

Decreased expression: 215

Increased expression: 1086

Total: 1301

**Table S5**  
**Transgenerational Female Hippocampus Regulated Genes**

**Apoptosis**

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio		
Bag3	163	103	<b>0.63</b>	AI231792	Bcl2-associated athanogene 3
Cbl27	16	160	<b>9.99</b>	AF275151	androgen receptor-related apoptosis-associated protein
Dapk1_predicte	55	149	<b>2.69</b>	BI281823	death associated protein kinase 1 (predicted)
Faim2	278	425	<b>1.53</b>	AF044201	Fas apoptotic inhibitory molecule 2
Pdcd6ip	457	706	<b>1.54</b>	BF389407	programmed cell death 6 interacting protein
RGD1306214_r	19	100	<b>5.32</b>	BE107277	similar to TGF-beta induced apoptosis protein 2 (predict
Tegt	358	559	<b>1.56</b>	NM_019381	testis enhanced gene transcript (Bax inhibitor-1)

**Cell Cycle**

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio		
Ccnd1	66	176	<b>2.67</b>	BI295861	cyclin D1
Ccnd2	41	546	<b>13.21</b>	L09752	cyclin D2
Ccnd3	87	139	<b>1.60</b>	NM_012766	cyclin D3
Cdc2l5	65	153	<b>2.37</b>	AI170800	cell division cycle 2-like 5 (cholinesterase-related cell di
Cdc2l6_predicte	275	480	<b>1.75</b>	BG376309	cell division cycle 2-like 6 (CDK8-like) (predicted)
Cdkn1c	66	105	<b>1.59</b>	AI013919	cyclin-dependent kinase inhibitor 1C (P57)
Ddb1	53	245	<b>4.58</b>	AJ277077	damage-specific DNA binding protein 1
Dab2	117	207	<b>1.77</b>	NM_024159	disabled homolog 2 (Drosophila)
Gas6	156	541	<b>3.47</b>	NM_057100	growth arrest specific 6
Numa1	100	168	<b>1.67</b>	AI599394	nuclear mitotic apparatus protein 1
RGD1560358_r	75	381	<b>5.11</b>	AW534965	similar to cell division cycle and apoptosis regulator 1 (f
Unc5b	54	108	<b>1.98</b>	BI303989	Unc-5 homolog B (C. elegans)

**Cytoskeleton-ECM**

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio		
Actb	1427	3416	<b>2.39</b>	NM_031144	actin, beta
Add1	60	176	<b>2.95</b>	NM_016990	adducin 1 (alpha)
Add3	46	269	<b>5.82</b>	AI407835	adducin 3 (gamma)
App	723	1257	<b>1.74</b>	BM986220	amyloid beta (A4) precursor protein
Actr1a_predicte	170	292	<b>1.71</b>	BI294346	ARP1 actin-related protein 1 homolog A (yeast) (predic
Actr2	1025	3030	<b>2.96</b>	BE107525	ARP2 actin-related protein 2 homolog (yeast)
Cdh13	139	525	<b>3.78</b>	BG381748	cadherin 13
Cdh2	147	230	<b>1.56</b>	AF097593	cadherin 2
Cdh22	36	526	<b>14.68</b>	NM_019161	cadherin 22
Chl1	211	848	<b>4.02</b>	BF524215	cell adhesion molecule with homology to L1CAM
Cspg5	33	99	<b>3.00</b>	AF292102	chondroitin sulfate proteoglycan 5
Cotl1_predicted	434	283	<b>0.65</b>	AI411057	coactosin-like 1 (Dictyostelium) (predicted)
Cntn1	82	127	<b>1.56</b>	NM_057118	contactin 1
Coro1c_predictr	23	91	<b>3.86</b>	BE120171	coronin, actin binding protein 1C (predicted)
Cttn	25	112	<b>4.45</b>	AF054618	cortactin
Dcn	142	230	<b>1.61</b>	BM390253	decorin
Dstn	305	483	<b>1.58</b>	AI170442	destrin
Dpy19l1_predic	145	315	<b>2.18</b>	BI296653	dpy-19-like 1 (C. elegans) (predicted)
Dbn1	978	551	<b>0.56</b>	NM_031024	drebrin 1
Dctn3_predicter	614	1050	<b>1.71</b>	AA893211	dynactin 3 (predicted)
Dst_predicted	72	162	<b>2.25</b>	AI235468	dystonin (predicted)
Dtna_predicted	458	878	<b>1.92</b>	BF396607	dystrobrevin alpha (predicted)
Dag1	52	138	<b>2.65</b>	AW251326	dystroglycan 1
Eml2	6	115	<b>17.88</b>	AF335571	<i>echinoderm microtubule associated protein like 2</i>
Eml4_predicted	146	91	<b>0.63</b>	BE111290	echinoderm microtubule associated protein like 4 (pred
Enc1	20	100	<b>4.90</b>	AA997271	Ectodermal-neural cortex 1
Epn2	40	88	<b>2.21</b>	NM_021852	epsin 2
Epb4.1l3	133	202	<b>1.52</b>	AB032828	erythrocyte protein band 4.1-like 3
Epb4.1l4a_pred	161	106	<b>0.66</b>	BE100811	erythrocyte protein band 4.1-like 4a (predicted)
Fndc3b_predict	22	115	<b>5.22</b>	AI176320	fibronectin type III domain containing 3B (predicted)
Gpiap1	108	216	<b>2.00</b>	BF546337	GPI-anchored membrane protein 1
Hook3	74	113	<b>1.53</b>	AI501458	hook homolog 3 (Drosophila)
Icam5_predicte	134	346	<b>2.58</b>	BE102418	intercellular adhesion molecule 5, telencephalin (predic
Kif1b	65	285	<b>4.42</b>	AB070355	kinesin family member 1B
Kif3a	29	122	<b>4.22</b>	BF397677	kinesin family member 3a
Kif5a	45	85	<b>1.88</b>	BF408765	kinesin family member 5A

Kif5c_predicted	21	148	7.17	BE104278	kinesin family member 5C (predicted)
KIFC2	541	332	0.61	AW433953	kinesin family member C2
Klc1	53	246	4.65	AI576961	kinesin light chain 1
Kifap3_predicte	16	187	11.78	AI227800	kinesin-associated protein 3 (predicted)
Lgals8	57	90	1.57	NM_053862	lectin, galactoside-binding, soluble 8
Lrln3_predicted	58	89	1.55	BE101066	leucine rich repeat and fibronectin type III domain conta
Map1lc3a	739	448	0.61	AI177372	microtubule-associated protein 1 light chain 3 alpha
Map1lc3b	276	418	1.52	AI233190	microtubule-associated protein 1 light chain 3 beta
Map1b	171	625	3.67	BG672052	microtubule-associated protein 1b
Mtap2	19	259	13.31	X74211	microtubule-associated protein 2
Mtap7_predicte	347	191	0.55	BG374192	Microtubule-associated protein 7 (predicted)
Mapre1	74	157	2.13	U75920	microtubule-associated protein, RP/EB family, member
Mtmr4_predicte	77	137	1.78	AI178892	myotubularin related protein 4 (predicted)
Mtmr9	109	411	3.77	BE111884	myotubularin related protein 9
LOC294446 /// I	85	156	1.84	M59859	myristoylated alanine rich protein kinase C substrate ///
Ncam1	275	427	1.56	AI409738	Neural cell adhesion molecule 1
Hnt	172	791	4.60	NM_017354	neurotrimin
Opcml	551	1095	1.99	M88709	opioid binding protein/cell adhesion molecule-like
Pclo	27	80	3.01	AF138789	piccolo (presynaptic cytomatrix protein)
Pkp4_predicted	54	153	2.83	AA819847	plakophilin 4 (predicted)
Podxl2_predicte	149	97	0.65	BI276946	podocalyxin-like 2 (predicted)
Col1a2	42	125	3.00	BM388837	procollagen, type I, alpha 2
Col11a2	92	56	0.61	BM391350	procollagen, type XI, alpha 2
Pfn2	255	446	1.75	AF228737	profilin 2
Pcdh17_predict	47	364	7.68	BF558981	protocadherin 17 (predicted)
Pcdh19_predict	76	227	2.97	AA997710	protocadherin 19 (predicted)
Pcdh7	38	97	2.54	AA956340	Protocadherin 7
Pcdha13	12	76	6.27	AW524833	protocadherin alpha 13
<b>Pcdhga1 /// Pcd</b>	<b>25</b>	<b>174</b>	<b>7.00</b>	<b>BE097805</b>	<b>protocadherin gamma subfamily C, 3 /// protocadhe</b>
Rdx	50	131	2.62	AW527313	radixin
Rsn	54	110	2.03	NM_031745	restin (Reed-Steinberg cell-expressed intermediate filar
Sparc	698	458	0.66	NM_012656	secreted acidic cysteine rich glycoprotein
Srrm2_predicte	143	908	6.33	BF408990	serine/arginine repetitive matrix 2 (predicted)
LOC679221	279	432	1.55	AA859614	similar to Microtubule-associated protein RP/EB family
Spnb2	123	342	2.78	BF559566	Spectrin beta 2
Spire1_predicte	117	182	1.55	BM387505	spire homolog 1 (Drosophila) (predicted)
Stmn4	465	713	1.53	AF026530	stathmin-like 4
Smap1l	79	141	1.79	BE097445	stromal membrane-associated protein 1-like
Sv2a	355	678	1.91	NM_057210	synaptic vesicle glycoprotein 2a
Sv2b	156	364	2.34	L10362	synaptic vesicle glycoprotein 2b
Tnr	34	229	6.80	NM_013045	tenascin R
Tspan5	42	88	2.10	AW251317	tetraspanin 5
Tmsb10	1044	636	0.61	NM_021261	thymosin, beta 10
Tmod2	110	327	2.98	BF567833	tropomodulin 2
Tpm1	178	280	1.57	M34135	tropomyosin 1, alpha
Tpm3	735	472	0.64	AI071098	tropomyosin 3, gamma
Vcpi1	56	300	5.37	BM383657	valosin containing protein (p97)/p47 complex interactin
Vamp1	20	95	4.79	M24104	vesicle-associated membrane protein 1
Vapb	13	83	6.65	NM_021847	vesicle-associated membrane protein, associated prote
Zyg11bl	80	207	2.59	BF567873	zyg-11 homolog B (C. elegans)-like

### Development

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio		
Apc	77	144	1.87	NM_012499	adenomatosis polyposis coli
Aph1a	53	82	1.56	BI275921	anterior pharynx defective 1a homolog (C. elegans)
Apeg1	143	86	0.60	NM_012905	aortic preferentially expressed gene 1
Atxn2_predicte	410	656	1.60	BE109041	ataxin 2 (predicted)
Atxn2l_predicte	52	88	1.71	BE349657	ataxin 2-like (predicted)
Bbx_predicted	42	227	5.40	BF392234	bobby sox homolog (Drosophila) (predicted)
Bai3_predicted	48	100	2.08	BF409866	Brain-specific angiogenesis inhibitor 3 (predicted)
Cops7a_predict	123	190	1.55	BE110772	COP9 (constitutive photomorphogenic) homolog, subur
Cpne8_predicte	18	119	6.75	AI059204	copine VIII (predicted)
Cryl1	241	135	0.56	BI293393	crystallin, lamda 1
<b>Crim1_predicte</b>	<b>31</b>	<b>160</b>	<b>5.07</b>	<b>BI289620</b>	<b>cysteine-rich motor neuron 1 (predicted)</b>
Cyfp2_predicte	131	242	1.84	BE105837	cytoplasmic FMR1 interacting protein 2 (predicted)
Dlgap3	17	99	5.80	U67139	discs, large (Drosophila) homolog-associated protein 3
Dlgh1	59	113	1.93	NM_012788	discs, large homolog 1 (Drosophila)
Dlgh2	26	81	3.11	BI296447	Discs, large homolog 2 (Drosophila)

Dlgh4	26	98	<b>3.85</b>	NM_019621	discs, large homolog 4 (Drosophila)
Dlgap4	334	537	<b>1.61</b>	U67140	discs, large homolog-associated protein 4 (Drosophila)
Elavl1_predicted	156	1107	<b>7.12</b>	BE104552	ELAV (embryonic lethal, abnormal vision, Drosophila)-li
Elavl2	413	1045	<b>2.53</b>	BF563441	ELAV (embryonic lethal, abnormal vision, Drosophila)-li
Fbn1	82	48	<b>0.59</b>	BM389019	fibrillin 1
Fcmd_predicted	232	473	<b>2.04</b>	AI101490	Fukuyama type congenital muscular dystrophy homolog
Grhl1_predicted	130	198	<b>1.52</b>	BM383749	Grainyhead-like 1 (Drosophila) (predicted)
Hdh	109	65	<b>0.60</b>	BI274329	Huntington disease gene homolog
Ibtk_predicted	135	89	<b>0.66</b>	AW525218	inhibitor of Bruton agammaglobulinemia tyrosine kinase
Vof16	7	122	<b>18.13</b>	BE107282	ischemia related factor vof-16
Kidins220	51	146	<b>2.86</b>	AF313464	kinase D-interacting substance 220
Lrrn6a	965	1821	<b>1.89</b>	AI711152	leucine rich repeat neuronal 6A
Mpv17_predicted	32	187	<b>5.78</b>	AA849966	Mpv17 transgene, kidney disease mutant-like (predicted)
Mbp	383	3170	<b>8.27</b>	NM_017026	myelin basic protein
Mog	43	88	<b>2.08</b>	BF410306	Myelin oligodendrocyte glycoprotein
Mobp	288	801	<b>2.78</b>	D28110	myelin-associated oligodendrocytic basic protein
Nsf	292	873	<b>2.99</b>	AF142097	N-ethylmaleimide sensitive fusion protein
Napg	80	174	<b>2.16</b>	AA956372	N-ethylmaleimide-sensitive factor attachment protein, g
Nedd4a	524	2373	<b>4.53</b>	BI284798	neural precursor cell expressed, developmentally down
Neurod2	39	116	<b>2.97</b>	NM_019326	neurogenic differentiation 2
LOC681423 /// I	18	117	<b>6.48</b>	BE117361	Nipped-B homolog (Drosophila) /// similar to delangin is
Odz2	122	264	<b>2.17</b>	NM_020088	odd Oz/ten-m homolog 2 (Drosophila)
Opa1	45	95	<b>2.12</b>	BF407962	Optic atrophy 1 homolog (human)
Pmp22	61	137	<b>2.23</b>	AW252810	peripheral myelin protein 22
Prg1	132	505	<b>3.84</b>	BF394800	plasticity related gene 1
Peo1_predicted	249	158	<b>0.64</b>	AA925995	progressive external ophthalmoplegia 1 homolog (hum
Rb1cc1_predict	196	449	<b>2.28</b>	BG671668	RB1-inducible coiled-coil 1 (predicted)
Rqcd1	93	160	<b>1.73</b>	AI044879	rcd1 (required for cell differentiation) homolog 1 (S. pon
Reln	43	89	<b>2.06</b>	NM_080394	reelin
Sema6a_predic	60	93	<b>1.55</b>	BM387083	sema domain, transmembrane domain (TM), and cytop
LOC682864	36	147	<b>4.08</b>	BE116720	similar to 82-kD FMRP Interacting Protein
LOC499716	47	78	<b>1.67</b>	AA818641	similar to DnaJ (Hsp40) homolog, subfamily B, member
LOC686892	44	188	<b>4.29</b>	AI180361	similar to muscleblind-like 1 isoform d
RGD1565589_f	151	273	<b>1.80</b>	BM390561	similar to myocardial ischemic preconditioning upregula
LOC682507	46	475	<b>10.31</b>	AI706673	similar to Neural Wiskott-Aldrich syndrome protein (N-V
RGD1566269_f	381	208	<b>0.55</b>	BE118896	Similar to Neuropilin- and tolloid-like protein 1 (predicte
RGD1310358_f	123	252	<b>2.06</b>	BF567255	similar to NNX3 (predicted)
RGD1559723_f	23	113	<b>4.82</b>	BE114807	similar to Spn protein (predicted)
LOC680647	36	86	<b>2.42</b>	BG667371	Similar to TFA2 protein
LOC687031	57	164	<b>2.90</b>	BE116590	Similar to transmembrane protein SHREW1
Slit3	171	88	<b>0.51</b>	BF386446	Slit homolog 3 (Drosophila)
Spg20	59	100	<b>1.71</b>	AI576427	spastic paraplegia 20, spartin (Troyer syndrome) homo
Smndc1	60	100	<b>1.67</b>	BI288396	survival motor neuron domain containing 1
Syn2	114	561	<b>4.92</b>	NM_019159	synapsin II
Sncb	135	228	<b>1.69</b>	NM_080777	synuclein, beta
Sncg	535	1003	<b>1.87</b>	NM_031688	synuclein, gamma
Tor1aip1	146	363	<b>2.49</b>	U19614	torsin A interacting protein 1
Trak2	101	844	<b>8.35</b>	BG378620	trafficking protein, kinesin binding 2
Trps1_predicted	62	307	<b>4.91</b>	BF398245	trichorhinophalangeal syndrome I (predicted)
Trps1_predicted	121	430	<b>3.55</b>	BF550315	trichorhinophalangeal syndrome I (predicted)
Wbscr1	16	142	<b>8.74</b>	H31800	Williams-Beuren syndrome chromosome region 1 hom
Wasip	626	377	<b>0.60</b>	NM_057192	Wiskott-Aldrich syndrome protein interacting protein
Whsc1l1_predic	94	239	<b>2.56</b>	AI234807	Wolf-Hirschhorn syndrome candidate 1-like 1 (predicte
Rbaf600	20	77	<b>3.85</b>	BF392966	ZUBR1

