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Jennifer D Stewart (jdstewart@wsu.edu) Andrew Storfer (astorfer@wsu.edu) Robert B Page (robert.page@uky.edu) Christopher K Beachy (christopher.beachy@minotstateu.edu) S R Voss (srvoss@uky.edu)

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### Transcriptional response of Mexican axolotls to *Ambystoma tigrinum* virus (ATV) infection

Jennifer D. Cotter<sup>1,\*</sup>, Andrew Storfer<sup>1</sup>, Robert B. Page<sup>2</sup>, Christopher K. Beachy<sup>3</sup>, and S. Randal Voss<sup>2</sup>

- <sup>1</sup> School of Biological Sciences, Washington State University, Pullman, WA 99164-4236
- <sup>2</sup> Department of Biology and Spinal Cord and Brain Injury Research Center, University
- of Kentucky, Lexington KY 40506
- <sup>3</sup> Department of Biology, Minot State University, Minot ND 58707
- <sup>\*</sup>To whom correspondence should be addressed
- JDS: jdstewart@wsu.edu
- AS: <u>astorfer@wsu.edu</u>
- RBP: <u>robert.page@uky.edu</u>
- CKB: <a href="mailto:christopher.beachy@minotstateu.edu">christopher.beachy@minotstateu.edu</a>
- SRV: <u>srvoss@uky.edu</u>

### Abstract

#### Background

Very little is known about the immunological responses of amphibians to pathogens that are causing global population declines. We used a custom microarray gene chip to characterize gene expression responses of axolotls (*Ambystoma mexicanum*) to an emerging viral pathogen, *Ambystoma tigrinum* virus (ATV).

#### Result

At 0, 24, 72, and 144 hours post-infection, spleen and lung samples were removed for estimation of host mRNA abundance and viral load. A total of 158 up-regulated and 105 down-regulated genes were identified across all time points using statistical and fold level criteria. The presumptive functions of these genes suggest a robust innate immune and antiviral gene expression response is initiated by *A. mexicanum* as early as 24 hours after ATV infection. At 24 hours, we observed transcript abundance changes for genes that are associated with phagocytosis and cytokine signaling, complement, and other general immune and defense responses. By 144 hours, we observed gene expression changes indicating host-mediated cell death, inflammation, and cytotoxicity.

#### Conclusion

Although *A. mexicanum* appears to mount a robust innate immune response, we did not observe gene expression changes indicative of lymphocyte proliferation in the spleen, which is associated with clearance of Frog 3 iridovirus in adult *Xenopus*. We speculate that ATV may be especially lethal to *A. mexicanum* and related tiger salamanders because they lack proliferative lymphocyte responses that are needed to clear highly virulent

iridoviruses. Genes identified from this study provide important new resources to investigate ATV disease pathology and host-pathogen dynamics in natural populations.

### Background

Emerging infectious diseases (EIDs) pose a serious threat to the health, stability, and persistence of human and wildlife populations [1-4]. Genetic and genomic tools have been incredibly useful for discovery of genes associated with host response and variation in resistance or susceptibility to a variety of pathogens [5-7]. The advent of genomic tools such as microarray analysis has offered new insights into host-pathogen systems. Additionally, their application to genomic response to host disease response allows rapid characterization of candidate genes for further research into control and eradication methods.

EIDs are a leading hypothesis for the global decline of amphibians and two pathogens in particular, *Batrachochytrium dendrobatidis* and Ranaviruses have been implicated in worldwide epizootics. Although studies are beginning to investigate possible mechanisms of resistance to these pathogens [8], in general, very little is known about the immune response of amphibians to EIDs. This is because most natural amphibian species are not used as laboratory models and we lack fundamental molecular tools to investigate disease pathology and host-pathogen interactions at the molecular level for all but a few species (e.g., *Ambystoma tigrinum spp., Xenopus spp.*).

Over the last 15 years, *Ranavirus* infections have been associated with marked increases in morbidity and mortality in fish, reptiles, and amphibians [9]. Ranaviruses are globally-distributed double-stranded, methylated DNA viruses of fish, amphibians and reptiles and are implicated in amphibian epizootics worldwide [9-11]. Both encapsulated and non-encapsulated forms can be infectious. The virus enters the cell via receptor mediated endocytosis or via fusion with the plasma membrane; and DNA and RNA synthesis occur in the nucleus, while protein synthesis occurs at morphologically specific assembly sites in the cytoplasm [9]. In North America, ranaviruses have been isolated from the majority of recent documented amphibian epizootics [12], including from tiger salamander (Ambystoma tigrinum) epizootics in Saskatchewan, Canada [13], Arizona [14], North Dakota, Utah, and Colorado, USA [15,16]. The viral variant that infects tiger salamanders, ATV, is transmitted either via direct contact with an infected animal or immersion in water that contains virus and infected individuals exhibit systemic hemorrhaging, edema, ulceration, and necrosis of the integument and internal organs [13,17,18]. In cases where ATV infection leads to mortality, it usually occurs within 2-3 weeks of exposure, with animals displaying symptoms often between 8-10 days post-exposure. Thus, ATV can rapidly overwhelm the tiger salamander immune response. However, mortality is not always a pathological endpoint because virulence and resistance are known to vary among ATV strains and tiger salamander populations, respectively, as indicated by both laboratory experiments and field observations [19]. Research characterizing the tiger salamander genomic response to ATV is needed to better understand the pathology, virulence, and possible mechanisms of resistance to this emerging disease.

The tiger salamander species complex includes *A. mexicanum* (Mexican axolotl), a model organism with a growing genomic and informatics resource base [20]. The immune system of the Mexican axolotl has been extensively studied using several classical approaches. Relative to other vertebrate models, the axolotl immune response has been described as immunodeficient [21,22]. There are several reasons for this characterization, including: production of only two immunoglobulin (Ig) classes, only one of which regulates the humoral response and neither of which is anamnestic [23,24]; no response to soluble antigens [25]; poor mixed lymphocyte reactions [26,27]; and lack of cellular cooperation during the humoral immune response as indicated by enhanced humoral immunity following thymectomy or X-ray irradiation [28,29]. Weak immune responses are known for salamanders in general, and the Mexican axolotl and related tiger salamanders are especially susceptible to ATV infections with high observed mortality rates both in the laboratory and in the field. Indeed, an outbreak of ATV in 2003 at the Indiana Axolotl Colony significantly reduced adult stocks before the virus was contained. By way of comparison, adult *Xenopus* effectively clear close-related FV3 *Ranavirus* with an immune response that includes an early T-cell proliferative phase in the spleen [30].

To further investigate the axolotl immune response to ATV, we used an Affymetrix custom microarray to identify genes that were significantly, differentially expressed in the spleen. We then compared these genes to a list of genes associated with regeneration that were previously identified from *A. mexicanum* using the same microarray platform. We reasoned that such a comparison would allow us to filter gene expression responses of humoral cells induced generally in response to injury and stress from those expressed specifically in response to ATV infection. Also, this comparison would potentially identify gene expression signatures associated with cell proliferation in response to ATV, as we have previously identified many cell proliferation probe sets on the *Ambystoma* genechip that are differentially regulated during spinal cord regeneration [31]. The genes that we describe provide mechanistic insights and new tools to investigate salamander antiviral responses in the laboratory and in natural populations.

### Methods

#### Animal care and surgery protocols

Inbred A. mexicanum eggs from a single full-sib mating were obtained from the Ambystoma Genetic Stock Center at the University of Kentucky. Each A. mexicanum egg and larva was reared in an individual container in aquifer water treated with ReptiSafe and changed weekly. Individuals were fed brine shrimp *ad libitum* for the first four weeks post-hatching and blackworms (Tubifex) ad libitum thereafter. Animals were reared in an environmental chamber on a 12:12 h light:dark cycle at 20°C. At 4.5 months of age, 12 individuals were injected with 100 $\mu$ l of 10<sup>6</sup> p.f.u./ ml of ATV isolated from the axolotl colony and suspended in cell culture medium. This amount of virus was determined to be the minimum lethal dose via injection in previous unpublished experiments (Storfer, unpublished data) and the strain utilized in the experiment was extracted from axolotls that had previously been infected and killed by the virus. Simultaneously, four uninfected (control) individuals were sacrificed in MS222 for spleen and lung removal. Spleens from all animals were flash frozen in liquid nitrogen. The same surgical procedure was performed on four infected individuals following 24, 72 and 144 hours of infection. Spleen tissue was utilized due to its previously noted importance in CD8+ T cell immune responses to Ranaviruses, particularly FV3, in frogs [30] Additionally, spleen is an important immune organ as antigens from the blood are processed in the spleen. Lung tissue was removed for viral quantification as it is an internal organ that can be utilized in early stage virus quantification (Stewart, unpublished data).

During the infection period behavioral observations were taken opportunistically. Total RNA was extracted from spleen with TRIzol (Invitrogen) according to the manufacturer's protocol. RNA isolations were further purified using RNeasy mini columns (Qiagen). The amount of RNA present in each isolate was determined via UV spectrophotometry, and RNA quality was inspected via a 2100 Agilent Bioanalyzer. Sixteen high quality isolates (four replicates at each of four sampling times: 0 (controls), 24, 72, and 144 hours post-infection) were used to make individual-specific pools of biotin labeled cRNA probes. Each of the 16 pools was then independently hybridized to an Amby\_001 custom Affymetrix GeneChip (for a more detailed description of the microarray platform see [31] and [32]). The University of Kentucky Microarray Core Facility generated cRNA probes and performed hybridizations according to standard Affymetrix protocols.

#### **Quality Control and Data Processing**

All quality control and processing analyses were done in R [33] (www.rproject.org). We used the Bioconductor package "affy" (www.bioconductor.org) to perform several quality control analyses at the individual probe level [34,35]. These analyses included: (1) viewing images of the log(intensity) values of the probes on each GeneChip to check for spatial artifacts, (2) investigating measures of central tendency and dispersion by viewing box-plots and histograms of all the GeneChips, (3) viewing pair-wise M versus A plot matrices for replicate GeneChips, and (4) viewing an RNA degradation plot [35] that enables the visualization of the 3' labeling bias associated with all GeneChips simultaneously. Upon conducting these probe level analyses, we background corrected, normalized, and summarized all sixteen GeneChips using the Robust Multi-array Average (RMA) algorithm [36]. Following this, we calculated correlation matrices for replicate GeneChips (four correlation matrices with four GeneChips per matrix; all r from replicate GeneChips > 0.980) on the summarized probeset level data. The strong correlations observed between replicate GeneChips suggests that we were able to obtain a high degree of repeatability within treatments.

#### Data Filtering

Microarrays may not accurately quantify the abundance of lowly expressed genes [37]. Calculating statistical tests for such genes adds to the multiple testing burden that is inherent to microarray studies. To address this issue, we filtered genes whose mean intensity across all 16 GeneChips was greater than the mean of the lowest quartiles ( $25^{th}$  percentiles) across all GeneChips (n = 16, mean = 5.83, SD = 0.06; data presented on a  $log_2$  scale). Upon imposing this filtering criterion, 3619 probe-sets were available for significance testing.

#### Identifying Differentially Expressed Genes

We used the Bioconductor package LIMMA [38,39] to generate moderated *t*statistics for all six of the possible pair-wise contrasts of the four sampling times investigated in our study. LIMMA employs an empirical Bayes methodology that effectively shrinks the sample variances towards a pooled estimate. This approach reduces the likelihood of obtaining large test statistics due to underestimation of the sample variances. The moderated *t*-statistics generated by LIMMA test the null hypothesis that the difference between the two groups being compared is zero (*i.e.*, group 1 – group 2 = 0). LIMMA also generates moderated *F*-statistics that test the null hypothesis that none of the contrasts within a family of contrasts are statistically significant. We corrected for multiple testing by applying the step-up algorithm [40] to the *P*-values of the moderated *F*-statistics associated with our six contrasts. Upon correcting for multiple testing, we identified 2322 genes (probe-sets) that were statistically significant. To prioritize amongst differentially expressed genes, we focused on probe-sets that exhibited two-fold or greater changes at any time-point relative to controls. Any gene that was non-significantly down-regulated but significantly up-regulated at one or more time points was considered up-regulated, and vice versa for classification of up- versus downregulation. We also required that these probe sets have moderated *F*-statistics greater than or equal to the 50<sup>th</sup> percentile of the 2322 *F*-statistics from the statistically significant probe-sets ( $F \ge 12.68$ ). We further limited our analysis to only those probe sets that exhibited significant sequence identity with a human reference sequence. We note that 263 probe-sets with no functional annotation were statistically significant, differentially expressed by  $\ge$  two-fold, and had *F*-values  $\ge 12.68$ .

#### Clustering

Hybridization intensities were averaged within treatment groups (0, 24, 72, and 144 hrs post-infection) and  $log_2$  ratios were calculated for each non-zero sampling time relative to 0 hours post-infection. Genesis v. 1.6.0 [41,42] was used cluster these  $log_2$  ratio data and to generate heat maps. Clustering was conducted using a Self Organizing Map (SOM) algorithm. Default conditions were used with the exception that the SOM was allowed to run for 263,000 iterations. The dimensions of the final SOM are  $2_x*1_y$ . These dimensions were determined by comparing output from several different combinations.

#### Enrichment Analyses

Functional annotation of genes by gene ontology was performed using the Database for Annotation, Visualization and Integrated Discovery (DAVID, [43]).

Functional annotation clustering was performed using the default settings with the exception of using the highest classification stringency.

#### Quantitative real-time PCR

We used quantitative real-time PCR (qPCR) to confirm the results of the microarrays. We estimated a fold change for 24 and 72 hr time points using the  $\Delta\Delta$ ct method of relative quantification [44], utilizing ribosomal protein L 19 as an endogenous control gene. The same total RNA that was used for microarray analysis was used to create cDNA for qPCR using the BioRad iScript cDNA synthesis kit, following manufacturer instruction. Primers for the qPCR were designed using Primer Express 2.0 (Applied Biosystems). Primers were designed to encompass the sequence of GeneChip probe sets (Additional file 1). qPCR was accomplished using SYBR Green chemistry.

To verify that exposed animals were infected and to quantify viral load and replication over time, we performed qPCR on lung tissue with TaqMan chemistry following the protocol detailed in [45]. ANOVA with a Tukey's HSD correction for all pairwise comparisons was performed to determine if viral loads were significantly different across time points.

### Results

#### Viral load and disease pathogenesis

Viral load for each animal was estimated using qPCR and then averaged for each time point (Fig 1). The significant increase in viral load across time points indicates that animals were infected and that viral replication was occurring. ANOVA with a Tukey's correction for multiple comparisons confirmed that viral load increased linearly between

24, 72, and 144 hours post-infection, and all time points were significantly different from all other time points ( $F_{3,44} = 242.56$ ;  $p \le 0.01$ ).

No animals displayed any gross symptoms of ATV infection in terms of hemorrhaging, lesions or edema, either externally or on any internal organs upon euthanasia and subsequent surgery. Similarly, there were no notable changes in behavior observed during the period of infection. This is likely due to the relatively short infection period utilized in this experiment. As noted in the introduction, infected animals often take 8-10 days, or more, to become symptomatic upon infection.

#### Gene clustering and functional annotation

We identified 263 probe sets with statistically significant differences in mRNA abundances between Day 0 and any other subsequent time point (Tables 1, 2). We assume that statistically significant probe sets correspond to genes that were differentially regulated after ATV infection. Cluster analysis of the statistically significant genes identified two groups that exhibited similar changes in mRNA abundance. After ATV infection at Day 0, 158 putative genes showed a significant increase in mRNA abundance at subsequent time points (Figure 2), while 105 transcripts showed a significant decrease (Figure 3). Thus, more genes were up-regulated than down-regulated in response to ATV infection. Overall, DAVID categorized statistically significant genes among 44 different groups that correspond to different biological processes. Eight of these groups contained more genes than would be expected by chance sampling of genes from the microarray (geometric mean p-value < 0.05); these groups were considered significantly enriched with candidate genes relative to other groups (Table 3).. Four of these significant groups contain gene ontologies related to immune response and pathogen response, including

innate immunity, complement activation, lysosome function, and antigen processing and presentation. The most enriched functional group contains genes primarily related to immune function and defense responses. The remaining four functional groups contain gene ontologies related to ion binding, ion transport, vitamin metabolism, and response to an unfolded protein. Many genes that were classified in broader biological process categories that are not directly immunity-related are nonetheless associated with immunity in vertebrates [e.g. 46-48].

