VAR_000262	P13569	1283	2.74	Q9CHL8	422	31.67	Yes
P13569	VAR_000264	P13569	1291	4.20	Q9CHL8	430	31.67	Yes
P13569	VAR_000265	P13569	1291	4.20	Q9CHL8	430	31.67	Yes
P13569	VAR_000266	P13569	1303	3.39	Q9CHL8	442	31.67	Yes
P13569	VAR_000267	P13569	1303	3.39	Q9CHL8	442	31.67	Yes
P13645	VAR_003833	P13645	442	3.26	P08670	392	36.16	No
P13645	VAR_010510	P13645	439	3.93	P08670	389	36.16	No
P13645	VAR_010511	P13645	446	4.09	P08670	396	36.16	No
P13647	VAR_003876	P13647	463	3.26	P08670	392	37.13	No
P13647	VAR_010466	P13647	467	4.09	P08670	396	37.13	No
P13647	VAR_023726	P13647	404	2.34	P08670	333	37.13	No
P13716	VAR_003635	P13716	240	2.28	P13716	240	100.00	No
P13804	VAR_002368	P13804	266	3.56	P13804	266	100.00	No
P13942	VAR_010655	P13942	808	3.76	P02452	133	40.35	No
P14136	VAR_017475	P14136	362	3.97	P08670	395	63.96	No
P14770	VAR_024997	P14770	24	5.79	P07359	24	34.62	No
P15153	VAR_017452	P15153	57	4.23	P15153	57	100.00	No
P15735	VAR_009518	P15735	189	3.47	P05132	200	33.06	No
P15735	VAR_009518	P15735	189	3.47	P00517	200	33.06	No
P15735	VAR_020854	P15735	157	2.62	P49137	190	35.77	No
P16144	VAR_011297	P16144	336	2.73	P05106	347	35.37	No
P16219	VAR_000316	P16219	383	3.62	P15651	383	94.63	No
P17661	VAR_007902	P17661	392	3.13	P08670	387	73.38	No
P17661	VAR_009189	P17661	344	2.79	P08670	339	73.38	No
P17661	VAR_018771	P17661	384	2.43	P08670	379	73.38	No
P17661	VAR_018772	P17661	388	3.89	P08670	383	73.38	Yes
P19013	VAR_016038	P19013	449	3.97	P08670	395	38.76	No
P19429	VAR_016078	P19429	81	2.49	P19429	81	100.00	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P19438	VAR_013410	P19438	59	5.87	P19438	102	33.33	No
P19438	VAR_013411	P19438	62	5.87	P19438	62	100.00	No
P19438	VAR_019302	P19438	59	5.87	P19438	102	33.33	No
P19438	VAR_019303	P19438	62	5.87	P19438	62	100.00	No
P19438	VAR_019304	P19438	99	5.87	P19438	99	100.00	Yes
P19438	VAR_019329	P19438	51	2.26	P19438	91	33.33	No
P20594	VAR_022584	P20594	115	3.94	P18910	128	41.89	No
P20807	VAR_001367	P20807	490	3.92	Q07009	416	56.86	No
P20807	VAR_009560	P20807	214	3.81	Q07009	189	55.44	Yes
P20807	VAR_009561	P20807	215	2.37	Q07009	190	55.44	Yes
P20807	VAR_009574	P20807	440	3.45	Q07009	366	56.86	No
P20807	VAR_009584	P20807	490	3.92	Q07009	416	56.86	No
P20807	VAR_009589	P20807	567	2.66	Q07009	494	56.86	No
P20807	VAR_009595	P20807	705	4.21	Q64537	154	57.14	No
P20807	VAR_009596	P20807	705	4.21	Q64537	154	57.14	No
P20823	VAR_003759	P20823	272	4.16	P40424	288	34.62	No
P20823	VAR_010537	P20823	12	2.61	P22361	12	94.86	No
P20823	VAR_010553	P20823	200	3.27	P06601	214	36.54	No
P20823	VAR_010556	P20823	229	3.65	P06601	243	36.54	No
P20823	VAR_010563	P20823	272	4.16	P40424	288	34.62	No
P20823	VAR_012483	P20823	20	3.33	P22361	20	94.86	No
P20933	VAR_005069	P20933	60	2.94	P20933	60	100.00	No
P20933	VAR_005071	P20933	101	2.95	P20933	101	100.00	No
P20933	VAR_005075	P20933	306	2.91	Q47898	304	37.41	No
P20933	VAR_015429	P20933	135	3.73	P20933	135	100.00	No
P20933	VAR_015432	P20933	257	4.01	P20933	257	100.00	No
P21439	VAR_023504	P21439	1161	3.24	Q9CHL8	473	46.15	Yes
P21953	VAR_004974	P21953	206	2.83	P84130	139	53.11	No
P22033	VAR_004416	P22033	368	2.01	P11653	345	63.48	No
P22033	VAR_004417	P22033	369	4.22	P11653	346	63.48	No
P22830	VAR_002385	P22830	267	2.22	P22830	267	100.00	No
P23760	VAR_003804	P23760	238	4.46	P40424	252	35.71	No
P23760	VAR_003805	P23760	265	2.61	P02836	500	37.50	No
P23760	VAR_003806	P23760	271	4.16	P40424	288	35.71	No
P23760	VAR_017537	P23760	271	4.16	P40424	288	35.71	No
P23760	VAR_017538	P23760	271	4.16	P40424	288	35.71	No
P24752	VAR_007500	P24752	158	2.24	P07097	120	43.14	No
P24752	VAR_007501	P24752	183	3.07	P07097	146	43.14	No
P25054	VAR_005040	P25054	1027	4.37	P25054	1027	100.00	Yes
P25054	VAR_005044	P25054	1176	4.29	P25054	1024	53.33	Yes
P26367	VAR_003812	P26367	44	3.43	Q02548	56	77.42	Yes
P26440	VAR_015966	P26440	411	2.65	P26440	411	100.00	No
P28069	VAR_003778	P28069	143	4.06	P14859	299	58.11	Yes
P28358	VAR_022582	P28358	319	2.82	Q6B2C0	185	37.04	Yes
P28360	VAR_003754	P28360	196	3.65	P06601	243	42.86	No
P29400	VAR_001915	P29400	129	3.76	P02452	133	47.37	No
P29400	VAR_001916	P29400	129	3.76	P02452	133	47.37	No
P29400	VAR_001923	P29400	325	3.86	P02452	145	42.11	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P29400	VAR_001924	P29400	325	3.86	P02452	145	42.11	No
P29400	VAR_001929	P29400	383	3.76	P02452	133	42.86	No
P29400	VAR_001930	P29400	400	3.86	P02452	151	42.86	No