#### Electron Transport

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio	Genbank	Gene Title
Cyp51	613	1238	<b>2.02</b>	NM_012941	cytochrome P450, subfamily 51
Glrx2	76	475	<b>6.27</b>	BG671304	Glutaredoxin 2 (thioltransferase)
Iag2	46	145	<b>3.13</b>	NM_053946	implantation-associated protein
Uqcrc2	241	418	<b>1.74</b>	BF290998	ubiquinol cytochrome c reductase core protein 2

#### Epigenetics/Gene Modifying Factors

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio	Genbank	Gene Title
Atrx	13	87	<b>6.48</b>	BF397805	alpha thalassemia/mental retardation syndrome X-link
Rere	96	61	<b>0.64</b>	BF412271	Arginine-glutamic acid dipeptide (RE) repeats
Ash1_predicted	20	143	<b>7.10</b>	BG663056	ash1 (absent, small, or homeotic)-like (Drosophila) (pre

Brd2	78	127	<b>1.64</b>	AA946361	bromodomain containing 2
Brd3_predicted	62	271	<b>4.37</b>	AI172375	bromodomain containing 3 (predicted)
Brd4	12	178	<b>14.61</b>	BM389207	bromodomain containing 4
Csnk2a1	19	268	<b>13.81</b>	BF288177	casein kinase II, alpha 1 polypeptide
Csnk2a2_predict	38	106	<b>2.81</b>	BI290750	casein kinase II, alpha 2, polypeptide (predicted)
Chd1_predicted	60	137	<b>2.28</b>	AA955721	chromodomain helicase DNA binding protein 1 (predict
<i>Chd2_predictec</i>	91	51	<b>0.56</b>	<i>BF396633</i>	<i>Chromodomain helicase DNA binding protein 2 (predic</i>
Cxxc5	276	564	<b>2.04</b>	BI281838	CXXC finger 5
Tia1	46	99	<b>2.14</b>	BI298817	Cytotoxic granule-associated RNA binding protein 1
Epc2_predicted	64	138	<b>2.17</b>	AW918173	enhancer of polycomb homolog 2 (Drosophila) (predict
H2afy	169	403	<b>2.38</b>	M99065	H2A histone family, member Y
Hdac4_predicte	46	120	<b>2.63</b>	BF419085	histone deacetylase 4 (predicted)
Impact	155	253	<b>1.63</b>	BG664101	imprinted and ancient
Id4	23	99	<b>4.38</b>	AI412150	inhibitor of DNA binding 4
Jmjd3_predicte	50	150	<b>3.02</b>	BE118720	jumonji domain containing 3 (predicted)
Metap1_predict	58	91	<b>1.59</b>	AI101470	methionyl aminopeptidase 1 (predicted)
Prdm2	33	168	<b>5.05</b>	BF388420	PR domain containing 2, with ZNF domain
Pcmt2_predict	709	1292	<b>1.82</b>	AA955163	protein-L-isoaspartate (D-aspartate) O-methyltransfera
RGD1566399_f	95	155	<b>1.63</b>	BE098769	similar to MYST histone acetyltransferase monocytic le
LOC314964	24	83	<b>3.51</b>	AA965250	similar to PHD finger protein 20-like 1 isoform 1
LOC681178 /// I	134	234	<b>1.75</b>	BI288196	similar to polycomb group ring finger 5
Smarca2	70	148	<b>2.12</b>	BE102268	SWI/SNF related, matrix associated, actin dependent re
Smarca4	24	223	<b>9.41</b>	BE111847	SWI/SNF related, matrix associated, actin dependent re

### Golgi Apparatus

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	<b>Ratio</b>	Genbank	Gene Title
Ap1s1_predicte	215	524	<b>2.44</b>	AA955775	adaptor protein complex AP-1, sigma 1 (predicted)
Ap2b1	26	95	<b>3.66</b>	M34176	adaptor-related protein complex 2, beta 1 subunit
Copz1_predicte	124	217	<b>1.75</b>	BF406339	coatamer protein complex, subunit zeta 1 (predicted)
Cbfa2t3_predict	45	93	<b>2.07</b>	BE115481	core-binding factor, runt domain, alpha subunit 2; trans
Gga3_predictec	320	114	<b>0.35</b>	BE112983	golgi associated, gamma adaptin ear containing, ARF t
Gla	60	112	<b>1.86</b>	BM385090	galactosidase, alpha
Golph3	58	101	<b>1.74</b>	NM_023977	golgi phosphoprotein 3
Hs2st1	226	439	<b>1.94</b>	AI714262	heparan sulfate 2-O-sulfotransferase 1
Pde4dip	54	114	<b>2.11</b>	NM_022382	phosphodiesterase 4D interacting protein (myomegalin
Pde4dip	450	278	<b>0.62</b>	AI763912	Phosphodiesterase 4D interacting protein (myomegalin
<i>RGD1560511_f</i>	79	189	<b>2.40</b>	<i>BI281965</i>	<i>similar to Vps41 protein (predicted)</i>
Surf4	146	97	<b>0.66</b>	AI227937	surfeit 4
Tgoln2	53	98	<b>1.87</b>	AW917192	trans-golgi network protein 2

### Growth Factors, Cyto- and Chemokines

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	<b>Ratio</b>	Genbank	Gene Title
Api5_predicted	64	108	<b>1.68</b>	H34636	apoptosis inhibitor 5 (predicted)
Bdnf	137	236	<b>1.72</b>	NM_012513	brain derived neurotrophic factor
Cxcl12	52	155	<b>2.97</b>	AF189724	chemokine (C-X-C motif) ligand 12
Crif1_predicted	44	81	<b>1.84</b>	AA866388	cytokine receptor-like factor 1 (predicted)
Igf2	131	308	<b>2.36</b>	NM_031511	insulin-like growth factor 2
Il6st	153	382	<b>2.49</b>	BM383427	interleukin 6 signal transducer
Gdf1_predicted	716	433	<b>0.60</b>	AI549010	longevity assurance homolog 1 (S. cerevisiae) (predict
Negr1	155	964	<b>6.21</b>	NM_021682	neuronal growth regulator 1
Ogn_predicted	55	117	<b>2.14</b>	BG664221	osteoglycin (predicted)
Sep3	88	183	<b>2.08</b>	NM_019375	septin 3
Sept6_predicte	117	262	<b>2.24</b>	AW532098	Septin 6 (predicted)
Sep7	294	569	<b>1.93</b>	NM_022616	septin 7
Sst	468	307	<b>0.66</b>	NM_012659	somatostatin
<b>Tgfb2</b>	<b>121</b>	<b>70</b>	<b>0.58</b>	<b>BE117736</b>	<b>Transforming growth factor, beta 2</b>
Tmeff1	74	177	<b>2.38</b>	NM_023020	transmembrane protein with EGF-like and two follistatin

### Immune Response

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	<b>Ratio</b>	Genbank	Gene Title
Cd2bp2_predict	32	76	<b>2.38</b>	BG379822	CD2 antigen (cytoplasmic tail) binding protein 2 (predic
Cd74	70	203	<b>2.89</b>	NM_013069	CD74 antigen (invariant polypeptide of major histocomp
Cd99	64	101	<b>1.59</b>	AI235284	CD99 antigen
C7 /// Tubb2c	51	89	<b>1.76</b>	BF284922	complement component 7 /// tubulin, beta 2c
Igsf4b_predicte	36	109	<b>3.06</b>	BF409960	immunoglobulin superfamily, member 4B (predicted)
Ke2	530	938	<b>1.77</b>	AW434268	MHC class II region expressed gene KE2

Pafah1b1	45	141	<b>3.11</b>	BG663460	platelet-activating factor acetylhydrolase, isoform Ib, al
<i>RT1-Ke4</i>	301	134	<b>0.44</b>	<i>BM389027</i>	<i>RT1 class I, locus Ke4</i>
RT1-Aw2	208	34	<b>0.16</b>	AA801218	RT1 class Ib, locus Aw2
<i>RT1-S3</i>	86	51	<b>0.59</b>	<i>AJ243974</i>	<i>RT1 class Ib, locus S3</i>
RT1-Da	45	153	<b>3.36</b>	Y00480	RT1 class II, locus Da
RT1-149	109	65	<b>0.60</b>	BI282965	RT1-149 protein
Sema4f	143	95	<b>0.66</b>	NM_019272	sema domain, immunoglobulin domain (Ig), transmemb
LOC690085	38	199	<b>5.29</b>	AW534002	Similar to B-cell CLL/lymphoma 7A
LOC680404 /// I	117	1507	<b>12.89</b>	BG374818	similar to Complement C1q-like protein 3 precursor (Gli
LOC296637	118	259	<b>2.20</b>	AA944347	similar to HLA-B associated transcript-2 isoform a
RGD1563429_f	61	463	<b>7.56</b>	BM391860	similar to T-cell activation leucine repeat-rich protein (pi
<i>Stag1_predicted</i>	100	49	<b>0.49</b>	<i>AI071210</i>	<i>Stromal antigen 1 (predicted)</i>
Thy1	213	597	<b>2.81</b>	NM_012673	thymus cell antigen 1, theta
Wrb	87	147	<b>1.69</b>	BI280216	tryptophan rich basic protein

### Metabolism & Transport

Sample	F3-Cont	F3-Vinc	Vin/Con	Gene Symbol	Gene Title
Agpat1	74	167	<b>2.24</b>	BF390804	1-acylglycerol-3-phosphate O-acyltransferase 1
Abhd8_predicted	10	227	<b>22.52</b>	AI137533	abhydrolase domain containing 8 (predicted)
Acs1	52	126	<b>2.45</b>	BI277523	acyl-CoA synthetase long-chain family member 1
Adpgk	65	170	<b>2.60</b>	BE102146	ADP-dependent glucokinase
Aldh1a2	59	119	<b>2.01</b>	NM_053896	aldehyde dehydrogenase family 1, subfamily A2
Aldh6a1	50	103	<b>2.05</b>	NM_031057	aldehyde dehydrogenase family 6, subfamily A1
Accn2	95	58	<b>0.61</b>	AJ309926	amiloride-sensitive cation channel 2, neuronal
<i>Aqp4</i>	84	255	<b>3.03</b>	<i>NM_012825</i>	<i>aquaporin 4</i>
Aqp9	86	53	<b>0.61</b>	NM_022960	aquaporin 9
<i>Arcn1</i>	55	153	<b>2.78</b>	<i>BF414061</i>	<i>archain 1</i>
<b>Arg2</b>	<b>125</b>	<b>245</b>	<b>1.96</b>	<b>AI574994</b>	<b>Arginase 2</b>
Asna1	194	308	<b>1.59</b>	AI406558	arsA arsenite transporter, ATP-binding, homolog 1 (bac
Arsb	72	114	<b>1.57</b>	AW917073	arylsulfatase B
Atp2a2	79	331	<b>4.19</b>	J04024	ATPase, Ca++ transporting, cardiac muscle, slow twitcl
Atp2b1	1829	3444	<b>1.88</b>	NM_053311	ATPase, Ca++ transporting, plasma membrane 1
Atp6v1a1_predi	636	1617	<b>2.54</b>	BI289589	ATPase, H transporting, lysosomal V1 subunit A (predic
Atp6v1b2	568	1031	<b>1.82</b>	NM_057213	ATPase, H transporting, lysosomal V1 subunit B2
Atp6v0a2	37	112	<b>3.03</b>	BE110564	ATPase, H+ transporting, lysosomal V0 subunit a isofo
Atp1a1	45	293	<b>6.56</b>	M74494	ATPase, Na+/K+ transporting, alpha 1 polypeptide
Atp1a3	1132	746	<b>0.66</b>	NM_012506	ATPase, Na+/K+ transporting, alpha 3 polypeptide
Atp1b2	15	194	<b>12.92</b>	U45946	ATPase, Na+/K+ transporting, beta 2 polypeptide
B3gat1	43	87	<b>2.03</b>	NM_054003	beta-1,3-glucuronyltransferase 1 (glucuronosyltransfera
Bace1	38	149	<b>3.88</b>	BG377970	Beta-site APP cleaving enzyme 1
Cacna2d3	20	99	<b>4.94</b>	BE108017	Calcium channel, voltage-dependent, alpha 2/delta 3 su
Cacnb1	79	144	<b>1.81</b>	NM_017346	calcium channel, voltage-dependent, beta 1 subunit
Cacng3	117	218	<b>1.87</b>	AF361340	calcium channel, voltage-dependent, gamma subunit 3
Cacng8	95	60	<b>0.63</b>	BE114019	Calcium channel, voltage-dependent, gamma subunit 8
Clstn2	24	268	<b>11.10</b>	BE109141	Calsyntenin 2
Chst2_predicted	290	585	<b>2.02</b>	AW520944	carbohydrate sulfotransferase 2 (predicted)
Car11	128	229	<b>1.78</b>	AW252112	carbonic anhydrase 11
Ca3	28	228	<b>8.01</b>	<i>AB030829</i>	<i>carbonic anhydrase 3</i>
Cs	136	269	<b>1.98</b>	NM_130755	citrate synthase
Cplx1	56	184	<b>3.29</b>	U35098	complexin 1
Cplx2	47	345	<b>7.29</b>	NM_053878	complexin 2
Ckmt1	1166	750	<b>0.64</b>	BI301453	creatine kinase, mitochondrial 1, ubiquitous
Capon	94	339	<b>3.60</b>	AF037071	C-terminal PDZ domain ligand of neuronal nitric oxide s
Dgkg	160	599	<b>3.75</b>	NM_013126	diacylglycerol kinase, gamma
Elovl6	54	271	<b>5.03</b>	BF396857	ELOVL family member 6, elongation of long chain fatty
Etnk1_predicted	83	133	<b>1.61</b>	BG668816	ethanolamine kinase 1 (predicted)
Xpo4_predicted	25	108	<b>4.31</b>	BF397646	exportin 4 (predicted)
Fmo1	58	87	<b>1.51</b>	NM_012792	flavin containing monooxygenase 1
Gja1	13	81	<b>6.11</b>	BG378227	gap junction membrane channel protein alpha 1
Got1	438	822	<b>1.87</b>	D00252	glutamate oxaloacetate transaminase 1
Glul	270	509	<b>1.88</b>	BI296610	glutamate-ammonia ligase (glutamine synthase)
Gls	67	145	<b>2.15</b>	M22586	glutaminase
Gapdh	1258	3188	<b>2.53</b>	X_Rat_GAPC	glyceraldehyde-3-phosphate dehydrogenase
Gpd1	112	66	<b>0.59</b>	BF399697	Glycerol-3-phosphate dehydrogenase 1 (soluble)
Gsk3b	16	115	<b>7.03</b>	BF287444	glycogen synthase kinase 3 beta
Gda	830	1718	<b>2.07</b>	AF245172	guanine deaminase
Hba-a1 /// LOC:	4846	2874	<b>0.59</b>	AI179404	hemoglobin alpha, adult chain 1 /// hemoglobin alpha 2
HK1	93	222	<b>2.40</b>	_Rat_Hexokir	hexokinase 1