#### Genes Up-regulated in Response to ATV

Across all time points, the majority of up-regulated genes were related to immune response or other related functions, such as inflammation and apoptosis. Other upregulated genes pertained to gene functions such as ion binding and transport, membrane related functions, and protein binding and modification. Twenty-three genes (represented by 26 probe sets) demonstrated 2-fold or greater changes at 24 hours post-infection, all of which were up-regulated. Ten of these 23 have functions pertaining to immune response. Of the remaining highly expressed genes, one was associated with inflammation, two to regulation of apoptosis, three to ion binding, three to protein binding and modification, one to transport, one to the extracellular constituent, and one to membrane and glycolipids. Many of these genes showed increasing transcript abundances over time. At 72 hours post infection, 43 genes had a greater than 5-fold change, and 40 genes had a greater than 5-fold change at 144 hours. The highest expression level, 91-fold increase at 144 hours, was observed for *interferon-induced protein with tetracopeptide repeats 5* (IFIT5).

#### Genes Down-regulated in Response to ATV

In contrast to the very high fold changes observed among up-regulated genes, the largest fold change observed among down-regulated genes was approximately 4.9-fold, in *chondroitin sulfate proteoglycan* (NCAN). Five down-regulated genes each code for regulation of transcription and translation. An additional 15 down-regulated genes correspond to 20 probe sets that have functions associated with cell division and mitosis, which was not observed in the up-regulated genes. Other notable down-regulated gene ontologies include one gene corresponding to pinocytosis and endocytosis, and one gene related to natural killer cell mediated cytotoxicity.

#### Validation of Microarray Results Using Quantitative Real-time PCR

We used qPCR to estimate fold changes for nine genes to verify our microarray data (Table 4). For five of the nine genes investigated (56%; *Myxovirus resistance 1, Macrophage receptor with collagenous structure, Complement component 3, Cyclin dependant kinase inhibitor 1B, Vaccinia related kinase 1*) there is good agreement between the microarray and qPCR data. In genes where the microarray estimates of fold change were modest (*Serine dehydratase like , Hemoglobin gamma alpha, Glycogen synthase kinase, Programmed cell death 8*) there is poorer agreement between fold change estimates from these two technologies. However, for this latter group of genes with modest fold change values, the microarray and qPCR data were always within four fold of each other. These results demonstrate that we were able to verify robust differences that were suggested by the microarray data.

#### Analyses to identify proliferation gene expression signatures

Comparison of gene expression after ATV and tail amputation identified 25 genes that are significantly up-regulated in both experimental frameworks (Table 5). No significantly down-regulated genes were identified in common. Several of the commonly up-regulated genes appear to be related to humoral immunity, and membrane and extracellular matrix related functions. Additionally, general stress response genes such as *heat shock 70kDa protein 5* were similarly regulated. None of the cell cycle genes that are significantly up-regulated during tail regeneration were identified in this study. Thus, there was no evidence of cell proliferation by spleen cells after ATV infection.

### **Discussion and conclusion**

Emerging infectious diseases are implicated in the global decline of amphibians and other animals [3, 49-51]. There is urgent need to develop understanding of amphibian immunological responses to pathogens and to identify host genes that may be important in disease resistance. Our study shows that functional genomics provides a means to rapidly meet these needs. We infected Mexican axolotls from the Ambystoma Genetic Stock Center with a viral pathogen that is clearly affecting tiger salamander populations in nature [10,13-15,-19]. Our results show that ATV infection induces transcriptional changes of genes that are known to function in vertebrate immunity. Below we discuss the transcriptional response in more detail and suggest hypotheses to explain why ATV is often lethal to axolotls and other tiger salamanders.

We detected significant gene expression changes 24 hours post infection. Many of these gene expression changes likely reflect transcription within lymphocytes, as they are the predominant cell type in the spleen of juvenile and adult axolotls [52]. Indeed, the

functions of many of these genes are associated with neutrophil, dendritic, and macrophage cell functions, including cytokine signaling (*chemokine (C-X-C motif*) receptor 4), phagocytosis and destruction of phagocytised particles (disabled homolog 2, mitogen-responsive phosphoprotein, neutrophil cytosolic factor 2, lysosomal-associated membrane protein 1, RAS homolog gene family, member B), complement (complement factor B, complement component 3), and inflammation (pentraxin related gene, rapidly induced by IL-1 beta, cytochrome B-245 beta polypeptide, n-myc and STAT interactor). Up-regulation of complement components that are known to function in the removal of viral particles, and up-regulation of the stress-associated transcription factor *jun-b*, clearly shows that ATV induced a humoral gene expression response in the axolotl. Further support for this idea was obtained by comparing ATV-induced gene expression changes to changes identified from a previous microarray experiment using A. mexicanum and the same microarray platform. Twenty-five genes that were up-regulated in response to ATV infection were also identified as significantly up-regulated during regeneration [31]. In both microarray studies, blood was not perfused from tissues prior to tissue collection and it is known that leukocyctes express genes during the early wound-healing phase of spinal cord and limb regeneration. Thus, it seems likely that many of the early gene expression changes that we observed in response to ATV-infection reflect a general, humoral transcriptional response to stress.

In addition to this general humoral response, the gene expression patterns that we observed suggest that the Mexican axolotl manifests an antiviral transcriptional response that is not unlike that observed in other vertebrates. For example, ATV infection clearly induces an interferon-mediated, antiviral response. Although probe sets for interferon

genes are not represented on the GeneChip, we estimate based upon literature surveys that at least 20% of the significant genes that we identified are known in other systems (*in vitro* and *in vivo*) to be involved in interferon-mediated transcription [53-55]. These genes exhibited some of the largest fold-changes and include two primary transcription factors that compete to activate (*interferon regulatory factor 1*, up-regulated) and repress (*interferon regulatory factor 2*, down-regulated) transcription of interferon-alpha and beta (Type 1 interferon), and inferon-inducible genes that recognize and degrade intra-cellular viral nucleic acid (*interferon induced with helicase C domain 1*). Considering further that four of the most highly enriched functional groups also contained genes relating to the immune response and pathogen response, the results show that axolotls mount a robust anti-viral response from 24-144 hours post-infection.

Given the robust immunological transcription response that we observed, it is curious why ATV is so virulent to tiger salamanders. In the closely related *Ranavirus* frog virus 3 (FV3), larval *Xenopus laevis* succomb to FV3 but adults effectively clear virons and develop lasting resistance to future infection [56]. Adult resistance in *X. laevis* is correlated with a significant proliferation of cytotoxic CD8<sup>+</sup> T cells in the spleen upon infection (within 6 days), as well as increased mortality upon CD8<sup>+</sup> T cell depletion [30,57]. Mortality events due to ATV are more significant among larvae in natural tiger salamander populations, however metamorphosed adult tiger salamanders are more susceptible than larvae to ATV infection in the lab [18]. It is well established that Mexican axolotls have a less complicated immune system and never develop the type of mature immune response typical of amniote vertebrates [21-29, 52]. We did not observe any gene expression changes that would indicate proliferative leukocyte responses in axolotl spleen. Perhaps this is because we used juvenile axolotls that are incapable of such a response. However, it is also possible that ATV maybe more resistant to the immune response mounted by *A. mexicanum* than FV3 is to the *Xenopus* immune response. Phylogenetic analyses indicate ATV is more closely related to iridoviruses found in fish than to FV3, which suggests a relatively recent host switch occurring with the introduction of sportfish to areas of the southwestern United States [15]. Iridoviruses found in sportfish have a larger genome and contain more ORFs related to immune evasion than FV3, which could also be related to improved performance of this virus on the salamander host [15]. Further studies are needed to better understand the ontogeny of immunological responses in axolotls, the virulence of different ranaviruses, and the role of innate versus adaptive immunity in ATV infection.

Our study has identified hundreds of new candidate genes for laboratory and field studies of stress and disease in tiger salamanders. Significantly more gene candidates will undoubtedly be discovered using a higher content, 2<sup>nd</sup> generation microarray that is currently under development. Genomic and bioinformatics tools make *Ambystoma* a powerful system for wildlife disease research. In particular, molecular information can be quickly cross-referenced from a genetically homogeneous strain that is available for laboratory studies (Mexican axolotl), to other closely related tiger salamander species in North America [20]. Such power is needed to quickly understand how ATV and other pathogens are overwhelming amphibian immune responses and causing population declines in nature.

### **Competing Interests**

No competing interests are declared.

### **Authors Contributions**

JDS performed infections and surgeries and drafted the manuscript. RBP performed microarray statistical and bioinformatic analyses. AS, CKB, and SRV contributed to experimental design and manuscript editing.

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### **Figure legends**

**Figure 1**. Log values of viral particles quantified with quantitative real-time PCR across all time points.

**Figure 2.** Expression profiles for cluster 1. Each row represents an individual gene, and each column a post-infection time point. Red coloration indicates increased expression of a gene relative to uninfected animals, and green indicates decreased expression. Genes (Cluster 1, n=158) identified to be significantly up-regulated in response to ATV infection.

**Figure 3**. Expression profiles for cluster 2. Each row represents an individual gene, and each column a post-infection time point. Red coloration indicates increased expression of a gene relative to uninfected animals, and green indicates decreased expression. Genes (Cluster 2, n=105) identified to be significantly down-regulated in response to ATV infection.

## Tables

Table 1.	Genes	which	were	signi	ificantly	y ur	o-regu	ılated	at	any	time	point.	Numbers

indicate fold change at that time point.

Gene ID	Gene Name	24 hr	72 hr	144 hr
immune response	Gene Name	24 11	7211	144 11
SRV_03329_at	INTERFERON-INDUCED PROTEIN WITH TETRATRICOPEPTIDE REPEATS 5	11.85	82.22	91.04
SRV 01342 at	INTERFERON REGULATORY FACTOR 1	1.94	12.71	12.79
SRV_01343_a_at	INTERFERON REGULATORY FACTOR 1	2.10	16.07	15.63
SRV_13637_a_at	INTERFERON INDUCED TRANSMEMBRANE PROTEIN 3 (1- 8U)	1.75	8.48	10.88
SRV_01199_a_at	CLUSTERIN	1.05	4.35	8.02
SRV_00442_at	SOLUTE CARRIER FAMILY 11 (PROTON-COUPLED DIVALENT METAL ION TRANSPORTERS), MEMBER 1	1.28	6.17	5.38
SRV_01303_a_at	GUANYLATE BINDING PROTEIN 1, INTERFERON- INDUCIBLE, 67KDA	1.10	2.35	2.49
SRV_02828_at SRV_02072_at	2'-5'-OLIGOADENYLATE SYNTHETASE 3, 100KDA CATHEPSIN S	2.47 1.54	13.56 5.51	9.35 6.43
SRV_02588_a_at	LECTIN, GALACTOSIDE-BINDING, SOLUBLE, 3 BINDING PROTEIN	4.04	26.37	24.40
SRV_02586_at	LECTIN, GALACTOSIDE-BINDING, SOLUBLE, 3 BINDING PROTEIN	3.88	21.30	18.03
response to virus				
SRV_03073_at	INTERFERON-INDUCED PROTEIN 44	2.05	9.44	13.16
SRV_01439_a_at	MYXOVIRUS (INFLUENZA VIRUS) RESISTANCE 1, INTERFERON-INDUCIBLE PROTEIN P78 (MOUSE) MYXOVIRUS (INFLUENZA VIRUS) RESISTANCE 1,	3.13	29.23	36.80
SRV_01441_at	INTERFERON-INDUCIBLE PROTEIN P78 (MOUSE)	3.26	28.24	35.84
SRV_04604_s_at SRV_04604_at	INTERFERON INDUCED WITH HELICASE C DOMAIN 1 INTERFERON INDUCED WITH HELICASE C DOMAIN 1	6.81 7.27	27.34 25.43	24.15 22.62
SRV_04518_at	RIBONUCLEASE L (2',5'-OLIGOISOADENYLATE SYNTHETASE-DEPENDENT)	3.30	7.38	8.66
cell defense response				
SRV_00353_at	NEUTROPHIL CYTOSOLIC FACTOR 2 (65KDA, CHRONIC GRANULOMATOUS DISEASE, AUTOSOMAL 2)	1.05	2.07	1.73
SRV_00264_a_at	MYELOPEROXIDASE	1.60	7.60	4.10
antigen processing and presentation				
SRV_05347_at	PROTEASOME (PROSOME, MACROPAIN) SUBUNIT, BETA TYPE, 9 (LARGE MULTIFUNCTIONAL PEPTIDASE 2)	1.27	3.73	4.25
lysosome/ endosome SRV 02581 a at	LYSOSOMAL-ASSOCIATED MEMBRANE PROTEIN 1	1.23	2.26	2.06
SRV_12596_at	EH-DOMAIN CONTAINING 1	1.22	4.16	3.47
inflammatory response				
SRV_10702_at	CYTOCHROME B-245, BETA POLYPEPTIDE (CHRONIC	1.69	5.58	4.40
 SRV 00330 at	GRANULOMATOUS DISEASE) CYTOCHROME B-245, BETA POLYPEPTIDE (CHRONIC	1.59	4.74	3.89
SRV_01877_at	GRANULOMATOUS DISEASE) CHEMOKINE (C-X-C MOTIF) RECEPTOR 4	1.29	5.70	8.08
SRV_01877_at	N-myc (and STAT) interactor	1.15	2.29	2.29
SRV_00453_a_at	TRANSFORMING GROWTH FACTOR, BETA 1 (CAMURATI- ENGELMANN DISEASE)	1.12	2.19	2.18
SRV_01617_a_at	PENTRAXIN-RELATED GENE, RAPIDLY INDUCED BY IL-1 BETA	2.83	12.82	17.76
cell cycle				
SRV_02067_at SRV_11882_s_at	POLO-LIKE KINASE 3 (DROSOPHILA) RAS HOMOLOG GENE FAMILY, MEMBER B	0.93 1.18	3.82 2.33	6.18 2.40