P29400	VAR_001939	P29400	521	3.86	P02452	136	38.60	No
P29400	VAR_001940	P29400	521	3.86	P02452	136	38.60	No
P29400	VAR_001942	P29400	609	3.86	P02452	151	43.86	No
P29400	VAR_001947	P29400	684	3.86	P02452	151	42.11	No
P29400	VAR_001950	P29400	796	3.86	P02452	142	47.37	No
P29400	VAR_001956	P29400	1104	3.86	P02452	142	40.35	Yes
P29400	VAR_001964	P29400	1421	3.76	P02452	133	43.86	Yes
P29400	VAR_001968	P29400	1517	4.30	Q7SIB2	61	58.23	Yes
P29400	VAR_001973	P29400	1649	2.97	Q7SIB2	193	58.02	Yes
P29400	VAR_001973	P29400	1649	2.97	P02462	1633	58.02	Yes
P29400	VAR_001974	P29400	1677	4.20	Q7SIB2	221	58.02	Yes
P29400	VAR_007992	P29400	331	3.86	P02452	151	42.11	No
P29400	VAR_008000	P29400	669	3.86	P02452	136	42.11	No
P29400	VAR_008008	P29400	1107	3.86	P02452	145	40.35	Yes
P29400	VAR_008009	P29400	1161	3.86	P02452	139	42.11	Yes
P29400	VAR_008011	P29400	1220	3.86	P02452	139	40.35	Yes
P29400	VAR_008012	P29400	1333	3.76	P02452	133	36.84	Yes
P29400	VAR_008013	P29400	1427	3.86	P02452	139	43.86	Yes
P29400	VAR_011221	P29400	192	3.86	P02452	136	54.39	No
P29400	VAR_011222	P29400	204	3.86	P02452	148	54.39	No
P29400	VAR_011229	P29400	319	3.86	P02452	139	42.11	No
P29400	VAR_011237	P29400	524	3.86	P02452	139	38.60	No
P29400	VAR_011241	P29400	603	3.86	P02452	145	43.86	No
P29400	VAR_011242	P29400	609	3.86	P02452	151	43.86	No
P29400	VAR_011249	P29400	681	3.86	P02452	148	42.11	No
P29400	VAR_011253	P29400	802	3.86	P02452	148	47.37	No
P29400	VAR_011269	P29400	1036	3.76	P02452	133	42.11	Yes
P29400	VAR_011270	P29400	1039	3.86	P02452	136	42.11	Yes
P29400	VAR_011271	P29400	1045	3.86	P02452	142	42.11	Yes
P29400	VAR_011275	P29400	1158	3.86	P02452	136	42.11	Yes
P29400	VAR_011276	P29400	1167	3.86	P02452	145	42.11	Yes
P29400	VAR_011277	P29400	1170	3.86	P02452	148	42.11	Yes
P29400	VAR_011281	P29400	1229	3.86	P02452	148	40.35	Yes
P29400	VAR_011290	P29400	1677	4.20	Q7SIB2	221	58.02	Yes
P29965	VAR_007524	P29965	227	2.84	P29965	227	100.00	No
P29965	VAR_017923	P29965	170	3.56	P29965	170	100.00	No
P29965	VAR_017927	P29965	174	4.04	P29965	174	100.00	No
P29965	VAR_017938	P29965	226	2.87	P29965	226	100.00	No
P30613	VAR_004042	P30613	337	4.23	P30613	337	100.00	No
P30613	VAR_004043	P30613	337	4.23	P30613	337	100.00	No
P30613	VAR_004044	P30613	339	4.19	P30613	339	100.00	No
P30613	VAR_004045	P30613	341	3.84	P30613	341	100.00	No
P30613	VAR_004052	P30613	384	4.05	P30613	384	100.00	No
P30613	VAR_004053	P30613	392	3.08	P30613	392	100.00	No
P30613	VAR_004054	P30613	393	4.41	P30613	393	100.00	No

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P30613	VAR_004055	P30613	393	4.41	P30613	393	100.00	No
P30613	VAR_004061	P30613	431	2.16	P30613	431	100.00	No
P30613	VAR_004075	P30613	559	2.22	P11974	515	59.17	Yes
P30613	VAR_004076	P30613	566	4.20	P30613	566	100.00	No
P30613	VAR_011445	P30613	222	3.84	P30613	222	100.00	No
P30613	VAR_011454	P30613	341	3.84	P30613	341	100.00	No
P30613	VAR_011455	P30613	342	2.93	P30613	342	100.00	No
P30613	VAR_011456	P30613	348	2.24	P30613	348	100.00	No
P30613	VAR_011459	P30613	376	3.70	P30613	376	100.00	No
P30613	VAR_011460	P30613	387	3.93	P30613	387	100.00	No
P30613	VAR_011461	P30613	390	4.19	P30613	390	100.00	No
P30613	VAR_011478	P30613	385	4.23	P30613	385	100.00	No
P30613	VAR_011480	P30613	479	2.41	P11974	435	59.17	Yes
P30613	VAR_011482	P30613	569	3.48	P30613	569	100.00	No
P30793	VAR_002638	P30793	134	2.33	P30793	134	100.00	No
P30793	VAR_002640	P30793	144	3.94	P30793	144	100.00	No
P30793	VAR_002644	P30793	186	3.10	P22288	177	97.12	No
P30793	VAR_002647	P30793	211	4.32	P22288	202	97.12	No
P30793	VAR_016896	P30793	135	2.72	P30793	135	100.00	No
P30793	VAR_016902	P30793	199	3.70	P22288	190	97.12	No
P30793	VAR_016903	P30793	211	4.32	P22288	202	97.12	No
P30793	VAR_016904	P30793	213	4.22	P30793	213	100.00	No
P31271	VAR_017775	P31271	371	4.16	P02836	503	37.50	No
P31271	VAR_017776	P31271	372	4.42	P02836	504	37.50	No
P33527	VAR_011489	P33527	671	3.57	P13569	451	42.69	No
P35240	VAR_000814	P35240	117	2.42	P11171	304	31.02	No
P35240	VAR_000825	P35240	535	2.05	P26038	517	38.76	No
P35240	VAR_000826	P35240	538	3.26	P26038	520	38.76	No
P35453	VAR_015953	P35453	314	2.38	P02836	500	33.93	No
P35520	VAR_002172	P35520	87	4.06	P35520	87	100.00	No
P35520	VAR_002176	P35520	130	2.28	P35520	130	100.00	Yes
P35520	VAR_008051	P35520	84	2.63	Q9WZD3	7	46.15	No
P35520	VAR_008064	P35520	151	2.48	Q9WZD3	66	46.15	No
P35520	VAR_021792	P35520	108	2.07	P35520	108	100.00	No