Hcn2	41	110	<b>2.67</b>	AW532988	hyperpolarization activated cyclic nucleotide-gated pota
Hprt	43	78	<b>1.81</b>	BE117366	Hypoxanthine guanine phosphoribosyl transferase
Kpnb1	18	92	<b>5.01</b>	NM_017063	karyopherin (importin) beta 1
Kcnp2	17	93	<b>5.40</b>	AF269283	Kv channel-interacting protein 2
Lypla2	63	132	<b>2.11</b>	NM_031342	lysophospholipase 2
Man1a_predicted	399	237	<b>0.59</b>	AA892549	mannosidase 1, alpha (predicted)
Man2a2_predic	295	192	<b>0.65</b>	BE099244	Mannosidase 2, alpha 2 (predicted)
Magi2	36	168	<b>4.69</b>	AF130819	membrane associated guanylate kinase, WW and PDZ
Maoa	235	752	<b>3.20</b>	D00688	monoamine oxidase A
Mgll	15	216	<b>14.29</b>	AY081195	monoglyceride lipase
Asah1	80	123	<b>1.54</b>	NM_053407	N-acylsphingosine amidohydrolase 1
Por	26	133	<b>5.12</b>	AI407454	P450 (cytochrome) oxidoreductase
Pex2	54	114	<b>2.13</b>	BF413248	Peroxin 2
Ppap2b	30	571	<b>18.84</b>	AW253995	phosphatidic acid phosphatase type 2B
Ptdss1	349	611	<b>1.75</b>	AA851302	phosphatidylserine synthase 1
Plcl1	1062	696	<b>0.66</b>	NM_053456	phospholipase C-like 1
Prps1	183	292	<b>1.59</b>	M29392	phosphoribosyl pyrophosphate synthetase 1
Prps2	124	203	<b>1.64</b>	NM_012634	phosphoribosyl pyrophosphate synthetase 2
Kctd1	212	140	<b>0.66</b>	AI501338	Potassium channel tetramerisation domain containing 1
Kctd12_predicted	105	162	<b>1.54</b>	AA800908	Potassium channel tetramerisation domain containing 1
Kcnj3	37	86	<b>2.35</b>	U09243	potassium inwardly-rectifying channel, subfamily J, mem
Kcnd2	326	900	<b>2.76</b>	AW528891	potassium voltage gated channel, Shal-related family, n
Kcnd3	177	274	<b>1.55</b>	NM_031739	potassium voltage gated channel, Shal-related family, n
Kcna1	129	511	<b>3.97</b>	M26161	potassium voltage-gated channel, shaker-related subfa
Kcng2	56	89	<b>1.58</b>	AI407741	potassium voltage-gated channel, subfamily G, membe
Kcnq3	43	108	<b>2.48</b>	BF400850	Potassium voltage-gated channel, subfamily Q, membe
Psap	523	937	<b>1.79</b>	NM_013013	prosaposin
Pin1_predicted	539	892	<b>1.65</b>	BM388765	protein (peptidyl-prolyl cis/trans isomerase) NIMA-inter
Ahcy11_predicted	199	486	<b>2.45</b>	BF397074	S-adenosylhomocysteine hydrolase-like 1 (predicted)
Srpk2_predicted	24	262	<b>10.93</b>	BE113419	serine/arginine-rich protein specific kinase 2 (predicted)
LOC365090	45	109	<b>2.44</b>	AW140661	similar to 5-nucleotidase, cytosolic II
LOC500282	173	306	<b>1.77</b>	BG671292	similar to ADP-ribosylation factor-like 10C
LOC682926 /// I	91	136	<b>1.50</b>	BF557618	similar to chromatin modifying protein 1B
RGD1311444_f	427	768	<b>1.80</b>	BF557088	similar to intracellular membrane-associated calcium-in
LOC679692 /// I	43	168	<b>3.89</b>	BE115518	similar to lysophosphatidylglycerol acyltransferase 1
LOC498358	48	128	<b>2.67</b>	AW918899	similar to solute carrier family 30 (zinc transporter), mem
LOC499587	276	1370	<b>4.96</b>	AW529244	similar to solute carrier family 7, member 14
RGD1564893_f	151	100	<b>0.66</b>	AI410604	similar to surfeit 5 isoform b (predicted)
Scn3a	37	158	<b>4.33</b>	NM_013119	sodium channel, voltage-gated, type III, alpha polypept
Slc1a3	129	477	<b>3.70</b>	AF265360	solute carrier family 1 (glial high affinity glutamate trans
Slc16a7	65	134	<b>2.05</b>	X97445	solute carrier family 16 (monocarboxylic acid transporte
Slc17a7	85	271	<b>3.19</b>	NM_053859	solute carrier family 17 (sodium-dependent inorganic pl
Slc2a13	68	279	<b>4.08</b>	NM_133611	solute carrier family 2 (facilitated glucose transporter), r
Slc24a2	233	1304	<b>5.60</b>	AF021923	solute carrier family 24 (sodium/potassium/calcium excl
Slc25a12_predi	200	124	<b>0.62</b>	AI578095	Solute carrier family 25 (mitochondrial carrier, Aralar), n
Slc27a4	161	97	<b>0.60</b>	BI286134	solute carrier family 27 (fatty acid transporter), member
Slc3a1	277	983	<b>3.55</b>	NM_017216	solute carrier family 3, member 1
Slc35a1_predic	56	88	<b>1.58</b>	BE103444	solute carrier family 35 (CMP-sialic acid transporter), m
Slc4a4	8	143	<b>19.07</b>	AF210250	solute carrier family 4, member 4
Slc5a3	765	455	<b>0.60</b>	BE116021	solute carrier family 5 (inositol transporters), member 3
Slco3a1	51	109	<b>2.15</b>	AF239219	solute carrier organic anion transporter family, member
St3gal4	34	87	<b>2.58</b>	BF289499	ST3 beta-galactoside alpha-2,3-sialyltransferase 4
Scd2	98	559	<b>5.73</b>	BE107760	stearoyl-Coenzyme A desaturase 2
Sc5d	151	290	<b>1.92</b>	AB052846	sterol-C5-desaturase (fungal ERG3, delta-5-desaturase
Snap91	173	95	<b>0.55</b>	AW531009	Synaptosomal-associated protein, 91kDa homolog (mo
Syt1	50	260	<b>5.22</b>	AI413003	synaptotagmin I
Syt4	27	312	<b>11.76</b>	L38247	synaptotagmin IV
Syt11	21	89	<b>4.15</b>	NM_031667	synaptotagmin XI
Txnrd1	53	86	<b>1.64</b>	U63923	thioredoxin reductase 1
---	77	221	<b>2.87</b>	AI029275	Transcribed locus, weakly similar to XP_001054001.1
Ugt8	82	154	<b>1.88</b>	L21698	UDP galactosyltransferase 8
Galnt1	41	104	<b>2.53</b>	NM_024373	UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-ac
Galnt2_predicted	156	282	<b>1.81</b>	BE111638	UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-ac
Vip	250	159	<b>0.64</b>	AI412212	vasoactive intestinal polypeptide
Vdac1	59	283	<b>4.79</b>	AF268467	voltage-dependent anion channel 1

#### Proteolysis

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio		

Cpd	174	846	<b>4.86</b>	NM_012836	carboxypeptidase D
Dcun1d4_predic	43	132	<b>3.05</b>	BF288456	DCN1, defective in cullin neddylation 1, domain contain
Dpp6	80	212	<b>2.66</b>	NM_022850	dipeptidylpeptidase 6
Dpp8_predicted	54	111	<b>2.06</b>	AI175505	dipeptidylpeptidase 8 (predicted)
Laptm4a	500	1185	<b>2.37</b>	AW525776	lysosomal-associated protein transmembrane 4A
LOC290864	56	93	<b>1.66</b>	AI408662	similar to Ubiquitin carboxyl-terminal hydrolase 4 (Ubiqui
LOC300472	48	78	<b>1.63</b>	BE097616	similar to Vacuolar protein sorting 26 homolog (VPS26)
LOC316395	63	191	<b>3.02</b>	BE119116	similar to HECT, C2 and WW domain containing E3 ubi
LOC679539 /// I	230	533	<b>2.31</b>	AW142658	ubiquitin-conjugating enzyme E2 variant 1 (predicted) /
<b>LOC686268 /// I</b>	<b>138</b>	<b>287</b>	<b>2.07</b>	<b>BE105606</b>	<b>similar to SUMO/sentrin specific protease 5 (predic</b>
Mmp14	22	81	<b>3.62</b>	X83537	matrix metalloproteinase 14 (membrane-inserted)
Mmp16	64	104	<b>1.64</b>	NM_080776	matrix metalloproteinase 16
Pja2	334	994	<b>2.98</b>	AI600019	praja 2, RING-H2 motif containing
Prss22_predicte	55	109	<b>1.97</b>	BM389391	protease, serine, 22 (predicted)
RGD1560364_f	147	77	<b>0.53</b>	BM384685	similar to vacuolar protein sorting 13C protein (predicte
RGD1561386_f	28	93	<b>3.31</b>	BE116890	similar to CBL E3 ubiquitin protein ligase (Signal transd
RGD1563278_f	18	148	<b>8.06</b>	BF402371	similar to Ubiquitin ligase protein DZIP3 (DAZ-interactir
RGD1565984_f	86	57	<b>0.66</b>	BE097603	similar to Ubiquitin carboxyl-terminal hydrolase 35 (Ubi
<b>Rnf187_predic</b>	<b>32</b>	<b>93</b>	<b>2.88</b>	<b>AI502527</b>	<b>ring finger protein 187 (predicted)</b>
<b>Rnf6_predicte</b>	<b>22</b>	<b>117</b>	<b>5.25</b>	<b>BI296352</b>	<b>ring finger protein (C3H2C3 type) 6 (predicted)</b>
Saps1_predicte	116	184	<b>1.59</b>	BF399333	SAPS domain family, member 1 (predicted)
Senp6_predicte	96	147	<b>1.54</b>	H32530	SUMO/sentrin specific peptidase 6 (predicted)
Siah1a	76	116	<b>1.53</b>	NM_080905	seven in absentia 1A
Smurf2_predict	9	106	<b>12.01</b>	BF393807	SMAD specific E3 ubiquitin protein ligase 2 (predicted)
Timp2	31	111	<b>3.54</b>	BF523128	tissue inhibitor of metalloproteinase 2
Tpp1	191	480	<b>2.51</b>	NM_031357	tripeptidyl peptidase I
Ube2d2	128	391	<b>3.05</b>	BF283846	Ubiquitin-conjugating enzyme E2D 2
Ube2g1	113	240	<b>2.12</b>	AF099093	ubiquitin-conjugating enzyme E2G 1 (UBC7 homolog, C
Ube2n	723	456	<b>0.63</b>	BI294702	ubiquitin-conjugating enzyme E2N
Ubl3	128	217	<b>1.69</b>	BG672206	ubiquitin-like 3
Usp1	16	110	<b>6.90</b>	AW525342	ubiquitin specific peptidase 1
Usp32_predicte	20	84	<b>4.28</b>	AI231505	ubiquitin specific protease 32 (predicted)
Usp33	156	333	<b>2.13</b>	BE108244	Ubiquitin specific peptidase 33
<i>Usp47_predicte</i>	<i>119</i>	<i>197</i>	<i>1.66</i>	<i>AI407830</i>	<i>ubiquitin specific protease 47 (predicted)</i>
Usp7	156	381	<b>2.45</b>	AI175593	Ubiquitin specific protease 7 (herpes virus-associated)
Usp8_predicted	224	361	<b>1.61</b>	AI137190	ubiquitin specific protease 8 (predicted)
Wsb2	106	211	<b>2.00</b>	AA848210	WD repeat and SOCS box-containing 2

### Receptors & Binding Proteins

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio		
Atrn	455	301	<b>0.66</b>	AB038388	attractin
<i>Abca1</i>	<i>35</i>	<i>124</i>	<i>3.58</i>	<i>AI502114</i>	<i>ATP-binding cassette, sub-family A (ABC1), member 1</i>
Aebp1_predicte	64	97	<b>1.51</b>	BI278482	AE binding protein 1 (predicted)
Agtrap	158	103	<b>0.65</b>	BE349785	angiotensin II, type I receptor-associated protein
Apba2	317	513	<b>1.62</b>	NM_031780	amyloid beta (A4) precursor protein-binding, family A, n
Appbp2	43	122	<b>2.84</b>	BE113611	amyloid beta precursor protein (cytoplasmic tail) bindin
Bmpr1a	70	124	<b>1.77</b>	NM_030849	bone morphogenetic protein receptor, type 1A
Bmpr2	8	176	<b>22.08</b>	BE118651	bone morphogenetic protein receptor, type II (serine/thre
Chp /// RGD156	58	129	<b>2.23</b>	AB070350	calcium binding protein p22 /// similar to calcium bindin
<i>Crebl2</i>	<i>56</i>	<i>163</i>	<i>2.91</i>	<i>BE102391</i>	<i>cAMP responsive element binding protein-like 2</i>
Ednrb	21	230	<b>11.00</b>	X57764	endothelin receptor type B
Egfr	8	87	<b>11.17</b>	M37394	epidermal growth factor receptor
Epha7	72	114	<b>1.58</b>	U21955	Eph receptor A7
Esrrg	47	97	<b>2.06</b>	AA996810	Estrogen-related receptor gamma
Fgfr1	103	233	<b>2.26</b>	S54008	Fibroblast growth factor receptor 1
Gabbr1	7	161	<b>22.23</b>	Y10369	gamma-aminobutyric acid (GABA) B receptor 1
Gabbr3	25	117	<b>4.69</b>	NM_017065	gamma-aminobutyric acid (GABA-A) receptor, subunit I
Gabrg1	48	103	<b>2.13</b>	NM_080586	gamma-aminobutyric acid A receptor, gamma 1
Gpr21_predicte	622	368	<b>0.59</b>	BG374688	G protein-coupled receptor 21 (predicted)
Gpr3711	356	632	<b>1.78</b>	AF087947	G protein-coupled receptor 37-like 1
Gpr85	232	367	<b>1.58</b>	AF203907	G protein-coupled receptor 85
Gpr89_predicte	128	213	<b>1.67</b>	BI275972	G protein-coupled receptor 89 (predicted)
Glrb	128	195	<b>1.53</b>	NM_053296	glycine receptor, beta subunit
Grm3	247	725	<b>2.93</b>	AW522430	glutamate receptor, metabotropic 3
Grm5	59	572	<b>9.76</b>	AW526330	Glutamate receptor, metabotropic 5
Htr4	253	148	<b>0.58</b>	BF401317	5-hydroxytryptamine (serotonin) receptor 4
<b>Igfbp2</b>	<b>156</b>	<b>266</b>	<b>1.71</b>	<b>NM_013122</b>	<b>insulin-like growth factor binding protein 2</b>
Itn2b	15	100	<b>6.75</b>	AW534352	Integral membrane protein 2B