SRV_02051_at         FAS HOMOLOG GENE FAMILY, MEMBER B         1.26         2.15         2.28           SRV_02050_at         FAS HOMOLOG GENE FAMILY, MEMBER B         1.23         2.51         2.55           SRV_01520_at         FAS HOMOLOG GENE FAMILY, MEMBER B         1.23         2.51         2.55           SRV_01520_at         Provestore for the second					
SRV_02050_atl         PAS HOMOLOG GENE FAMILY, MEMBER B         1.23         2.51         2.56           SRV_01520_atl         EXOSTOSES (MULTPLE)1         0.90         2.07         1.90           glycolysis         pyruvate kinase, muscle         1.41         4.60         4.22           SRV_01519_atl         pyruvate kinase, muscle         1.65         5.94         5.43           B_s_atl         ALDOLASE A, FRUCTOSE BISPHOSPHATE         1.22         3.14         2.86           SRV_01351_att         JUN B PROTO-ONCOGENE         1.76         7.82         7.54           SRV_03210_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR I         9.52         7.0         2.45           SRV_02251_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR I         1.55         2.70         2.45           SRV_02230_at         COMPLEMENT FACTOR B         3.40         10.37         13.66           SRV_02399_a_st         COMPLEMENT FACTOR B         3.46         10.33         13.66           SRV_02399_a_st         CASPASE ECQUITIMENT DOMAIN FAMILY, MEMBER 6         1.35         4.73         4.80           SRV_02399_a_st         CASPASE ECQUITIMENT DOMAIN FAMILY, MEMBER 6         1.35         4.76         1.81         3.46         10.33         3.26         3.76	SBV 02051 at	BAS HOMOLOG GENE FAMILY, MEMBER B	1.26	2.15	2.28
SRV_00154_att         EXOSTOSES (MULTIPLE) 1         0.50         2.07         1.90           glycolysis SRV_01520_at         pyruvate kinase, muscle         1.41         4.60         4.22           SRV_01520_at         pyruvate kinase, muscle         1.41         4.60         4.22           SRV_01519_a_att         pyruvate kinase, muscle         1.41         4.60         4.22           SRV_01336_a_at         INHEIDTO ONCOGENE         1.76         7.82         7.54           SRV_01336_a_at         INHEIDTO ONCOGENE         1.76         7.82         7.54           SRV_02310_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR 1         0.55         2.70         2.46           SRV_02316_at         SPE HOMOLOG FALLYOP HELLY PROTEIN         1.56         3.14         3.57           SRV_03646_at         SPE HOMOLOG FALLYOP HELLY PROTEIN         1.23         4.52         2.30           sRV_0037_a_at         COMPLEMENT COMPONENT 3         1.80         9.79         1.366           SRV_0239_a_at         CASPASE RECOUNTERN TOWAIN FAMILY, MERER 6         1.35         4.78           SRV_02312_at         HEAT SHOCK PROTEIN GRAD RETAIN (GULCOSE-REGULATED PROTEIN 38/LOCOSE-REGULATED PROTEIN 38/LOCOSE-REGULATED CINCOSE REGULATED CINCOSE REGULATED CINCOSE REGULATED CINCOSE REGULATED PROTEIN 38/LOCOSE REGULATED CINCOSE REGULATED CINCOSE R					
glycolysis SRV_01520_at         pyruvate kinase, muscle         1.41         4.60         4.22           B, S, at         ALDOLASE A, FRUCTOSE BISPHOSPHATE         1.22         3.14         2.266           transcription SRV_01351_at         JUN B PROTO ONCOGENE         1.76         7.82         7.54           SRV_0136_a_at         JUN B PROTO ONCOGENE         1.66         3.14         2.26           SRV_02310_at         EUKAPOTIC TANASCATOR TO TANGATINATION FACTOR I         0.95         2.27         2.45           SRV_02571_at         INHIBITOR OF DIAL BINDING S DOMINANT NEGATIVE         1.66         3.14         2.22         2.30           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_0299_a_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           SRV_02905.at         HEAT SHOCK 7X0AP PROTEIN 5 (GLUCOSE REGULATED D         2.70         10.28         7.60           SRV_0313.a, at         CASPASE RECRUITINEND COMAIN FAMICINALIN-2)         1.18         3.45         2.71           SRV_0314.a, at         HEAT SHOCK 7X0AP APROTEIN 5 (GLUCOSE REGULATED D         2.70         10.28         2.71           SRV_0315.a, at         CASPASE RECRUITINEND COMAIN FAMICINALIN-2)					
SRV_01520_at         pyruxate kinase, muscle         1.41         4.60         4.22           B_S,at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.22         3.14         2.96           Itranscription         JUN B PROTO-ONCOGENE         1.76         7.82         7.54           SRV_01531_at         JUN B PROTO-ONCOGENE         1.76         7.82         7.54           SRV_0364_at         INHIBITOR OF DNA BINDING 3, DOMINANT NEGATIVE         1.56         3.14         3.57           SRV_03646_at         SPEN HOMOLOG (IROSOPHILA) NI TERMINAT NEGATIVE         1.56         3.14         3.57           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_02399_a.g.tl         MATRIX METALLOPFEDIASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_02390_a.g.tl         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.76           SRV_02390_a.g.tl         CASPASE RECOUNTENT TO MAIN FAMILY, MEMBER 6         1.36         4.77         4.80           SRV_0237_a.g.tl         CASPASE RECOUNTENT TO MAIN FAMILY, MEMBER 6         1.36         4.78         4.80           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7	SRV_00154_at	EXOSTOSES (MULTIPLE) T	0.90	2.07	1.90
SRV_01520_at         pyruxate kinase, muscle         1.41         4.60         4.22           B_S,at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.22         3.14         2.96           Itranscription         JUN B PROTO-ONCOGENE         1.76         7.82         7.54           SRV_01531_at         JUN B PROTO-ONCOGENE         1.76         7.82         7.54           SRV_0364_at         INHIBITOR OF DNA BINDING 3, DOMINANT NEGATIVE         1.56         3.14         3.57           SRV_03646_at         SPEN HOMOLOG (IROSOPHILA) NI TERMINAT NEGATIVE         1.56         3.14         3.57           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_02399_a.g.tl         MATRIX METALLOPFEDIASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_02390_a.g.tl         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.76           SRV_02390_a.g.tl         CASPASE RECOUNTENT TO MAIN FAMILY, MEMBER 6         1.36         4.77         4.80           SRV_0237_a.g.tl         CASPASE RECOUNTENT TO MAIN FAMILY, MEMBER 6         1.36         4.78         4.80           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7					
SRV_01519_a_at         private kinase, muscle         1.56         5.54         5.48           B_s_at         ALDOLASE A, FRUCTOSE-BISHOSPHATE         1.22         3.14         2.36           Itranscription         JUN B PROTO-ONCOGENE         1.76         7.82         7.54           SRV_01351_at         JUN B PROTO-ONCOGENE         1.76         7.82         7.54           SRV_02310_at         EUKARVOTIC TRANSLATION TREMINATION FACTOR I         0.95         2.70         2.45           SRV_03846_at         SPEN HOMOLOG, TRANSCRIPTIONAL REGULATOR         1.15         2.22         2.30           Omplement         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           SRV_02517_at         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           SRV_02516_at         CASPASE RECORUTIMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04970_a_at         CASPASE RECORUTIMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04370_a_at         CASPASE RECORUTIMENT DOMAIN FAMILY, MEMBER 1         1.3         3.53         3.26           SRV_04970_a_at         CASPASE RECORUTIMENT DOMAIN FAMILY, MEMBER 1         1.21         3.56         3.38           SRV_04970_a_at         <					
Tell_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.22         3.14         2.96           transcription SRV_01351_at         JUN B PROTO-ONCOGENE         1.76         7.82         7.54           SRV_01351_at         IMHBITO OF DNA BINDING 3, DOMINANT NEGATIVE         1.56         3.14         3.57           SRV_02310_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR 1         0.95         2.70         2.45           SRV_02310_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR 1         0.95         2.70         2.45           SRV_02517_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_0145_a,at         COMPLEMENT FACTOR B         3.46         18.33         20.05           apoptosis         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_02399_a,at         CAMPAGE REPTIDASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_04300,a,at         CASPASE RECRUITEMENT TOMAIN FAMILY, MEMBER 6         1.35         4.76         4.80           SRV_04300,a,at         CASPASE RECRUITEMENT TOMAGE- NOCES-REGULATED         1.30         2.56         3.36           SRV_0132,a,at         GROWTH AREST AND DNA-DAMAGE- NOLOGE- REGULATED         1.	SRV_01520_at	pyruvate kinase, muscle	1.41	4.60	4.22
Tell_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.22         3.14         2.96           transcription SRV_01351_at         JUN B PROTO-ONCOGENE         1.76         7.82         7.54           SRV_01351_at         IMHBITO OF DNA BINDING 3, DOMINANT NEGATIVE         1.56         3.14         3.57           SRV_02310_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR 1         0.95         2.70         2.45           SRV_02310_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR 1         0.95         2.70         2.45           SRV_02517_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_0145_a,at         COMPLEMENT FACTOR B         3.46         18.33         20.05           apoptosis         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_02399_a,at         CAMPAGE REPTIDASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_04300,a,at         CASPASE RECRUITEMENT TOMAIN FAMILY, MEMBER 6         1.35         4.76         4.80           SRV_04300,a,at         CASPASE RECRUITEMENT TOMAGE- NOCES-REGULATED         1.30         2.56         3.36           SRV_0132,a,at         GROWTH AREST AND DNA-DAMAGE- NOLOGE- REGULATED         1.	SBV 01519 a at	pyruvate kinase, muscle	1.56	5.94	5.48
transcription SRV_01351_at SRV_01351_at         JUN B PROTO-ONCOGENE HELIX-LOOP-HELIX PROTEIN         1.76         7.82         7.54           SRV_01351_at         INHIBITOR OF DNA BINDING 3, DOMINANT NEGATIVE HELIX-LOOP-HELIX PROTEIN         1.56         3.14         3.57           SRV_02310_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR 1         0.95         2.70         2.45           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SPK_0137_a, at         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           SRV_02399_a_atl         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA GELATINASE, 92KDA PROTEIN 5 (GULCOSE-REGULATED PROTEIN, 78KDA)         2.70         10.28         7.60           SRV_0216_a_t         HEAT SHOCK 7KOA PROTEIN 5 (GULCOSE-REGULATED PROTEIN, 78KDA)         1.83         4.84         2.70         10.28         7.60           SRV_03023_a_t         CASPASE RECOLUMENT ON MORTALIN-2)         1.18         3.45         2.71         3.48         4.80           SRV_0132_a_t         EVKARYOTIC TRANSLAWIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_0213_a_t         GRAWA         EPSILON 1         1.21         3.56         3.38           SRV_0213_a_t         EVANDARO NANDANA					
SRV_01351_at         JUN B PROTO-ONCOGENE         1.76         7.82         7.82           SRV_01308_a_at         INHISTOR OF DNA BINDING S, DOMINANT NEGATIVE HELIX-LOOP-HELIX PROTEIN         1.56         3.14         3.57           SRV_03646_at         SPEN HOMOLOG, TRANSCRIPTIONAL REGULATOR (DROSOPHILA)         1.52         2.22         2.30           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_0137_a_at         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           SRV_02390_a_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           apoptosis         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA (GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.19         6.79         8.78           SRV_02390_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.36         4.78         4.80           SRV_0470_a_at         CASPASE RECRUITMENT DOMAIN FACTOR 1         1.21         3.56         3.38           SRV_0212_at         EUKARYOTIC TRANSLATION NAMAGE-INDUCIBLE, BETA         1.30         2.264         2.71           SRV_0213_a_at         GADWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.33         3.26         3.33         3.26           SRV_01812_a_at	D_3_4(	ALDOLAGE A, THOUTOBE DIGITIOUT HATE	1.22	0.14	2.50
SRV_01351_at         JUN B PROTO-ONCOGENE         1.76         7.82         7.82           SRV_01308_a_at         INHISTOR OF DNA BINDING S, DOMINANT NEGATIVE HELIX-LOOP-HELIX PROTEIN         1.56         3.14         3.57           SRV_03646_at         SPEN HOMOLOG, TRANSCRIPTIONAL REGULATOR (DROSOPHILA)         1.52         2.22         2.30           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_0137_a_at         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           SRV_02390_a_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           apoptosis         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA (GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.19         6.79         8.78           SRV_02390_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.36         4.78         4.80           SRV_0470_a_at         CASPASE RECRUITMENT DOMAIN FACTOR 1         1.21         3.56         3.38           SRV_0212_at         EUKARYOTIC TRANSLATION NAMAGE-INDUCIBLE, BETA         1.30         2.264         2.71           SRV_0213_a_at         GADWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.33         3.26         3.33         3.26           SRV_01812_a_at	the second sector the sec				
SRV_01396_a_at         INHIBITOR OF DNA BINDING 3, DOMINANT NEGATIVE HELLX.COP-HELLY PROTEIN NELX.COP-HELLY PROTEIN         1.56         3.14         3.57           SRV_02310_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR 1         0.95         2.70         2.45           SRV_02310_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR 1         0.95         2.70         2.45           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_0137_a_at         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           SRV_0239_a_at         CAMPLEMENT COMPONENT 3         1.80         3.46         18.33         20.05           SRV_0239_a_at         CASPASE RECRUITMENT DOMAIN FACTOR B         3.46         18.33         20.05           SRV_0216_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.36         3.46         2.80         2.64           SRV_01212_at         EVARAVICIC TRANSLATION FACTOR 1         1.21         3.56         3.38         2.80         2.84         2.80         2.80         2.84         2.80         2.84         2.85         3.36         2.85					
SHV_00135.8_a         HELIX-LOOP-HELIX PROTEIN         1.56         3.14         3.57           SRV_03046_at         EUKARVOTIC TRANSLATION TERMINATION FACTOR 1         0.95         2.70         2.45           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           sRV_01145_a_at         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           SRV_02516_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           SRV_02516_at         GASPASE RECONTINE IN COMPONENT 3         1.00         7.70         10.28         7.60           SRV_02132_at         GASPASE RECONTINENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         GASPASE RECONTIVENT DOMAIN FAMILY, MEMBER 1         1.31         3.53         3.26           SRV_03753_at         GASPASE RECONTRANDON ADAMAGE-INDUCIBLE, ETA         1.39         7.54         8.89           SRV_03003_a,at         FUCOHFORE C, SOMATIC         1.30         3.53         3.26           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, ETA         1.39         7.53	SRV_01351_at		1.76	7.82	7.54
SRV_02310_at         EUKAPUOTIC TRANSLATION IF RATION FACTOR 1         0.95         2.70         2.45           SRV_02571_at         SPEN HOMOLOG, TRANSCRIPTIONAL REGULATOR         1.15         2.22         2.30           complement         SPEN HOMOLOG, TRANSCRIPTIONAL REGULATOR         1.15         2.22         2.30           complement         SRV_0137_att         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           sRV_02399_a_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           sRV_02390_a_tt         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_02390_a_tt         CASPASE RECRUMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04970_a_tt         CASPASE RECRUMENT DOMAIN FAMILY, MEMBER 6         1.30         2.80         2.64           SRV_01232_at         EUKARYOTIC TRANSLATION LLONGATION FACTOR 1         1.21         3.56         3.38           SRV_02132_at         HEAT SHOCK YORD PROTEIN 90KDA BETA (GRP94), MEMBER 1         1.31         3.53         3.26           SRV_03023_a_tt         HEAT SHOCK NONT PANTERIX         S.99         7.44         2.85         3.35           SRV_03023_a_tt         HEAT SHOCK NONT PANTERIX         S.99         <		INHIBITOR OF DNA BINDING 3, DOMINANT NEGATIVE	1 50	0.14	0.57
SRV_02310_at         EUKARYOTIC TRANSLATION TERMINATION FACTOR 1         0.95         2.70         2.45           SRV_03646_at         SPEN HOMOLOG, TRANSCHIPTIONAL REGULATOR (DROSOPHILA)         1.15         2.22         2.30           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_00137_a_at         COMPLEMENT COMPONENT 3         1.80         7.79         13.66           sRV_01145_a_at         COMPLEMENT COMPONENT 3         3.46         18.33         20.05           apoptosis         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.70         10.28         7.60           SRV_0239a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.30         2.80         2.64           SRV_02132_at         EUKARYOTIC TRANSLATION FLONGATION FACTOR 1         1.21         3.56         3.38           SRV_01612_at         HEAT SHOCK 70KDA PROTEIN 9(MORTALIN-2)         1.18         3.32.66         3.38           SRV_01612_at         HEAT SHOCK PROTEIN 90KDA BETA (GRP94), MEMBER 1         1.31         3.59         7.44         2.70         3.73         3.26	SRV_01336_a_at	HELIX-LOOP-HELIX PROTEIN	1.56	3.14	3.57
SRV_03046_at         SPEN HOMOLOG, TRANSCRIPTIONAL REGULATOR (DROSOPHILA)         1.15         2.22         2.30           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_0173_at         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           SRV_02571_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           apoptosis         SRV_02516_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           SRV_02516_at         GASPASE RECUTIVENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_0373_at         GASPASE RECUTIVENT DOMAIN FAMILY, MEMBER 6         1.38         3.45         2.70         10.28         7.60           SRV_03703_at         GASPASE RECUTIVENT DOMAIN FAMILY, MEMBER 6         1.38         3.45         2.73         2.80         2.64           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE INDUGIELE DETA         1.39         7.34         8.89           SRV_00157_a_at         HEAT SHOCK PROTEIN 90KDA BETA (GRPA), MEMBER 1         1.31         3.53         3.26           SRV_00128_a         HEAT SHOCK PROTEIN 90KDA BETA (GRPA), MEMBER 1         1.39         7.34         8.89	SBV 02310 at		0.95	2 70	2 4 5
SHV_00849_at         (DROSOPHILA)         I.13         2.22         2.30           SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement         SRV_0137_a_at         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           apoptosis         SRV_01145_a_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           SRV_02399_a_at         GELATINASE 9/GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_0239_a_at         GELATINASE 9/GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.76         4.80           SRV_02132_at         HEAT SHOCK 70KDA PROTEIN 96 (MORTALIN-2)         1.18         3.45         2.80         2.64           SRV_02132_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, ETA         1.39         7.34         8.49           SRV_00302_a_at         GAMMA         GAMMA         1.27         4.78         4.42           SRV_00282_a_at         SUDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78 </td <td>0111_02010_01</td> <td></td> <td>0.00</td> <td>2.70</td> <td>2.40</td>	0111_02010_01		0.00	2.70	2.40
SRV_02571_at         HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)         1.23         4.52         6.10           complement SRV_0145_a_at         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           apoptosis         SRV_0239_a_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           spoptosis         SRV_0239_a_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           sRV_02516_at         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.70         10.28         7.60           SRV_02164_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04300_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.30         2.80         2.64           EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.31         3.53         3.28         3.38           SRV_0323_a.t         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_05162_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_0506_a_at         SYNDECAN BINDING PROTEIN S(WATENIN)         1.27         4.78         4.42           SRV_00202_a         UDP-NACTEYLGUCOSAMINE PYROPHOS	SRV 03646 at		1.15	2.22	2.30
complement SRV_00137_a_atl         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           apoptosis         SRV_01145_a_atl         COMPLEMENT FACTOR B         3.46         18.33         20.05           SRV_02399_a_atl         MATRIX METALLOPEPTIDASE 9 (GELATINASE 9.92KDA GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.19         6.79         8.78           SRV_02399_a_atl         MATRIX METALLOPEPTIDASE 9 (GELATINASE 9.92KDA CASPASE RECULATED PROTEIN, 78KDA)         2.70         10.28         7.60           SRV_02164_a_atl         HEAT SHOCK YORDA PROTEIN 96 (MORTALIN-2)         1.13         3.45         2.17           SRV_02132_atl         CASPASE RECULTMENT DOWAIN FAMILY, MEMBER 6         1.30         2.80         2.64           SRV_02132_a_atl         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03023_a_atl         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BITA         1.31         3.53         3.26           SRV_05147_a_atl         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00138_a_atl         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BITA         1.18         2.18         2.55           SRV_0137_a_atl         UDP-N-ACTEYLGUCOSAMINE PYROPHOSPHORYLASE 1         1.44         2.14         2.14         2.14					
SRV_0137_a_att         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           apoptosis         SRV_01145_a_att         COMPLEMENT FACTOR B         3.46         18.33         20.05           sRV_02399_a_att         MATRIX METALLOPEPTIDASE 9 (GELATINASE, 92KDA         2.19         6.79         8.78           SRV_02396_a_tt         MATRIX METALLOPEPTIDASE 9 (GELATINASE, 92KDA         2.19         6.79         8.78           SRV_04970_a_tt         GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.10         10.28         7.60           SRV_06154_a_tt         HEAT SHOCK 70KDA PROTEIN 96 (MORTALIN-2)         1.18         3.45         2.70         10.28         7.60           SRV_02132_att         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.33         4.78         4.80           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.30         3.53         3.26           SRV_03023_a_att         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.32         4.55         3.35           SRV_03147_a_att         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_0322_a_att         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_0137_a_att         UDP-N-ACTEVLGLUCOSAMINE	SRV_02571_at	HAIRY AND ENHANCER OF SPLIT 1, (DROSOPHILA)	1.23	4.52	6.10
SRV_0137_a_att         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           apoptosis         SRV_01145_a_att         COMPLEMENT FACTOR B         3.46         18.33         20.05           sRV_02399_a_att         MATRIX METALLOPEPTIDASE 9 (GELATINASE, 92KDA         2.19         6.79         8.78           SRV_02396_a_tt         MATRIX METALLOPEPTIDASE 9 (GELATINASE, 92KDA         2.19         6.79         8.78           SRV_04970_a_tt         GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.10         10.28         7.60           SRV_06154_a_tt         HEAT SHOCK 70KDA PROTEIN 96 (MORTALIN-2)         1.18         3.45         2.70         10.28         7.60           SRV_02132_att         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.33         4.78         4.80           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.30         3.53         3.26           SRV_03023_a_att         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.32         4.55         3.35           SRV_03147_a_att         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_0322_a_att         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_0137_a_att         UDP-N-ACTEVLGLUCOSAMINE					
SRV_0137_a_att         COMPLEMENT COMPONENT 3         1.80         9.79         13.66           apoptosis         SRV_01145_a_att         COMPLEMENT FACTOR B         3.46         18.33         20.05           sRV_02399_a_att         MATRIX METALLOPEPTIDASE 9 (GELATINASE, 92KDA         2.19         6.79         8.78           SRV_02396_a_tt         MATRIX METALLOPEPTIDASE 9 (GELATINASE, 92KDA         2.19         6.79         8.78           SRV_04970_a_tt         GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.10         10.28         7.60           SRV_06154_a_tt         HEAT SHOCK 70KDA PROTEIN 96 (MORTALIN-2)         1.18         3.45         2.70         10.28         7.60           SRV_02132_att         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.33         4.78         4.80           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.30         3.53         3.26           SRV_03023_a_att         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.32         4.55         3.35           SRV_03147_a_att         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_0322_a_att         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_0137_a_att         UDP-N-ACTEVLGLUCOSAMINE	complement				
SRV_01145_a_at         COMPLEMENT FACTOR B         3.46         18.33         20.05           apoptosis         SRV_02399_a_at         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.19         6.79         8.78           SRV_02516_at         HEAT SHOCK 70KDA PROTEIN 5 (GLUCOSE-REGULATED PROTEIN 78KDA)         2.70         10.28         7.60           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_0412_a_at         HEAT SHOCK 70KDA PROTEIN 98 (MORTALIN-2)         1.18         3.45         2.71           SRV_01232_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_0323_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_0323_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, 1.00         3.73         5.99           SRV_05107_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           C s_at         pHOSPHOSMANE PYROPHONSPHORYLASE 1         1.84         2.61         2.65           SRV_0129_a_at         GANMA         1.00         3.73         5.99           SRV_0202_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27		COMPLEMENT COMPONENT 3	1.80	9 79	13.66
apoptosis         MATRIX METALLOPEPTIDASE 9 (GELATINASE B, 92KDA         2.19         6.79         8.78           SRV_02516_at         HEAT SHOCK 70KDA PROTEIN, 78KDA)         2.70         10.28         7.60           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA 1.39         7.34         8.89           SRV_03147_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00132_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00132_a_at         GAMMA         1.21         2.64         2.59         3.55           SRV_00132_a_at         GAMMA         1.22         4.75         3.42         2.59           SRV_00138_a.at         C_5.at         ALDOLASE A, FRUCTOSE BISPHOSPHATE         1.24         2.64 <td< td=""><td></td><td></td><td></td><td></td><td></td></td<>					
SRV_02399_a_at         MATRIX METALLOPEPTIDASE 9 (GELATINASE 8, 92KDA GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.19         6.79         8.78           SRV_02516_at         GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.70         10.28         7.60           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.76         3.73           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, DETA         1.00         3.73         5.99           SRV_03023_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_03023_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.32           SRV_03024_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.32           SRV_03026_at         SYNDECAN BINDING PROTEIN (SYN	SRV_01145_a_at	COMPLEMENT FACTOR B	3.46	18.33	20.05
SRV_02399_a_at         MATRIX METALLOPEPTIDASE 9 (GELATINASE 8, 92KDA GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.19         6.79         8.78           SRV_02516_at         GELATINASE, 92KDA TYPE IV COLLAGENASE)         2.70         10.28         7.60           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.76         3.73           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, DETA         1.00         3.73         5.99           SRV_03023_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_03023_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.32           SRV_03024_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.32           SRV_03026_at         SYNDECAN BINDING PROTEIN (SYN					
SRV_02516_at         GELATINASE, 92KDA TYPE IV COLLAGENASE.)         2.19         6.79         6.76           SRV_02516_at         HEAT SHOCK 70KDA PROTEINS, 5(GLUCOSE-REGULATED PROTEIN, 78KDA)         2.70         10.28         7.60           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04300_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.30         2.80         2.64           SRV_0132_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03735_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.30         3.73         5.99           metabolic process         SRV_05147_a_at         glucose-6-phosphate dehydrogenase         1.32         4.78         4.42           SRV_00108_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.42         2.61         2.65           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.42         2.62         2.45         3.35           SRV_00129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.42         2.64         2.30         2.13	apoptosis				
SRV_02516_at         GELATINASE, 92KDA TYPE IV COLLAGENASE.)         2.19         6.79         6.76           SRV_02516_at         HEAT SHOCK 70KDA PROTEINS, 5(GLUCOSE-REGULATED PROTEIN, 78KDA)         2.70         10.28         7.60           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04300_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.30         2.80         2.64           SRV_0132_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03735_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.30         3.73         5.99           metabolic process         SRV_05147_a_at         glucose-6-phosphate dehydrogenase         1.32         4.78         4.42           SRV_00108_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.42         2.61         2.65           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.42         2.62         2.45         3.35           SRV_00129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.42         2.64         2.30         2.13		MATRIX METALLOPEPTIDASE 9 (GELATINASE B. 92KDA			
SRV_02516_at         HEAT SHOCK 70KDA PROTEINS (GLUCOSE-REGULATED PROTEIN, 78KDA)         2.70         10.28         7.60           SRV_04970_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_02132_at         CHORNME C, SOMATIC         1.30         2.80         2.64           SRV_02132_at         EUKARVOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, 1.00         3.73         5.99         3.73         5.99           SRV_0322_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_0322_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.32           SRV_05108_at         DPA-NAGE-SINDORANG SINDARY         1.84         2.18         2.59           SRV_05108_at         DPA-NACTEVLGULCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_005108_at         C_s.at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.12         2.2	SRV_02399_a_at		2.19	6.79	8.78
SRV_02516_at         PROTEIN, 78KDA)         2.70         10.25         7.80           SRV_04370_a_at         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04300_a_at         CASPASE RECRUITMENT DOMAIND FACTOR 1         1.30         2.80         2.64           SRV_02132_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.00         3.73         5.99           metabolic process         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_0308_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHORYLASE 1         1.84         2.18         2.59           SRV_0129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.24         3.64         3.39           SRV_02020_at         CARBOXYLESTERASE 1         1.12         1.97         2.48         3.86           SRV_03054_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.24         2.36         2.33					
-         -         PHOTEIN, 78/04/0           SRU_04970_a_att         CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6         1.35         4.78         4.80           SRV_04154_a_att         HEAT SHOCK 70KDA PROTEIN 98 (MORTALIN-2)         1.18         3.45         2.71           SRV_04300_a_att         CYTOCHROME C, SOMATIC         1.30         2.80         2.66           SRV_01312_a_att         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_01312_a_att         HEAT SHOCK PROTEIN 90KDA BETA (GRP94), MEMBER 1         1.31         3.53         3.26           GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_0323_a_att         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_0323_a_att         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.30         3.75         5.99           metabolic process         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_0322_a_att         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00222_a_att         DPN-ACTEVICQUOSAMINE PYREPOHOSPHORYLASE 1         1.84         2.18         2.59           SRV_00202_att         CAR	SRV 02516 at		2.70	10.28	7.60
SRV_08154_a_at         HEAT SHOCK 70KDA PROTEIN 9B (MORTALIN-2)         1.18         3.45         2.71           SRV_02132_at         CYTOCHROME C, SOMATIC         1.30         2.80         2.64           SRV_02132_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_05147_a_att         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00132_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_01737_a_at         UDP-N-ACTEYLGUCOSAMINE PYROPHOSPHATE         1.00         3.73         5.99           SRV_0108_at         DPHOSENTER ANINOTRANSFERASE 1         1.42         2.18         2.59           SRV_0129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.24         2.42         2.27           A_s.at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13 </td <td></td> <td></td> <td></td> <td></td> <td></td>					