P35548	VAR_010201	P35548	172	3.65	P06601	243	46.43	No
P35555	VAR_002278	P35555	129	5.88	P00740	108	50.00	No
P35555	VAR_002291	P35555	723	4.22	P07204	441	42.42	No
P35555	VAR_002323	P35555	1249	5.87	Q9JJS8	152	37.50	Yes
P35555	VAR_002331	P35555	1893	2.77	P07204	443	44.83	Yes
P35555	VAR_002339	P35555	2258	5.87	Q9JJS8	152	40.63	Yes
P35555	VAR_017971	P35555	154	5.88	P05106	562	34.78	No
P35555	VAR_017974	P35555	560	3.48	P07204	469	45.16	Yes
P35555	VAR_017985	P35555	723	4.22	P07204	441	42.42	No
P35555	VAR_017986	P35555	734	5.87	Q9JJS8	152	38.24	No
P35555	VAR_017988	P35555	776	5.87	Q9JJS8	152	34.38	No
P35555	VAR_017989	P35555	776	5.87	Q9JJS8	152	34.38	No
P35555	VAR_017991	P35555	816	5.88	P09871	143	37.50	No
P35555	VAR_017995	P35555	921	5.87	Q9JJS8	152	30.77	No

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P35555	VAR_018007	P35555	1374	5.87	Q9JJS8	152	40.63	Yes
P35555	VAR_018019	P35555	1796	3.48	P07204	469	36.84	Yes
P35555	VAR_023865	P35555	541	5.87	Q9JJS8	152	34.38	No
P35555	VAR_023871	P35555	832	5.88	P08709	132	52.00	No
P35555	VAR_023873	P35555	1058	3.48	P07204	469	40.63	Yes
P35555	VAR_023881	P35555	1333	5.87	Q9JJS8	152	35.14	Yes
P35555	VAR_023884	P35555	1475	3.48	P07204	469	51.61	Yes
P35555	VAR_023885	P35555	1475	3.48	P07204	469	51.61	Yes
P35555	VAR_023895	P35555	1900	5.87	Q9JJS8	152	40.00	Yes
P35556	VAR_002350	P35556	1252	5.87	Q9JJS8	152	41.03	Yes
P35556	VAR_010741	P35556	1252	5.87	Q9JJS8	152	41.03	Yes
P35557	VAR_003698	P35557	175	2.67	P19367	179	48.04	No
P35557	VAR_003709	P35557	279	2.31	P05708	283	51.88	No
P35557	VAR_003711	P35557	300	3.58	P05708	304	51.88	No
P35557	VAR_003712	P35557	300	3.58	P05708	304	51.88	No
P35557	VAR_010586	P35557	108	4.34	P19367	560	55.61	No
P35557	VAR_010587	P35557	137	2.30	P19367	589	55.61	No
P35625	VAR_007509	P35625	191	2.22	P16035	200	45.88	No
P35908	VAR_009186	P35908	482	3.97	P08670	395	35.18	No
P35908	VAR_009187	P35908	485	3.45	P08670	398	35.18	No
P35908	VAR_010516	P35908	490	3.39	P08670	403	35.18	No
P35916	VAR_018413	P35916	1041	4.05	Q07912	256	35.04	Yes
P35916	VAR_018415	P35916	1114	4.23	Q06187	596	36.69	Yes
P35916	VAR_018416	P35916	1137	4.13	P11362	722	52.38	Yes
P36897	VAR_022344	P36897	200	3.76	P36897	200	100.00	No
P37173	VAR_022352	P37173	336	2.75	P36897	291	41.05	Yes
P37231	VAR_010728	P37231	495	2.39	Q07869	458	69.23	Yes
P38117	VAR_002369	P38117	163	3.92	P38117	164	100.00	No
P38117	VAR_025804	P38117	127	3.97	P38117	127	100.00	No
P38117	VAR_025804	P38117	127	3.97	P38117	128	100.00	No
P40337	VAR_005742	P40337	155	2.93	P40337	155	100.00	No
P40337	VAR_005743	P40337	156	4.15	P40337	156	100.00	No
P40337	VAR_005744	P40337	156	4.15	P40337	156	100.00	No
P40337	VAR_005746	P40337	157	3.07	P40337	157	100.00	No
P40337	VAR_005748	P40337	158	2.61	P40337	158	100.00	No
P40337	VAR_005749	P40337	158	2.61	P40337	158	100.00	No
P40337	VAR_008101	P40337	155	2.93	P40337	155	100.00	No
P40692	VAR_004438	P40692	64	2.54	P54278	71	37.11	No
P42771	VAR_001412	P42771	23	2.65	P42771	23	100.00	No
P42771	VAR_001440	P42771	74	2.61	Q60773	71	51.61	No
P42771	VAR_001441	P42771	74	2.61	Q60773	71	51.61	No
P42771	VAR_001453	P42771	89	2.65	P42771	89	100.00	No
P42771	VAR_001454	P42771	89	2.65	P42771	89	100.00	No
P43034	VAR_007724	P43034	148	4.99	P62871	53	33.33	No
P43034	VAR_015398	P43034	30	3.95	P63005	30	100.00	No
P43246	VAR_004488	P43246	834	3.36	P23909	779	48.91	No
P43403	VAR_015538	P43403	465	4.05	Q07912	256	38.98	Yes
P43681	VAR_000295	P43681	280	2.61	P02711	272	50.00	No

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P43681	VAR_017531	P43681	280	2.61	P02711	272	50.00	No
P43699	VAR_015189	P43699	213	4.16	P40424	288	32.14	No
P49748	VAR_000349	P49748	366	3.75	P15651	297	38.10	No
P49748	VAR_000350	P49748	366	3.75	P15651	297	38.10	No
P49748	VAR_000356	P49748	453	3.62	P15651	383	38.10	No
P49748	VAR_000357	P49748	454	3.23	P15651	384	38.10	No
P49748	VAR_000358	P49748	456	2.95	P15651	386	38.10	No
P49748	VAR_000359	P49748	459	2.02	P15651	389	38.10	No
P49748	VAR_000361	P49748	469	2.65	Q06319	374	36.49	No
P49748	VAR_000362	P49748	469	2.65	Q06319	374	36.49	No
P50219	VAR_017876	P50219	248	3.39	P02836	459	50.00	Yes
P50219	VAR_017879	P50219	292	4.16	P02836	503	50.00	No
P50219	VAR_017881	P50219	295	4.16	P40424	288	35.71	No
P50219	VAR_017882	P50219	295	4.16	P40424	288	35.71	No
P51149	VAR_018722	P51149	129	2.78	P62825	126	32.91	No
P51149	VAR_018723	P51149	162	3.47	P62826	156	32.91	No
P51149	VAR_018723	P51149	162	3.47	P62826	157	32.91	No