LOC498912	170	288	<b>1.69</b>	AA859108	similar to Alpha- and gamma-adaptin-binding protein p3
LOC681927 /// I	66	109	<b>1.65</b>	BF419095	similar to SEC24 related gene family, member C
LOC688018	74	506	<b>6.82</b>	BM389412	similar to SH3-domain binding protein 3
LOC689560 /// I	9	80	<b>9.30</b>	BF418135	ryanodine receptor 2, cardiac /// ryanodine receptor 2 (c
Lphn1	181	501	<b>2.76</b>	AF111099	latrophilin 1
Lrp6_predicted	18	125	<b>6.86</b>	BM391816	low density lipoprotein receptor-related protein 6 (predi
Nab1	51	78	<b>1.53</b>	NM_022856	Ngfi-A binding protein 1
Narg1_predicte	20	249	<b>12.33</b>	AW527638	NMDA receptor-regulated gene 1 (predicted)
Ncald	85	239	<b>2.82</b>	BF420172	neurocalcin delta
Nisch	182	342	<b>1.88</b>	BE108587	nischarin
Nrp2	593	364	<b>0.61</b>	AA859669	Neuropilin 2
Nrxn1	230	993	<b>4.32</b>	NM_021767	neurexin 1
Nrxn3	271	170	<b>0.63</b>	NM_053817	neurexin 3
Oprl1	123	81	<b>0.66</b>	NM_031569	opioid receptor-like 1
Osbp2_predicte	140	213	<b>1.53</b>	AI112698	oxysterol binding protein 2 (predicted)
Osbp1a	35	121	<b>3.41</b>	AI137224	oxysterol binding protein-like 1A
Otub1_predicte	155	331	<b>2.13</b>	BI274378	OTU domain, ubiquitin aldehyde binding 1 (predicted)
P34	41	102	<b>2.47</b>	NM_134398	p34 protein
Pgrmc2	84	179	<b>2.14</b>	AI137488	progesterone receptor membrane component 2
Plxdc2_predicte	34	242	<b>7.14</b>	BE116194	plexin domain containing 2 (predicted)
Plxnb1_predicte	58	95	<b>1.64</b>	BI294858	plexin B1 (predicted)
RGD1308847_f	37	81	<b>2.19</b>	AI716277	similar to SPla/Ryanodine receptor SPRY (1J970) (pre
RGD1559552_f	24	243	<b>10.01</b>	BF396678	similar to peroxisome proliferator-activated receptor bin
RGD1560070_f	92	901	<b>9.75</b>	BG380409	similar to ataxin 2-binding protein 1 isoform 2 (predicte
RGD1560587_f	238	590	<b>2.48</b>	AW525048	similar to Eph receptor A4 (predicted)
Rimbp2	66	170	<b>2.59</b>	BE102826	RIM binding protein 2
Rora_predicted	153	230	<b>1.50</b>	AI235414	RAR-related orphan receptor alpha (predicted)
Rtn4r	46	101	<b>2.20</b>	NM_053613	reticulon 4 receptor
Sec14l2	32	95	<b>2.96</b>	NM_053801	SEC14-like 2 (S. cerevisiae)
Sec61a2_predic	840	532	<b>0.63</b>	BG381529	Sec61, alpha subunit 2 (S. cerevisiae) (predicted)
Sh3bgr1_predict	261	411	<b>1.57</b>	BG672648	SH3-binding domain glutamic acid-rich protein like (pre
<b>Stxbp1</b>	<b>65</b>	<b>162</b>	<b>2.48</b>	<b>U06069</b>	<b>syntaxin binding protein 1</b>
Tbl1xr1_predict	27	112	<b>4.12</b>	AA944938	transducin (beta)-like 1X-linked receptor 1 (predicted)
Thra	15	97	<b>6.44</b>	M18028	thyroid hormone receptor alpha
Thrap2_predicte	132	436	<b>3.31</b>	BE109224	thyroid hormone receptor associated protein 2 (predicte
Wbp4	32	146	<b>4.58</b>	BG663444	VW domain binding protein 4

### Signaling

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio	Genbank	Gene Title
Abi1	311	1190	<b>3.83</b>	NM_024397	abl-interactor 1
Adnp	37	83	<b>2.25</b>	NM_022681	activity-dependent neuroprotective protein
Adrbk1	65	156	<b>2.40</b>	NM_012776	adrenergic receptor kinase, beta 1
<b>Agtpbp1_predi</b>	<b>48</b>	<b>103</b>	<b>2.13</b>	<b>BF553179</b>	<b>ATP/GTP binding protein 1 (predicted)</b>
Ak1	47	82	<b>1.75</b>	NM_024349	adenylate kinase 1
Akap13	72	178	<b>2.45</b>	AW526712	A kinase (PRKA) anchor protein 13
Anp32a	80	164	<b>2.05</b>	NM_012903	acidic (leucine-rich) nuclear phosphoprotein 32 family, r
Anp32e	98	149	<b>1.53</b>	AI008642	acidic (leucine-rich) nuclear phosphoprotein 32 family, r
Apln	44	121	<b>2.73</b>	NM_031612	apelin, AGTRL1 ligand
Arhgap1_predic	24	80	<b>3.37</b>	AI535143	Rho GTPase activating protein 1 (predicted)
Arhgap5	23	162	<b>7.07</b>	BM384008	Rho GTPase activating protein 5
Ascc3l1	81	143	<b>1.77</b>	AI502051	Activating signal cointegrator 1 complex subunit 3-like
Brinp2	89	154	<b>1.73</b>	BF551863	BMP/retinoic acid-inducible neural-specific protein 2
<b>Camk2a</b>	<b>565</b>	<b>867</b>	<b>1.53</b>	<b>BM384558</b>	<b>calcium/calmodulin-dependent protein kinase II, al</b>
Camk2b	1041	2482	<b>2.38</b>	NM_021739	calcium/calmodulin-dependent protein kinase II, beta
Camk2g	165	930	<b>5.62</b>	NM_133605	calcium/calmodulin-dependent protein kinase II gamma
Camk2n1	191	400	<b>2.09</b>	BE111167	Calcium/calmodulin-dependent protein kinase II inhibi
Canx	93	167	<b>1.79</b>	BI297860	calnexin
Cap1	172	434	<b>2.53</b>	NM_022383	CAP, adenylate cyclase-associated protein 1 (yeast)
Cap2	272	806	<b>2.96</b>	NM_053874	CAP, adenylate cyclase-associated protein, 2 (yeast)
Cd47	247	385	<b>1.56</b>	NM_019195	CD47 antigen (Rh-related antigen, integrin-associated
Cdc42bbp	23	78	<b>3.35</b>	AF021936	Cdc42 binding protein kinase beta
Centg2_predicte	275	650	<b>2.37</b>	BF558819	centaurin, gamma 2 (predicted)
Centg3_predicte	4	88	<b>24.82</b>	AW252124	centaurin, gamma 3 (predicted)
<b>Cit</b>	<b>98</b>	<b>269</b>	<b>2.75</b>	<b>AA957183</b>	<b>Citron</b>
Cnksr2	60	124	<b>2.06</b>	AF102853	connector enhancer of kinase suppressor of Ras 2
Csda	162	87	<b>0.54</b>	AA866458	cold shock domain protein A
Csnk1a1	158	253	<b>1.60</b>	U77583	casein kinase 1, alpha 1
Csnk1d	83	141	<b>1.71</b>	L07578	casein kinase 1, delta



Ctnnb1	70	162	<b>2.33</b>	NM_053357	catenin (cadherin associated protein), beta 1
Ehd4	62	103	<b>1.67</b>	BF395171	EH-domain containing 4
Ensa	117	311	<b>2.65</b>	BI289922	endosulfine alpha
Epha5	27	175	<b>6.49</b>	BF416432	EphA5
Farp1_predicted	16	118	<b>7.16</b>	AI547942	FERM, RhoGEF (Arhgef) and pleckstrin domain protein
Freq	25	77	<b>3.11</b>	NM_024366	frequenin homolog (Drosophila)
Gapvd1_predicti	161	247	<b>1.54</b>	BE102596	GTPase activating protein and VPS9 domains 1 (predic
Garnl1	92	140	<b>1.52</b>	AF041107	GTPase activating RANGAP domain-like 1
Gnao	189	726	<b>3.85</b>	AF413212	guanine nucleotide binding protein, alpha o
Gnb1	1029	526	<b>0.51</b>	NM_030987	guanine nucleotide binding protein, beta 1
Gnb2	384	657	<b>1.71</b>	AF277892	guanine nucleotide binding protein, beta polypeptide 2
Gria2	568	1541	<b>2.72</b>	M85035	glutamate receptor, ionotropic, AMPA2
Gria3	17	84	<b>4.93</b>	M85036	glutamate receptor, ionotropic, AMPA3 (alpha 3)
Grin1a	104	197	<b>1.90</b>	BG664035	glutamate receptor, ionotropic, N-methyl D-aspartate-lik
Grip1	77	51	<b>0.67</b>	AI408747	Glutamate receptor interacting protein 1
Hipk2_predicted	73	123	<b>1.69</b>	BM392321	homeodomain interacting protein kinase 2 (predicted)
Hpcal4	82	171	<b>2.09</b>	BI296503	Hippocalcin-like 4
Hspa2	194	129	<b>0.67</b>	BF410146	heat shock 70kDa protein 2
Hspa4	62	153	<b>2.46</b>	AF077354	heat shock protein 4
Hspca	295	2016	<b>6.82</b>	BG671521	heat shock protein 1, alpha
Hspcb	249	904	<b>3.63</b>	BI285700	heat shock 90kDa protein 1, beta
Ifi271	1002	585	<b>0.58</b>	NM_130743	interferon, alpha-inducible protein 27-like
Kalrn	60	194	<b>3.22</b>	NM_032062	kalirin, RhoGEF kinase
Khdrbs1	70	131	<b>1.87</b>	AF393783	KH domain containing, RNA binding, signal transductio
LOC289606	164	328	<b>2.00</b>	BF404116	similar to Gamma-aminobutyric-acid receptor alpha-2 s
LOC365949 /// I	75	247	<b>3.29</b>	AI111738	similar to nemo like kinase /// similar to nemo like kinas
LOC497978	103	234	<b>2.26</b>	AI101639	similar to diacylglycerol kinase epsilon
LOC498351	295	190	<b>0.64</b>	BG670238	similar to signal recognition particle,72 kDa subunit
LOC678826 /// I	548	883	<b>1.61</b>	BI295776	similar to sorbin and SH3 domain containing 1 isoform :
LOC679252	16	136	<b>8.73</b>	AI070638	similar to Myeloid/lymphoid or mixed-lineage leukemia
LOC680419	218	132	<b>0.61</b>	AA963364	Similar to Ras suppressor protein 1
LOC682488 /// I	87	178	<b>2.04</b>	AI711306	similar to Ras-related protein Rab-1B
LOC683917 /// I	95	353	<b>3.70</b>	AI102520	similar to gamma-aminobutyric acid (GABA(A)) recepto
Lsg1	116	30	<b>0.26</b>	AA899663	large subunit GTPase 1 homolog (S. cerevisiae)
Pygm	33	90	<b>2.70</b>	AI717476	muscle glycogen phosphorylase
Map4k4_predicti	162	91	<b>0.56</b>	BE117850	mitogen-activated protein kinase kinase kinase
Mapk1	101	221	<b>2.18</b>	NM_053842	mitogen activated protein kinase 1
Mapk10	168	263	<b>1.56</b>	NM_012806	mitogen activated protein kinase 10
Mapk3	207	355	<b>1.71</b>	AF155236	mitogen activated protein kinase 3
Mapk8ip	144	282	<b>1.96</b>	AF092450	mitogen activated protein kinase 8 interacting protein
Mapk8ip3	82	224	<b>2.72</b>	BF553848	mitogen-activated protein kinase 8 interacting protein 3
Mapk9	40	136	<b>3.38</b>	NM_017322	mitogen-activated protein kinase 9
MGC125015	55	125	<b>2.28</b>	AA996838	similar to PAK/PLC-interacting protein 1
Nek7_predicted	69	182	<b>2.65</b>	BE117215	NIMA (never in mitosis gene a)-related expressed kinas
Nkiras1_predict	284	622	<b>2.19</b>	AA997472	NFKB inhibitor interacting Ras-like protein 1 (predicted)
Notch1	55	85	<b>1.55</b>	BF389398	Notch gene homolog 1 (Drosophila)
Ntrk2	1283	766	<b>0.60</b>	BE102996	Neurotrophic tyrosine kinase, receptor, type 2
Nucks	66	273	<b>4.11</b>	NM_022799	nuclear ubiquitous casein kinase and cyclin-dependent
Pacsin1	31	430	<b>13.84</b>	NM_017294	protein kinase C and casein kinase substrate in neuron
Pak3	220	421	<b>1.91</b>	NM_019210	p21 (CDKN1A)-activated kinase 3
Pde10a	260	587	<b>2.26</b>	NM_022236	phosphodiesterase 10A
Pde4b	129	296	<b>2.29</b>	AF202733	phosphodiesterase 4B, cAMP specific
Pik3c2a_predicti	198	321	<b>1.62</b>	AI059449	phosphatidylinositol 3-kinase, C2 domain containing, al
Pik3r2	122	235	<b>1.93</b>	NM_022185	phosphatidylinositol 3-kinase, regulatory subunit, polyp
Pip5k1a_predict	41	107	<b>2.65</b>	BG153281	phosphatidylinositol-4-phosphate 5-kinase, type 1, alph
Pip5k2a	40	87	<b>2.14</b>	NM_053926	phosphatidylinositol-4-phosphate 5-kinase, type II, alph
Pkia	123	199	<b>1.62</b>	AA996685	protein kinase inhibitor, alpha
Plaa	110	213	<b>1.95</b>	NM_053866	phospholipase A2, activating protein
<i>Plekhb2_predicti</i>	10	113	<b>11.39</b>	<i>AW254369</i>	<i>pleckstrin homology domain containing, family B (evect</i>
<b>Plekhh1_predi</b>	<b>78</b>	<b>25</b>	<b>0.32</b>	<b>BI275435</b>	<b>pleckstrin homology domain containing, family H (v</b>
Plekhj1	137	83	<b>0.60</b>	AA848429	pleckstrin homology domain containing, family J membe
Ppp1ca	906	1416	<b>1.56</b>	NM_031527	protein phosphatase 1, catalytic subunit, alpha isoform
Ppp1r12a	19	168	<b>8.94</b>	BF406594	protein phosphatase 1, regulatory (inhibitor) subunit 12
Ppp1r3c	69	159	<b>2.30</b>	AW530361	protein phosphatase 1, regulatory (inhibitor) subunit 3C
Ppp1r7	139	248	<b>1.78</b>	H32309	protein phosphatase 1, regulatory (inhibitor) subunit 7
Ppp2r1a	57	226	<b>3.97</b>	BF394544	protein phosphatase 2 (formerly 2A), regulatory subunit
Ppp2r5e_predict	105	211	<b>2.01</b>	BG664036	protein phosphatase 2, regulatory subunit B (B56), epsi
<i>Ppp3ca</i>	20	76	<b>3.90</b>	<i>BF388224</i>	<i>Protein phosphatase 3, catalytic subunit, alpha isoform</i>
Ppp3r1	87	379	<b>4.37</b>	NM_017309	protein phosphatase 3, regulatory subunit B, alpha isof