SRV_04300_a_at         CYTOCHROME C, SOMATIC         1.30         2.80         2.64           SRV_02132_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.31         3.53         3.26           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.00         3.73         5.99           metabolic process         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.00         3.73         5.99           SRV_03032_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_01737_a, at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_00129_a, at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.12         1.97         2.48           SRV_00202_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE         1.28         3.86         2.38           SRV_003054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS         2.17         8.20         6.82           SRV_01480_at         THIOREDOXIN         1.19         3.29         3.51           SRV_0	SRV_04970_a_at	CASPASE RECRUITMENT DOMAIN FAMILY, MEMBER 6	1.35	4.78	4.80
SRV_04300_a_at         CYTOCHROME C, SOMATIC         1.30         2.80         2.64           SRV_02132_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1         1.21         3.56         3.38           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.31         3.53         3.26           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.00         3.73         5.99           metabolic process         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.00         3.73         5.99           SRV_03032_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_01737_a, at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_00129_a, at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.12         1.97         2.48           SRV_00202_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE         1.28         3.86         2.38           SRV_003054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS         2.17         8.20         6.82           SRV_01480_at         THIOREDOXIN         1.19         3.29         3.51           SRV_0	SRV 08154 a at	HEAT SHOCK 70KDA PROTEIN 9B (MORTALIN-2)	1.18	3.45	2.71
SRV_02132_at         EUKARYOTIC TRANSLATION ELONGATION FACTOR 1 EPSILON 1         1.21         3.56         3.38           SRV_01812_a_at         HEAT SHOCK PROTEIN 90KDA BETA (GRP94), MEMBER 1         1.31         3.53         3.26           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           metabolic process         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_05147_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00292_a         UDP-N-ACTEYLIGULCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.55           SRV_01737_a_at         UDP-N-ACTEYLIGULCOSAMINE PYROPHOSPHORYLASE 1         1.24         2.48         2.42           A_sat         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_00202_at         CARBOXYLESTERASE 1         1.61         7.61         7.01           SRV_01313_a_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_0186_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE<					
SRV_02132_at         EPSILON 1         1.21         3.56         3.38           SRV_01812_att         HEAT SHOCK PROTEIN 90KDA BETA (GRP94), MEMBER 1         1.31         3.53         3.26           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.31         3.53         3.26           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, I.00         3.73         5.99           metabolic process         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00322_a.t         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00292_a_att         PHOSPHOMANNOMUTASE 2         1.14         2.61         2.65           SRV_0108_a.at         DUP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_00129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.24         2.64         3.39           SRV_00202_at         CARBOXYLESTERASE 1 (MONCYTE/MACROPHAGE         1.28         3.86         2.38           SRV_01818_at         THIOREDOXIN         1.9         3.29         3.61         3.72           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE         1.59         3.67         2.77           SRV_0148	0111_04000_u_u		1.00	2.00	2.04
SRV_01812_a.at         HEAT SHOCK PROTEIN 90KDA BETA (GRP94), MEMBER 1         1.31         3.53         3.26           SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_03023_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA         1.39         7.34         8.89           SRV_05147_a_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, I.00         3.73         5.99           metabolic process         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_05147_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHONLASE 1         1.84         2.18         2.59           SRV_0108_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.12         1.97         2.48           SRV_00129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.24         3.64         3.39           SRV_00202_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE         1.28         3.86         2.38           SRV_01313_a_at         THIOREDOXIN         1.19         3.29         3.51           SRV_03064_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE         1.59         3.67         2.77	SRV 02132 at		1.21	3.56	3.38
SRV_03753_at         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, SRV_0332 a_at         1.00         3.73         5.99           metabolic process SRV_01737_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_01018_a_at         PHOSPHOSEAHSTE MAINOTRANSFERASE 1         1.12         1.97         2.48           SRV_00129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_00129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_02026Af_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_011818_at         THIOREDOXIN         1.19         3.29         3.51					
SRV_03023_a_att         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, GAMMA         1.00         3.73         5.99           metabolic process SRV_05147_a_att         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00322_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_00323_a_at         PHOSPHOMANNOMUTASE 2         1.14         2.61         2.65           SRV_05108_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOSERINE AMINOTRANSFERASE 1         1.12         1.97         2.48           C_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_0179_a_att         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_0202_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE SRV_01313_a_at         SERINE ESTERASE 1)         1.61         7.61         7.01           SRV_01313_a_at         THOREDOXIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF RAPPA LIGHT POLYPEPTIDE GENE SRV_01480_at         1.45         3.72         3.40           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.15	SRV_01812_a_at	HEAT SHOCK PROTEIN 90KDA BETA (GRP94), MEMBER 1	1.31	3.53	3.26
SRV_03023_a_att         GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, GAMMA         1.00         3.73         5.99           metabolic process SRV_05147_a_att         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00322_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_00323_a_at         PHOSPHOMANNOMUTASE 2         1.14         2.61         2.65           SRV_05108_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOSERINE AMINOTRANSFERASE 1         1.12         1.97         2.48           C_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_0179_a_att         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_0202_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE SRV_01313_a_at         SERINE ESTERASE 1)         1.61         7.61         7.01           SRV_01313_a_at         THOREDOXIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF RAPPA LIGHT POLYPEPTIDE GENE SRV_01480_at         1.45         3.72         3.40           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.15	SRV 03753 at	GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE, BETA	1.39	7.34	8.89
SRV_0323_a_at         GAMMA         1.00         3.73         5.99           metabolic process SRV_05147_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_0032_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_001737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_0129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.15         2.42         2.27           A_s.at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_00129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_00202_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE         1.28         3.86         2.38           signal transduction         SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE					
metabolic process SRV_05147_a_at SRV_00332_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00332_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_00292_a_at         PHOSPHOMANNOMUTASE 2         1.14         2.61         2.65           SRV_05108_a_at         DPI-N-ACTEYLGUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_00129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.24         2.46         3.39           SRV_00129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_00129_a_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE         1.28         3.86         2.38           SRV_0202_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE         1.28         3.86         2.38           SRV_01313_a_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01462_at         SRV_01313_a_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01482_at         GRANULIN         1.45         3.72         3.40           SRV_01294_aat         CONTAINING GTASE ACTIVATING PROTEIN 2         1.23         2.59         2.69	SRV_03023_a_at		1.00	3.73	5.99
SRV_05147_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00322_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOMANNOMUTANSE2 2         1.14         2.61         2.65           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOMANNOMUTANSEPRASE 1         1.12         1.97         2.48           C_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_00129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_02002_at         CARBOXYLESTERASE 1         (MONOCYTE/MACROPHAGE         2.17         8.20         6.82           SRV_01318_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01313_a_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE         1.59         3.67         2.77           SRV_02894_a_at					
SRV_05147_a_at         SYNDECAN BINDING PROTEIN (SYNTENIN)         1.27         4.78         4.42           SRV_00322_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOMANNOMUTANSE2 2         1.14         2.61         2.65           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOMANNOMUTANSEPRASE 1         1.12         1.97         2.48           C_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_00129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_02002_at         CARBOXYLESTERASE 1         (MONOCYTE/MACROPHAGE         2.17         8.20         6.82           SRV_01318_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01313_a_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE         1.59         3.67         2.77           SRV_02894_a_at					
SRV_00332_a_at         glucose-6-phosphate dehydrogenase         1.32         4.55         3.35           SRV_00292_a_at         UDP-N-ACTEVLGUCOSAMINE PYROPHOSPHORYLASE 1         1.14         2.61         2.65           SRV_01737_a_at         UDP-N-ACTEVLGUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOSERINE AMINOTRANSFERASE 1         1.12         1.97         2.48           C_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.24         3.64         3.39           SRV_00129_a_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.28         3.86         2.30           SRV_00202_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE SERINE ESTERASE 1)         1.61         7.61         7.01           signal transduction SRV_02657_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_011313_a_at         GRANULIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE SRV_01480_at         1.9         3.67         2.77           SRV_002894_a_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 1         1.10					
SRV_00292_a_at         PHOSPHOMANNOMUTASE 2         1.14         2.61         2.65           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOSERINE AMINOTRANSFERASE 1         1.12         1.97         2.48           C_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.24         3.64         3.39           SRV_00129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_02002_at         CARBOXYLESTERASE 1         1.01         7.61         7.01           signal transduction         SRV_01313_a_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01313_a_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01462_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01313_a_at         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE         1.59         3.67         2.77           SRV_02894_a_at         CAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1         1.02         2.147         3.17         2.44		SYNDECAN BINDING PROTEIN (SYNTENIN)	1.27	4.78	4.42
SRV_00292_a_at         PHOSPHOMANNOMUTASE 2         1.14         2.61         2.65           SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOSERINE AMINOTRANSFERASE 1         1.12         1.97         2.48           C_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.24         3.64         3.39           SRV_00129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_02002_at         CARBOXYLESTERASE 1         1.01         7.61         7.01           signal transduction         SRV_01313_a_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01313_a_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01462_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01313_a_at         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE         1.59         3.67         2.77           SRV_02894_a_at         CAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1         1.02         2.147         3.17         2.44	SRV 00332 a at	glucose-6-phosphate dehydrogenase	1.32	4.55	3.35
SRV_01737_a_at         UDP-N-ACTEYLGLUCOSAMINE PYROPHOSPHORYLASE 1         1.84         2.18         2.59           SRV_05108_a_at         PHOSPHOSERINE AMINOTRANSFERASE 1         1.12         1.97         2.48           C_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.15         2.42         2.27           A_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_00129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_02002_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE         1.28         3.86         2.38           signal transduction         SERINE ESTERASE 1)         1.61         7.61         7.01           SRV_03054_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01313_a_at         CRANULIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE         1.59         3.67         2.77           SRV_01480_at         IO MOTIF CONTAINING GTASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         IO MOTIF CONTAINING GTASE ACTIVATING PROTEIN 1         1.00         2.09         1.76           SRV_02894_a					
SRV_05108_a_at         PHOSPHOSERINE AMINOTRANSFERASE 1         1.12         1.97         2.48           C_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.15         2.42         2.27           A_s_at         ALDOLASE A, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_0129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_02002_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE SERINE ESTERASE 1)         1.28         3.86         2.38           signal transduction SRV_02657_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01313_a_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01313_a_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_01313_a_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.59         3.67         2.77           SRV_02894_a_at         CHORTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.10         2.09					
C_satALDOLASE A, FRUCTOSE-BISPHOSPHATE1.152.422.27A_satALDOLASE A, FRUCTOSE-BISPHOSPHATE1.243.643.39SRV_00129_a_atALDOLASE B, FRUCTOSE-BISPHOSPHATE1.262.302.13SRV_02002_atCARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE SERINE ESTERASE 1)1.283.862.38signal transduction SRV_03054_atPRE-B-CELL COLONY ENHANCING FACTOR 11.617.617.01SRV_03054_atMACROPHAGE RECEPTOR WITH COLLAGENOUS SRV_01313_a_at2.178.206.82SRV_01313_a_atTHIOREDOXIN1.193.293.51SRV_01462_atNUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)1.593.672.77SRV_01480_atPURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 21.473.172.44SRV_02894_a_atCAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1 (YEAST)1.102.091.76SRV_03227_at SRV_01179_a_atNIDOGEN 2 (OSTEONIDOGEN)1.423.694.29SRV_02687_s_atUDP-GLCNAC:BETAGAL BETA-1,3-N-1.4443.50					
A_s_atALDOLASE A, FRUCTOSE-BISPHOSPHATE1.243.643.39SRV_00129_a_atALDOLASE B, FRUCTOSE-BISPHOSPHATE1.262.302.13SRV_02002_atCARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE SERINE ESTERASE 1)1.262.302.13signal transduction SRV_02657_atPRE-B-CELL COLONY ENHANCING FACTOR 11.617.617.01SRV_03054_atPRE-B-CELL COLONY ENHANCING FACTOR 11.617.617.01SRV_03054_atMACROPHAGE RECEPTOR WITH COLLAGENOUS SRV_01313_a_at2.178.206.82SRV_01818_at SRV_01462_atTHIOREDOXIN1.193.293.51SRV_00306_a_at SRV_01480_atNUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)1.593.672.77SRV_03206_a_at SRV_01480_atIQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 21.232.592.69SRV_02894_a_at SRV_03227_at SRV_01179_a_atOHLORIDE INTRACELLULAR CHANNEL 11.151.952.10membrane SRV_02687_s_atNIDOGEN 2 (OSTEONIDOGEN) MANNOSE-6-PHOSPHATE RECPTOR BINDING PROTEIN 11.304.823.55SRV_04888_a_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.414.443.50					
SRV_00129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_02002_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE SERINE ESTERASE 1)         1.28         3.86         2.38           signal transduction SRV_02657_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_03054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS SRV_01818_at         2.17         8.20         6.82           SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.45         3.72         3.40           SRV_01462_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.47         3.17         2.44           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.10         2.09         1.76           SRV_03227_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_02687_s_at         NIDOGEN 2 (OSTEONIDOGEN)         1.34         3.82         3.97           SRV_02687_s_at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41	C_s_at	ALDOLASE A, FRUCTOSE-BISPHOSPHATE	1.15	2.42	2.27
SRV_00129_a_at         ALDOLASE B, FRUCTOSE-BISPHOSPHATE         1.26         2.30         2.13           SRV_02002_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE SERINE ESTERASE 1)         1.28         3.86         2.38           signal transduction SRV_02657_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_03054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS SRV_01818_at         2.17         8.20         6.82           SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.45         3.72         3.40           SRV_01462_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.47         3.17         2.44           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.10         2.09         1.76           SRV_03227_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_02687_s_at         NIDOGEN 2 (OSTEONIDOGEN)         1.34         3.82         3.97           SRV_02687_s_at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41	A s at	ALDOLASE A. FRUCTOSE-BISPHOSPHATE	1.24	3.64	3.39
SRV_02002_at         CARBOXYLESTERASE 1 (MONOCYTE/MACROPHAGE SERINE ESTERASE 1)         1.28         3.86         2.38           signal transduction SRV_02657_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_03054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS STRUCTURE         2.17         8.20         6.82           SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01313_a_at         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.59         3.67         2.77           SRV_03006_a_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2         1.47         3.17         2.44           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.10         2.09         1.76           SRV_03227_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_02687_s_at         NIDOGEN 2 (OSTEONIDOGEN)         1.34         3.82         3.97           SRV_02687_s_at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44 <t< td=""><td></td><td></td><td></td><td></td><td></td></t<>					
SRV_02002_at         SERINE ESTERASE 1)         1.28         3.86         2.38           signal transduction SRV_02657_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_03054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS SRV_01818_at         2.17         8.20         6.82           SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.59         3.67         2.77           SRV_03006_a_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2         1.47         3.17         2.44           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.10         2.09         1.76           SRV_03227_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_01179_a_at         NIDOGEN 2 (OSTEONIDOGEN)         1.34         3.82         3.97           SRV_02687_s_at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44         3.50	0111_00120_4_4		1.20	2.00	2.10
Signal transduction SRV_02657_atPRE-B-CELL COLONY ENHANCING FACTOR 11.617.617.01SRV_03054_atMACROPHAGE RECEPTOR WITH COLLAGENOUS SRV_01818_at2.178.206.82SRV_01818_atSTRUCTURE2.178.206.82SRV_01313_a_atGRANULIN1.453.723.40SRV_01462_atNUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)1.593.672.77SRV_03006_a_atIQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 21.232.592.69SRV_04480_atPURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 21.473.172.44SRV_02894_a_atCAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 11.102.091.76SRV_03227_atCHLORIDE INTRACELLULAR CHANNEL 11.151.952.10membrane SRV_01179_a_atNIDOGEN 2 (OSTEONIDOGEN) CD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.423.694.29SRV_02887_s_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.414.443.50	SRV 02002 at		1.28	3.86	2.38
SRV_02657_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_03054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS STRUCTURE         2.17         8.20         6.82           SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01313_a_at         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.59         3.67         2.77           SRV_03006_a_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2         1.47         3.17         2.44           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.10         2.09         1.76           SRV_03227_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_01179_a_at         NIDOGEN 2 (OSTEONIDOGEN)         1.34         3.82         3.97           MANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN         1.30         4.82         3.55           SRV_04888_a at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44         3.50		SERINE ESTERASE 1)			
SRV_02657_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_03054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS STRUCTURE         2.17         8.20         6.82           SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01313_a_at         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.59         3.67         2.77           SRV_03006_a_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2         1.47         3.17         2.44           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.10         2.09         1.76           SRV_03227_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_01179_a_at         NIDOGEN 2 (OSTEONIDOGEN)         1.34         3.82         3.97           MANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN         1.30         4.82         3.55           SRV_04888_a at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44         3.50					
SRV_02657_at         PRE-B-CELL COLONY ENHANCING FACTOR 1         1.61         7.61         7.01           SRV_03054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS STRUCTURE         2.17         8.20         6.82           SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01313_a_at         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.59         3.67         2.77           SRV_03006_a_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2         1.47         3.17         2.44           SRV_02894_a_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.10         2.09         1.76           SRV_03227_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_01179_a_at         NIDOGEN 2 (OSTEONIDOGEN)         1.34         3.82         3.97           MANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN         1.30         4.82         3.55           SRV_04888_a at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44         3.50	signal transduction				
SRV_03054_at         MACROPHAGE RECEPTOR WITH COLLAGENOUS STRUCTURE         2.17         8.20         6.82           SRV_03054_at         STRUCTURE         2.17         8.20         6.82           SRV_01313_a_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01313_a_at         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.59         3.67         2.77           SRV_01480_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2         1.47         3.17         2.44           SRV_02894_a_at         CAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1 (YEAST)         1.10         2.09         1.76           SRV_03227_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.15         1.95         2.10           membrane         SRV_01179_a_at         NIDOGEN 2 (OSTEONIDOGEN) CD63 ANTIGEN (MELANOMA 1 ANTIGEN)         1.34         3.82         3.97           SRV_02687_s_at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44         3.50	SRV 02657 at	PRE-B-CELL COLONY ENHANCING FACTOR 1	1.61	7.61	7.01
SRV_03054_at         STRUCTURE         2.17         8.20         6.82           SRV_01818_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01313_a_at         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.59         3.67         2.77           SRV_03006_a_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2         1.23         2.59         2.69           SRV_01480_at         PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2         1.47         3.17         2.44           SRV_02894_a_at         CAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1         1.10         2.09         1.76           SRV_030227_at         CHLORIDE INTRACELLULAR CHANNEL 1         1.15         1.95         2.10           membrane         SRV_01179_a_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_02687_s_at         NIDOGEN 2 (OSTEONIDOGEN)         1.34         3.82         3.97           SRV_02687_s_at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44         3.50					
SRV_01818_at SRV_01313_a_at         THIOREDOXIN         1.19         3.29         3.51           SRV_01313_a_at         GRANULIN         1.45         3.72         3.40           SRV_01462_at         NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)         1.59         3.67         2.77           SRV_03006_a_at SRV_01480_at         IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2 PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2         1.47         3.17         2.44           SRV_02894_a_at         CAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1 (YEAST)         1.10         2.09         1.76           SRV_03227_at SRV_01179_a_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_02687_s_at         NIDOGEN 2 (OSTEONIDOGEN)         1.34         3.82         3.97           SRV_04888 a_at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44         3.50	SRV_03054_at		2.17	8.20	6.82
SRV_01313_a_atGRANULIN1.453.723.40SRV_01462_atNUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)1.593.672.77SRV_03006_a_at SRV_01480_atIQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2 PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 21.473.172.44SRV_02894_a_at SRV_00844_a_atCAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1 (YEAST)1.102.091.76SRV_03227_at SRV_01179_a_atNIDOGEN 2 (OSTEONIDOGEN) CD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.423.694.29SRV_02687_s_atUDP-GLCNAC:BETAGAL BETA-1,3-N-1.304.823.55				0.00	0.54
SRV_01462_atNUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE ENHANCER IN B-CELLS 2 (P49/P100)1.593.672.77SRV_03006_a_at SRV_01480_atIQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2 PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2 CAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1 (YEAST)1.232.592.69SRV_00844_a_at SRV_00844_a_atCHLORIDE INTRACELLULAR CHANNEL 11.102.091.76membrane SRV_03227_at SRV_01179_a_atNIDOGEN 2 (OSTEONIDOGEN) CD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.423.694.29SRV_02687_s_atUDP-GLCNAC:BETAGAL BETA-1,3-N-1.304.823.55					
SRV_01462_at       ENHANCER IN B-CELLS 2 (P49/P100)       1.59       3.67       2.77         SRV_03006_a_at       IQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2       1.23       2.59       2.69         SRV_01480_at       PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2       1.47       3.17       2.44         SRV_02894_a_at       CAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1       1.10       2.09       1.76         SRV_00844_a_at       CHLORIDE INTRACELLULAR CHANNEL 1       1.15       1.95       2.10         membrane       SRV_03227_at       NIDOGEN 2 (OSTEONIDOGEN)       1.42       3.69       4.29         SRV_01179_a_at       CD63 ANTIGEN (MELANOMA 1 ANTIGEN)       1.34       3.82       3.97         SRV_02687_s_at       UDP-GLCNAC:BETAGAL BETA-1,3-N-       2.41       4.44       3.50	SRV_01313_a_at	GRANULIN	1.45	3.72	3.40
SRV_03006_a_at SRV_01480_atIQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2 PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2 PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2 (YEAST)1.23 1.472.59 3.172.69 2.44SRV_02894_a_at SRV_00844_a_atIQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN COUPLED, 2 (YEAST)1.47 1.103.17 2.44SRV_00844_a_at SRV_03227_at SRV_01179_a_at SRV_02687_s_atIQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 1 (YEAST)1.10 2.092.09 1.76MANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN 11.34 3.82 3.973.82 3.97SRV_04888 a_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.41 4.444.44		NUCLEAR FACTOR OF KAPPA LIGHT POLYPEPTIDE GENE	4 50	0.07	0 77
SRV_03006_a_at SRV_01480_atIQ MOTIF CONTAINING GTPASE ACTIVATING PROTEIN 2 PURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 2 CAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 1 (YEAST)1.23 1.472.59 3.172.69 2.44SRV_02894_a_at SRV_00844_a_atCHLORIDE INTRACELLULAR CHANNEL 11.102.091.76membrane SRV_03227_at SRV_01179_a_atNIDOGEN 2 (OSTEONIDOGEN) CD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.423.694.29SRV_02687_s_atNIDOGEN 2 (OSTEONIDOGEN) CD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.343.823.97SRV_02687_s_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.414.443.50	SRV_01462_at	ENHANCER IN B-CELLS 2 (P49/P100)	1.59	3.67	2.77
SRV_01480_atPURINERGIC RECEPTOR P2Y, G-PROTEIN COUPLED, 21.473.172.44SRV_02894_a_atCAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 11.102.091.76SRV_00844_a_atCHLORIDE INTRACELLULAR CHANNEL 11.151.952.10membraneSRV_03227_atNIDOGEN 2 (OSTEONIDOGEN)1.423.694.29SRV_01179_a_atCD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.343.823.97SRV_02687_s_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.414.443.50	SBV 02006 a at		1 22	2 50	2 60
CAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 11.102.091.76SRV_02894_a_atCAP, ADENYLATE CYCLASE-ASSOCIATED PROTEIN 11.102.091.76SRV_00844_a_atCHLORIDE INTRACELLULAR CHANNEL 11.151.952.10membraneSRV_03227_atNIDOGEN 2 (OSTEONIDOGEN)1.423.694.29SRV_01179_a_atCD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.343.823.97SRV_02687_s_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.414.443.50					
SRV_02894_a_at       (YEAST)       1.10       2.09       1.76         SRV_00844_a_at       CHLORIDE INTRACELLULAR CHANNEL 1       1.15       1.95       2.10         membrane       SRV_03227_at       NIDOGEN 2 (OSTEONIDOGEN)       1.42       3.69       4.29         SRV_01179_a_at       CD63 ANTIGEN (MELANOMA 1 ANTIGEN)       1.34       3.82       3.97         SRV_02687_s_at       MANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN       1.30       4.82       3.55         SRV_04888_a_at       UDP-GLCNAC:BETAGAL BETA-1,3-N-       2.41       4.44       3.50	SRV_01480_at		1.47	3.17	2.44
SRV_00844_a_atCHLORIDE INTRACELLULAR CHANNEL 11.151.952.10membrane SRV_03227_at SRV_01179_a_at SRV_02687_s_atNIDOGEN 2 (OSTEONIDOGEN)1.423.694.29SRV_02687_s_atCD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.343.823.97SRV_02687_s_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.414.443.50	SRV 02894 a at		1 10	2 09	1 76
membraneSRV_03227_atNIDOGEN 2 (OSTEONIDOGEN)1.423.694.29SRV_01179_a_atCD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.343.823.97SRV_02687_s_atMANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN1.304.823.55SRV_04888_a_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.414.443.50	0110_02004_a_at	(YEAST)	1.10	2.00	1.70
membraneSRV_03227_atNIDOGEN 2 (OSTEONIDOGEN)1.423.694.29SRV_01179_a_atCD63 ANTIGEN (MELANOMA 1 ANTIGEN)1.343.823.97SRV_02687_s_atMANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN1.304.823.55SRV_04888_a_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.414.443.50	SBV 00844 a at	CHLOBIDE INTRACELLULAB CHANNEL 1	1.15	1.95	2.10
SRV_03227_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_01179_a_at         CD63 ANTIGEN (MELANOMA 1 ANTIGEN)         1.34         3.82         3.97           SRV_02687_s_at         MANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN         1.30         4.82         3.55           SRV_04888_a_at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44         3.50	0111_00011_a_a				
SRV_03227_at         NIDOGEN 2 (OSTEONIDOGEN)         1.42         3.69         4.29           SRV_01179_a_at         CD63 ANTIGEN (MELANOMA 1 ANTIGEN)         1.34         3.82         3.97           SRV_02687_s_at         MANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN         1.30         4.82         3.55           SRV_04888_a_at         UDP-GLCNAC:BETAGAL BETA-1,3-N-         2.41         4.44         3.50	mombrana				
SRV_01179_a_atCD63 ANTIGEN (MELANOMA 1 ANTÍGEN)1.343.823.97SRV_02687_s_atMANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN1.304.823.55SRV_04888_a_atUDP-GLCNAC:BETAGAL BETA-1,3-N-2.414.443.50			4.40	0.00	4.00
SRV_04888_3_st UDP-GLCNAC:BETAGAL BETA-1,3-N- 2.41 4.44 3.50					
SRV_02687_s_at 1 1.30 4.82 3.55 SRV_04888_a_at UDP-GLCNAC:BETAGAL BETA-1,3-N- 2.41 4.44 3.50	SRV_01179_a_at		1.34	3.82	3.97
SRV_02687_s_at 1 1.30 4.82 3.55 SRV_04888_a_at UDP-GLCNAC:BETAGAL BETA-1,3-N- 2.41 4.44 3.50		MANNOSE-6-PHOSPHATE RECEPTOR BINDING PROTEIN	1 00	4.00	0 55
SBV 04888 a at UDP-GLCNAC:BETAGAL BETA-1,3-N-	SHV_02687_s_at		1.30	4.82	3.55
	SRV_04888 a at		2.41	4.44	3.50
		AUE I I LULUUUJAIVIIN I LI KANSFEKASE 3			