P51159	VAR_010654	P51159	73	5.75	P63012	76	46.88	No
P51159	VAR_011335	P51159	152	3.49	P01112	134	33.96	Yes
P51587	VAR_020718	P51587	1524	4.41	P51587	1524	100.00	Yes
P51587	VAR_020725	P51587	2072	2.87	P51587	1538	44.12	Yes
P51812	VAR_006196	P51812	431	3.87	P05132	52	33.75	No
P52333	VAR_010498	P52333	910	3.26	Q06187	481	32.26	Yes
P52952	VAR_003752	P52952	178	2.30	P02836	494	48.21	Yes
P52952	VAR_010117	P52952	188	4.42	P02836	504	48.21	No
P53634	VAR_009541	P53634	249	3.15	P53634	249	100.00	No
P53634	VAR_009542	P53634	252	4.57	P53634	252	100.00	No
P53634	VAR_009544	P53634	301	3.78	P07711	181	38.54	No
P53634	VAR_016936	P53634	429	6.32	P53634	429	100.00	No
P53634	VAR_019038	P53634	236	4.19	P53634	236	100.00	No
P53634	VAR_019041	P53634	300	3.79	O46427	183	42.79	No
P53634	VAR_019042	P53634	300	3.79	O46427	183	42.79	No
P53634	VAR_019043	P53634	301	3.78	P07711	181	38.54	No
P53634	VAR_019046	P53634	319	3.20	P07711	199	38.54	No
P53634	VAR_019047	P53634	412	4.46	P53634	412	100.00	No
P55084	VAR_021130	P55084	118	4.26	P28790	73	37.50	No
P55084	VAR_021131	P55084	122	2.70	P28790	77	37.50	No
P55084	VAR_021132	P55084	134	3.58	P28790	89	37.50	No
P57727	VAR_011678	P57727	251	5.08	P00761	42	40.38	No
P57727	VAR_011679	P57727	404	3.88	P07338	216	41.28	No
P57727	VAR_013495	P57727	407	4.23	P07338	219	41.28	No
P58304	VAR_011618	P58304	200	4.16	P40424	288	32.14	No
P58304	VAR_011619	P58304	200	4.16	P40424	288	32.14	No
P60174	VAR_007535	P60174	72	3.80	P00939	72	98.32	No
P60174	VAR_007539	P60174	170	3.99	P04789	172	53.59	No
P61457	VAR_005530	P61457	96	2.39	P61459	96	100.00	No
P61626	VAR_004281	P61626	85	2.78	P61626	85	100.00	No
P62070	VAR_006848	P62070	72	4.56	P01112	61	61.88	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P63092	VAR_003441	P63092	201	4.27	P63096	177	41.62	No
P63092	VAR_003442	P63092	201	4.27	P63096	177	41.62	No
P63092	VAR_003443	P63092	227	4.62	P10824	203	41.91	No
P63092	VAR_017844	P63092	201	4.27	P63096	177	41.62	No
P63092	VAR_017845	P63092	201	4.27	P63096	177	41.62	No
P63092	VAR_017846	P63092	201	4.27	P63096	177	41.62	No
P63092	VAR_017847	P63092	227	4.62	P10824	203	41.91	No
P63092	VAR_017848	P63092	231	4.27	P04896	231	99.74	No
P68032	VAR_012857	P68032	101	2.27	P68135	101	98.93	No
P68032	VAR_012861	P68032	333	2.90	P68139	333	98.93	No
P68032	VAR_012862	P68032	363	3.71	P68135	363	98.93	Yes
P68133	VAR_011682	P68133	117	2.62	P68135	117	100.00	No
P68133	VAR_015579	P68133	42	2.20	P68135	42	100.00	No
P68133	VAR_015583	P68133	258	2.20	P68135	258	100.00	No
P68133	VAR_015586	P68133	288	3.61	P68135	288	100.00	No
P68133	VAR_015587	P68133	359	2.63	P68135	359	100.00	Yes
P68871	VAR_002878	P68871	15	5.78	P02118	15	69.40	Yes
P68871	VAR_002879	P68871	15	5.78	P02118	15	69.40	Yes
P68871	VAR_002882	P68871	17	2.30	P68871	17	100.00	No
P68871	VAR_002883	P68871	17	2.30	P68871	17	100.00	No
P68871	VAR_002884	P68871	17	2.30	P68871	17	100.00	No
P68871	VAR_002885	P68871	18	2.32	P02118	18	69.40	Yes
P68871	VAR_002886	P68871	19	2.34	P02118	19	69.40	Yes
P68871	VAR_002887	P68871	19	2.34	P02118	19	69.40	Yes
P68871	VAR_002888	P68871	19	2.34	P02118	19	69.40	Yes
P68871	VAR_002907	P68871	26	2.10	P68871	26	100.00	No
P68871	VAR_002908	P68871	26	2.10	P68871	26	100.00	No
P68871	VAR_002914	P68871	30	2.86	P68871	30	100.00	No
P68871	VAR_002919	P68871	35	2.99	P68871	35	100.00	No
P68871	VAR_002920	P68871	36	3.98	P68871	36	100.00	No
P68871	VAR_002921	P68871	36	3.98	P68871	36	100.00	No
P68871	VAR_002922	P68871	36	3.98	P68871	36	100.00	No
P68871	VAR_002923	P68871	37	3.09	P68871	37	100.00	No
P68871	VAR_002924	P68871	37	3.09	P68871	37	100.00	No
P68871	VAR_002925	P68871	37	3.09	P68871	37	100.00	No
P68871	VAR_002943	P68871	52	2.02	P02118	52	69.40	No
P68871	VAR_002944	P68871	52	2.02	P02118	52	69.40	No
P68871	VAR_002961	P68871	66	2.40	P02089	66	79.85	No
P68871	VAR_002962	P68871	67	3.12	P02089	67	79.85	No
P68871	VAR_002963	P68871	67	3.12	P02089	67	79.85	No
P68871	VAR_002969	P68871	70	2.18	P02089	70	79.85	No
P68871	VAR_002979	P68871	77	3.17	P02089	77	79.85	No
P68871	VAR_002980	P68871	77	3.17	P02089	77	79.85	No
P68871	VAR_002981	P68871	77	3.17	P02089	77	79.85	No
P68871	VAR_002982	P68871	78	2.52	P02089	78	79.85	No
P68871	VAR_002983	P68871	79	2.25	P02089	79	79.85	No
P68871	VAR_002993	P68871	88	2.86	P68871	88	100.00	No
P68871	VAR_002994	P68871	88	2.86	P68871	88	100.00	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P68871	VAR_003001	P68871	92	5.44	P02089	92	79.85	No