Ppp5c	114	185	<b>1.62</b>	NM_031729	protein phosphatase 5, catalytic subunit
Prkaa2	22	187	<b>8.41</b>	NM_023991	protein kinase, AMP-activated, alpha 2 catalytic subunit
Prkcbp1	230	602	<b>2.62</b>	BM391419	protein kinase C binding protein 1
Prkce	618	1719	<b>2.78</b>	AA799421	protein kinase C, epsilon
<i>Prkce</i>	89	34	<b>0.38</b>	BI301465	<i>Protein kinase C, epsilon</i>
Prkwk1	52	81	<b>1.55</b>	NM_053794	protein kinase, lysine deficient 1
<b>Ptgds</b>	<b>2735</b>	<b>4345</b>	<b>1.59</b>	<b>J04488</b>	<b>prostaglandin D2 synthase</b>
Ptp4a2	327	985	<b>3.02</b>	NM_053475	protein tyrosine phosphatase 4a2
Ptpn11	81	45	<b>0.55</b>	AI535048	Protein tyrosine phosphatase, non-receptor type 11
Ptpns1	34	99	<b>2.96</b>	D38468	protein tyrosine phosphatase, non-receptor type substr
Ptprd	48	289	<b>6.08</b>	L19933	protein tyrosine phosphatase, receptor type, D
Ptprf	51	263	<b>5.16</b>	X83505	protein tyrosine phosphatase, receptor type, F
<i>Ptprj</i>	139	231	<b>1.67</b>	<i>NM_017269</i>	<i>protein tyrosine phosphatase, receptor type, J</i>
Rab10	136	219	<b>1.62</b>	NM_017359	RAB10, member RAS oncogene family
Rab11b	139	298	<b>2.15</b>	D01046	RAB11B, member RAS oncogene family
Rab14	96	706	<b>7.37</b>	NM_053589	RAB14, member RAS oncogene family
Rab15	41	96	<b>2.31</b>	BF554320	RAB15, member RAS oncogene family
Rab5a	80	189	<b>2.36</b>	NM_022692	RAB5A, member RAS oncogene family
Rab5b_predicte	71	129	<b>1.82</b>	AA996576	RAB5B, member RAS oncogene family (predicted)
Rab6a	112	271	<b>2.43</b>	AA686757	RAB6A, member RAS oncogene family
Rab7	175	314	<b>1.79</b>	NM_023950	RAB7, member RAS oncogene family
Ralgs2	21	94	<b>4.51</b>	BF525299	Ral GEF with PH domain and SH3 binding motif 2
Ramp2	252	166	<b>0.66</b>	NM_031646	receptor (calcitonin) activity modifying protein 2
Rapgef6_predic	29	146	<b>5.04</b>	BI289486	Rap guanine nucleotide exchange factor (GEF) 6 (pred
Rasa1	126	217	<b>1.73</b>	NM_013135	RAS p21 protein activator 1
Rasgrf2	69	183	<b>2.64</b>	NM_053721	RAS protein-specific guanine nucleotide-releasing fact
Rasgrp1	82	241	<b>2.95</b>	AF081196	RAS guanyl releasing protein 1
Rasl10b_predic	58	102	<b>1.76</b>	AW919370	RAS-like, family 10, member B (predicted)
Rfng	61	116	<b>1.89</b>	NM_021849	radical fringe gene homolog (Drosophila)
RGD1309388_f	127	561	<b>4.43</b>	BF544981	similar to DIP13 alpha (predicted)
RGD1311455_f	90	230	<b>2.55</b>	BE108260	Similar to MAPK-interacting and spindle-stabilizing prot
RGD1560049_f	57	529	<b>9.24</b>	AW520764	similar to Dual specificity protein phosphatase 3 (T-DSF
RGD1561090_f	86	659	<b>7.62</b>	AI412803	similar to protein tyrosine phosphatase, receptor type, I
RGD1561817_f	85	129	<b>1.51</b>	BG666454	similar to Traf2 and NCK interacting kinase, splice vari
RGD1564560_f	27	342	<b>12.73</b>	AA850780	similar to RCK (predicted)
Rgs17_predicte	124	476	<b>3.83</b>	AI229118	regulator of G-protein signaling 17 (predicted)
Rgs4	68	176	<b>2.58</b>	U27767	regulator of G-protein signaling 4
<i>RICS_predictec</i>	95	248	<b>2.61</b>	<i>BE097238</i>	<i>RhoGAP involved in beta-catenin-N-cadherin and NMD</i>
Rims1	114	224	<b>1.97</b>	NM_052829	regulating synaptic membrane exocytosis 1
Riok3_predictec	40	79	<b>1.98</b>	BE126478	RIO kinase 3 (yeast) (predicted)
Rnd3	29	98	<b>3.37</b>	AI598323	Rho family GTPase 3
Sbf1_predicted	36	156	<b>4.39</b>	BF554377	SET binding factor 1 (predicted)
Set_predicted	601	1057	<b>1.76</b>	BG381708	SET translocation (predicted)
Sipa111	78	180	<b>2.30</b>	AF026504	signal-induced proliferation-associated 1 like 1
Snx10	39	105	<b>2.73</b>	AI007889	sorting nexin 10
Snx12_predicte	63	229	<b>3.65</b>	BF389280	Sorting nexin 12 (predicted)
Snx27	34	489	<b>14.57</b>	AA900057	sorting nexin family member 27
Snx4_predicted	67	179	<b>2.66</b>	AI029221	Sorting nexin 4 (predicted)
Shoc2	36	167	<b>4.66</b>	AI112329	soc-2 (suppressor of clear) homolog (C. elegans)
Sorcs3_predicte	35	95	<b>2.73</b>	BF409626	sortilin-related VPS10 domain containing receptor 3 (pr
Synj2	75	138	<b>1.86</b>	AY034051	synaptojanin 2
Taok1	34	145	<b>4.21</b>	AF084205	TAO kinase 1
Tbl1x_predictec	449	294	<b>0.65</b>	BE108225	transducin (beta)-like 1 X-linked (predicted)
Tiam1	22	87	<b>3.91</b>	BM389265	T-cell lymphoma invasion and metastasis 1
Tollip_predicted	85	143	<b>1.68</b>	BE112982	toll interacting protein (predicted)
Ywhab	214	493	<b>2.31</b>	BG380730	tyrosine 3-monooxygenase/tryptophan 5-monooxygena

### Transcription

Sample	F3-Cont	F3-Vinc	Vin/Con	Gene Symbol	Gene Title
Ahctf1_predicte	49	78	<b>1.60</b>	BF396355	AT hook containing transcription factor 1 (predicted)
Ahdc1_predicte	32	109	<b>3.40</b>	BG376923	AT hook, DNA binding motif, containing 1 (predicted)
Ank3	116	194	<b>1.68</b>	AJ428573	ankyrin 3, epithelial
Ankrd52_predic	52	294	<b>5.60</b>	AA964250	ankyrin repeat domain 52 (predicted)
Arid2_predicted	46	110	<b>2.37</b>	BI289741	AT rich interactive domain 2 (Arid-rfx like) (predicted)
Arid4a_predicte	24	138	<b>5.70</b>	BF391127	AT rich interactive domain 4A (Rbp1 like) (predicted)
Arid4b	66	135	<b>2.07</b>	NM_053421	AT rich interactive domain 4B (Rbp1 like)
Basp1	749	1185	<b>1.58</b>	NM_022300	brain abundant, membrane attached signal protein 1
Bcl11b_predicte	14	120	<b>8.85</b>	BF398531	B-cell leukemia/lymphoma 11B (predicted)

Bcr_predicted	91	627	<b>6.87</b>	BE120831	breakpoint cluster region (predicted)
Bhlhb2	498	299	<b>0.60</b>	AI548256	Basic helix-loop-helix domain containing, class B2
Bicap	53	101	<b>1.89</b>	NM_133582	bladder cancer associated protein homolog (human)
Cbfa211_predict	55	107	<b>1.96</b>	BE105678	CBFA2T1 identified gene homolog (human) (predicted)
Cdc42se1	252	163	<b>0.65</b>	BI284508	CDC42 small effector 1
Ddx17	67	141	<b>2.11</b>	BI296754	DEAD (Asp-Glu-Ala-Asp) box polypeptide 17
Ddx24	70	652	<b>9.32</b>	AW252511	DEAD (Asp-Glu-Ala-Asp) box polypeptide 24
Ddx3x	32	117	<b>3.64</b>	BF391513	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 3, X-irr
Dhx36_predicte	27	832	<b>31.27</b>	BE114021	DEAH (Asp-Glu-Ala-His) box polypeptide 36 (predicted)
Dmtf1	807	529	<b>0.66</b>	BE098802	cyclin D binding myb-like transcription factor 1
<b>Dnaja4</b>	<b>17</b>	<b>79</b>	<b>4.64</b>	<b>BF565278</b>	<b>DnaJ (Hsp40) homolog, subfamily A, member 4</b>
Dnajb1_predicte	137	226	<b>1.65</b>	BM384926	DnaJ (Hsp40) homolog, subfamily B, member 1 (predic
Dnajb6	10	75	<b>7.88</b>	AW528827	DnaJ (Hsp40) homolog, subfamily B, member 6
Dnajc7	126	367	<b>2.92</b>	BI285682	DnaJ (Hsp40) homolog, subfamily C, member 7
Dnd1	36	89	<b>2.48</b>	BF552813	dead end homolog 1 (zebrafish)
Etv1_predicted	72	131	<b>1.83</b>	AI112936	Ets variant gene 1 (predicted)
Fbxo9	160	252	<b>1.58</b>	BG669135	F-box only protein 9
Fbxw2_predicte	49	79	<b>1.63</b>	AW915859	F-box and WD-40 domain protein 2 (predicted)
Foxo1a	13	153	<b>11.98</b>	BI295511	forkhead box O1A
Fubp1	121	77	<b>0.64</b>	AI233709	Far upstream element (FUSE) binding protein 1
Fyn	191	358	<b>1.88</b>	NM_012755	fyn proto-oncogene
Gtf2i	83	146	<b>1.76</b>	AI715835	general transcription factor II I
Hip1	282	159	<b>0.56</b>	BE110671	huntingtin interacting protein 1
Impad1	119	287	<b>2.41</b>	BF396528	inositol monophosphatase domain containing 1
Ing3	58	169	<b>2.92</b>	BI296751	inhibitor of growth family, member 3
Jun	179	331	<b>1.84</b>	BI288619	Jun oncogene
Kit	33	76	<b>2.29</b>	NM_022264	v-kit Hardy-Zuckerman 4 feline sarcoma viral oncogene
Klf6	109	186	<b>1.71</b>	NM_031642	Kruppel-like factor 6
Klhl24	116	451	<b>3.90</b>	BF550390	kelch-like 24 (Drosophila)
LOC303057	31	116	<b>3.71</b>	AI411326	similar to step II splicing factor SLU7; DNA segment, CI
LOC309957	81	437	<b>5.41</b>	BI281877	similar to myocyte enhancer factor 2C
LOC362154 /// I	215	749	<b>3.49</b>	BF550329	brain zinc finger protein /// similar to brain Zn-finger pro
LOC500591	127	720	<b>5.66</b>	BF564825	similar to calmodulin-binding transcription activator 1
LOC678886 /// I	405	687	<b>1.70</b>	BE109666	similar to Enhancer of rudimentary homolog
LOC679725 /// I	35	133	<b>3.81</b>	BI296626	similar to MASK-4E-BP3 protein
LOC680802 /// I	20	83	<b>4.11</b>	BE101875	similar to Zinc finger protein 45 (BRC1744)
LOC685277	131	721	<b>5.50</b>	BE104102	Similar to liver-specific bHLH-Zip transcription factor
<b>LOC685374</b>	<b>100</b>	<b>151</b>	<b>1.52</b>	<b>BG671371</b>	<b>similar to ankyrin repeat domain 13c</b>
LOC685707 /// I	31	83	<b>2.64</b>	BF387266	neuron navigator 1 (predicted) /// similar to neuron navi
LOC685865	280	178	<b>0.64</b>	BG371744	Similar to zinc finger protein 458
LOC686480 /// I	237	91	<b>0.38</b>	AW528215	similar to Protein C12orf11 (Sarcoma antigen NY-SAR-1
LOC687516	112	180	<b>1.61</b>	BI296193	similar to zinc finger protein 146
LOC688133	255	170	<b>0.67</b>	BE114154	similar to mbt domain containing 1
LOC688144 /// I	65	137	<b>2.12</b>	BE105705	similar to ankyrin repeat domain 40
LOC690038	65	113	<b>1.74</b>	AI009167	Similar to Bcl2-associated athanogene 2
Mcf2l	20	80	<b>4.00</b>	NM_053951	mcf.2 transforming sequence-like
Mll	74	120	<b>1.62</b>	BG671918	myeloid/lymphoid or mixed-lineage leukemia
Mll5	33	227	<b>6.77</b>	BE117891	myeloid/lymphoid or mixed-lineage leukemia 5 (trithora:
Mpp5_predictec	39	235	<b>6.08</b>	AI072027	membrane protein, palmitoylated 5 (MAGUK p55 subfa
Mpp6_predictec	89	193	<b>2.17</b>	BE097259	membrane protein, palmitoylated 6 (MAGUK p55 subfa
Mta1	53	103	<b>1.93</b>	AJ132046	metastasis associated 1
Mtpn	129	334	<b>2.59</b>	AI711244	myotrophin
Mxd4_predictec	34	113	<b>3.35</b>	BI274201	Max dimerization protein 4 (predicted)
Mxi1	52	103	<b>1.96</b>	NM_013160	Max interacting protein 1
Mycl1	108	186	<b>1.72</b>	BI300996	v-myc myelocytomatosis viral oncogene homolog 1, lur
Ncor1	37	240	<b>6.54</b>	BE100543	nuclear receptor co-repressor 1
Nfia	36	80	<b>2.23</b>	D78019	nuclear factor I/A
Nfib	178	842	<b>4.73</b>	BE099050	nuclear factor I/B
Nfix	48	290	<b>6.09</b>	BF420722	nuclear factor I/X
Nfyc	38	84	<b>2.23</b>	NM_012866	nuclear transcription factor-Y gamma
Nol6_predicted	74	114	<b>1.53</b>	AI409184	nucleolar protein family 6 (RNA-associated) (predicted)
Nolc1	114	175	<b>1.54</b>	M94288	nucleolar and coiled-body phosphoprotein 1
Nono	121	222	<b>1.84</b>	BF555394	non-POU domain containing, octamer-binding
Npat_predicted	101	67	<b>0.66</b>	AI014116	nuclear protein in the AT region (predicted)
Npat_predicted	122	64	<b>0.52</b>	BF391717	Nuclear protein in the AT region (predicted)
Nr1h2	62	114	<b>1.84</b>	AI169222	nuclear receptor subfamily 1, group H, member 2
Nr2f2	10	79	<b>7.83</b>	NM_080778	nuclear receptor subfamily 2, group F, member 2
Nr2f6	56	91	<b>1.63</b>	NM_139113	nuclear receptor subfamily 2, group F, member 6
Nr4a2	61	118	<b>1.94</b>	U72345	nuclear receptor subfamily 4, group A, member 2

Nr4a3	525	1096	<b>2.09</b>	NM_031628	nuclear receptor subfamily 4, group A, member 3
<b>Nrip3_predicted</b>	<b>135</b>	<b>237</b>	<b>1.76</b>	<b>BI288541</b>	<b>Nuclear receptor interacting protein 3 (predicted)</b>
NTF2	70	117	<b>1.67</b>	BE109988	nuclear transport factor 2
Nucb1	259	397	<b>1.53</b>	Z36277	nucleobindin 1
Os-9	132	88	<b>0.66</b>	AI169058	amplified in osteosarcoma
Plag1	69	115	<b>1.66</b>	BI290063	pleiomorphic adenoma gene 1
Pogz_predicted	34	75	<b>2.23</b>	AI177507	pogo transposable element with ZNF domain (predicted)
Psip1	72	111	<b>1.54</b>	BF397366	PC4 and SFRS1 interacting protein 1
pur-beta	52	92	<b>1.79</b>	BI284455	transcription factor Pur-beta
Rala	87	344	<b>3.96</b>	NM_031093	v-ral simian leukemia viral oncogene homolog A (ras re
Recc1	137	476	<b>3.47</b>	BM391856	replication factor C 1
Rexo4	44	102	<b>2.31</b>	AI237143	REX4, RNA exonuclease 4 homolog (S. cerevisiae)
RGD1305133	36	352	<b>9.69</b>	BE116128	similar to Ab2-008
RGD1306327	169	457	<b>2.71</b>	AW530286	similar to downregulated in renal cell carcinoma
RGD1307526	292	190	<b>0.65</b>	BE114469	similar to modulator of estrogen induced transcription
RGD1310712_f	68	127	<b>1.87</b>	BI297740	similar to EMSY protein (predicted)
RGD1562348_f	338	811	<b>2.40</b>	BE117929	similar to ankyrin repeat domain protein 17 isoform b (p
RGD1562639_f	49	86	<b>1.74</b>	BG670537	similar to c-myc promoter binding protein (predicted)
RGD1562686_f	37	163	<b>4.41</b>	BI288731	similar to genetic suppressor element 1 (predicted)
<i>RGD1563119_f</i>	<i>125</i>	<i>841</i>	<b>6.75</b>	<i>BE104219</i>	<i>similar to MADS box transcription enhancer factor 2, pc</i>
RGD1564056_f	2	77	<b>51.63</b>	AI500735	similar to cellular repressor of E1A-stimulated genes 2
RGD1565099_f	10	174	<b>17.42</b>	AI045857	similar to BTEB3 protein (predicted)
RGD1565496_f	180	401	<b>2.22</b>	BF283779	similar to Butyrate-induced transcript 1 (predicted)
RGD1565549_f	88	307	<b>3.47</b>	BE113614	similar to polybromo-1 (predicted)
RGD1565591_f	140	262	<b>1.87</b>	BG380847	similar to Ski protein (predicted)
RGD1565646_f	32	121	<b>3.74</b>	BF548737	similar to SOX2 protein (predicted)
Rtel1	83	53	<b>0.64</b>	BI274548	regulator of telomere elongation helicase 1
Sdccag33_pred	131	62	<b>0.48</b>	BF411100	serologically defined colon cancer antigen 33 (predicted)
Sertad2	59	146	<b>2.46</b>	BI295862	SERTA domain containing 2
Sp1	39	163	<b>4.22</b>	AI705174	Sp1 transcription factor
Sreb1	37	100	<b>2.74</b>	AF286470	sterol regulatory element binding factor 1
Sreb2	763	482	<b>0.63</b>	AI170663	sterol regulatory element binding factor 2 (predicted)
Sox4_predicted	66	101	<b>1.53</b>	BI297183	SRY-box containing gene 4 (predicted)
Tcf20	108	168	<b>1.55</b>	BF394639	transcription factor 20
Tcf4	269	1307	<b>4.86</b>	NM_053369	transcription factor 4
Thoc2_predicted	26	139	<b>5.26</b>	AI556851	THO complex 2 (predicted)
Tnks2_predicted	51	121	<b>2.39</b>	H32233	tankyrase, TRF1-interacting ankyrin-related ADP-ribose
Tox_predicted	64	107	<b>1.69</b>	AI101139	thymocyte selection-associated HMG box gene (predic
Tpr	9	98	<b>11.44</b>	BE118639	translocated promoter region
Tspyl4	629	1359	<b>2.16</b>	BI281738	TSPY-like 4
Ubn1_predicted	10	108	<b>10.94</b>	AI578566	ubinnuclein 1 (predicted)
Ubtf	51	90	<b>1.77</b>	AI407688	upstream binding transcription factor, RNA polymerase
Usf1	59	129	<b>2.20</b>	NM_031777	upstream transcription factor 1
VeZF1_predicted	41	77	<b>1.86</b>	BI284190	vascular endothelial zinc finger 1 (predicted)
Zbtb20_predicted	155	543	<b>3.50</b>	BF415114	zinc finger and BTB domain containing 20 (predicted)
Zbtb4_predicted	1082	681	<b>0.63</b>	BM384146	zinc finger and BTB domain containing 4 (predicted)
Zcchc7_predicted	74	285	<b>3.84</b>	AI535316	Zinc finger, CCHC domain containing 7 (predicted)
Zfhx1b	28	176	<b>6.28</b>	BG377397	zinc finger homeobox 1b
Zfhx2	10	83	<b>7.96</b>	BI284196	zinc finger homeobox 2
Zfp179	553	368	<b>0.67</b>	AF054586	zinc finger protein 179
Zfp180	542	880	<b>1.62</b>	U41164	zinc finger protein 180
Zfp238	29	112	<b>3.89</b>	NM_022678	zinc finger protein 238
Zfp265	111	182	<b>1.64</b>	AF013965	zinc finger protein 265
Zfp365	70	157	<b>2.23</b>	BF408799	zinc finger protein 365
Zfp91	60	710	<b>11.78</b>	BE111631	zinc finger protein 91
Znf142_predicted	46	93	<b>2.00</b>	BE117672	Zinc finger protein 142 (clone pHZ-49) (predicted)
Znf292	11	218	<b>19.22</b>	BF388585	zinc finger protein 292
---	104	427	<b>4.13</b>	BE099838	Non-coding RNA expressed in the brain, repeat sequer
---	1036	3621	<b>3.49</b>	AA963228	Non-coding RNA expressed in the brain, repeat sequer