SRV 04819 a at	TRANSMEMBRANE PROTEIN 49	1.46	3.40	3.20
SRV_04070_a_at	LEUCINE RICH REPEAT CONTAINING 59	1.22	2.48	2.78
SRV 03407 at	FER-1-LIKE 3, MYOFERLIN (C. ELEGANS)	1.22	2.67	3.81
SRV 05439 at	HYPOTHETICAL PROTEIN LOC441168	1.40	3.43	2.79
3HV_03439_al		1.40	5.45	2.19
SRV 02874 a at	BRAIN ABUNDANT, MEMBRANE ATTACHED SIGNAL	1.33	2.80	2.31
	PROTEIN 1			
SRV_04226_a_at	GTPASE, IMAP FAMILY MEMBER 5	1.13	2.05	2.02
transport				
SRV 00744 a at	ADIPOSE DIFFERENTIATION-RELATED PROTEIN	3.49	12.49	7.34
ent_eent_a_a	PROTECTIVE PROTEIN FOR BETA-GALACTOSIDASE	0.10	12:10	7.01
SRV 00294 s at		1.53	5.23	4.73
	(GALACTOSIALIDOSIS)			
SRV_02592_a_at	LYSYL OXIDASE-LIKE 1	1.18	2.81	4.59
SRV_03991_a_at	SOLUTE CARRIER FAMILY 15, MEMBER 3	1.36	5.33	3.88
SRV 03562 at	ERO1-LIKE (S. CEREVISIAE)	1.00	2.24	1.77
	SOLUTE CARRIER FAMILY 7 (CATIONIC AMINO ACID			
SRV_04996_at	TRANSPORTER, Y+ SYSTEM), MEMBER 3	1.07	1.75	2.11
SRV 01134 at	ATPASE, H+ TRANSPORTING, LYSOSOMAL 56/58KDA, V1	1.04	2.17	1.59
	SUBUNIT B2			
SBV 01220 at	CYTOCHROME C OXIDASE SUBUNIT VIB POLYPEPTIDE 1	1.15	4.51	4.18
SRV_01220_at	(UBIQUITOUS)	1.15	4.51	4.10
SRV_02133_a_at	GOLGI SNAP RECEPTOR COMPLEX MEMBER 2	1.05	2.24	1.88
0111_02100_u_u		1.00		1.00
and the second second second second				
extracellular region	<i></i>			
SRV_02948_a_at	fibulin 1	0.93	1.65	2.24
SRV 01275 at	fibulin 1	0.81	1.54	2.10
SRV_02965_at	TISSUE FACTOR PATHWAY INHIBITOR 2	3.56	3.04	2.56
SRV 03142 at	ELASTIN MICROFIBRIL INTERFACER 1	1.11	3.37	3.73
0110_00142_at		1.11	0.07	0.70
ion binding				
SRV 00713 a at	TRANSCOBALAMIN I (VITAMIN B12 BINDING PROTEIN, R	2.55	23.02	12.98
011v_00715_a_at	BINDER FAMILY)	2.55	20.02	12.30
SRV 02456 at	GASTRIC INTRINSIC FACTOR (VITAMIN B SYNTHESIS)	3.36	20.67	11.63
SRV 07722 at	CHROMOSOME 17 OPEN READING FRAME 27	0.94	4.46	4.63
SRV_05460_at	tripartite motif-containing 39			4.07
		1.58	3.81	
SRV_05065_at	tripartite motif-containing 17	1.53	2.86	3.46
SRV_00371_a_at	FRUCTOSE-1,6-BISPHOSPHATASE 1	2.40	5.01	3.14
SRV 00741 a at	ADENOSINE DEAMINASE, RNA-SPECIFIC	1.44	3.76	2.86
	PROSTAGLANDIN-ENDOPEROXIDE SYNTHASE 1			
SRV 00562 a at	(PROSTAGLANDIN G/H SYNTHASE AND	1.29	2.05	2.54
011v_00002_a_at		1.23	2.05	2.54
	CYCLOOXYGENASE)			
SRV_05448_at	ATPase, Ca++ transporting, cardiac muscle, slow twitch 2	0.87	2.16	2.13
SRV_02724_at	METHIONINE ADENOSYLTRANSFERASE II, ALPHA	1.66	2.06	1.56
	LATENT TRANSFORMING GROWTH FACTOR BETA	4.05	4 70	0.45
SRV_00449_a_at	BINDING PROTEIN 1	1.05	1.79	3.15
	SECRETED PROTEIN, ACIDIC, CYSTEINE-RICH			
SRV 00131 a at		1.19	1.99	2.74
	(OSTEONECTIN)			
SRV_04896_a_at	TRANSKETOLASE (WERNICKE-KORSAKOFF SYNDROME)	1.44	4.02	2.52
SRV_02047_at	ANNEXIN A2	1.29	2.15	2.45
SRV 01345 at	INOSITOL 1,4,5-TRIPHOSPHATE RECEPTOR, TYPE 3	1.19	2.80	2.35
SRV_12418_at	MYOSIN, LIGHT POLYPEPTIDE 9, REGULATORY	1.01	1.99	2.20
SRV_01649_at	RETICULOCALBIN 1, EF-HAND CALCIUM BINDING DOMAIN	1.16	3.01	2.68
511V_01049_at		1.10	5.01	2.00
protein binding/transport				
SRV_03206_at	V-SET AND IMMUNOGLOBULIN DOMAIN CONTAINING 4	1.90	13.14	9.17
	SERPIN PEPTIDASE INHIBITOR, CLADE H (HEAT SHOCK			
SRV 00797 a at	PROTEIN 47), MEMBER 1, (COLLAGEN BINDING PROTEIN	1.77	4.67	4.79
0110_00707_a_at		1.77	4.07	4.75
		1 0 0	0.50	0.55
SRV_04964_a_at	TUBULIN, BETA 6	1.06	2.53	2.55
SRV_03477_a_at	TRANSMEMBRANE PROTEIN 4	1.59	2.22	2.39
SRV_02085_at	FK506 BINDING PROTEIN 1B, 12.6 KDA	1.29	1.71	2.18
SRV 02652 a at	PROTEIN DISULFIDE ISOMERASE FAMILY A, MEMBER 6	1.28	2.16	1.99
SRV 02814 at	DNAJ (HSP40) HOMOLOG, SUBFAMILY B, MEMBER 1	2.35	4.29	1.73
011v_02014_at		2.00	7.23	1.75
SRV 03255 a at	AHA1, ACTIVATOR OF HEAT SHOCK 90KDA PROTEIN	1.07	2.14	1.53
	ATPASE HOMOLOG 1 (YEAST)			
SRV_05534_at	HYPOTHETICAL PROTEIN MGC16212	1.74	4.05	3.73
SRV 00840 a at	COLD INDUCIBLE RNA BINDING PROTEIN	1.03	2.46	3.20
SRV_01147_a_at	BIGLYCAN	0.99	2.01	2.90
SRV 05461 x at	tripartite motif-containing 39	1.32	2.53	2.30
SRV_02070_a_at	CYSTEINE AND GLYCINE-RICH PROTEIN 1	1.27	2.22	2.46