P68871	VAR_003002	P68871	92	5.44	P02089	92	79.85	No
P68871	VAR_003003	P68871	92	5.44	P02089	92	79.85	No
P68871	VAR_003004	P68871	92	5.44	P02089	92	79.85	No
P68871	VAR_003013	P68871	97	3.47	P68871	97	100.00	No
P68871	VAR_003014	P68871	97	3.47	P68871	97	100.00	No
P68871	VAR_003015	P68871	97	3.47	P68871	97	100.00	No
P68871	VAR_003016	P68871	97	3.47	P68871	97	100.00	No
P68871	VAR_003017	P68871	98	2.88	P68871	98	100.00	No
P68871	VAR_003018	P68871	99	2.45	P68871	99	100.00	No
P68871	VAR_003019	P68871	100	3.08	P68871	100	100.00	No
P68871	VAR_003020	P68871	100	3.08	P68871	100	100.00	No
P68871	VAR_003025	P68871	102	2.60	P68871	102	100.00	No
P68871	VAR_003026	P68871	102	2.60	P68871	102	100.00	No
P68871	VAR_003040	P68871	117	3.30	P68871	117	100.00	No
P68871	VAR_003041	P68871	117	3.30	P68871	117	100.00	No
P68871	VAR_003042	P68871	119	2.23	P68871	119	100.00	No
P68871	VAR_003051	P68871	123	2.16	P68871	123	100.00	No
P68871	VAR_003052	P68871	124	2.47	P68871	124	100.00	No
P68871	VAR_003053	P68871	124	2.47	P68871	124	100.00	No
P68871	VAR_003054	P68871	124	2.47	P68871	124	100.00	No
P68871	VAR_003058	P68871	127	2.16	P68871	127	100.00	No
P68871	VAR_003059	P68871	127	2.16	P68871	127	100.00	No
P68871	VAR_003060	P68871	128	2.12	P68871	128	100.00	No
P68871	VAR_003069	P68871	132	2.60	P68871	132	100.00	No
P68871	VAR_003070	P68871	132	2.60	P68871	132	100.00	No
P68871	VAR_010144	P68871	114	2.56	P02089	114	79.85	No
P68871	VAR_010145	P68871	114	2.56	P02089	114	79.85	No
P68871	VAR_025399	P68871	117	3.30	P68871	117	100.00	No
P69891	VAR_003141	P69891	36	3.98	P02070	35	74.63	No
P69891	VAR_003141	P69891	36	3.98	P68871	36	74.63	No
P69891	VAR_003142	P69891	37	3.09	P02070	36	74.63	No
P69891	VAR_003142	P69891	37	3.09	P68871	37	74.63	No
P69891	VAR_003163	P69891	79	2.25	P02089	79	70.15	No
P69891	VAR_003168	P69891	97	3.47	P02070	96	74.63	No
P69891	VAR_003168	P69891	97	3.47	P68871	97	74.63	No
P69891	VAR_003175	P69891	128	2.12	P69891	128	100.00	No
P69892	VAR_003131	P69892	15	5.78	P02118	15	73.13	Yes
P69892	VAR_003139	P69892	26	2.10	P68871	26	75.37	No
P69892	VAR_003156	P69892	66	2.40	P02089	66	70.90	No
P69892	VAR_003157	P69892	66	2.40	P02089	66	70.90	No
P69892	VAR_003162	P69892	77	3.17	P02089	77	70.90	No
P69892	VAR_003166	P69892	92	5.44	P02089	92	70.90	No
P69892	VAR_003171	P69892	117	3.30	P68871	117	75.37	No
P69892	VAR_003174	P69892	125	2.06	P69891	125	99.25	No
P69892	VAR_020646	P69892	17	2.30	P68871	17	75.37	No
P69892	VAR_020647	P69892	19	2.34	P02118	19	73.13	Yes
P69892	VAR_020651	P69892	75	2.38	P02089	75	70.90	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P69905	VAR_002729	P69905	11	2.28	P02118	12	41.41	Yes
P69905	VAR_002731	P69905	14	5.78	P02118	15	41.41	Yes
P69905	VAR_002733	P69905	16	2.30	P01990	16	67.69	Yes
P69905	VAR_002734	P69905	16	2.30	P01990	16	67.69	Yes
P69905	VAR_002739	P69905	20	2.28	P02118	19	41.41	Yes
P69905	VAR_002740	P69905	20	2.28	P02118	19	41.41	Yes
P69905	VAR_002748	P69905	27	2.10	P01958	27	87.69	No
P69905	VAR_002749	P69905	27	2.10	P01958	27	87.69	No
P69905	VAR_002750	P69905	27	2.10	P01958	27	87.69	No
P69905	VAR_002752	P69905	31	2.86	P69905	31	100.00	No
P69905	VAR_002754	P69905	37	3.98	P01965	37	83.85	No
P69905	VAR_002756	P69905	40	2.44	P02074	38	46.09	No
P69905	VAR_002756	P69905	40	2.44	P02070	38	46.09	No
P69905	VAR_002759	P69905	44	2.17	P69905	44	100.00	No
P69905	VAR_002760	P69905	44	2.17	P69905	44	100.00	No
P69905	VAR_002761	P69905	45	3.01	P02208	54	30.77	Yes
P69905	VAR_002762	P69905	45	3.01	P02208	54	30.77	Yes
P69905	VAR_002763	P69905	47	2.60	P02208	57	30.77	Yes
P69905	VAR_002764	P69905	47	2.60	P02208	57	30.77	Yes
P69905	VAR_002765	P69905	47	2.60	P02208	57	30.77	Yes
P69905	VAR_002766	P69905	47	2.60	P02208	57	30.77	Yes
P69905	VAR_002774	P69905	56	2.63	P02208	71	30.77	No
P69905	VAR_002775	P69905	56	2.63	P02208	71	30.77	No
P69905	VAR_002779	P69905	59	2.52	P02089	64	42.19	No
P69905	VAR_002782	P69905	61	2.40	P02089	66	42.19	No
P69905	VAR_002783	P69905	61	2.40	P02089	66	42.19	No
P69905	VAR_002784	P69905	62	3.12	P02089	67	42.19	No
P69905	VAR_002790	P69905	72	3.17	P02089	77	42.19	No
P69905	VAR_002791	P69905	74	2.25	P02089	79	42.19	No
P69905	VAR_002792	P69905	74	2.25	P02089	79	42.19	No
P69905	VAR_002793	P69905	74	2.25	P02089	79	42.19	No
P69905	VAR_002801	P69905	80	2.20	P02089	85	42.19	No
P69905	VAR_002808	P69905	87	5.44	P02089	92	42.19	No
P69905	VAR_002809	P69905	87	5.44	P02089	92	42.19	No
P69905	VAR_002814	P69905	94	2.45	P69905	94	100.00	No