#### Translation & Protein Modification

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio		
Adarb1	32	117	<b>3.67</b>	NM_012894	adenosine deaminase, RNA-specific, B1
Arf1	478	955	<b>2.00</b>	NM_022518	ADP-ribosylation factor 1
Arf3	129	298	<b>2.32</b>	NM_080904	ADP-ribosylation factor 3
Arl1	350	554	<b>1.58</b>	NM_022385	ADP-ribosylation factor-like 1
Arl2bp	81	223	<b>2.74</b>	BF567478	ADP-ribosylation factor-like 2 binding protein
Arl6ip5	322	584	<b>1.82</b>	NM_023972	ADP-ribosylation factor-like 6 interacting protein 5

<i>Brunol4_predict</i>	107	55	<b>0.52</b>	AW524497	<i>Bruno-like 4, RNA binding protein (Drosophila) (predict</i>
Cct6a	33	76	<b>2.34</b>	AA875047	Chaperonin subunit 6a (zeta)
Cugbp1	271	474	<b>1.75</b>	BF408840	CUG triplet repeat, RNA binding protein 1
Cugbp2	94	209	<b>2.23</b>	AF090695	CUG triplet repeat, RNA binding protein 2
Dazap2	263	425	<b>1.62</b>	AI102612	DAZ associated protein 2
Eif2s3x	32	97	<b>3.07</b>	BG662620	eukaryotic translation initiation factor 2, subunit 3, struc
Eif4a1	340	597	<b>1.76</b>	BI284436	eukaryotic translation initiation factor 4A1
Eif4ebp2	189	311	<b>1.64</b>	BF419161	eukaryotic translation initiation factor 4E binding protei
Eif4g2 /// Eif4g2	230	406	<b>1.77</b>	BG672378	eukaryotic translation initiation factor 4 gamma, 2 /// eul
Eif5	116	928	<b>8.03</b>	BE107346	eukaryotic translation initiation factor 5
Eif5a	590	1119	<b>1.90</b>	BI283681	eukaryotic translation initiation factor 5A
Eif5b /// LOC68	77	147	<b>1.90</b>	BE117773	eukaryotic translation initiation factor 5B /// similar to Eu
Eprs	49	282	<b>5.79</b>	BF553211	glutamyl-prolyl-tRNA synthetase
Etf1	183	293	<b>1.60</b>	BF285301	eukaryotic translation termination factor 1
Hnrpa2b1_pred	64	213	<b>3.33</b>	BG666358	heterogeneous nuclear ribonucleoprotein A2/B1 (predic
Hnrpa3	103	181	<b>1.76</b>	AI409455	Heterogeneous nuclear ribonucleoprotein A3
Hnrpu	829	1264	<b>1.52</b>	AI177494	heterogeneous nuclear ribonucleoprotein U
Hnrpu1_predict	69	108	<b>1.56</b>	BG381219	heterogeneous nuclear ribonucleoprotein U-like 1 (pred
Khsrp	251	750	<b>2.98</b>	BI295086	KH-type splicing regulatory protein
Larp5_predictec	266	170	<b>0.64</b>	AI228249	La ribonucleoprotein domain family, member 5 (predicte
LOC297481	106	238	<b>2.24</b>	AI575608	similar to eukaryotic translation initiation factor 4E mem
LOC502603	38	146	<b>3.86</b>	AI145768	similar to splicing factor p54
LOC680726	163	248	<b>1.52</b>	AW144676	similar to RNA binding motif, single stranded interacting
LOC686858 /// I	73	110	<b>1.51</b>	AW531791	similar to small nuclear RNA activating complex, polype
Pabpn1	358	555	<b>1.55</b>	AW525563	poly(A) binding protein, nuclear 1
Papola_predicte	61	299	<b>4.88</b>	BE108853	poly (A) polymerase alpha (predicted)
Pum1_predictec	54	140	<b>2.58</b>	BI282129	pumilio 1 (Drosophila) (predicted)
Qki	143	240	<b>1.68</b>	BE102226	Quaking homolog, KH domain RNA binding (mouse)
RAMP4	79	138	<b>1.75</b>	AI103695	ribosome associated membrane protein 4
Rbm14	27	75	<b>2.76</b>	BG670091	RNA binding motif protein 14
Rbm25_predicte	54	210	<b>3.92</b>	AW434972	RNA binding motif protein 25 (predicted) /// similar to RI
Rbm5	261	777	<b>2.98</b>	BI293987	RNA binding motif protein 5
Rbm9_predictec	90	543	<b>6.04</b>	BE111095	RNA binding motif protein 9 (predicted)
RGD1560397_f	447	808	<b>1.81</b>	BF284305	similar to RNA-binding protein Musashi2-S (predicted)
Rnase4	191	290	<b>1.52</b>	NM_020082	ribonuclease, RNase A family 4
Rnpc2	185	390	<b>2.11</b>	BG372903	RNA-binding region (RNP1, RRM) containing 2
Rps6ka5_predic	21	154	<b>7.26</b>	AI045780	ribosomal protein S6 kinase, polypeptide 5 (predicted)
Serpbp1	1544	2586	<b>1.68</b>	AF388527	Serpine1 mRNA binding protein 1
Sf3b1	66	1024	<b>15.56</b>	BM384277	splicing factor 3b, subunit 1
Sf3b2_predictec	39	99	<b>2.52</b>	BE098910	splicing factor 3b, subunit 2 (predicted)
Tsn	90	181	<b>2.02</b>	NM_021762	translin

#### Miscellaneous & Unknown

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	<b>Ratio</b>	Genbank	Gene Title
---	93	200	<b>2.14</b>	AW535280	Retroviral-like ovarian specific transcript 30-1 mRNA
---	249	468	<b>1.88</b>	BI303933	Transcribed locus, strongly similar to XP_226779.4 zin
Aff4_predicted	5	86	<b>16.45</b>	AI556957	AF4/FMR2 family, member 4 (predicted)
Dmxl1_predicte	37	100	<b>2.68</b>	AA924048	Dmx-like 1 (predicted)
Fytd1	16	107	<b>6.53</b>	AI501447	forty-two-three domain containing 1
Gramd3	101	260	<b>2.58</b>	C06752	GRAM domain containing 3
LOC289378	117	70	<b>0.60</b>	BI288621	similar to B0432.8
LOC308398	49	101	<b>2.06</b>	BE107128	similar to F28C1.3a
LOC363544	379	241	<b>0.64</b>	AI012506	similar to 2610111M03Rik protein
LOC498962	118	74	<b>0.63</b>	BM387366	similar to C50H11.1
LOC681715 /// I	48	177	<b>3.67</b>	BF416086	similar to CG9047-PA, isoform A
LOC683839 /// I	98	217	<b>2.22</b>	AW921279	hypothetical protein LOC683839 /// hypothetical protein
LOC686503 /// I	88	200	<b>2.27</b>	AA944330	similar to CDC42 small effector 2 (predicted) /// hypothe
LOC687166 /// I	51	120	<b>2.37</b>	BF420090	similar to gene trap ROSA b-geo 22
LOC690789 /// I	81	210	<b>2.59</b>	AI104518	similar to product is unknown~seizure-related gene (pr
Lysmd3	78	175	<b>2.26</b>	AI071181	LysM, putative peptidoglycan-binding, domain containir
Mfhas1_predict	49	140	<b>2.88</b>	AI136864	malignant fibrous histiocytoma amplified sequence 1 (p
MGC94335	45	78	<b>1.72</b>	BF404452	similar to hypothetical protein FLJ22555
Nudcd1_predict	72	116	<b>1.61</b>	AI408455	NudC domain containing 1 (predicted)
Ociad1	99	59	<b>0.60</b>	BI275477	OClA domain containing 1
Perq1_predictec	16	101	<b>6.13</b>	AW526967	PERQ amino acid rich, with GYF domain 1 (predicted)
PNAS-4	154	500	<b>3.25</b>	AW913871	CGI-146 protein
RGD1306101_f	37	233	<b>6.27</b>	BE095778	similar to 4933407C03Rik protein (predicted)
RGD1306694_f	87	142	<b>1.64</b>	BF523573	similar to hypothetical protein (predicted)

RGD1307615_f	25	142	<b>5.80</b>	AI407061	similar to hypothetical protein FLJ13045 (predicted)
RGD1307729_f	68	113	<b>1.65</b>	BF419602	similar to KIAA0853 protein (predicted)
RGD1307907_f	158	99	<b>0.63</b>	AI146080	similar to hypothetical protein FLJ14681 (predicted)
RGD1307981_f	78	144	<b>1.85</b>	BF408881	similar to cisplatin resistance-associated overexpresser
RGD1308329_f	178	288	<b>1.62</b>	BM390702	similar to KIAA0869 protein (predicted)
RGD1308795_f	42	129	<b>3.05</b>	AW143311	similar to hypothetical protein FLJ12994 (predicted)
RGD1309054_f	137	248	<b>1.82</b>	AI231159	similar to FKSG26 protein (predicted)
RGD1309450_f	58	91	<b>1.57</b>	BI291600	similar to KIAA2010 protein (predicted)
RGD1309550	222	340	<b>1.53</b>	BI295047	similar to hypothetical protein D12Erd771e
RGD1309752	15	159	<b>10.91</b>	AW524173	Similar to hypothetical protein D630010C10
RGD1309995_f	22	113	<b>5.20</b>	BF398054	similar to CG13957-PA (predicted)
RGD1310052_f	34	77	<b>2.26</b>	BF561001	similar to hypothetical protein FLJ40362 (predicted)
RGD1310139_f	167	107	<b>0.64</b>	AI144739	Similar to KIAA0303 (predicted)
RGD1310433_f	50	105	<b>2.11</b>	AW254450	similar to mKIAA1757 protein (predicted)
RGD1310474_f	9	110	<b>12.14</b>	BF567629	similar to KIAA0423 (predicted)
RGD1311595	138	569	<b>4.13</b>	BE107859	similar to KIAA2026 protein
RGD1311678	187	674	<b>3.61</b>	AW524670	Similar to 4921517L17Rik protein
<b>RGD1311958_f</b>	<b>104</b>	<b>388</b>	<b>3.74</b>	<b>BF563206</b>	<b>similar to 6430514L14Rik protein (predicted)</b>
RGD1312005_f	72	132	<b>1.83</b>	BF392349	similar to DD1 (predicted)
<b>RGD1559605_f</b>	<b>17</b>	<b>409</b>	<b>24.57</b>	<b>BE101933</b>	<b>Similar to hypothetical protein FLJ25477 isoform 2</b>
RGD1559693_f	69	147	<b>2.11</b>	BF400799	similar to Hypothetical protein 6330514E13 (predicted)
RGD1559930	20	151	<b>7.50</b>	AW920445	similar to mKIAA0256 protein (predicted)
RGD1560834_f	115	256	<b>2.23</b>	BE108367	similar to FRBZ1 protein (FRBZ1) (predicted)
RGD1560924_f	37	161	<b>4.38</b>	AI011501	similar to C230080I20Rik protein (predicted)
RGD1561500_f	259	171	<b>0.66</b>	BM390695	similar to hypothetical protein FLJ14800 (predicted)
RGD1561597_f	56	174	<b>3.13</b>	BE118049	similar to mKIAA0518 protein (predicted)
RGD1561653_f	60	706	<b>11.79</b>	AI059295	similar to HECT domain containing 1 (predicted)
RGD1562123_f	100	247	<b>2.47</b>	BE117893	similar to hypothetical protein (predicted)
RGD1562407_f	93	193	<b>2.07</b>	BG665671	similar to WAC (predicted)
RGD1563072_f	126	237	<b>1.88</b>	BI290787	similar to hypothetical protein FLJ38984 (predicted)
RGD1563612_f	174	344	<b>1.98</b>	BE108162	similar to testymin (predicted)
RGD1563838_f	34	77	<b>2.29</b>	BG665533	similar to leucine zipper protein 2 (predicted)
RGD1563977_f	113	69	<b>0.61</b>	AI145746	Similar to protein 4.1G (predicted)
RGD1564625_f	87	139	<b>1.61</b>	BF397371	similar to transmembrane protein TM9SF3 (predicted)
RGD1564852_f	73	119	<b>1.63</b>	AI575906	similar to hypothetical protein FLJ14503 (predicted)
RGD1564943_f	96	243	<b>2.52</b>	BE118876	similar to 4930429A08Rik protein (predicted)
RGD1564964_f	18	266	<b>15.04</b>	BE116698	similar to WD repeat domain 11 protein (predicted)
RGD1564983_f	29	77	<b>2.67</b>	BE107672	similar to leucine rich repeat containing 10 (predicted)
RGD1565095_f	82	669	<b>8.14</b>	AA893212	Similar to hypothetical protein MGC52110 (predicted)
RGD1565267_f	48	147	<b>3.07</b>	BM387829	similar to PS1D protein (predicted)
RGD1565556_f	374	1171	<b>3.13</b>	AI409823	similar to cajalin 2 isoform a (predicted)
RGD1565602_f	76	182	<b>2.38</b>	BM383996	similar to PLU1 (predicted)
RGD1566064_f	69	282	<b>4.08</b>	BI289641	similar to KIAA1096 protein (predicted)
RGD1566117_f	135	928	<b>6.87</b>	AI555865	similar to hypothetical protein FLJ23033 (predicted)
RGD1566201_f	38	150	<b>3.96</b>	BE118739	similar to mKIAA0960 protein (predicted)
Rtn4	68	505	<b>7.48</b>	AF051335	reticulon 4
Scoc	223	399	<b>1.79</b>	AI029749	short coiled-coil protein
Sel1h	36	86	<b>2.38</b>	BI285936	Sel1 (suppressor of lin-12) 1 homolog (C. elegans)
Sgta	113	202	<b>1.79</b>	NM_022703	small glutamine-rich tetratricopeptide repeat (TPR)-con
Slitrk1_predicte	41	94	<b>2.29</b>	AW523107	SLIT and NTRK-like family, member 1 (predicted)
Slitrk3_predicte	35	105	<b>2.96</b>	AA819827	SLIT and NTRK-like family, member 3 (predicted)
Svop	143	255	<b>1.79</b>	NM_134404	SV2 related protein
Tmem16c_pred	25	98	<b>3.98</b>	AI144648	transmembrane protein 16C (predicted)
<b>Tmem24</b>	<b>537</b>	<b>295</b>	<b>0.55</b>	<b>BG378195</b>	<b>transmembrane protein 24</b>
Trim23	79	126	<b>1.60</b>	L04760	tripartite motif protein 23
Ttc3_predicted	242	989	<b>4.08</b>	BF388771	tetratricopeptide repeat domain 3 (predicted)
Ttc9c	75	139	<b>1.86</b>	BG668930	Tetratricopeptide repeat domain 9C
Ttyh1_predictec	1113	2975	<b>2.67</b>	AI412746	weety homolog 1 (Drosophila) (predicted)
Wdr37_predicte	105	712	<b>6.77</b>	BG662814	WD repeat domain 37 (predicted)
Wdr48_predicte	82	134	<b>1.63</b>	BE101118	WD repeat domain 48 (predicted)
Wdr59	81	135	<b>1.66</b>	AW527783	WD repeat domain 59