SRV_00257_at		1.12	1.93	2.44
SRV_04005_at	KDEL (LYS-ASP-GLU-LEU) ENDOPLASMIC RETICULUM PROTEIN RETENTION RECEPTOR 3	0.90	2.07	2.06
SRV_02647_a_at	ARP2 ACTIN-RELATED PROTEIN 2 HOMOLOG (YEAST)	1.23	2.33	1.93
SRV_01234_a_at	CATENIN (CADHERIN-ASSOCIATED PROTEIN), ALPHA 1, 102KDA	0.98	1.43	2.05
SRV_01504_a_at	PHOSPHOGLUCONATE DEHYDROGENASE	1.19	2.20	1.50
SRV_05174_a_at	THREE PRIME REPAIR EXONUCLEASE 2	1.57	5.55	5.02
protein modification				
SRV 04305 a at	GLYCINE N-METHYLTRANSFERASE	1.24	1.84	3.49
SRV_01832_a_at	UBIQUITIN-CONJUGATING ENZYME E2L 3	1.04	2.09	1.99
	TRANSGLUTAMINASE 1 (K POLYPEPTIDE EPIDERMAL			
SRV_00309_at	TYPE I, PROTEIN-GLUTAMINE-GAMMA-	3.22	16.60	14.36
	GLUTAMYLTRANSFERASE)			
SRV_02093_a_at	HEAT SHOCK 70KDA PROTEIN 9B (MORTALIN-2)	1.35	3.91	2.79
SRV_02989_at	HEAT SHOCK 70KDA PROTEIN 8	2.28	6.93	2.73
SRV_01225_at	CRYSTALLIN, ALPHA B	1.27	3.35	2.38
SRV_05456_a_at	UBIQUITIN SPECIFIC PEPTIDASE 2	1.27	2.56	2.00
SRV_05457_a_at	UBIQUITIN SPECIFIC PEPTIDASE 2	1.26	2.28	1.89
 SRV_11417_a_at	MATRIX METALLOPEPTIDASE 1 (INTERSTITIAL COLLAGENASE)	5.26	13.96	3.65
SRV_04306_at	GLYCINE N-METHYLTRANSFERASE	1.03	2.15	4.66
SRV_00327_a_at	CATHEPSIN K (PYCNODYSOSTOSIS)	1.37	3.23	3.94
cellular process		1.00	04.74	00.01
SRV_11663_a_at	TIMP METALLOPEPTIDASE INHIBITOR 1	1.96	24.74	23.31
SRV_04387_at	RETINOIC ACID RECEPTOR RESPONDER (TAZAROTENE INDUCED) 1	1.64	6.63	9.22
SRV_03285_at	RELATED RAS VIRAL (R-RAS) ONCOGENE HOMOLOG 2	1.76	5.07	4.97
SRV_04911_at	SIMILAR TO THIOREDOXIN DOMAIN-CONTAINING 2 PROTEIN PHOSPHATASE 1, CATALYTIC SUBUNIT, ALPHA	1.34	3.15	3.20
SRV_01534_at	ISOFORM	1.15	2.04	2.18
SRV_04858_at	POLY (ADP-RIBOSE) POLYMERASE FAMILY, MEMBER 9	0.97	2.36	2.15
SRV_03421_a_at	LR8 PROTEIN	1.31	1.95	2.02
	V-YES-1 YAMAGUCHI SARCOMA VIRAL RELATED	1.37	2.46	2.12
SRV_11406_at	ONCOGENE HOMOLOG	1.37	2.40	2.12
other				
other SRV_03887_at	DYNEIN, CYTOPLASMIC 1, LIGHT INTERMEDIATE CHAIN 1	1.31	2.42	2.20
SRV_03007_at	poly (ADP-ribose) glycohydrolase	1.31	2.42	2.20
SRV_01920_at SRV 00155 a at	COAGULATION FACTOR XIII, A1 POLYPEPTIDE	1.31	2.46	2.23
SRV_00155_a_at	KERATIN 8	1.20	2.46 2.90	2.35
		1.19	2.90 4.08	
SRV_00775_at SRV 11767 a at	ARGINASE, TYPE II INTERFERON INDUCED TRANSMEMBRANE PROTEIN 5	1.44	4.08 2.74	6.06 4.24
	MACROPHAGE EXPRESSED GENE 1			
SRV_07726_a_at		1.69	4.25	3.18
SRV_01302_at		1.07	2.39	2.64
SRV_02761_at	ARGININE-RICH, MUTATED IN EARLY STAGE TUMORS	1.10	2.17	1.99
SRV_03758_a_at	ARRESTIN DOMAIN CONTAINING 2	0.90	5.48	8.72