P69905	VAR_002815	P69905	95	3.08	P69905	95	100.00	No
P69905	VAR_002816	P69905	95	3.08	P69905	95	100.00	No
P69905	VAR_002817	P69905	97	2.60	P69905	97	100.00	No
P69905	VAR_002821	P69905	109	2.56	P02089	114	42.19	No
P69905	VAR_002823	P69905	112	3.30	P01958	112	87.69	No
P69905	VAR_002823	P69905	112	3.30	P01966	112	87.69	No
P69905	VAR_002825	P69905	114	2.07	P69905	114	100.00	No
P69905	VAR_002826	P69905	114	2.07	P69905	114	100.00	No
P69905	VAR_002827	P69905	114	2.07	P69905	114	100.00	No
P69905	VAR_002835	P69905	122	2.46	P69905	122	100.00	No
P69905	VAR_002837	P69905	126	2.03	P69905	126	100.00	No
P69905	VAR_002838	P69905	126	2.03	P69905	126	100.00	No
P69905	VAR_002839	P69905	127	2.60	P01958	127	87.69	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
P69905	VAR_025002	P69905	31	2.86	P69905	31	100.00	No
P69905	VAR_025389	P69905	94	2.45	P69905	94	100.00	No
P69905	VAR_025392	P69905	126	2.03	P69905	126	100.00	No
P78363	VAR_008430	P78363	965	3.62	Q9YGA6	38	32.72	No
P78363	VAR_008431	P78363	978	3.11	Q9YGA6	51	32.72	No
P78363	VAR_008436	P78363	1087	3.85	Q9YGA6	165	32.72	Yes
P78363	VAR_012547	P78363	971	3.73	Q9YGA6	44	32.72	No
P78363	VAR_012558	P78363	1063	3.70	Q9YGA6	141	32.72	Yes
P78363	VAR_012559	P78363	1087	3.85	Q9YGA6	165	32.72	Yes
P78385	VAR_023052	P78385	407	3.97	P08670	395	37.66	No
P78504	VAR_013203	P78504	386	2.12	P00740	105	41.94	No
P80365	VAR_015639	P80365	227	2.70	P19992	147	30.52	No
P80365	VAR_015640	P80365	237	2.81	P19992	157	30.52	No
P80365	VAR_015642	P80365	250	2.52	P19992	170	30.52	No
P80404	VAR_008883	P80404	220	3.11	P80147	220	95.95	No
P82279	VAR_011642	P82279	250	5.88	P08709	132	48.39	No
P82279	VAR_022943	P82279	195	5.88	P09871	143	41.38	No
P82279	VAR_022946	P82279	383	5.88	P08709	130	41.94	No
P82279	VAR_022954	P82279	681	5.88	P09871	143	31.03	No
P82279	VAR_022966	P82279	894	2.65	P00740	100	46.67	No
P82279	VAR_022977	P82279	1205	3.29	P09871	161	34.48	Yes
P82279	VAR_022980	P82279	1321	5.88	P08709	130	51.61	Yes
P98172	VAR_023131	P98172	111	2.41	P52800	114	61.15	No
P98172	VAR_023132	P98172	115	4.26	P52800	118	61.15	No
P98172	VAR_023133	P98172	119	4.10	P52800	122	61.15	No
P98172	VAR_023134	P98172	119	4.10	P52800	122	61.15	No
P98172	VAR_023135	P98172	119	4.10	P52800	122	61.15	No
Q00266	VAR_006935	Q00266	55	2.81	P13444	56	97.98	No
Q00266	VAR_006937	Q00266	264	3.43	P13444	265	97.08	No
Q00266	VAR_006939	Q00266	322	3.04	P13444	323	97.08	No
Q01955	VAR_011212	Q01955	1207	3.86	P02452	139	36.84	Yes
Q01955	VAR_011217	Q01955	1334	3.86	P02452	148	43.86	Yes
Q01955	VAR_011219	Q01955	1661	4.20	Q7SIB2	221	49.38	Yes
Q01974	VAR_010771	Q01974	620	4.44	Q06187	525	37.75	Yes
Q02388	VAR_001825	Q02388	2073	3.86	P02452	145	40.35	Yes
Q02388	VAR_001826	Q02388	2076	3.86	P02452	148	40.35	Yes
Q02388	VAR_001827	Q02388	2079	3.86	P02452	151	40.35	Yes
Q02388	VAR_001830	Q02388	2569	3.86	P02452	148	43.86	Yes
Q02388	VAR_001832	Q02388	2623	3.86	P02452	142	33.33	Yes
Q02388	VAR_001836	Q02388	2749	3.86	P02452	139	42.86	Yes
Q02388	VAR_011169	Q02388	1812	3.86	P02452	139	47.37	Yes
Q02388	VAR_011184	Q02388	2064	3.86	P02452	136	40.35	Yes
Q02388	VAR_011185	Q02388	2079	3.86	P02452	151	40.35	Yes
Q02388	VAR_011188	Q02388	2207	3.86	P02452	139	42.86	Yes
Q02388	VAR_011190	Q02388	2263	3.86	P02452	136	42.11	Yes
Q02388	VAR_011194	Q02388	2366	3.86	P02452	142	42.86	Yes
Q02388	VAR_011195	Q02388	2369	3.86	P02452	145	42.86	Yes
Q02388	VAR_015520	Q02388	1815	3.86	P02452	142	47.37	Yes

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
Q03692	VAR_001844	Q03692	598	3.71	Q00780	661	61.29	No
Q06124	VAR_015601	Q06124	42	2.34	P35235	42	100.00	No
Q06124	VAR_015613	Q06124	139	2.38	O89100	84	42.47	No
Q06187	VAR_006220	Q06187	27	3.31	Q06187	27	100.00	No
Q06187	VAR_006221	Q06187	27	3.31	Q06187	27	100.00	No
Q06187	VAR_006227	Q06187	287	3.80	O60880	13	30.14	No
Q06187	VAR_006231	Q06187	306	4.27	P35235	32	32.43	No
Q06187	VAR_006232	Q06187	333	3.75	P27986	670	33.80	No
Q06187	VAR_006239	Q06187	407	2.92	Q06187	407	100.00	Yes
Q06187	VAR_006249	Q06187	508	5.05	P54763	742	40.16	Yes
Q06187	VAR_006251	Q06187	519	4.10	P08069	1134	34.94	Yes
Q06187	VAR_006254	Q06187	524	4.05	Q07912	256	41.30	Yes
Q06187	VAR_006255	Q06187	524	4.05	Q07912	256	41.30	Yes
Q06187	VAR_006256	Q06187	525	4.44	Q06187	525	100.00	Yes
Q06187	VAR_006267	Q06187	591	2.29	Q07912	325	41.30	Yes
Q06187	VAR_006268	Q06187	593	3.57	Q07912	327	41.30	Yes
Q06187	VAR_006269	Q06187	593	3.57	Q07912	327	41.30	Yes