#### EST's

Sample	F3-Cont	F3-Vinc	Vin/Con	Genbank	Gene Title
Gene Symbol	Raw	Raw	Ratio	Genbank	Gene Title
---	180	667	<b>3.71</b>	BF406608	Transcribed locus, strongly similar to XP_223397.3 sin
---	12	84	<b>7.08</b>	BE109363	Transcribed locus, strongly similar to XP_579796.1 hy
Fam31b_predic	24	134	<b>5.57</b>	BM385853	Family with sequence similarity 31, member B (predicte
LOC366300	598	1122	<b>1.88</b>	AI170706	hypothetical LOC366300

LOC682182 /// I	51	77	<b>1.52</b>	BE095833	similar to RIKEN cDNA 1110067D22 (predicted) /// simi
RGD1304711_f	354	222	<b>0.63</b>	AW526713	Similar to RIKEN cDNA 4921521J11 (predicted)
RGD1305387	76	120	<b>1.58</b>	BI288527	similar to RIKEN cDNA 2610207I16
RGD1305492_f	48	96	<b>2.01</b>	BG670905	similar to RIKEN cDNA 1700034P14 (predicted)
RGD1305671_f	29	186	<b>6.46</b>	BE116226	Similar to expressed sequence AI317237 (predicted)
<b>RGD1305755</b>	<b>51</b>	<b>285</b>	<b>5.62</b>	<b>AI179665</b>	<b>Similar to RIKEN cDNA 5033406L14</b>
RGD1305903_f	31	125	<b>4.05</b>	AI556398	similar to RIKEN cDNA G630024C07 gene (predicted)
RGD1306053	49	91	<b>1.85</b>	BF392629	similar to RIKEN cDNA 1200016B17
RGD1306067	61	243	<b>4.00</b>	AW141642	similar to chromosome 20 open reading frame 6
RGD1306284	273	180	<b>0.66</b>	AA866227	similar to RIKEN cDNA 1110005A03
RGD1306739_f	135	83	<b>0.61</b>	AI171288	similar to RIKEN cDNA 1700040L02 (predicted)
RGD1306873	33	137	<b>4.16</b>	AI177725	similar to RIKEN cDNA 2210010N04 gene
RGD1308290_f	53	96	<b>1.81</b>	AA894335	similar to RIKEN cDNA 5730454B08 (predicted)
RGD1309104_f	116	395	<b>3.40</b>	BF388757	similar to RIKEN cDNA 1700025G04 gene (predicted)
RGD1309385_f	39	238	<b>6.14</b>	BG372587	similar to RIKEN cDNA E030034P13 (predicted)
RGD1310127	139	88	<b>0.64</b>	AA894060	similar to cDNA sequence BC017158
RGD1310351_f	47	238	<b>5.04</b>	BI296537	similar to RIKEN cDNA 4732418C07 (predicted)
RGD1311686	223	347	<b>1.56</b>	BI294949	similar to chromosome 16 open reading frame 5
RGD1310722_f	42	690	<b>16.34</b>	AW534218	similar to RIKEN cDNA D130059P03 gene (predicted)
RGD1311086	221	146	<b>0.66</b>	BI288424	similar to RIKEN cDNA 2610029K21
RGD1311456_f	55	266	<b>4.83</b>	AI045965	similar to RIKEN cDNA B230380D07 (predicted)
RGD1311783_f	265	469	<b>1.77</b>	BG668881	similar to RIKEN cDNA 2010012O05 (predicted)
<i>RGD1311835</i>	<i>81</i>	<i>136</i>	<i>1.69</i>	<i>BI284801</i>	<i>similar to RIKEN cDNA 1110021N07</i>
<b>RGD1359529</b>	<b>67</b>	<b>166</b>	<b>2.48</b>	<b>BE096504</b>	<b>similar to chromosome 1 open reading frame 63</b>
RGD1359691	137	48	<b>0.35</b>	BF398271	hypothetical LOC287534
RGD1559896_f	31	85	<b>2.77</b>	AW253651	similar to RIKEN cDNA 2310022B05 (predicted)
RGD1560479_f	23	211	<b>9.24</b>	BF394331	similar to RIKEN cDNA 5330439J01 (predicted)
<i>RGD1560913_f</i>	<i>5</i>	<i>164</i>	<b>30.87</b>	<i>AA799328</i>	<i>similar to expressed sequence AW413625 (predicted)</i>
RGD1560957_f	137	85	<b>0.62</b>	BE119096	similar to RIKEN cDNA 5730466H23 (predicted)
RGD1562563_f	150	283	<b>1.89</b>	BE116089	Similar to RIKEN cDNA G430041M01 (predicted)
RGD1562618_f	74	136	<b>1.85</b>	BG381652	similar to RIKEN cDNA 6030419C18 gene (predicted)
RGD1563001_f	70	343	<b>4.90</b>	BI304125	similar to RIKEN cDNA 2010106G01 (predicted)
RGD1563441_f	28	77	<b>2.74</b>	BF406306	similar to RIKEN cDNA A030009H04 (predicted)
RGD1563912_f	441	868	<b>1.97</b>	BI282028	RGD1563912 (predicted)
RGD1564227_f	39	158	<b>4.06</b>	BI296242	similar to RIKEN cDNA 5730557B15 (predicted)
RGD1564379_f	66	100	<b>1.52</b>	BF284939	RGD1564379 (predicted)
RGD1564778_f	131	259	<b>1.97</b>	AI168933	similar to RIKEN cDNA 4121402D02 (predicted)
RGD1564957_f	36	138	<b>3.89</b>	BF387898	similar to RIKEN cDNA 3110007P09 (predicted)
RGD1565957_f	514	325	<b>0.63</b>	BI294688	similar to RIKEN cDNA 1110061N23 (predicted)
---	200	126	<b>0.63</b>	AF023090	Transcribed locus
---	131	885	<b>6.75</b>	BF398122	Transcribed locus
---	31	76	<b>2.48</b>	AI029767	Transcribed locus
---	66	125	<b>1.91</b>	AA956307	Transcribed locus
---	25	84	<b>3.35</b>	BF398673	Transcribed locus
---	9	115	<b>13.55</b>	BE117514	Transcribed locus
---	189	120	<b>0.63</b>	AI146156	Transcribed locus
---	954	583	<b>0.61</b>	BM389272	Transcribed locus
---	252	469	<b>1.86</b>	BF420311	Transcribed locus
---	327	892	<b>2.72</b>	BM384203	Transcribed locus
---	43	232	<b>5.43</b>	BF394499	Transcribed locus
---	17	90	<b>5.26</b>	BF403330	Transcribed locus
---	87	51	<b>0.59</b>	BE106592	Transcribed locus
---	88	177	<b>2.01</b>	H34328	Transcribed locus
---	45	220	<b>4.93</b>	BG668689	Transcribed locus
---	44	167	<b>3.79</b>	BF394561	Transcribed locus
---	81	36	<b>0.44</b>	BI300158	Transcribed locus
---	205	1136	<b>5.56</b>	BF546710	Transcribed locus
---	70	183	<b>2.60</b>	BF522208	Transcribed locus
---	32	109	<b>3.38</b>	AA848370	Transcribed locus
---	217	135	<b>0.62</b>	BF390904	Transcribed locus
---	122	77	<b>0.64</b>	AW524239	Transcribed locus
---	79	156	<b>1.98</b>	BE119960	Transcribed locus
---	106	64	<b>0.60</b>	BF415784	Transcribed locus
---	54	247	<b>4.59</b>	BF404490	Transcribed locus
---	32	187	<b>5.85</b>	BF404446	Transcribed locus
---	335	200	<b>0.60</b>	BF391580	Transcribed locus
---	41	90	<b>2.21</b>	BF405168	Transcribed locus
---	83	258	<b>3.11</b>	BF393011	Transcribed locus
---	206	93	<b>0.45</b>	BE120509	Transcribed locus

---	313	204	<b>0.65</b>	AW521804	Transcribed locus
---	73	113	<b>1.54</b>	BF411608	Transcribed locus
---	145	289	<b>1.99</b>	AW526127	Transcribed locus
---	48	247	<b>5.16</b>	AW529817	Transcribed locus
---	26	76	<b>2.93</b>	BM958510	Transcribed locus
---	27	81	<b>2.97</b>	BF396580	Transcribed locus
---	<b>11</b>	<b>78</b>	<b>6.85</b>	<b>BE105421</b>	<b>Transcribed locus</b>
---	45	83	<b>1.83</b>	AI113308	Transcribed locus
---	263	476	<b>1.81</b>	BE098012	Transcribed locus
---	600	314	<b>0.52</b>	H34760	Transcribed locus
---	43	252	<b>5.82</b>	<i>BF545930</i>	<i>Transcribed locus</i>
---	23	153	<b>6.80</b>	BF548061	Transcribed locus
---	1301	856	<b>0.66</b>	BE099398	Transcribed locus
---	106	42	<b>0.40</b>	AW920217	Transcribed locus
---	38	81	<b>2.14</b>	BF291053	Transcribed locus
---	1181	1892	<b>1.60</b>	BF544005	Transcribed locus
---	110	271	<b>2.47</b>	AW522341	Transcribed locus
---	360	574	<b>1.59</b>	BF390608	Transcribed locus
---	66	141	<b>2.15</b>	AI102173	Transcribed locus
---	26	105	<b>4.00</b>	AW531516	Transcribed locus
---	32	86	<b>2.70</b>	AI044545	Transcribed locus
---	36	94	<b>2.63</b>	AI601993	Transcribed locus
---	254	149	<b>0.59</b>	BE096857	Transcribed locus
---	388	225	<b>0.58</b>	BG380768	Transcribed locus
---	112	297	<b>2.65</b>	AW523015	Transcribed locus
---	196	311	<b>1.59</b>	AW529108	Transcribed locus
---	955	608	<b>0.64</b>	BG668719	Transcribed locus
---	62	108	<b>1.73</b>	H31734	Transcribed locus
---	34	691	<b>20.25</b>	AI145951	Transcribed locus
---	173	115	<b>0.66</b>	AW529067	Transcribed locus
---	71	127	<b>1.80</b>	AI575856	Transcribed locus
---	74	196	<b>2.65</b>	AI412813	Transcribed locus
---	53	142	<b>2.68</b>	BG664080	Transcribed locus
---	127	265	<b>2.08</b>	AI575519	Transcribed locus
---	113	300	<b>2.66</b>	AA956982	Transcribed locus
---	7	172	<b>26.30</b>	AI030231	Transcribed locus
---	328	188	<b>0.57</b>	AI071202	Transcribed locus
---	89	35	<b>0.39</b>	BF392359	Transcribed locus
---	8	197	<b>23.43</b>	BF400601	Transcribed locus
---	33	336	<b>10.30</b>	BE120522	Transcribed locus
---	42	81	<b>1.92</b>	AA956085	Transcribed locus
---	23	136	<b>5.82</b>	BI303858	Transcribed locus
---	192	86	<b>0.45</b>	BE106361	Transcribed locus
---	64	331	<b>5.16</b>	BF549454	Transcribed locus
---	13	112	<b>8.85</b>	BE096515	Transcribed locus
---	12	79	<b>6.83</b>	BE111692	Transcribed locus
---	153	361	<b>2.36</b>	AI556642	Transcribed locus
---	26	124	<b>4.76</b>	BI295733	Transcribed locus
---	84	143	<b>1.71</b>	AA850595	Transcribed locus
---	79	161	<b>2.05</b>	BF406435	Transcribed locus
---	29	139	<b>4.79</b>	AA956757	Transcribed locus
---	82	287	<b>3.51</b>	BM391661	Transcribed locus
---	135	343	<b>2.54</b>	BI296013	Transcribed locus
---	135	89	<b>0.66</b>	AI227988	Transcribed locus
---	75	46	<b>0.61</b>	BI280337	Transcribed locus
---	30	93	<b>3.12</b>	BE096618	Transcribed locus
---	200	122	<b>0.61</b>	BF389889	Transcribed locus
---	54	115	<b>2.14</b>	BI291162	Transcribed locus
---	59	118	<b>2.00</b>	AI043817	Transcribed locus
---	173	84	<b>0.48</b>	<i>BF399309</i>	<i>Transcribed locus</i>
---	81	54	<b>0.66</b>	BF405616	Transcribed locus
---	20	107	<b>5.45</b>	BI294889	Transcribed locus
---	23	80	<b>3.53</b>	BE107619	Transcribed locus
---	89	170	<b>1.91</b>	BF419818	Transcribed locus
---	34	354	<b>10.29</b>	BI300794	Transcribed locus
---	183	113	<b>0.62</b>	BF283381	Transcribed locus
---	131	228	<b>1.74</b>	AI231225	Transcribed locus
---	78	41	<b>0.53</b>	<i>BF391128</i>	<i>Transcribed locus</i>
---	34	92	<b>2.74</b>	BI295124	Transcribed locus