### **Table 2**. Genes which were significantly down-regulated at any time point. Numbers

indicate fold change at that time point.

transport

ID	NAME	21 hr	70 hr	111 br
transcription	INAIVIE	24 hr	72 hr	144 hr
SRV 04230 at	CHROMOSOME X OPEN READING FRAME 15	0.73	0.44	0.50
SRV 01344 a at	INTERFERON REGULATORY FACTOR 2	1.02	0.44	0.50
	TAF9 RNA POLYMERASE II, TATA BOX BINDING			
SRV_01768_a_at	PROTEIN (TBP)-ASSOCIATED FACTOR, 32KDA	0.84	0.44	0.50
SRV_03843_a_at	MEDIATOR OF RNA POLYMERASE II TRANSCRIPTION, SUBUNIT 31 HOMOLOG (YEAST)	0.95	0.50	0.45
SRV_01892_at	ZINC FINGER PROTEIN 282	0.98	0.39	0.42
translation				
SRV 03800 a at	MITOCHONDRIAL RIBOSOMAL PROTEIN S7	1.13	0.50	0.54
SRV 03598 at	MITOCHONDRIAL RIBOSOMAL PROTEIN L19	0.73	0.49	0.53
SRV 04607 at	PEPTIDE DEFORMYLASE-LIKE PROTEIN	1.01	0.46	0.50
SRV 04925 a at	HYPOTHETICAL PROTEIN MGC11102	1.01	0.43	0.48
	EUKARYOTIC TRANSLATION INITIATION FACTOR 4E			
SRV_01958_at	BINDING PROTEIN 3	1.05	0.32	0.31
Natural Killer cell mediated				
cytotoxicity				
AE at	TUBULIN, BETA 2C	0.88	0.52	0.49
, <u>,</u> ,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	TODOLIN, DETAZO	0.00	0.02	0.40
apoptosis				
SRV 11815 at	CASP2 AND RIPK1 DOMAIN CONTAINING ADAPTOR	0.70	0.45	0.52
	WITH DEATH DOMAIN			
SRV_01489_at	PRKC, APOPTOSIS, WT1, REGULATOR	0.88	0.43	0.44
ion binding/ transport				
SRV_03020_at	TRAF-TYPE ZINC FINGER DOMAIN CONTAINING 1	0.85	0.49	0.53
 SRV 01742 at	SPECTRIN, ALPHA, NON-ERYTHROCYTIC 1 (ALPHA-	0.82	0.46	0.52
	FODRIN)			
SRV_02131_a_at	PEPTIDASE (MITOCHONDRIAL PROCESSING) BETA	0.80	0.48	0.51
SRV_02733_at	MITOCHONDRIAL INTERMEDIATE PEPTIDASE	0.59	0.39	0.40
SRV_04112_at	HYPOTHETICAL PROTEIN FLJ20699	0.93	0.42	0.39
SRV_00559_a_at	PRIMASE, POLYPEPTIDE 1, 49KDA	0.79	0.42	0.39
SRV_03126_at	RING FINGER PROTEIN 113A	0.95	0.39	0.39
SRV 03759 at	ATP SYNTHASE, H+ TRANSPORTING, MITOCHONDRIAL	0.91	0.30	0.25
0.11_00700_at	F0 COMPLEX, SUBUNIT S (FACTOR B)	0.01	0.00	0.20
SRV_01177_a_at		1.19	0.52	0.49
	DIPHOSPHOHYDROLASE 1			
SRV_04638_a_at	MEMBRANE-ASSOCIATED RING FINGER (C3HC4) 7	0.78	0.44	0.47
SRV_03403_at	MAKORIN, RING FINGER PROTEIN, 1	1.02	0.55	0.48
SRV_02137_at	GUANINE DEAMINASE	0.93	0.36	0.36
SRV_12156_at	TUMOR PROTEIN D52 CHONDROITIN SULFATE PROTEOGLYCAN 3	0.93	0.58	0.45
SRV_02173_at	(NEUROCAN)	1.38	0.57	0.20
motobolio proceso				
metabolic process SRV 02346 a at	GLUTATHIONE S-TRANSFERASE OMEGA 1	1.04	0 54	0 47
SRV_02346_a_at SRV 04215 at	PHOSPHOGLUCOMUTASE 2		0.54	0.47
SRV_04215_at SRV_04397_at		1.08 0.86	0.57	0.46 0.46
	NITRILASE FAMILY, MEMBER 2 NON-METASTATIC CELLS 7, PROTEIN EXPRESSED IN	0.00	0.49	
SRV_03369_at	(NUCLEOSIDE-DIPHOSPHATE KINASE)	1.00	0.55	0.43
SRV_00123_at	AMINOLEVULINATE, DELTA-, DEHYDRATASE	0.71	0.49	0.31
SRV 00160 s at	FUMARYLACETOACETATE HYDROLASE	0.83	0.61	0.49
SRV_00135_at	ASPARTOACYLASE (CANAVAN DISEASE) 6-PHOSPHOFRUCTO-2-KINASE/FRUCTOSE-2,6-	0.92	0.30	0.22
SRV_01499_at	6-PHOSPHOFROCIO-2-KINASE/FROCIOSE-2,6- BIPHOSPHATASE 1	0.88	0.65	0.47
SRV_11745_at	ACYL-COENZYME A OXIDASE 3, PRISTANOYL	0.87	0.51	0.47
SRV_03094_at	LIPOIC ACID SYNTHETASE	0.83	0.47	0.47
SRV_05217_a_at	SERINE DEHYDRATASE-LIKE	1.03	0.52	0.33

SRV_03906_at	HEMATOPOIETIC STEM/PROGENITOR CELLS 176 HYPOTHETICAL PROTEIN FLJ22028	0.81 0.98	0.49 0.54	0.58 0.50
SRV_04743_a_at SRV 02065 a at	ADAPTOR-RELATED PROTEIN COMPLEX 2, MU 1	0.98	0.54 0.59	0.50 0.48
SRV 03218 a at	SUBUNIT GABA(A) RECEPTOR-ASSOCIATED PROTEIN-LIKE 2	0.90	0.28	0.40
SRV 05300 a at	SFT2 DOMAIN CONTAINING 2	0.93	0.28	0.37
SRV 05537 a at	TRAFFICKING PROTEIN PARTICLE COMPLEX 6B	0.95	0.49	0.23
SRV_02033_a_at	SELENIUM BINDING PROTEIN 1	1.05	0.58	0.45
protein binding/modification	humethatical protein FL 111000	0.69	0.44	0.50
SRV_04235_at SRV 02678 a at	hypothetical protein FLJ11280 M-PHASE PHOSPHOPROTEIN 6	0.68 0.84	0.44 0.45	0.56 0.51
SRV 02198 a at	GLUTAMYL-PROLYL-TRNA SYNTHETASE	0.84	0.43	0.31
SRV 01481 at	PHOSPHATIDYLETHANOLAMINE BINDING PROTEIN 1	1.05	0.56	0.49
SRV 01495 at	PYRUVATE DEHYDROGENASE KINASE, ISOZYME 2	1.04	0.41	0.31
SRV_04077_a_at	UBIQUITIN-CONJUGATING ENZYME E2R 2	0.78	0.51	0.47
SRV_04977_s_at	CNDP DIPEPTIDASE 2 (METALLOPEPTIDASE M20 FAMILY)	1.27	0.49	0.22
SRV 04977 at	CNDP DIPEPTIDASE 2 (METÁLLOPEPTIDASE M20	1.20	0.46	0.22
	FAMILY) UBIQUITIN-CONJUGATING ENZYME E2B (RAD6			
SRV_01825_at	HOMOLOG)	0.90	0.47	0.51
RNA binding/processing		0 00	0 47	0.54
SRV_03823_at	RNA BINDING MOTIF PROTEIN, X-LINKED 2 SYF2 HOMOLOG, RNA SPLICING FACTOR (S.	0.80	0.47	0.54
SRV_03721_at	CEREVISIAE)	0.96	0.46	0.49
SRV_03417_at	MITOCHONDRIAL RIBOSOMAL PROTEIN S28	0.86	0.51	0.47
SRV_03836_at	EXOSOME COMPONENT 1	1.03	0.46	0.46
cell cycle/ cell division	COILED-COIL DOMAIN CONTAINING 5 (SPINDLE			
SRV_05218_a_at	ASSOCIATED)	0.68	0.36	0.46
SRV_03244_a_at	FREQUENTLY REARRANGED IN ADVANCED T-CELL LYMPHOMAS 2	0.89	0.38	0.46
SRV_05024_at	ZW10 INTERACTOR	0.96	0.41	0.44
SRV_00804_at	CDC6 CELL DIVISION CYCLE 6 HOMOLOG (S. CEREVISIAE)	0.70	0.50	0.42
SRV_03256_at	TPX2, MICROTUBULE-ASSOCIATED, HOMOLOG (XENOPUS LAEVIS)	0.91	0.41	0.51
SRV 04156 at	CELL DIVISION CYCLE ASSOCIATED 8	0.71	0.43	0.50
SRV_03593_at	DISCS, LARGE HOMOLOG 7 (DROSOPHILA)	0.66	0.39	0.49
SRV_14350_at	NIMA (NEVER IN MITOSIS GENE A)-RELATED KINASE 3	0.96	0.42	0.30
SRV_02556_at	SMC4 STRUCTURAL MAINTENANCE OF CHROMOSOMES 4-LIKE 1 (YEAST)	0.69	0.43	0.47
SRV_03257_at	TPX2, MICROTUBULE-ASSOCIATED, HOMOLOG (XENOPUS LAEVIS)	0.79	0.35	0.45
SRV_02235_at	KINESIN FAMILY MEMBER 11	0.53	0.31	0.44
SRV_01290_at	FERRITIN, HEAVY POLYPEPTIDE 1	1.00	0.54	0.42
SRV_02151_a_at	CENTRIN, EF-HAND PROTEIN, 2	0.65	0.39	0.39
SRV_04253_a_at	NUCLEOLAR AND SPINDLE ASSOCIATED PROTEIN 1 CYCLIN-DEPENDENT KINASE INHIBITOR 2C (P18,	0.62	0.28	0.37
SRV_05141_at	INHIBITS CDK4)	0.85	0.57	0.37
SRV_00033_copy4_at	T-cell acute lymphocytic leukemia 1	1.02	0.47	0.36
SRV_00033_at SRV_00033_copy2_at	T-cell acute lymphocytic leukemia 1 T-cell acute lymphocytic leukemia 1	0.97 0.96	0.44 0.46	0.35 0.34
SRV_00033_copy2_at SRV_00033_copy1_at	T-cell acute lymphocytic leukemia 1	0.96	0.46 0.44	0.34 0.33
SRV_00033_copy3_at	T-cell acute lymphocytic leukemia 1	0.94	0.44	0.33
membrane				
SRV_04260_at	CHROMOSOME 9 OPEN READING FRAME 46	1.05	0.56	0.47
SRV_04763_at	CHROMOSOME 10 OPEN READING FRAME 57	1.24	0.46	0.42
SRV_04650_a_at	TRANSMEMBRANE 6 SUPERFAMILY MEMBER 1	1.00	0.47	0.32
SRV_05571_a_at	OXIDATION RESISTANCE 1 TRANSLOCASE OF OUTER MITOCHONDRIAL	0.77	0.37	0.42
SRV_03611_a_at	MEMBRANE 70 HOMOLOG A (YEAST)	0.97	0.55	0.48
kinase activity				
SRV_05333_at	RIO kinase 3 (yeast)	1.06	0.47	0.52
SRV 05450 a at	INTEGRIN BETA 1 BINDING PROTEIN 3	0.87	0.53	0.46

SRV_01863_at	VACCINIA RELATED KINASE 1	0.61	0.39	0.51
pinocytosis/ endocytosis	DISABLED HOMOLOG 2, MITOGEN-RESPONSIVE			
SRV_00866_at	PHOSPHOPROTEIN (DROSOPHILA)	0.84	0.49	0.43
	· · · · · · · · · · · · · · · · · · ·			
DNA damage		0.75	0.44	0.07
SRV_04199_at SRV 02222 at	NEI ENDONUCLEASE VIII-LIKE 3 (E. COLI) HUS1 CHECKPOINT HOMOLOG (S. POMBE)	0.75 0.72	0.41 0.45	0.37 0.57
3HV_02222_at	HOST CHECKFOINT HOMOLOG (S. FOMBE)	0.72	0.45	0.57
muscle development/				
contraction				
SRV_01033_a_at	INTERFERON-RELATED DEVELOPMENTAL REGULATOR 1	1.18	0.38	0.53
	MYOSIN, LIGHT POLYPEPTIDE 1, ALKALI; SKELETAL,			
SRV_05143_a_at	FAST	0.79	0.50	0.49
SRV_00058_s_at	tropomyosin	0.89	0.53	0.48
other				
	FICOLIN (COLLAGEN/FIBRINOGEN DOMAIN			
SRV_01932_a_at	CONTAINING) 3 (HAKATA ANTIGEN)	1.03	0.58	0.46
SRV_05356_s_at	FAMILY WITH SEQUENCE SIMILARITY 58, MEMBER A	0.89	0.46	0.47
SRV_02972_at	GLUTAREDOXIN 5 HOMOLOG (S. CEREVISIAE)	0.97	0.49	0.49
SRV_05356_at	FAMILY WITH SEQUENCE SIMILARITY 58, MEMBER A	1.09	0.58	0.58
SRV_05263_at	SOLUTE CARRIER FAMILY 39 (ZINC TRANSPORTER), MEMBER 3	0.73	0.45	0.51
SRV 04739 a at	ZINC FINGER PROTEIN 403	0.97	0.49	0.51
SRV 03830 at	SHWACHMAN-BODIAN-DIAMOND SYNDROME	0.94	0.47	0.46
SRV 03448 at	COILED-COIL DOMAIN CONTAINING 59	0.89	0.42	0.44
SRV 02223 a at	ISOPENTENYL-DIPHOSPHATE DELTA ISOMERASE 1	0.89	0.37	0.43
SRV 04160 at	SDA1 DOMAIN CONTAINING 1	0.88	0.48	0.42
SRV_05216_at	SERUM AMYLOID A-LIKE 1	0.82	0.42	0.42
SRV_04991_a_at	MYC INDUCED NUCLEAR ANTIGEN	0.90	0.41	0.39
SRV_00134_a_at	ARGININOSUCCINATE LYASE	1.10	0.44	0.36
SRV_05376_at	WILLIAMS BEUREN SYNDROME CHROMOSOME REGION 27	0.84	0.36	0.31

### **Table 3**. Significant ( $p \le 0.05$ geometric mean p-value) functional groups obtained from

functional annotation using DAVID.