Q06187	VAR_006270	Q06187	597	4.47	Q07912	331	41.30	Yes
Q06187	VAR_006272	Q06187	612	3.26	P00520	455	48.19	No
Q06187	VAR_006272	Q06187	612	3.26	P00519	455	48.19	Yes
Q06187	VAR_006273	Q06187	618	4.13	P08631	478	41.53	Yes
Q06187	VAR_006276	Q06187	632	5.76	P08631	492	41.53	Yes
Q06187	VAR_006277	Q06187	640	4.27	P08631	500	41.53	Yes
Q06187	VAR_006278	Q06187	640	4.27	P08631	500	41.53	Yes
Q06187	VAR_006280	Q06187	646	2.31	P08631	506	41.53	Yes
Q06187	VAR_008293	Q06187	27	3.31	Q06187	27	100.00	No
Q06187	VAR_008305	Q06187	287	3.80	O60880	13	30.14	No
Q06187	VAR_008307	Q06187	306	4.27	P35235	32	32.43	No
Q06187	VAR_008319	Q06187	508	5.05	P54763	742	40.16	Yes
Q06187	VAR_008323	Q06187	524	4.05	Q07912	256	41.30	Yes
Q06187	VAR_008326	Q06187	562	6.26	Q06187	562	100.00	Yes
Q06187	VAR_008330	Q06187	618	4.13	P08631	478	41.53	Yes
Q06187	VAR_008331	Q06187	618	4.13	P08631	478	41.53	Yes
Q07001	VAR_021211	Q07001	271	3.54	P02711	260	34.31	No
Q09428	VAR_000100	Q09428	715	3.84	P68187	39	33.72	No
Q09428	VAR_008540	Q09428	1492	3.35	Q9CHL8	499	35.75	Yes
Q09428	VAR_015009	Q09428	1505	3.85	Q9KQW9	506	33.52	Yes
Q13253	VAR_011361	Q13253	35	4.16	Q13253	35	100.00	No
Q13253	VAR_018324	Q13253	35	4.16	Q13253	35	100.00	No
Q13402	VAR_009328	Q13402	503	3.77	P13538	529	41.12	No
Q13402	VAR_024047	Q13402	519	2.04	P13538	545	41.12	No
Q13402	VAR_024048	Q13402	756	4.13	P10587	803	40.00	No
Q13485	VAR_011380	Q13485	493	2.48	Q13485	493	100.00	No
Q13485	VAR_019571	Q13485	352	3.51	Q13485	352	100.00	No
Q13608	VAR_007918	Q13608	812	3.51	Q01853	585	52.46	No
Q13608	VAR_007919	Q13608	812	3.51	Q01853	585	52.46	No
Q13950	VAR_012132	Q13950	113	2.98	Q01196	62	91.04	No
Q13950	VAR_012133	Q13950	118	2.83	Q01196	67	91.04	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
Q13950	VAR_012137	Q13950	169	3.35	Q01196	118	91.04	No
Q13950	VAR_012142	Q13950	193	3.89	Q01196	142	91.04	Yes
Q13950	VAR_012145	Q13950	200	3.21	Q01196	149	91.04	No
Q13950	VAR_012146	Q13950	205	3.07	Q01196	154	91.04	No
Q13950	VAR_012147	Q13950	209	2.48	Q01196	158	91.04	No
Q14533	VAR_018116	Q14533	402	3.97	P08670	395	37.66	No
Q15672	VAR_004496	Q15672	131	3.34	P01106	377	44.90	No
Q15672	VAR_015219	Q15672	156	3.54	P01106	403	44.90	No
Q16667	VAR_013849	Q16667	187	3.46	Q16667	187	100.00	No
Q16836	VAR_024081	Q16836	258	3.56	Q16836	258	100.00	No
Q5LJ48	VAR_022986	Q5LJ48	116	2.77	P01135	54	43.33	No
Q6XZB0	VAR_023760	Q6XZB0	55	3.82	P29183	62	35.37	No
Q8NBP7	VAR_025453	Q8NBP7	253	3.05	P04072	104	33.04	No
Q92838	VAR_011080	Q92838	332	3.13	Q92838	332	100.00	No
Q92838	VAR_013487	Q92838	302	3.03	Q92838	302	100.00	No
Q92887	VAR_000099	Q92887	768	3.65	Q58206	153	31.65	Yes
Q92887	VAR_010756	Q92887	1382	4.20	Q9CHL8	430	37.99	Yes
Q92947	VAR_000394	Q92947	309	2.30	Q06319	262	31.03	No
Q92947	VAR_000396	Q92947	333	2.79	Q06319	286	31.03	No
Q92947	VAR_000408	Q92947	390	3.77	P15651	368	31.51	No
Q92947	VAR_000409	Q92947	390	3.77	P15651	368	31.51	No
Q92968	VAR_009306	Q92968	326	2.44	P08631	127	32.08	Yes
Q99456	VAR_008528	Q99456	429	4.94	P08670	399	33.11	No
Q99497	VAR_020496	Q99497	149	3.28	Q99497	149	100.00	Yes
Q99574	VAR_008520	Q99574	49	3.58	O35684	49	86.93	No
Q99574	VAR_008521	Q99574	52	3.19	O35684	52	86.93	No
Q99684	VAR_016213	Q99684	403	2.74	P03001	166	31.82	No
Q99697	VAR_003765	Q99697	115	3.65	P06601	243	62.50	No
Q99697	VAR_003766	Q99697	137	4.16	P40424	288	33.93	No
Q99758	VAR_023498	Q99758	568	3.62	P68187	38	33.33	No
Q9GZU5	VAR_013876	Q9GZU5	264	3.75	P41391	132	38.10	No
Q9H3D4	VAR_020870	Q9H3D4	243	3.89	P02340	172	57.22	Yes
Q9H3D4	VAR_020871	Q9H3D4	243	3.89	P02340	172	57.22	Yes
Q9H3D4	VAR_020873	Q9H3D4	318	3.89	P04637	248	56.19	No
Q9H3D4	VAR_020874	Q9H3D4	319	3.89	P04637	249	56.19	Yes
Q9HCC0	VAR_012792	Q9HCC0	99	3.09	Q8GBW6	61	35.11	No
Q9HCC0	VAR_012793	Q9HCC0	155	2.11	Q9X4K7	123	32.40	No
Q9NZR4	VAR_014246	Q9NZR4	166	3.12	P06601	215	62.50	No
Q9UBP0	VAR_010198	Q9UBP0	499	3.83	Q01853	637	40.66	No
Q9UBX0	VAR_010225	Q9UBX0	160	4.16	P40424	288	33.93	No
Q9UBX5	VAR_017153	Q9UBX5	227	2.97	Q9JJS8	162	41.94	No
Q9ULV5	VAR_017558	Q9ULV5	20	2.65	P22121	196	36.98	Yes
Q9UM47	VAR_012878	Q9UM47	146	5.88	P08709	132	38.71	No
Q9UM47	VAR_012886	Q9UM47	222	5.88	P08709	130	58.06	No
Q9UM47	VAR_012887	Q9UM47	224	5.88	P08709	132	58.06	No
Q9UM47	VAR_012900	Q9UM47	1261	5.88	P00740	108	38.71	Yes
Q9Y458	VAR_021832	Q9Y458	183	3.80	O15119	191	50.55	Yes
Q9Y5X4	VAR_010025	Q9Y5X4	97	4.26	P03372	234	44.00	No

Mut acc	Variant	Prot Acc	Resid	Cons	Templ acc	Templ resid	% id	Cryst Cont?