---	70	253	<b>3.62</b>	AA899937	<i>Transcribed locus</i>
---	186	287	<b>1.55</b>	BF393945	Transcribed locus
---	253	161	<b>0.64</b>	BE107167	Transcribed locus
---	62	96	<b>1.53</b>	AI232217	Transcribed locus
---	84	55	<b>0.66</b>	BF283404	Transcribed locus
---	302	171	<b>0.57</b>	AI229933	Transcribed locus
---	273	144	<b>0.53</b>	AW915115	Transcribed locus
---	15	134	<b>8.86</b>	BF406304	<i>Transcribed locus</i>
---	<b>112</b>	<b>322</b>	<b>2.88</b>	<b>AA944136</b>	<b><i>Transcribed locus</i></b>
---	166	105	<b>0.63</b>	BF562962	Transcribed locus
---	257	603	<b>2.35</b>	BF546770	Transcribed locus
---	348	198	<b>0.57</b>	BF284027	Transcribed locus
---	144	314	<b>2.18</b>	AI229321	Transcribed locus
---	15	90	<b>6.07</b>	H31285	Transcribed locus
---	176	315	<b>1.79</b>	AA874903	Transcribed locus
---	59	188	<b>3.21</b>	BE100612	Transcribed locus
---	204	129	<b>0.63</b>	AI146262	Transcribed locus
---	55	121	<b>2.19</b>	BF416395	Transcribed locus
---	127	61	<b>0.48</b>	AA817920	Transcribed locus
---	3	122	<b>40.67</b>	BG668477	Transcribed locus
---	81	135	<b>1.66</b>	AI072068	Transcribed locus
---	55	178	<b>3.25</b>	AW532389	Transcribed locus
---	51	253	<b>4.95</b>	BF409092	Transcribed locus
---	38	81	<b>2.14</b>	AA997253	Transcribed locus
---	21	81	<b>3.87</b>	BE096277	Transcribed locus
---	48	93	<b>1.94</b>	BE120904	Transcribed locus
---	159	597	<b>3.75</b>	AA925373	Transcribed locus
---	36	145	<b>3.99</b>	BM390588	Transcribed locus
---	28	106	<b>3.77</b>	AI229409	Transcribed locus
---	250	156	<b>0.62</b>	BE108597	Transcribed locus
---	92	147	<b>1.59</b>	BF559356	Transcribed locus
---	115	1006	<b>8.77</b>	BE109132	Transcribed locus
---	170	292	<b>1.72</b>	BG667918	Transcribed locus
---	74	133	<b>1.80</b>	BG671630	Transcribed locus
---	79	158	<b>1.99</b>	BF555825	Transcribed locus
---	140	89	<b>0.64</b>	AI072798	Transcribed locus
---	<b>85</b>	<b>213</b>	<b>2.50</b>	<b>AA925807</b>	<b><i>Transcribed locus</i></b>
---	20	156	<b>7.90</b>	AI071698	Transcribed locus
---	90	47	<b>0.53</b>	BG665568	Transcribed locus
---	544	352	<b>0.65</b>	AI230360	Transcribed locus
---	<b>190</b>	<b>76</b>	<b>0.40</b>	<b>AI102821</b>	<b><i>Transcribed locus</i></b>
---	175	111	<b>0.64</b>	AA926109	<i>Transcribed locus</i>
---	556	1131	<b>2.03</b>	BM958512	Transcribed locus
---	117	191	<b>1.63</b>	AW921244	Transcribed locus
---	38	348	<b>9.25</b>	BI281615	Transcribed locus
---	206	865	<b>4.19</b>	BG372598	Transcribed locus
---	339	219	<b>0.65</b>	BF563716	Transcribed locus
---	868	1987	<b>2.29</b>	BF555795	Transcribed locus
---	80	52	<b>0.65</b>	AW523077	Transcribed locus
---	90	375	<b>4.16</b>	BE108047	Transcribed locus
---	87	141	<b>1.63</b>	AI237079	Transcribed locus
---	232	361	<b>1.55</b>	AI176342	Transcribed locus
---	46	100	<b>2.17</b>	AA859319	<i>Transcribed locus</i>
---	74	212	<b>2.86</b>	BE120748	Transcribed locus
---	107	603	<b>5.66</b>	BE105050	Transcribed locus
---	155	83	<b>0.54</b>	AA875457	Transcribed locus
---	55	260	<b>4.77</b>	AI717668	Transcribed locus
---	110	59	<b>0.54</b>	AI070489	<i>Transcribed locus</i>
---	843	1377	<b>1.63</b>	BF285731	Transcribed locus
---	162	92	<b>0.57</b>	BI280114	Transcribed locus
---	171	91	<b>0.53</b>	BE105488	Transcribed locus
---	70	124	<b>1.78</b>	BE110067	<i>Transcribed locus</i>
---	372	764	<b>2.05</b>	AW530378	Transcribed locus
---	47	115	<b>2.45</b>	BE121056	Transcribed locus
---	26	189	<b>7.40</b>	AI555166	Transcribed locus
---	28	97	<b>3.51</b>	AI060117	Transcribed locus
---	15	83	<b>5.37</b>	AI145015	<i>Transcribed locus</i>
---	120	315	<b>2.63</b>	AA819045	Transcribed locus
---	60	170	<b>2.86</b>	AW524106	Transcribed locus

---	305	144	<b>0.47</b>	BF409213	Transcribed locus
---	16	392	<b>24.17</b>	BE108751	Transcribed locus
---	74	322	<b>4.37</b>	BE114458	Transcribed locus
---	31	94	<b>3.02</b>	BF391155	Transcribed locus
---	12	146	<b>12.61</b>	AW534519	Transcribed locus
---	47	96	<b>2.04</b>	AW534466	Transcribed locus
---	94	728	<b>7.72</b>	AI145433	Transcribed locus
---	18	87	<b>4.84</b>	BF416420	Transcribed locus
---	28	94	<b>3.38</b>	BF390757	Transcribed locus
---	81	132	<b>1.63</b>	AI406475	Transcribed locus
---	8	148	<b>18.44</b>	AW530527	Transcribed locus
---	19	138	<b>7.30</b>	BM386352	Transcribed locus
---	63	130	<b>2.08</b>	AI103408	Transcribed locus
---	<b>68</b>	<b>1546</b>	<b>22.67</b>	<b>AI555855</b>	<b>Transcribed locus</b>
---	551	335	<b>0.61</b>	H31701	Transcribed locus
---	97	203	<b>2.09</b>	AI229240	Transcribed locus
---	10	198	<b>19.54</b>	BF402498	Transcribed locus
---	84	55	<b>0.65</b>	AI058315	Transcribed locus
---	81	146	<b>1.79</b>	BE102139	Transcribed locus
---	84	53	<b>0.63</b>	AI137306	Transcribed locus
---	150	243	<b>1.62</b>	BM386302	Transcribed locus
---	158	430	<b>2.72</b>	AI535567	Transcribed locus
---	96	413	<b>4.29</b>	BF420785	Transcribed locus
---	123	193	<b>1.57</b>	AI011930	Transcribed locus
---	38	78	<b>2.06</b>	AI233902	Transcribed locus
---	34	81	<b>2.40</b>	BI288184	Transcribed locus
---	79	163	<b>2.07</b>	BM383595	Transcribed locus
---	28	108	<b>3.91</b>	BF392753	Transcribed locus
---	79	167	<b>2.11</b>	BE108208	Transcribed locus
---	86	140	<b>1.63</b>	AI717163	Transcribed locus
---	204	355	<b>1.74</b>	AW534671	Transcribed locus
---	37	170	<b>4.55</b>	BF397054	Transcribed locus
---	41	120	<b>2.92</b>	AA900904	Transcribed locus
---	61	164	<b>2.71</b>	AA818967	Transcribed locus
---	45	119	<b>2.64</b>	BF407470	Transcribed locus
---	85	134	<b>1.57</b>	BE107074	Transcribed locus
---	146	234	<b>1.60</b>	AI179450	Transcribed locus
---	7	79	<b>10.61</b>	BG379394	Transcribed locus
---	190	330	<b>1.74</b>	BF411826	Transcribed locus
---	116	1031	<b>8.89</b>	AI511069	Transcribed locus
---	48	434	<b>9.05</b>	AW252020	Transcribed locus
---	195	129	<b>0.66</b>	BI276118	Transcribed locus
---	<b>263</b>	<b>706</b>	<b>2.69</b>	<b>AI112113</b>	<b>Transcribed locus</b>
---	73	161	<b>2.21</b>	AW531387	Transcribed locus
---	55	139	<b>2.51</b>	BF399121	Transcribed locus
---	21	110	<b>5.16</b>	BF284914	Transcribed locus
---	657	1034	<b>1.57</b>	BI292687	Transcribed locus
---	43	86	<b>2.01</b>	AA859337	Transcribed locus
---	465	1380	<b>2.97</b>	AW535380	Transcribed locus
---	96	157	<b>1.64</b>	BE106526	Transcribed locus
---	46	78	<b>1.68</b>	BF548480	Transcribed locus
---	84	51	<b>0.60</b>	BI291457	Transcribed locus
---	103	62	<b>0.60</b>	BF418563	Transcribed locus
---	51	94	<b>1.86</b>	BE099568	Transcribed locus
---	204	344	<b>1.69</b>	BI295869	Transcribed locus
---	470	1446	<b>3.08</b>	AI170377	Transcribed locus
---	140	330	<b>2.37</b>	BM386385	Transcribed locus
---	136	241	<b>1.77</b>	BF412303	Transcribed locus
---	53	589	<b>11.20</b>	AW532489	Transcribed locus
---	87	232	<b>2.66</b>	BF283340	Transcribed locus
---	113	333	<b>2.96</b>	BF550404	Transcribed locus
---	163	92	<b>0.57</b>	BF416276	Transcribed locus
---	40	85	<b>2.14</b>	BG378933	Transcribed locus
---	116	281	<b>2.42</b>	AA955579	Transcribed locus
---	21	96	<b>4.67</b>	BF396151	Transcribed locus
---	622	406	<b>0.65</b>	BG378070	Transcribed locus
---	100	66	<b>0.66</b>	BI286851	Transcribed locus
---	51	117	<b>2.29</b>	BE108174	Transcribed locus
---	25	112	<b>4.59</b>	BE108246	Transcribed locus

---	288	153	<b>0.53</b>	AW143156	Transcribed locus
---	66	129	<b>1.96</b>	AI101372	Transcribed locus
---	194	359	<b>1.86</b>	BF415701	Transcribed locus
---	315	190	<b>0.60</b>	BI282767	Transcribed locus
---	23	200	<b>8.64</b>	BF402633	Transcribed locus
---	56	221	<b>3.98</b>	BE109509	Transcribed locus
---	23	168	<b>7.45</b>	AI385171	Transcribed locus
---	77	371	<b>4.81</b>	BF408438	Transcribed locus
---	63	382	<b>6.06</b>	BM389190	Transcribed locus
---	24	90	<b>3.72</b>	BG372400	Transcribed locus
---	40	98	<b>2.44</b>	AI101245	Transcribed locus
---	82	193	<b>2.34</b>	BE111820	Transcribed locus
---	264	141	<b>0.53</b>	AI407719	Transcribed locus
---	53	198	<b>3.73</b>	BF400811	Transcribed locus
---	333	517	<b>1.55</b>	BM384026	Transcribed locus
---	333	646	<b>1.94</b>	BM388843	Transcribed locus
---	75	251	<b>3.34</b>	BF400779	Transcribed locus
---	119	76	<b>0.64</b>	AI232806	Transcribed locus
---	30	1207	<b>39.95</b>	AI228978	Transcribed locus
---	117	62	<b>0.53</b>	BG381647	Transcribed locus
---	202	135	<b>0.67</b>	AA859010	Transcribed locus
---	716	473	<b>0.66</b>	AI171776	Transcribed locus
---	212	607	<b>2.87</b>	BE109208	Transcribed locus
---	250	156	<b>0.62</b>	BM386212	Transcribed locus
---	206	129	<b>0.63</b>	AI598550	Transcribed locus
---	104	232	<b>2.23</b>	AA799420	Transcribed locus
---	111	70	<b>0.63</b>	BG373057	Transcribed locus
---	<b>147</b>	<b>364</b>	<b>2.48</b>	<b>AI103530</b>	<b>Transcribed locus</b>
---	57	342	<b>5.95</b>	AI555608	Transcribed locus
---	81	123	<b>1.52</b>	AW914907	Transcribed locus
---	122	193	<b>1.58</b>	BF544403	Transcribed locus
---	52	83	<b>1.60</b>	BF542239	Transcribed locus
---	375	206	<b>0.55</b>	BI303277	Transcribed locus
---	74	137	<b>1.86</b>	AW920828	Transcribed locus
---	67	115	<b>1.71</b>	AI407047	Transcribed locus, moderately similar to XP_001144552
---	70	171	<b>2.46</b>	BE104676	Transcribed locus, strongly similar to NP_001029332.1
---	693	405	<b>0.58</b>	AI454332	Transcribed locus, strongly similar to XP_001081628.1
---	281	498	<b>1.77</b>	BF390195	CDNA clone IMAGE:7320582
---	51	135	<b>2.68</b>	BF562934	CDNA clone IMAGE:7320582
---	77	962	<b>12.45</b>	AI008409	CDNA clone IMAGE:7321089
---	237	145	<b>0.61</b>	AI237047	CDNA clone IMAGE:7365681
---	126	80	<b>0.63</b>	BF398091	CDNA clone IMAGE:7374368
---	37	203	<b>5.48</b>	BE106331	CDNA clone IMAGE:7461178
---	295	469	<b>1.59</b>	BI294768	---
---	381	161	<b>0.42</b>	BM384537	---
---	46	86	<b>1.85</b>	BF547596	---
---	55	99	<b>1.79</b>	AI547718	---
---	23	147	<b>6.29</b>	AI705744	---
---	195	370	<b>1.90</b>	H33235	---
---	120	390	<b>3.24</b>	AI103026	---
---	294	688	<b>2.34</b>	BE116953	---
---	162	258	<b>1.59</b>	BI282114	---
---	302	461	<b>1.53</b>	BI275155	---
---	74	271	<b>3.67</b>	AW527270	---
---	236	151	<b>0.64</b>	BM385286	---
---	109	283	<b>2.60</b>	BG372713	---
---	207	849	<b>4.10</b>	AA997406	---
---	248	493	<b>1.99</b>	BF400907	---
---	1710	3508	<b>2.05</b>	{_Rat_beta-ac	---
---	240	784	<b>3.27</b>	{_Rat_beta-ac	---
---	543	1920	<b>3.54</b>	_Rat_beta-ac	---
---	126	211	<b>1.68</b>	BF556405	---
---	<b>49</b>	<b>82</b>	<b>1.67</b>	<b>AI172311</b>	---
---	2376	1570	<b>0.66</b>	AI145313	---
---	14	77	<b>5.36</b>	AI112375	---
---	94	61	<b>0.65</b>	AA858748	---
---	143	70	<b>0.49</b>	BF522861	---
---	382	181	<b>0.47</b>	BF545849	---
---	133	200	<b>1.50</b>	BE120211	---

---	539	347	<b>0.64</b>	BG380279	---
---	51	1212	<b>23.99</b>	AI103917	---
---	84	52	<b>0.62</b>	AI030449	---
---	102	712	<b>6.97</b>	AI101194	---
---	51	139	<b>2.75</b>	BI289762	---
---	1779	1185	<b>0.67</b>	BE115454	---
---	11	127	<b>11.37</b>	BF398431	---
---	37	114	<b>3.04</b>	BG668744	---
---	66	100	<b>1.51</b>	BE097409	---
---	45	85	<b>1.89</b>	AW535011	---
---	42	167	<b>4.02</b>	AW527250	---
---	131	84	<b>0.64</b>	BE116860	---
---	51	153	<b>2.98</b>	BE119914	---
---	45	85	<b>1.91</b>	<i>BI288579</i>	---
---	170	345	<b>2.02</b>	AW921158	---
---	100	152	<b>1.53</b>	BF396974	---
---	129	75	<b>0.58</b>	AA859524	---
---	177	96	<b>0.54</b>	BE117273	---
---	77	129	<b>1.68</b>	AI715113	---
---	246	152	<b>0.62</b>	BG373537	---
---	14	78	<b>5.60</b>	BF415798	---
---	53	158	<b>2.97</b>	BI293026	---
---	47	104	<b>2.23</b>	BF397998	---
---	50	80	<b>1.60</b>	BF400933	---
---	105	58	<b>0.55</b>	BI280367	---
---	131	271	<b>2.06</b>	BF544968	---
---	19	106	<b>5.58</b>	AA851046	---
---	22	283	<b>12.82</b>	BF548081	---
---	403	249	<b>0.62</b>	BM383406	---
---	89	53	<b>0.60</b>	BG380561	---
---	573	1349	<b>2.36</b>	BE101133	---
---	131	86	<b>0.66</b>	AI574734	---
---	238	146	<b>0.61</b>	BE105500	---
---	27	95	<b>3.48</b>	BF386160	---
---	29	171	<b>5.84</b>	BE115821	---
---	160	106	<b>0.66</b>	BF413298	---
---	76	18	<b>0.24</b>	BE111542	---
---	35	134	<b>3.82</b>	AI501579	---
---	<i>101</i>	<i>59</i>	<i>0.58</i>	<i>BF410240</i>	---
---	87	53	<b>0.60</b>	AI013683	---
---	178	277	<b>1.56</b>	BF397301	---
---	282	180	<b>0.64</b>	AI145935	---
---	88	43	<b>0.49</b>	BF419655	---
---	54	192	<b>3.56</b>	BF419406	---
---	28	76	<b>2.73</b>	BF390754	---
---	<b>55</b>	<b>97</b>	<b>1.76</b>	<b>BF403869</b>	---
---	107	950	<b>8.92</b>	AI575254	---
---	45	930	<b>20.86</b>	AI710284	---
---	420	279	<b>0.67</b>	BF420262	---
---	670	433	<b>0.65</b>	BF403383	---
---	29	376	<b>12.95</b>	AI045904	---
---	240	151	<b>0.63</b>	AI104117	---
---	29	97	<b>3.37</b>	AW920849	---
---	306	1144	<b>3.74</b>	BF567766	---
---	52	116	<b>2.21</b>	AA684862	---
---	18	112	<b>6.11</b>	AI059349	---
---	18	105	<b>5.94</b>	AI705744	---
---	17	81	<b>4.74</b>	BF559198	---
---	5	86	<b>15.98</b>	AI234943	---

**Note - The bolded genes are similar between male and female gene sets.  
The italic genes are similar within the same sex between amygdala and hippocampus.**