		Number				Numbe	
		of				r of	p-
	Ontology	Genes	p-value		Ontology	Genes	value
Functional				Functional	water-soluble vitamin		
Group 1	response to biotic stimulus	25	<0.001	Group 5	metabolism	4	0.02
<0.001	immune response	21	<0.001	0.033	vitamin metabolism pyridine nucleotide	4	0.03
	defense response	22	<0.001		metabolism	3	0.07
Functional				Functional	di-, tri-valent inorganic cation		
Group 2	cation binding	34	0.001	Group 6	transport	5	0.006
0.002	ion binding	36	0.004	0.042	metal ion transport	5	0.06
	metal ion binding	36	0.004		cation transport	6	0.22
Functional				Functional			
Group 3	innate immunity	4	0.001	Group 7	bcr protein	3	0.02
0.005	immune response	5	0.004	0.045	molecular chaperone	4	0.02
	innate immune response	4	0.007		Heat shock protein Hsp70	3	0.03
	complement activation	3	0.02		Heat shock protein 70 antigen processing and	3	0.03
					presentation	4	0.09
					cell surface	3	0.20
Functional						-	
Group 4	lysosome	7	0.002	Functional			
0.008	lysosome	7	0.009	Group 8	response to unfolded protein	5	0.02
	lytic vacuole	7	0.009	0.048	response to protein stimulus	5	0.02
	vacuole	7	0.02		chaperone	6	0.26

Gene name	Micro	oarray	qP	CR
	24	72	24	72
Myxovirus resistance 1 MX1	3.13	29.23	5.97	24.44
Macrophage receptor with collagenous structure MARCO	2.17	8.20	3.36	15.24
Complement component 3 C3	1.80	9.79	2.14	14.78
Cyclin dependant kinase inhibitor 1B CDKN1B	-1.17	-1.74	-2.26	-3.16
Vaccinia related kinase 1 VRK1	-1.64	-2.58	-1.23	-1.88
Serine dehydratase like SDSL	1.03	-1.92	-1.35	-1.01
Hemoglobin gamma alpha HBG1	-1.01	-1.02	-1.54	-1.87
Glycogen synthase kinase GSK3A	-1.07	-1.08	-1.13	2.76
Programmed cell death 8 PDCD8	1.13	-1.10	1.30	1.85

**Table 4**. Fold changes obtained from microarray and from quantitative real-time PCR.

Gene ID	Gene Name	Gene Ontology
	protective protein for beta-galactosidase	
SRV_00294_s_at	(galactosialidosis)	proteolysis, protein transport
	transglutaminase 1 (K polypeptide epidermal type	membrane, cell envelope, protein
SRV_00309_at	I, protein-glutamine-gamma-glutamyltransferase)	modification
SRV_00327_a_at	cathepsin K (pycnodysostosis)	proteolysis
	cytochrome b-245, beta polypeptide (chronic	humoral response, inflammatory
SRV_00330_at	granulomatous disease)	response
SRV_00371_a_at	fructose-1,6-bisphosphatase 1	metal ion binding (zinc)
51( V_005 / 1_u_u	solute carrier family 11 (proton-coupled divalent	metar fon omding (zme)
SRV_00442_at	metal ion transporters), member 1	immune response, ion transport
SK v _00++2_at	transcobalamin I (vitamin B12 binding protein, R	minute response, for transport
SDV 00712 a at	binder family)	ion transport/hinding (ashalt)
SRV_00713_a_at	Under family)	ion transport/binding (cobalt)
CDV 00744 a at	a dimage differentiation related motoin	fatty acid transport, extracellular
SRV_00744_a_at	adipose differentiation-related protein	region
SRV_01179_a_at	CD63 antigen (melanoma 1 antigen)	endosome, membrane
SRV_01342_at	interferon regulatory factor 1	immune response, transcription
SRV_01351_at	jun B proto-oncogene	transcription
SRV_01818_at	thioredoxin	signal transduction
	matrix metalloproteinase 9 (gelatinase B, 92kDa	extracellular matrix, apoptosis,
SRV_02399_a_at	gelatinase, 92kDa type IV collagenase)	proteolysis
SRV_02456_at	gastric intrinsic factor (vitamin B synthesis)	ion transport/binding (cobalt)
	heat shock 70kDa protein 5 (glucose-regulated	anti-apoptosis, endoplasmic
SRV_02516_at	protein, 78kDa)	reticulum
	lectin, galactoside-binding, soluble, 3 binding	cell adhesion, cellular defense
SRV_02586_at	protein	response, signal transduction
SRV_03054_at	macrophage receptor with collagenous structure	signal transduction
		innate immune response, regulation
SRV_04604_at	interferon induced with helicase c domain 1	of apoptosis, response to virus
SRV_04819_a_at	transmembrane protein 49	membrane, endoplasmic reticulum
	UDP-GlcNAc:betaGal beta-1,3-N-	,,
SRV_04888_a_at	acetylglucosaminyltransferase 5	membrane, CNS development
SRV 04911 at	thioredoxin domain containing 2 (spermatozoa)	cell redox homeostasis
5 <b>K</b> v _04711_at	unoredoxin domain containing 2 (spermatozoa)	nucleotide binding, protein
<b>SPV</b> 04064 a at	tubulin beta MGC4083	polymerization
SRV_07726_a_at	macrophage expressed gene 1	none
CDV 11417 - 4	matrix metalloproteinase 1 (interstitial	anoto classic ion kin dia ( -in -)
SRV_11417_a_at	collagenase)	proteolysis, ion binding (zinc)
0011 11/22	tissue inhibitor of metalloproteinase 1 (erythroid	
SRV_11663_a_at	potentiating activity, collagenase inhibitor)	enzyme inhibitor, cell proliferation
	protective protein for beta-galactosidase	
SRV_00294_s_at	(galactosialidosis)	proteolysis, protein transport

**Table 5**. Genes expressed in both ATV infection and spinal cord injury

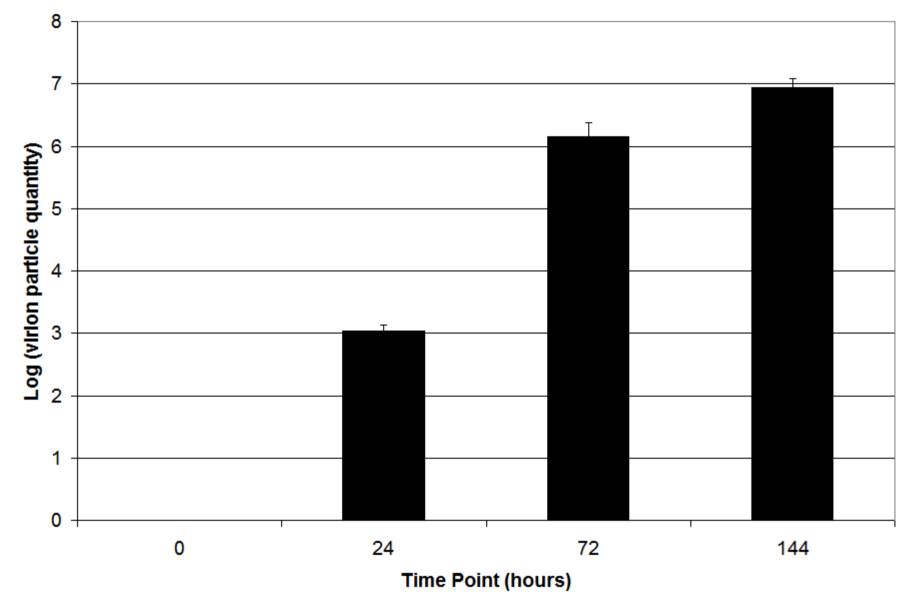
### **Additional files**

Additional file 1

Format: doc

Title: Appendix A

**Description:** Primer sequences used for qPCR verification of microarray data.



-3.0 1:1

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24 hours 72 hours hours

Figure

4

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#### -3.0 1:1 3.0 hours 24 t 72 h 144 SOM Vector

Figure 3

hours

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PHOSPHOGLUCONATE DEHYDROGENASE "AHA1, ACTIVATOR OF HEAT SHOCK 90KDA PROTEIN ATPASE HOMOLOG 1 (YEAST)" "ATPASE, H+ TRANSPORTING, LYSOSOMAL 56/58KDA, V1 SUBUNIT B2" MITOCHONDRIAL RIBOSOMAL PROTEIN S7 INTERFERON-RELATED DEVELOPMENTAL REGULATOR 1 RIO kinase 3 (yeast) HEMATOPOIETIC STEM/PROGENITOR CELLS 176 RIO kinase 3 (yeast) HEMATOPOIETIC STEM/PROGENITOR CELLS 176 PEPTIDE DEFORMYLASE-LIKE PROTEIN ZINC FINGER PROTEIN 403 INTERFERON REGULATORY FACTOR 2 ECTONUCLEOSIDE TRIPHOSPHATE DIPHOSPHOHYDROLASE 1 HYPOTHETICAL PROTEIN FL32028 PHOSPHATIDYLETHANOLAMINE BINDING PROTEIN 1 "TPX2. MICROTUBULE-ASSOCIATED, HOMOLOG (XENOPUS LAEVIS)" UBIQUITIN-CONJUGATING ENZYME 228 (RAD6 HOMOLOG) TRAF-TYPE ZINC FINGER POMAIN CONTAINING 1 GLUTAREDOXIN 5 HOMOLOG (S. CEREVISIAE) "MAKORIN, RING FINGER PROTEIN, 1" "SYE2 HOMOLOG, RNA SPLICING FACTOR (S. CEREVISIAE)" "RNA BINDING MOTIF PROTEIN, X-LINKED 2" "RNA BINDING MOTIF PROTEIN, X-LINKED 2" "RNA BINDING MOTIF PROTEIN MICOTTAINOT 1 GLUTATHIONE S-TRANSFERASE OMEGA 1 "SPECTRIN, ALPHA, NON-ERYTHROCYTIC 1 (ALPHA-FODRIN)" M-PHASE PHOSPHOPROTEIN 8 TRANSLOCASE OF OUTER MITOCHONDRIAL MEMBRANE 70 HOMOLOG A (YEAST) PHOSPHOGULCOMUTASE 2 GLUTATMICAL PROLYL-TRNA SYNTHETASE EXOSOME 200FPONENT 1 FIGOLIN (COLLAGEN/FIBRINOGEN DOMAIN CONTAINING) 3 (HAKATA ANTIGEN) FICOLIN (COLLAGEN/FIBRINOGEN DOMAIN CONTAINING) 3 (HAKATA ANTIGEN) PIDULIN (BULLAGENPIDENINGEN DOMAIN CONTAINING (CLASSING) (CLASSING tropomyosin MITRILASE FAMILY, MEMBER 2" "INTRILASE FAMILY, MEMBER 2" "MYOSIN, LIGHT POLYPEPTIDE 1, ALKALI; SKELETAL, FAST" "NON-METASTATIC CELLS 7, PROTEIN EXPRESSED IN (NUCLEOSIDE-DIPHOSPHATE KINASE)" ZW10 INTERACTOR "NON-METASTATIC CELLS 7, PROTEIN EXPRESSED IN (NUCLEOSIDE-DIPHOSPHATE K ZW10 INTERACTOR LIPOIC ACID SYNTHETASE COILED-COLL DOMAIN CONTAINING 59 hypothetical protein FLJ11280 "PRKC, APOPTOSIS, W11, REGULATOR" CHROMOSOME 10 OPEN READING FRAME 57 "SOLUTE CARRIER FAMILY 39 (ZINC TRANSPORTER), MEMBER 3" ZINC FINGER PROTEIN 282 CHROMOSOME X OPEN READING FRAME 15 UBIQUITIN-CONJUGATING ENZYME E2R 2 "FERRITIN, HEAVY POLYPEPTIDE 1" CASP2 AND RIPKI DOMAIN CONTAINING ADAPTOR WITH DEATH DOMAIN MEMBRANE-ASSOCIATED RING FINGER (C3HC4)7 ISOPENTENYL-DIPHOSPHATE DELTA ISOMERASE 1 CELL DIVISION CYCLE ASSOCIATED, HOMOLOG (XENOPUS LAEVIS)" SDA1 DOMAIN CONTAINING EN SDA1 DOMAIN CONTAINING 1 "DISABLED HOMOLOG 2, MITOGEN-RESPONSIVE PHOSPHOPROTEIN (DROSOPHILA)" SERUM AWYLOID ALIKE 1 HYPOTHETICAL PROTEIN FLI20899 RING FINGER PROTEIN 113A SMC4.STRUCTURAL MAINTEMANCE OF CHROMOSOMES 4-LIKE 1 (YEAST) RING FINGER FROTEIN TI3A SMC4 STRUCTURAL MAINTENANCE OF CHROMOSOMES 4-LIKE 1 (YEAST) MYC INDUCED NUCLEAR ANTIGEN "DISCS, LARGE HOMOLOG 7 (DROSOPHILA)" OXIDATION RESISTANCE 1 COILED-COIL DOMAIN CONTAINING 5 (SPINDLE ASSOCIATED) Toell active lymphocytic leukemia 1 GABA(A) RECEPTOR-ASSOCIATED PROTEIN-LIKE 2 ARGININOSUCCINATE LYASE VACCINIA RELATED KINASE 1 GUANINE DEAMINASE VACCINIA RELATED KINASE 1 GUANINE DEAMINASE "PRIMASE, POLYPEPTIDE 1, 40KDA" "CYCLIN-DEPENDENT KINASE INHIBITOR 2C (P18, INHIBITS CDK49" T-cell acute lymphocytic leukemia 1 CDC8 CELL DIVISION CYCLE 6 HOMOLOG (S. CEREVISIAE) T-cell acute lymphocytic leukemia 1 TRAFFICKING PROTEIN PARTICLE COMPLEX 6B T-cell acute lymphocytic leukemia 1 SERINE DEHYDRATASE-LIKE NEI ENDONUCLEASE VIILLIKE 3 (E. COLI) TRANSMEMBRANE 6 SUPERFAMILY MEMBER 1 T-cell acute lymphocytic leukemia 1 "PYRUVATE DEHYDROGENASE KINASE, ISOZYME 2" EUKARYOTIC TRANSLATION INITIATION FACTOR 4E BINDING PROTEIN 3 "CENTRIN, EF-HAND PROTEIN, 2" NIMA (NEVER IN MITOSIS GENE A); RELATED KINASE 3 MITOCHONDRIAL, INTERMEDIATE PEPTIDASE WILLIAMS BEUREN SYNDROME CHROMOSOME REGION 27 SFT2 DOMAIN CONTAINING 2 NUCLEOLAR AND SPINDLE ASSOCIATED PROTEIN 1 KINSIN FAMILY MEMBER 11 "AMINOLEVULIATE, DELTA, DEHYDRATASE" "ATP SYNTHASE, H+ TRANSPORTING, MITOCHONDRIAL FD COMPLEX, SUBUNIT S (FACTOR B)" CNDP DIPEPTIDASE 2 (METALLOPEPTIDASE M20 FAMILY) ASPARTOACYLASE (CANAVAN DISEASE)

### Additional files provided with this submission:

Additional file 1: supp1.doc, 21K http://www.biomedcentral.com/imedia/2004261874218210/supp1.doc