Q9Y6D9	VAR_019714	Q9Y6D9	516	2.08	Q9Y6D9	516	100.00	No

Appendix H

Table H.1: List of diseases and dosage sensitive genes compiled by the Baylor College of Medicine Medical Genetics Laboratory.

Disease description	Gene
1q41q42 deletion	DISP1
van der Woude syndrome	IRF6
Short stature, pituitary and cerebellar defects, and small sella turcica	LHX4
Pituitary anomalies with holoprosencephaly-like features	GLI2
Synpolydactyly/Syndactyly II//Split hand foot malformation 5 (SHFM 5)	HOXD13
Feingold	MYCN
nephronophthisis	NPHP1
SATB2, cleft palate	SATB2
Severe myoclonic epilepsy of infancy (SMEI) or Dravet syndrome	SCN1A
Holoprosencephaly 2, SIX3	SIX3
ASHG 2006	SUMO1
Mowat-Wilson	ZEB2
Noonan	SOS1
Heterotaxy 2	CFC1
Hypertension with CHD	BMPR2
Blepharophimosis	FOXL2
Waardenburg syndrome type II (WS2A)	MITF
3q29 microdeletion	PAK2
microphthalmia	SOX2
forebrain defects, left-right laterality defects	TDGF1
	TGFBR2
TP73L, split foot/split hand 4	TP63
Dandy-Walker syndrome	ZIC1, ZIC4
Noonan	RAF1
Rieger	PITX2
alfa synuclein	SCNA
Cornelia de Lange	NIPBL
microcephaly, CHD	NKX2-5
microcephaly, CHD	NPM1
Sotos	NSD1

Disease description	Gene
Treacher Collins syndrome	TCOF1
ADLD adult onset aut. dom. leukodystrophy	LMNB1
	EGR2
Chronic pancreatitis	SPINK1
Congenital 21-alpha hydroxylase deficiency	CYP21A2
Cleidocranial dysplasia	RUNX2
Prader-Willi-like phenotype	SIM1
VEGF	VEGF
Transient neonatal diabetes loci on 6q24 (OMIM 601410)	ZAC
Iridogoniodysgenesis anomaly, Axenfeld-Rieger syndrome	FKHL7 (FOXC1)
COL1A2	COL1A2
Williams	ELN
speech delay	FOXP2
Greig	GLI3
Williams	LIMK1
Split hand/foot	SHFM1
Holoprosencephaly 3, SHH	SHH
Saethre Chotzen	TWIST1
Hereditary pancreatitis	PRSS1
Schizophrenia & epilepsy	CNTNAP2
CHARGE	CHD7
Langer-Giedion	EXT1
Branchiootorenal (BOR)/Melnick-Fraser/Oto-facio-cervical)OFC)	EYA1
Congenital heart disease	GATA4
Bipolar disorder	IMPA1
Langer Giedion	TRPS1
Tetralogy of Fallot	ZFPM2/FOG2
9q34 microdeletion	EHMT1
GPR51, overgrowth	GABBR2
Nail-Patella	LMX1B
9q34 microdeletion	NOTCH1
Gorlin syndrome/Holoprosencephaly 7	PTCH1
Robinow/brachydactyly 1 Olivieri et al	ROR2
Sex reversal - Steroidogenic factor SF-1	SF-1
Loeys-Dietz syndrome	TGFBR1
Tuberous sclerosis	TSC1
Split food split hand 3	FBXW4
hypoparathyroidism, sensorineural deafness, and renal disease, HDR	GATA3
GRID1, 10q22q23 deletion	GRID1
Nebulette	NEBL
NRG3, 10q22q23 deletion	NRG3
PTEN, Cowden syndrome, Bannayan-Zonana syndrome	PTEN
Hirschsprung	RET
Potocki-Shaffer	ALX4
behavioral problems and autistic spectrum disorder. (OMIM 114130)	CALC1
behavioral problems and autistic spectrum disorder. (OMIM 114130)	CALC2
Potocki-Shaffer	EXT2
Beckwith-Wiedeman	H19

Disease description	Gene
Beckwith-Wiedeman	IGF2
Beckwith-Wiedeman	KCNQ1
Mitochondrial complex 1 deficiency	NDUFV1
Beckwith-Wiedeman	p57 (CDKN1C)
WAGR, Aniridia, PAX6	PAX6
Craniosynostosis	SOX6
WAGR, Wilms tumor, WT1	WT1
Stickler syndrome	COL2A1
Osteopoikilosis, short stature and MR	HMGA2
Osteopoikilosis, short stature and MR	LEMD3
Microduplication, Ruiter et al 2007	NOS1
Noonan	PTPN11
Microduplication, Ruiter et al 2008	RFC5
Microduplication, Ruiter et al 2006	THRAP2
Timothy	CACNA1C
Holt-Oram	TBX5
ulnar-mammary syndrome	TBX3
GPC5, brachydactyly and other skeletal anomalies	GPC5
GPC6, brachydactyly and other skeletal anomalies	GPC6
Retinoblastoma	RB
Holoprosencephaly 5, ZIC2	ZIC2
Hirschsprung	EDNRB
Anophthalmia, pituitary hypoplasia, and ear anomalies	BMP4
14q11.2 deletion syndrome	CHD8
FOXP1B	FOXP1B
14q11.2 deletion syndrome	SUPT16H
Branchiootic syndrome-3	SIX1
Oculoauriculovertebral spectrum (?)	SIX6
15q13.3 microdeletion	CHRNA7
Marfan	FBN1
Severe IUGR, developmental delay, postnatal growth retardation	IGF1R
NR2F2, Diaphragmatic hernia	NR2F2
PML	PML
PWS/AS	SNRPN
PWS/AS	UBE3A
Rubinstein-Taybi	CREBBP
Rubinstein-Taybi	DNASE1
alpha thalasemia-MR syndrome	HBA1
alpha thalasemia-MR syndrome	HBA2
Tuberous sclerosis	PKD1
Polycystic kidney disease	TSC2
Townes-Brocks	SALL1
Osteogenesis imperfecta type IV	COL1A1
17q21.31 microdeletion	CRHR1
Cystinosis	CTNS
Miller-Dieker	LIS1
17q21.31 microdeletion	MAPT
NF1	NF1

Disease description	Gene
CMT1A	PMP22
SMS	RAI1
Campomelic dysplasia	SOX9
TCF2, renal cysts and diabetes	TCF2
Miller-Dieker	YWHAE
Dyggve Melchior Clausen	DYM
Holoprosencephaly 4	TGIF1
Pitt-Hopkins	TCF4
BMP2	BMP2
Brachydactyly C	GDF5
Alagille	JAG1
Coloboma	SNAP25
Alzheimer - early onset	APP
SIM2	SIM2
Holoprosencephaly 1	TMEM1
Metachromatic leukodystrophy	ARSA
NF2	NF2
22q13.3 deletion	SHANK3
DGS	TBX1